

Behavioural responses to ectoparasites in pied flycatchers *Ficedula hypoleuca*: an experimental study

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Nests of cavity-nesting birds usually harbor some species of haematophagous ectoparasites that feed on the incubating adults and nestlings. Given the negative impact of ectoparasites on nestlings there will be selection on hosts to reduce parasite infestations through behavioural means. We have experimentally reduced the abundance of all ectoparasites in nests of pied flycatchers *Ficedula hypoleuca* to explore both whether there are changes in the frequency and duration of putative anti-parasite behaviours by tending adults, as well as whether such anti-parasite behaviours are able to compensate for the deleterious effects that parasites may have on nestlings. Heat treatment of nests substantially decreased the density of ectoparasites, and thereby positively affected nestling growth. The frequency and intensity of female grooming and nest sanitation behaviours during the incubation and nestling periods decreased as a consequence of the experimental reduction of ectoparasite infestation. Although nestlings begged more intensely in infested nests, the experiment had no significant effect on parental provisioning effort. Reduction of parasites resulted in larger nestlings shortly before fledging and increased fledging success. This study shows a clear effect of a complete natural nest ectoparasite fauna on parental behaviour at the nest and nestling growth in a cavity-nesting bird. Although ectoparasites induce anti-parasite behaviours in females, these behaviours are not able to fully remove parasite's deleterious effects on nestling growth and survival.

Parasites have been proposed as an important ecological and evolutionary force affecting avian life histories and behaviour (Atkinson and van Riper 1991, Møller 1997). Cavity-nesting birds have been traditionally associated with selective pressures arising from the thermal environment and the impact of nest predation (Hansell 2000). However, the microclimatically stable environment of cavity nests and the presence of an abundant food supply may offer excellent breeding conditions for ectoparasites, and it is thought that many nest ectoparasitic arthropods have evolved specifically in such nesting environments (Waage 1979, Marshall 1981). Thus, ectoparasites in the nest may be an additional important evolutionary factor modulating adaptations of cavitynesting birds (Heeb et al. 1999, Tripet et al. 2002a).

Ectoparasites cause removal of nutritional and energy resources from hosts that could otherwise be used for growth, maintenance, or reproduction (Møller 1993). They may also induce costly immune and inflammatory responses (Møller et al. 2005, Owen et al. 2009). Conversely the immature immune systems of altricial nestlings result in stronger direct impacts from ectoparasitism faced by nestlings with the need to assign sufficient nutritional resources to growth (Saino et al. 1998, Szep and Møller 1999). Given the negative impact of ectoparasites on host fitness there will be selection on hosts to avoid parasite infestations through behavioural, physiological and immunological responses (Møller and Erritzoe 1996, Hart 1997, Heeb et al. 1998, Cantarero et al. 2013). All these responses are complementary and may be induced in adults, nestlings or both (Keymer and Read 1991, Hart 1992, Simon et al. 2005). Nesting adults may avoid nest sites with high ectoparasite loads (Moore 2002), due to the association between old nest material and higher abundance of certain types of ectoparasites (Mazgajski 2007, Tomás et al. 2007, López-Arrabé et al. 2012) and bacteria (González-Braojos et al. 2012). Adults may also take measures to indirectly minimize the effects of nest parasites through incorporation of fresh plant material containing compounds that either directly affect the development of parasites (Clark and Mason 1988, Lafuma et al. 2001, Malan et al. 2002, Tomás et al. 2012) or stimulate elements of the immune system of chicks that help them to cope better with the harmful activities of ectoparasites (Mennerat et al. 2008). Nevertheless, adult cavity-nesting birds are faced with the presence of nest ectoparasites, and likely have a suite of behaviours directed a minimizing the impacts of parasites (Keymer and Read 1991, Loye and Zuk 1991, Hart 1992, Mazgajski 2009).

The main behavioural defenses against ectoparasites are grooming and nest sanitation (Christe et al. 1996). Grooming behaviour may be operationally defined as manipulation of the plumage with the bill (Nelson et al. 1977, Murray 1990). One of its functions may be to dislodge ectoparasites hiding or residing among feathers (Cotgreave and Clayton 1994, Waite et al. 2012). Thus both adults and nestlings may groom themselves in the presence of ectoparasites (O'Connor et al. 2010). Nest sanitation (Welty 1982) refers to behaviours by parents in altricial species tending to remove ectoparasites on nestlings or nest material (Hurtrez-Boussès et al. 2000), removing from the nest both these as well as eggshells (Montevecchi 1974), fecal material (Blair 1941) or dead nestlings (Skutch 1976). Parents are expected to allocate time to nest sanitation in order to control the load of harmful ectoparasites in the nest materials and on the nestlings. Such anti-parasite behaviours may be time-consuming (Cotgreave and Clayton 1994) and therefore may reduce the time that a parent bird can devote to foraging and provisioning offspring.

Nest ectoparasites are a community of species, and the entire community must be considered when examining the influences of these ectoparasites on host behaviour and fitness. For example the nests of pied flycatchers Ficedula hypoleuca in Iberian populations usually harbour three species of haematophagous ectoparasites, namely mites Dermanyssus gallinoides, blowflies larvae Protocalliphora azurea and hen fleas Ceratophyllys gallinae (Merino and Potti 1995, 1996). Nests may also contain haematophagous flying insects such as blackflies and biting midges (Martínez-de la Puente et al. 2009). In mites, adult and nymphal stages are haematophagous, while their short generation times allow the build-up of very large populations with detrimental effects on host reproductive success (Merino and Potti 1995, 1996, Moreno et al. 2009). Mites may be present in nest materials even before nestlings hatch and may feed on incubating females (Pacejka et al. 1996). They have been shown to be the most virulent ectoparasite of Iberian pied flycatcher populations (Merino and Potti 1995, Moreno et al. 2009). Blowfly larvae live in bird nests and feed intermittently on nestling blood (Bennett and Whitworth 1991, Remeš and Krist 2005). Larvae of fleas are not haematophagous, but adult fleas need blood to produce eggs (Tripet and Richner 1997). Therefore, the number of flea larvae in nests indicates the fecundity of adult fleas (Eeva et al. 1994). Fleas may be present in nest materials already during incubation (Harper et al. 1992).

To explore behavioural anti-parasite strategies it is necessary to conduct field experiments where the levels of infestation are strictly controlled in all treatments (Christe et al. 1996, Heeb et al. 1998, Tripet et al. 2002b, Fitze et al. 2004) or experiments in which nests with a reduced ectoparasite loads are compared with natural controls (Allander 1998, Martínez-de la Puente et al. 2010). Each approach has its advantages and disadvantages. Controlled levels of infestation are useful when dealing with a single parasite and reduce environmentally induced variation. On the other hand, natural controls versus experimental reductions allow manipulations of complete ectoparasite faunas while retaining natural levels of infestation as controls, and are especially useful when the effects of whole ectoparasite faunas with their natural interactions are of interest. Moreover, natural controls reflect the effects of whole ectoparasite faunas on nestlings in the wild.

Our goal is to examine the impacts of an entire nest ectoparasite community on reproductive behaviour of their hosts. In our study of pied flycatchers in central Spain, we have reduced the abundance of all ectoparasites by a heat treatment of nestboxes. We have assumed that experimental reduction would have a negative impact on ectoparasite abundance and a positive impact on nestling growth and survival. We have then compared control and experimental host behaviour within the nest-box using data from video films (see Hurtrez-Boussès et al. 2000 for a similar approach). Video-recordings inside the nest-box were made during the incubation and at two stages of the nestling period (nestlings of 3 and 9 d of age). Our objectives were to explore changes in the frequency and duration of parental grooming and nest sanitation behaviours as a consequence of the abundance of ectoparasites, and to examine the impacts of these behaviours of adult birds. We have hypothesized that: 1) behavioural responses to ectoparasites should be more frequent in control nests than in experimental nests. This pattern should occur during both the incubation and nestling periods; 2) there should be a trade-off between brooding nestlings and nest sanitation behaviours at the early nestling stage; 3) nestlings should beg more intensely in control nests due to the increased food demand induced by ectoparasites; 4) parents should respond to higher begging levels in control nests by increasing provisioning rates only if time consumed by anti-parasite behaviours does not compromise that available for foraging.

Material and methods

General field methods

The study was conducted during the spring of 2012 in a montane forest of Pyrenean oak *Quercus pyrenaica*, at 1200 m a.s.l. in Valsaín, central Spain (40°54′N, 4°01′W) where pied flycatchers breeding in nest-boxes have been studied since 1991 (see Sanz et al. 2003 for general description). Of 552 nest-boxes, 91 were occupied by pied flycatchers and the rest by other species, mainly great tits, nuthatches and blue tits (see Lambrechts et al. 2010 for dimensions, structure and placement of nest-boxes).

Egg laying in the pied flycatcher population under study typically begins in late May, and modal clutch size is six. The female incubates and broods alone and receives part of her food from her mate (Moreno et al. 2011). No brooding is observed after nestlings attain 7 d of age (Sanz and Moreno 1995). Breeding activities are followed routinely every year and laying and hatching dates and brood sizes at hatching and fledging are determined.

On day 3 (hatching day = day 1), we weighed all nestlings in each brood together with a digital scale to the nearest 0.1 g to give an average nestling mass when divided by brood size. On day 13 (hatching day = day 1), we ringed nestlings and measured their tarsus lengths with a digital callipers to the nearest 0.01 mm and their wing lengths with a stopped ruler to the nearest mm. Nestlings were also weighed with a Pesola spring balance to the nearest 0.25 g. Nestlings from one nest flew before being measured on day 13.

Experimental reduction of ectoparasites protocol

Of the 91 nest boxes occupied by pied flycatchers we selected those whose laying date was between dates 45 and 51 (1 April = day 1). We assigned 56 nests randomly to two

groups, to minimize any possible confounding effects such as variation in microclimate among nest-boxes. The first group was left unmanipulated (control group, n = 37). In the second experimental group (n = 19), we reduced the number of ectoparasites by a heat-treatment for 30 s at 750 W using a microwave oven. For the time that the original nests were treated (around 30 min), a fresh substitute nest was introduced into the nest-box (these nests had been collected in previous seasons after being abandoned prior to laying and kept frozen at -20° C until use). This treatment ensured that experimental nests did not contain live arthropods when placed in the nest-box (Rendell and Verbeek 1996), although some arthropods may colonize the nest material after the treatment. To avoid the loss of water during the heat-treatment, the nests were placed into a hermetic plastic container. To prevent recurrence of ectoparasite colonization a total of three heat-treatments were made in the experimental group: 1) 7 d after clutch completion, 2) when nestlings were 2 d old (hatching day = day 1) and 3) when nestlings were 8 d old. Furthermore, before returning the nest the flame from a butane jet torch lighter (Microtorch GT-3000) was passed across the walls of the nest-box to kill ectoparasites that might remain there. Nests in the control group were visited on the same days and handled in a similar way to experimental ones.

Ectoparasite abundance estimation

One or two days after nestlings fledged (17 d after hatching), all nests were removed in sealed plastic bags and taken to the laboratory, where they were subjected to arthropod removal in Berlese funnels for 48 h until nests were thoroughly dried and no arthropods were moving in the nest material. Ectoparasite identification was made with the aid of a stereoscopic microscope (Olympus SZX7). We assume that all mites are hematophagous given their red color as a consequence of the ingested blood (for arthropods collection and abundance estimations see Moreno et al. 2009).

Video recordings

Seven days after clutch completion (day 7 of incubation), we recorded nest activity inside nest-boxes for about 90 min $(91.45 \pm SE \ 24.63 \text{ min}, n = 58)$ with a cold white light (LED 5 mm) powered by a 3 V battery and a camera (GoPro HD Hero1) mounted on the roof inside the nest-box. Video recordings were made one day after experimental treatment and nest handling. Nest-boxes were again recorded two days after the day of hatching of the young $(88.63 \pm 13.01 \text{ min})$, n = 57) and 8 d after hatching of the young (85.04 \pm 20.01 min, n = 55). In two nests all chicks died after day 3 but we have included records taken during incubation and day 3. All films were recorded between 08:00-17:00 h, and no differences between experimental groups with respect to time of filming were found (incubation period: U=297.0, p = 0.346; nestling period day 3: U = 314.5, p = 0.522; nestling period day 9: U = 272.5, p = 0.277). We excluded the time until the first nest visit by parents (14.31 ± 11.55) min, n = 164). No evidence of stress or unnatural behaviour like extremely long absence periods from the nest or trying to peck at the camera system were observed after the first visit.

Because of technical problems, we failed to record the behaviour at two different nests in the control group, one from young nestlings and another from older nestlings.

Behavioural data analysis

From recordings taken during incubation we estimated the proportion of time spent by the female inside the nest-box or 'egg attendance' which includes the time allocated to incubating and turning the eggs, and the mean duration of incubation sessions and recesses. Furthermore, we monitored two specific types of female behaviour: 'grooming' and 'nest sanitation'. 'Grooming' is the combined time in which female spends preening or scratching herself (Cotgreave and Clayton 1994) while 'nest sanitation' is a period of active search with the head buried, sometimes deeply, into the nest material (Cotgreave and Clayton 1994). In our study, we define nest sanitation as burying the bill in the nest material or carrying out nest materials. As scratching by females resting on the nest could not be observed accurately, grooming refers mostly to preening with the bill. We assume that these behaviours in our study population have the functions implied by the terms derived from the literature, although our experiment intends precisely to confirm these functional interpretations. We obtained the proportion and the mean duration of these behaviours over the time that the female was inside the box. In addition, we also counted the number of incubation feedings by males.

From recordings during the early nestling phase we obtained hourly provisioning rates by males and females and the amount of time spent by females in 'nest attendance', 'brooding', 'grooming' or 'nest sanitation'. 'Nestlings attendance' includes the proportion of time spent by the female inside the nest-box. 'Brooding' activity is defined as the proportion of time spent by the female inside the nest-box covering young nestlings in relation to the total time spent inside the nest-box. 'Nest sanitation' and 'grooming' were calculated in the same way as during the incubation stage. Pied flycatcher females do not exhibit 'sleeping' behaviours during the incubation or nestling periods like in other species (Tripet et al. 2002b).

From recordings during the late nestling phase we obtained hourly provisioning rates by males and females and nest sanitation behaviour. Nest sanitation at this stage only considers removal of nest material from nest-boxes as the chicks do not need brooding and female visits to the nestbox are just for feeding. We also recorded the posture during begging of nestlings. Nestling postures were assigned based on a modification of the scale used by Leonard et al. (2003) following a scale of increasing intensity: 0 = head down, no gaping; 1 = head down, gaping, sitting on tarsi; 2 = head up, gaping, sitting on tarsi; 3 = same as 2, plus neck stretched upward; and 4 = same as 3, but body lifted off tarsi. On each visit of an adult to the nest with food, we scored the maximum postural begging intensity of each nestling. We then estimated the average value of the maximum begging intensity at each visit for the whole brood.

Statistical analyses

Many response variables were normally distributed or successfully normalized through logarithmic transformation

Table 1. Differences in breeding variables and ectoparasite abundances (means \pm SE, n in parenthesis) and results of GLM analyses and Mann–Whitney U-tests.

	Control	Experimental	Statistic	р
Breeding data				
Laying date	48.162 ± 1.642 (37)	47.386 ± 1.012 (19)	$F_1 = 3.70$	0.060
Hatching date	66.351 ± 1.230 (37)	66.947 ± 1.311 (19)	$F_1 = 1.30$	0.260
Clutch size	5.622 ± 0.594 (37)	5.84 ± 0.501 (19)	Wald = 0.107	0.743
Brood size 13 d	4.722 ± 1.446 (36)	5.263 ± 0.733 (19)	Wald = 0.741	0.389
Ectoparasites				
Blowflies	6.162 ± 8.748 (37)	0.684 ± 1.887 (19)	$U_1 = 172.0$	< 0.005
Mites	3347.57 ± 4543.55 (37)	274.053 ± 906.913 (19)	$F_1 = 17.76$	< 0.001
Fleas	24.946 ± 88.329 (37)	0.000 ± 0.000 (19)	$U_1 = 247.0$	< 0.01

(Kolmogorov-Smirnov, p > 0.20) and were therefore analyzed with GLM models (STATISTICA package) assuming a normal error with treatment as explanatory factor. Hatching success (proportion eggs that hatched) was not normally distributed even when transformed but its residuals were, so it was analyzed with a GLM with treatment as explanatory factor and laying date and clutch size as continuous predictors. Clutch size and brood size were analyzed with GLM models assuming a Poisson distribution with treatment as explanatory factor. Grooming and nest sanitation variables could not be normalized and were analyzed with nonparametric tests (Mann-Whitney U test). Nest sanitation (nestling day 9) and fledging success (proportion hatched chicks that fledged) were analyzed as frequencies (Yes-1/ No-0 observation of sanitation in the nest and Yes-1/ No-0 cases of all chicks hatched becoming fledglings) with Chi-squared contingency tables.

All parametric behavioural variables were analyzed with treatment as explanatory factor and hatching date, brood size, date and time of filming as continuous predictors. Nonsignificant predictors were sequentially removed until only significant effects remained in the final model. Only the effects of treatment are presented in all cases, even when non-significant.

Results

The two treatments did not differ with respect to laying date, hatching date, clutch size or brood size (Table 1). In order to assess the efficiency of our manipulation, we compared the ectoparasite abundances of the two treatments. The experiment was successful because the experimental nests differed from controls in the abundances of ectoparasites sampled in the predicted direction (Table 1).

The control (3.36 ± 0.49) and the experimental group (3.76 ± 0.61) differed in the mean nestling mass (g) on day 3 (*F*=6.87, p=0.011). We then tested for the effects of

treatment on brood-averaged nestling morphometric measurements and mass near fledging (tarsus length, wing length and body mass at day 13), controlling for hatching date and brood size. Nestlings in experimental nests were significantly larger with respect to wing length and tarsus length (Table 2), while there were no differences between treatments in body mass (Table 2). Tarsus length was negatively related to hatching date (adjusted R² = 0.25). The control (90.09 ± 14.13) and the experimental (90.30 ± 11.61) groups did not differ in hatching success (*F* = 0.003, p > 0.90) when controlling for laying date and clutch size (both p > 0.20), while fledging success was marginally lower (χ^2 = 2.82, p = 0.093) in control nests (0.89 ± 0.32) than in experimental nests (1.00 ± 0.00).

Treatment did not significantly affect incubation attendance (% incubation time) or the mean of recess and incubation session durations of females (Table 3). Female grooming behaviour was less frequent and the mean duration of grooming sessions were significantly shorter in the experimental group than in the control group (Table 3). Nest sanitation behaviour time was also more intensive in the control group (Table 3).

In relation to the second hypothesis, the proportion of brooding time on day 3 was significantly higher in the experimental group than in the control one as predicted (Table 3). Grooming variables showed the same pattern between treatments as during incubation (Table 3). Nest sanitation time was again higher in the control group (Table 3). There was no experimental effect on provisioning rates of males and females (all p > 0.2). Control nests where nest sanitation behaviours occurred showed higher mite infestations than control nests where these behaviours did not occur (Fig. 1). This relationship was not found for blowfly larvae or fleas. There were marginally more nest sanitation events in control nests than in experimental nests (Table 3).

In accordance with our third hypothesis, begging intensity of nestlings was higher in control nests (Table 3) and

Table 2. Differences in brood-averaged nestling morphology and mass (means \pm SE, number of broods in parenthesis) and results of GLM models on nestling condition parameters with treatment as explanatory factor and hatching date and brood size as continuous predictors (**p < 0.01, *p < 0.05).

	Control	Experimental	Treatment F	Hatching date F	Brood size F
Nestling day 13					
Tarsus length (mm)	17.42 ± 0.47 (34)	17.79 ± 0.38 (19)	6.615*	9.027**	0.133
Body mass (g)	13.98±0.94 (34)	14.17 ± 0.92 (19)	0.510	0.231	0.478
Wing length (mm)	46.85 ± 2.40 (34)	48.57 ± 2.32 (19)	7.126*	1.050	0.072

Table 3. Differences (means + SE, n in parenthesis) in behavioural variables between the two treatments and results of GLM analyse	ŝ
(significant p-values in bold), Mann–Whitney U-tests and Chi-squared contingency tables (sanitation present or absent).	

	Control	Experimental	Statistic	р
Incubation				
Grooming (%)	0.400 ± 0.586 (37)	0.122 ± 0.240 (19)	U = 182	0.012
Mean grooming (s)	3.00 ± 3.00 (37)	2.00 ± 3.00 (19)	U = 185	0.015
Egg attendance (%)	63.844 ± 11.74 (37)	60.750 ± 12.645 (19)	$F_1 = 0.81$	0.372
Mean session (min)	10.516 ± 4.433 (37)	9.317 ± 6. 167 (19)	$F_1 = 0.68$	0.412
Mean recess (min)	5.950 ± 2.000 (37)	4.217 ± 2.217 (19)	$F_1 = 1.55$	0.218
Nest sanitation (%)	1.905 ± 1.596 (37)	1.041 ± 1.289 (19)	$F_1 = 5.50$	0.029
Nestling day 3				
Nestling attendance (%)	53.274 ± 14.59 (36)	48.358 ± 13.106 (19)	$F_1 = 1.34$	0.254
Brooding (%)	93.045 ± 3.127 (36)	95.997 ± 3.981 (19)	$F_1 = 7.60$	0.008
Mean grooming duration (s)	2.00 ± 2.00 (36)	1.00 ± 2.00 (19)	$\dot{U} = 172$	0.048
Grooming (%)	0.108±0.143 (36)	0.047 ± 0.121 (19)	U = 168	0.039
Nest sanitation (NS) (%)	5.367 ± 2.562 (36)	2.879 ± 3.659 (19)	$F_1 = 7.80$	0.007
Mean NS duration (s)	5.00 ± 3.00 (36)	3.00 ± 2.00 (19)	$\dot{U} = 121$	< 0.005
Male provisioning (h^{-1})	9.298 ± 5.102 (36)	$11.108 \pm 4.566 \ (19)$	$F_1 = 1.62$	0.209
Female provisioning (h^{-1})	5.950 ± 3.076 (36)	5.791 ± 2.557 (19)	$F_1 = 0.03$	0.853
Nestling day 9				
Nest sanitation (yes/no)	0.176 ± 0.387 (34)	0.000 ± 0.000 (19)	$\chi^2 = 3.78$	0.052
Male provisioning (h^{-1})	11.934 ± 5.685 (34)	11.290 ± 5.268 (19)	$F_1 = 0.16$	0.688
Female provisioning (h ⁻¹)	11.384 ± 5.541 (34)	10.345 ± 5.074 (19)	$F_1 = 0.45$	0.503
Begging intensity score	1.149 ± 0.636 (34)	0.741 ± 0.376 (19)	$F_1 = 6.16$	0.016

was positively associated with parental provisioning rates (Fig. 2).

Contrary to our fourth hypothesis and despite the association with begging intensity male and female provisioning rates on day 9 were not related to treatment (Table 3).

Discussion

This study shows that the behaviour of pied flycatcher females inside the nest-box was clearly affected by ectoparasite abundance. The heat treatment of the nests decreased strongly the nest density of blowflies, mites and fleas, and

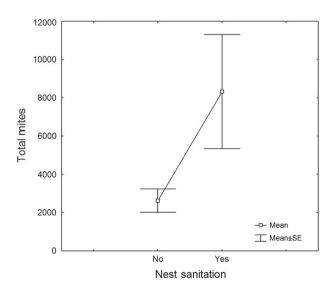


Figure 1. Total mites (\pm SE) in relation to the presence (Yes) or absence (No) of nest sanitation behaviour in the control group (p = 0.0045) in the late nestling phase.

thereby positively affected nestling growth. Experimental nests resulted in larger nestlings shortly before fledging. The frequency and intensity of female grooming and nest sanitation behaviours during the incubation and nestling periods decreased as a consequence of the experimental reduction of ectoparasite infestation. The experimental treatment did not affect incubation attendance and there were no effects on male or female provisioning rates to the chicks at any stage.

There is mixed evidence concerning the impact of ectoparasites on reproductive success in altricial cavitynesting birds. While some experimental studies have found strong deleterious effects on nestling growth and survival (Richner et al. 1993, Heeb et al. 1998), others have only found weak effects or none at all (Tripet et al. 2002b). These differences among host species and populations may

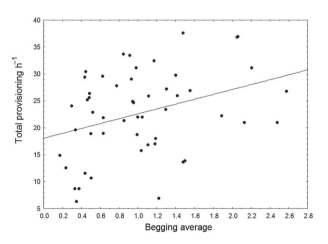


Figure 2. Association between hourly provisioning rates (male and female) and begging average intensity in the late nestling phase (Spearman correlation: r = 0.48, p < 0.005).

depend on the absolute levels of infestation found in different regions or habitats (Eeva et al. 1994, Fitze et al. 2004). Strong effects are thus mostly found in areas where climate is favourable for arthropod survival and dispersal during and between avian breeding seasons (Dufva and Allander 1996, Merino and Potti 1996). Reproductive success in Iberian pied flycatcher populations has been previously shown to suffer the impacts of nest-dwelling ectoparasites (Merino and Potti 1995, 1996, 1998, Merino et al. 1998, Moreno et al. 2009).

In agreement with several previous experimental studies we found marked effects of ectoparasites on nestling growth (Heeb et al. 2000, Tomás et al. 2008, Brommer et al. 2011). Tarsus and wing length of nestlings were negatively affected by ectoparasite abundance although we did not find an effect on body mass. Tarsus length of pied flycatcher nestlings has been related to their recruitment probability from fledging until breeding (Alatalo and Lundberg 1986), so the effects of ectoparasites may affect the future fitness of nestlings. For the observation that we found no effect of treatment on nestling body mass, there are two non-mutually exclusive alternative interpretations of our results. On the one hand, nestling growth improves under favourable conditions for breeding (Sanz 1995). Conditions during the year of study (2012) must have been especially favourable as nestlings attained their largest masses since the inception of the study (1991), which may explain why we found no effect of treatment on body mass. Additionally, control nestlings could reduce ectoparasite effects on body mass by increasing self preening or standing on top of one another (O'Connor et al. 2010).

Control nestlings showed as expected an increase in begging intensity, which is positively associated with parental provisioning rates, as was found in the great tit Parus major (Christe et al. 1996). Older nestlings suffering from higher ectoparasite loads begged more intensely as a response to their higher nutritional needs. Parental provisioning frequency depends on begging intensity as found in other studies (Kedar et al. 2000, Kölliker et al. 2000, Wright et al. 2002). Like Fitze et al. (2004) we noticed no effect of ectoparasite reduction on parental provisioning rates at any nestling age despite the strong correlation between begging intensity and provisioning rates. However, we found no effects of the experiment on parental provisioning frequency which could explain the lack of difference in the body mass of nestlings between treatments and it could have been caused by factors for which we did not control such as prey quality, ectoparasite virulence, nestling resistance or environmental constraints (Lehmann 1993, Møller 1994). Rogers et al. (1991) also showed no effect in parental provisioning frequencies in response to ectoparasites (see also Tripet et al. 2002b). There is also evidence that parental effort in pied flycatchers is energetically tightly constrained thereby precluding responses to variation in brood demand (Moreno et al. 1997, 1999). This lack of parental response may explain why their food provisioning was incapable of compensating for ectoparasite effects leading to smaller size at fledging in control nests. It is also possible that increased dedication to nest sanitation in control nests contributed to reduce the capacity of parents to augment their provisioning rates sufficiently to be detectable. That fledging success was marginally higher in experimental nests supports the existence of ectoparasite effects on nestling survival (Lundberg and Alatalo 1992, Moreno et al. 1999) although other fitness costs such as the probability of recruitment could be expressed after fledging (Thomas et al. 2007).

It is assumed that as a consequence of the negative impact of ectoparasites on nestlings, hosts have evolved behavioural responses (Cantarero et al. 2013). Ectoparasites present during incubation in pied flycatcher nests are mites and fleas. Females groom themselves more in control nests which may imply a direct response to the attachment of these ectoparasites on their skin and plumage. Nevertheless, grooming activity may not occupy sufficient time to constrain incubation attendance in females (Hurtrez-Boussès et al. 2000). However, our results on grooming behaviour indicate that tending females may suffer some costs induced by nest ectoparasites through attachment and possibly blood-sucking even before nestlings hatch (see also Tomás et al. 2008). When the nestlings hatch, the blowflies lay their eggs on their skin, and emerging larvae then begin feeding on nestling blood. The blood-sucking larvae of blowflies feed intermittently on the blood of nestling birds (Rognes 1991), although they may try to attach also to brooding females (Bennett and Whitworth 1991) given that their belly skin is naked at this stage. The combined effect of nest ectoparasites induced a lower body mass of chicks in control nests compared with treated nests already two days after hatching. This indicates that ectoparasites impair the growth of nestlings from hatching, a cost for which parents are apparently not able to compensate. If variation in parasite abundance is obvious to attending parents, we should expect that females in the control group compared to those in the treated group should allocate more time to anti-parasite behaviours and restrict the time spent on brooding chicks, sleeping (Tripet et al. 2002b) or foraging and provisioning nestlings (Christe et al. 1996). We found that control females reduced their proportion of time spent in the nest-box brooding compared to experimental females, but not with respect to total nestling attendance. The fact that females from control nests increased anti-parasite behaviours but maintained similar brooding attendances and provisioning rates as at experimental nests suggests that the time costs of these behaviours are not sufficiently important to reduce time available for foraging at this early stage of nestling development.

The function of nest sanitation behaviour by introducing the bill in the nest material has been debated (Haftorn 1994). One possibility is that birds actually destroy and even consume ectoparasitic arthropods (Rothschild and Clay 1952). This behaviour may also occur in pied flycatchers because we observed females swallowing some collected items on videorecordings of control nests. Nest sanitation could also be used to chase blowfly larvae or adult fleas away from their own body or that of their nestlings, thereby preventing them from biting the incubating female or the nestlings. We also observed one female attacking an adult blowfly that entered the nest-box while she was brooding which could prevent oviposition in the nest. The difference in the time invested in behavioural defences indicates that females may be able to choose to increase the amount of time allocated to control of nest ectoparasites.

Changes in the frequency and duration of grooming or nest sanitation may be interpreted as responses to ectoparasites. Our results are consistent with several previous studies in great tits (Richner et al. 1993) and blue tits (Christe et al. 1996, Hurtrez-Boussès et al. 2000, Tripet et al. 2002b) that showed that females spent more time on nest sanitation when the nest was infested with fleas. The number and duration of grooming sessions also increased in control nests. This suggests that this behaviour may have evolved in response to ectoparasites and that females could thereby minimize the fitness costs associated with ectoparasite infestations (Richner et al. 1993).

This is the first study showing a clear effect of a complete natural ectoparasite fauna on parental behaviour and nestling growth in a cavity-nesting bird. Our test of effects of ectoparasites is conservative as we were not able to completely remove all ectoparasites and as the study was performed under especially good conditions for breeding. Ectoparasites induce significant changes in female grooming and nest sanitation behaviours which are not able to fully remove their natural deleterious effects on nestling growth and survival.

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