

Daily foraging routines and feeding effort of a small bird feeding on a predictable resource

Olsson, Ola; Wiktander, Ulf; Nilsson, Sven

Published in:

Royal Society of London. Proceedings B. Biological Sciences

10.1098/rspb.2000.1164

2000

Link to publication

Citation for published version (APA):

Olsson, O., Wiktander, U., & Nilsson, S. (2000). Daily foraging routines and feeding effort of a small bird feeding on a predictable resource. Royal Society of London. Proceedings B. Biological Sciences, 267(1451), 1457-1461. https://doi.org/10.1098/rspb.2000.1164

Total number of authors:

General rights

Unless other specific re-use rights are stated the following general rights apply: Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights

- Users may download and print one copy of any publication from the public portal for the purpose of private study or research.

 • You may not further distribute the material or use it for any profit-making activity or commercial gain

 • You may freely distribute the URL identifying the publication in the public portal

Read more about Creative commons licenses: https://creativecommons.org/licenses/

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

LUND UNIVERSITY

Download date: 17. Dec. 2025



Daily foraging routines and feeding effort of a small bird feeding on a predictable resource

O. Olsson*†, U. Wiktander and S. G. Nilsson

Department of Ecology, Animal Ecology, Lund University, Ecology Building, SE-223 62 Lund, Sweden

According to theory, a small diurnal bird living in a predictable environment should have the highest feeding effort as late as possible in the day in order to minimize the time it carries large and costly reserves. The feeding effort should also decline with increasing food availability. We tested both these ideas with the lesser spotted woodpecker (*Dendrocopos minor*). For most of the year, this bird feeds on woodliving insects in dead tree branches. This food supply is likely to be highly predictable on a daily scale. Our results corroborated the theory. We found that the proportion of time spent actively feeding was lower in the mornings (before noon) than in the afternoons. We also found that woodpeckers spent less time feeding the higher their food availability. However, for a given food availability they spent more time feeding in the afternoons. This supports the idea that feeding is less and other activities are more valuable in the mornings given a predictable food resource. This is the first demonstration of daily routines in small birds concordant with a predictable environment. In spring, males but not females reduced their feeding time. This difference between the sexes may be related to their sex-specific reproductive effort.

Keywords: daily routines; foraging; giving-up density

1. INTRODUCTION

The optimal foraging behaviour of animals should be a trade-off between the benefits of gaining energy and the risk of being killed by predation (e.g. Lima 1986; McNamara & Houston 1990; Houston & McNamara 1993; Lilliendahl *et al.* 1996). For a small bird whose life may depend on their ability to outmanoeuvre a raptor, the risk of predation may increase severely with increasing body weight (Rogers & Smith 1993; Witter & Cuthill 1993; Metcalfe & Ure 1995).

Models of optimal daily foraging routines for birds (Bednekoff & Houston 1994; McNamara et al. 1994) predict that those having predictable access to food should postpone their feeding activity until late in the day. This prediction is dependent on two assumptions: (i) that birds carrying a lot of reserves suffer from a higher predation risk (Bednekoff 1996; Kullberg et al. 1996), and (ii) that reserves are essential in order to avoid starvation at night. In order to maximize survival, the optimal trade-off between starvation and predation will therefore be to feed late in the day and, hence, reduce the time spent carrying high reserves.

In contrast, birds with unpredictable access to food must feed early in the day. This is because they must ensure sufficient reserves by the end of the day so that they survive the night (Bednekoff & Houston 1994; McNamara *et al.* 1994). Once sufficient reserves are gained they may then spend time idle in order to reduce their predation risk.

Other models of foraging strategies (McNamara & Houston 1987; Werner & Anholt 1993; Anholt & Werner 1998; Brown 1999; Olsson & Holmgren 1999) predict that

animals in rich environments should spend less time feeding than those in poor environments. The necessary assumption for generating these predictions is that the predation risk increases with feeding activity and reserve levels and the benefit from harvesting energy is a decelerating function of a cumulative harvest (Houston *et al.* 1997)

The food availability of an environment is often difficult to estimate based solely on direct sampling (Hutto 1990; Poulin & Lefebvre 1997). A more efficient means of estimating environmental quality is to consult the forager itself by the use of behavioural indicators (Brown 1988; Olsson & Holmgren 1999; Olsson et al. 1999). Foragers using depletable food patches should abandon these when their value to the forager corresponds to the average value of the environment (Charnov 1976; Brown 1988, 1999; Olsson & Holmgren 1998). The amount of food left in the patches at that time, the giving-up density (GUD), is one of the essential components of the behavioural indicator. By joining GUDs with data on patch residence times and a fitness measure, one has a full behavioural indicator. That is, these parameters may jointly estimate the cost of predation and the value of energy (Brown 1999; Olsson & Holmgren 1999). This can be used to determine what factors generate differences in quality between environments and to estimate that quality itself (Olsson & Holmgren 1999).

In a previous study (Olsson et al. 1999) of lesser spotted woodpeckers ($Dendrocopos\ minor$), we used these behavioural indicators and determined that food availability is the main factor varying between territories. Consequently, the average GUD of an individual, \overline{GUD} , is a good estimate of the food availability that an individual has in its territory. Within individuals, however, GUDs may change, even at constant food availability, in response to changes in the costs and benefits associated with foraging (Brown 1988).

Here we will present data on the daily foraging routines, feeding activity and food availability (in terms

^{*}Author for correspondence (ola.olsson@zooekol.lu.se).

[†] Present address: Department of Biological Sciences (MC 066), University of Illinois at Chicago, 845 West Taylor Street, Chicago, IL 60607-7060, USA.

of GUD) of lesser spotted woodpeckers during both winter and spring. During this time the lesser spotted woodpecker feeds almost exclusively on wood-boring insect larvae (mainly long-horn beetles, Cerambycidae) in thin (1-5 cm) dead branches of living trees (Cramp 1985; Olsson 1998). An exception may be the moth Argyresthia goedarthella which occurs abundantly in the trunk bark or dead branches of Betulaceae trees (birches and alders) in the years when these tree species bloom (Agassiz 1987). The larvae usually become accessible to the woodpeckers in late March or early April, but during 1996 they appeared later than usual and also in higher abundance (Olsson 1998; Olsson et al. 1999).

As the woodpeckers' prey are incapable of moving distances of more than a few millimetres, we were confident that the diurnal patterns of foraging by woodpeckers do not depend on the daily routines of their prey.

During the winter, the woodpeckers live practically solitarily in vast home ranges which may partly overlap with those of other individuals (Wiktander 1998). From the end of March until mid-May they form pairs, defend territories and excavate nest holes in dead trees (Olsson 1998; Wiktander 1998). Thus, in the winter most nonforaging time is devoted to resting or preening, whereas territory defence, breeding interactions and nest excavation become more frequent in spring.

2. METHODS

The study was conducted between 1990 and 1996 in a 125 km² study area around Lake Möckeln in southern Sweden. The area is a mixture of open land and forest and the forests are both coniferous and deciduous. The woodpeckers mainly use the deciduous parts of the forest during spring.

We studied the woodpeckers using binoculars and noted their behaviour. In total, 85 individuals were studied and all individuals were observed for at least 15 min (range 15.2-979 min and mean 113 min). The observations were made between 1 September each year and the day when oaks came into leaf the following spring (range 3-20 May in the years studied). This study halted at leaf break because at that time new food resources, such as aphids and caterpillars, become available to the woodpeckers and the woodpeckers abandon their wood-living food resources. By ending the observations at leaf break, the behavioural observations were also not confounded by direct breeding behaviours as egg laying starts after leaf break (Wiktander et al. 2000). The territorial activities and nest excavation start fairly synchronously in the population at the end of March. We chose 21 March as the division between winter and spring.

The majority of all non-foraging activities were either nest building or perching. We defined perching as either resting or actively preening, as both activities provide ample possibilities for predator scanning. Perching did not include the 'freezing' behaviour which the woodpeckers performed in the presence of avian predators. During perching, resting and preening were often mixed within short time-periods and were sometimes ambiguous to separate. Several activities, such as drumming, territorial interactions and mating, were also recorded, but the duration of these activities was too short for productive analysis.

All observations were grouped as being made in the mornings (before 12.00) or in the afternoons (after 12.00). Preliminary analyses showed that moving the division by an hour in either

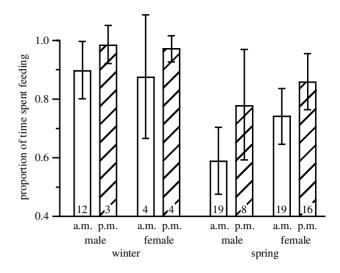


Figure 1. The proportion of the total observed time which was spent actively foraging in winter and spring in males and females: before versus after 12.00. Mean \pm s.e. The number of individuals observed is shown inside each bar.

direction did not alter the conclusions. The observations were made throughout the woodpeckers' active time, i.e. from approximately 30 min after sunrise until 30 min before sunset (Wiktander 1998), although, in any given day, a single woodpecker was only followed for several hours.

We used the proportion of time engaged in foraging, nest excavation and perching for analysis of the behavioural data. The proportion of time spent foraging is the ratio of the total time spent foraging to the total time observed during the given period of a day (i.e. not per foraging patch). The statistical distribution of this proportion was normalized by arcsine transformation. However, the proportions of time spent on nonforaging activities included many observations of zero value and, hence, could not be normalized. We therefore first used the frequencies of individuals observed performing these activities in Fisher's exact tests for the analyses. For the individuals engaged in these activities, we also analysed the respective proportions using ANOVAs.

The GUDs were measured in the thin dead branches which the woodpeckers had foraged on. They were calculated as the number of prey remaining in the branches per square decimetre of surface. The measure was achieved by cutting down the branches and using X-rays to count the numbers of remaining larvae (see Olsson et al. (1999) for a complete description of the method). We log transformed the GUD values before analysis to normalize the distribution. Data on the GUDs were collected from 1993 to 1996 from a total of 35 individuals with at least two branches sampled per individual.

Each individual occurs only once in each analysis. In the cases where individuals were observed in more than one period, one of these periods was selected for analysis at random.

3. RESULTS

The lesser spotted woodpeckers foraged more actively in the afternoons than in the mornings $(F_{180} = 11.22)$ and p = 0.001) (figure 1). Furthermore, they spent less time actively foraging during the spring than during the winter period ($F_{1,80} = 18.52$ and p < 0.0005). The males

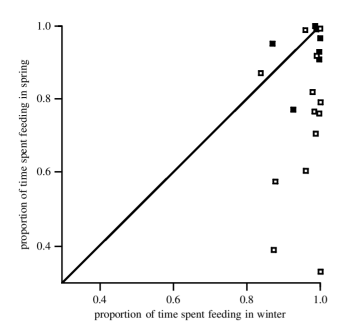


Figure 2. The proportion of time spent feeding by 20 individual woodpeckers during winter (horizontal axis) and spring (vertical axis). Open symbols are males and filled symbols are females. The diagonal line indicates the 1:1 line, i.e. where the points would be expected to be if the feeding times were equal in the two seasons.

and females spent similar proportions of their time feeding $(F_{1.80} = 0.015 \text{ and } p = 0.6)$. However, there was a tendency for the males to decrease their feeding times more in spring than did the females (interaction term $F_{1.80} = 3.36$ and p = 0.070) (figure 1). All other interaction terms were removed from this ANOVA as they were nonsignificant (p > 0.6 in all cases).

We made a repeated-measures ANOVA within individuals across seasons in order to further investigate the seasonal change in foraging between the sexes. Only 20 individuals were observed during both seasons and, hence, the analysis has to be confined to these. As in the former analysis, the individuals fed less in spring than in winter $(F_{1.18} = 22.20 \text{ and } p < 0.0005)$. However, the males reduced their feeding time significantly more than did the females $(F_{1,18} = 6.48 \text{ and } p = 0.020)$ (figure 2). As in the previous analysis, when combining seasons the males and females fed for similar proportions of time $(F_{1,18} = 2.61)$ and p = 0.12).

During the spring period, the proportion of time spent foraging was negatively correlated with the GUD (figure 3). This relationship was influenced by the time of day such that, in the mornings, the time devoted to foraging was lower than in the afternoon for a given GUD (figure 3). In an ANCOVA of the proportion of foraging time using the GUD as the covariate and time of day as a factor, both were significant $(F_{1.22} = 5.64)$ and p = 0.027, and $F_{1,22} = 7.71$ and p = 0.011, respectively and $r^2 = 0.45$), whereas their interaction was not (partial r = 0.10 and p = 0.3). During the winter period the $\overline{\rm GUD}$ was measured on too few individuals for productively making the same analysis.

During the Argyresthia presence in 1996, the $\overline{\text{GUD}}$ was higher than during previous periods (Olsson et al. 1999).

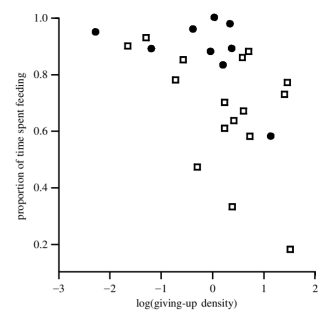


Figure 3. The proportion of time spent foraging as a function of the mean giving-up density of prey in patches which the same individuals were observed on. Squares indicate individuals observed in the mornings and circles those observed in the afternoons.

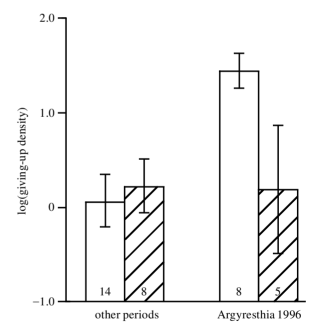


Figure 4. The mean (\pm s.e.) giving-up density (log transformed) of prey in patches before and during the peak occurrence of Argyresthia in 1996. All years before 1996 and 1996 before the Argyresthia appeared are thus included in the 'other periods'. Open bars are mornings and hatched bars are afternoons. The number of individuals observed is shown inside each bar.

Separating the mornings and afternoons, it appears that most of this difference was due to an exceptionally high GUD in the mornings during the Argyresthia presence (figure 4). This was formally shown by an ANOVA with the season, time of day and sex as factors. Only the season and the interaction between the season and time of day were close to significance (before versus during Argyresthia in 1996, $F_{1.31} = 3.53$ and p = 0.070, mornings versus afternoons, $F_{1.31} = 2.44$ and p = 0.13, and interaction, $F_{1.31} = 4.01$ and p = 0.054). All other terms had pvalues above 0.15. When testing mornings and afternoons separately, the difference between seasons was strong in the mornings $(F_{1,20} = 12.1 \text{ and } p = 0.002)$ but not in the afternoons ($F_{1.11} = 0.005$ and p = 0.9).

The woodpeckers spent a similar proportion of time foraging during the Argyresthia presence and the other period of spring $(F_{1.81} = 1.29 \text{ and } p = 0.26)$. The feeding effort depended on the time of day and sex of the individuals only $(F_{1.81} = 8.62 \text{ and } p = 0.004, \text{ and } F_{1.81} = 6.55 \text{ and}$ p = 0.012).

The proportion of perching individuals did not differ between winter and spring: 15 out of 23 versus 43 out of 62 (Fisher's exact p = 0.8).

There was a similar proportion of individuals observed perching in the mornings and afternoons in both seasons combined (Fisher's exact p = 0.15) and the proportion of individuals observed perching did not differ between males and females (Fisher's exact p = 0.6). However, among these perching individuals the time spent perching was higher in the mornings ($F_{1,55} = 5.37$ and p = 0.026), but similar between the sexes (two-way ANOVA, $F_{1.55} = 0.024$ and p = 0.9).

Out of the 23 individuals observed during the winter period, only one spent some time excavating a nesting hole, whereas during the spring period 36 out of the 62 observed did so (Fisher's exact p < 0.0005).

Excavation of nest holes was predominantly performed during mornings in the spring (Fisher's exact p = 0.035). In total, the frequency of excavation was approximately the same across the sexes (Fisher's exact p = 0.17). However, out of the 36 individuals observed to build nests, males spent significantly more time at this activity, but there was no difference between mornings and afternoons $(F_{1.33} = 6.01 \text{ and } p = 0.020, \text{ and } F_{1.33} = 0.00 \text{ and}$ p = 1.00).

4. DISCUSSION

Our major and novel findings were that lesser spotted woodpeckers feed less actively in the mornings than in the afternoons, that individuals in territories with high food availability feed less than those with low food availability and that females feed more than males during spring.

Previous studies have shown routines expected for unpredictable feeding conditions in several bird species (great tits Parus major, Lilliendahl et al. 1996; blackbirds Turdus merula, Cresswell 1998). That is, most of the mass increase took place in the mornings, but was also dynamic such that previous mass loss could be rapidly regained. The species studied spent a large fraction of their time in winter feeding on the ground. Therefore, the weather conditions may often make their feeding gains unpredictable (Rogers & Smith 1993).

Our study is the first to show the opposite pattern, i.e. more feeding in the afternoon. This is the predicted optimal daily routine for birds with predictable food resources (Bednekoff & Houston 1994; McNamara et al. 1994), i.e. when their starvation risk is negligible. It seems justifiable to argue that this is the case for woodpeckers.

During the course of a year, their prey density distribution does not vary stochastically over time (except for the rapid increase in the Argyresthia density). The accessibility of the wood-living prey should also not be severely influenced by weather conditions, etc. The physical properties of the branches and also the activity of the prey may to an extent vary with factors such as temperature and humidity. However, as these factors vary much more between days than within days, we find it unlikely that they will influence diurnal foraging patterns. Furthermore, the woodpeckers visited over 100 branches (patches) per hour (arithmetic mean 116) (Olsson 1998; O. Olsson and U. Wiktander, personal observations). Thus, although their gain from single patch visits is clearly stochastic and should have a large standard deviation, the standard error of the expectation over a few hours or a whole day will be small. Hence, our study together with previous studies (Lilliendahl et al. 1996; Cresswell 1998) fit the patterns predicted by theory.

The lesser spotted woodpeckers' propensity to engage in activities other than gaining energy was influenced by the availability of food in the environment, as measured by the GUD. This result in itself indicates the existence of trade-offs between energetic gain and other factors which may enhance fitness, as assumed in a number of theoretical models (McNamara & Houston 1987; Werner & Anholt 1993; Anholt & Werner 1998; Brown 1999). Results in accordance with such models have been shown in the wild in only a few cases (e.g. Davies & Lundberg 1985). Woodpeckers with abundant food spend much of their time keeping their plumage in good condition and being vigilant against predators and they can afford to invest more into excavating high-quality nests. All of these activities may obviously have a positive influence on fitness without increasing their energy gain.

During the peak occurrence of Argyresthia in 1996, the GUD was higher in the mornings, both compared to the previous period and compared to the afternoons in the same period. This higher intake rate (during the active foraging time) in the mornings is obviously chosen by the woodpeckers as their food availability does not change (so dramatically) between mornings and afternoons. As such, this strongly indicates that the woodpeckers' foraging is dynamic and possibly state dependent, as has also been demonstrated, for example, in great tits (Lilliendahl et al. 1996). Having higher GUDs but shorter feeding times in the mornings than in the afternoons implies that the woodpeckers found activities alternative to feeding more valuable in the mornings than in the afternoons (Brown 1988). Such a conclusion is in complete agreement with models of daily routines (Bednekoff & Houston 1994; McNamara et al. 1994), although it is only implicit in them.

The total time that woodpeckers are outside their roosting hole in spring is much longer than during the winter (O. Olsson and U. Wiktander, personal observations). This factor alone may be responsible for why the woodpeckers fed for a lesser proportion of their time during spring than in winter. It is interesting though that males and females respond differently to seasonal change. Males reduce their feeding effort in order to invest more time into nest excavation instead. Females do not. This is probably related to the fact that the approaching breeding may present different challenges and opportunities for females than for males (e.g. Askenmo et al. 1992). The most obvious difference is of course that the female is the one to produce the eggs which may be energetically rather costly (see the review by Monaghan & Nager 1997). Females may therefore value energy higher than males. Our result that males and females spent similar proportions of time feeding in winter but that females fed more in spring supports this idea. Given that foraging is the most dangerous activity this may explain the observation (Wiktander 1998) that females suffer higher mortality prior to egg laying than do males. The females may thus pay a high cost for their increased energy gains.

We thank Anders Stagen, Krister Wahlström, Fredrik Östrand, Leif Appelgren, Fredrik Haas and Martin Stjernman for assistance in the field. We also thank Jan-Åke Nilsson for encouragement and valuable discussions throughout this project and Joel Brown for discussions and ideas. Peter Bednekoff, Ron Ydenberg, Wendy Jackson and two anonymous reviewers made very valuable comments to previous versions of the paper. Most of the fieldwork was financed by Worldwide Fund for Nature Sweden.

REFERENCES

- Agassiz, D. J. L. 1987 The British Argyresthiinae and Yponomeutinae. Proc. Trans. Br. Entomol. Nat. Hist. Soc. 20, 1-26.
- Anholt, B. R. & Werner, E. E. 1998 Predictable changes in predation mortality as a consequence of changes in food availability and predation risk. Evol. Ecol. 12, 729-738.
- Askenmo, C., Neergaard, R. & Arvidsson, B. L. 1992 Prelaying time budget in rock-pipits: priority rules of males and females. Anim. Behav. 44, 957-965.
- Bednekoff, P. A. 1996 Translating mass dependent flight performance into predation risk: an extension of Metcalfe & Ure. Proc. R. Soc. Lond. B 263, 887-889.
- Bednekoff, P. A. & Houston, A. I. 1994 Avian daily foraging patterns: effects of digestive constraints and variability. Evol. Ecol. 8, 36-52.
- Brown, J. S. 1988 Patch use as an indicator of habitat preference, predation risk, and competition. Behav. Ecol. Sociobiol.
- Brown, J. S. 1999 Vigilance, patch use and habitat selection: foraging under predation risk. Evol. Ecol. Res. 1, 49-71.
- Charnov, E. L. 1976 Optimal foraging, the marginal value theorem. Theor. Pop. Biol. 9, 129-136.
- Cramp, S. 1985 The birds of the Western Palearctic, vol.4. Oxford University Press.
- Cresswell, W. 1998 Diurnal and seasonal mass variation in blackbirds Turdus merula: consequences for mass-dependent predation risk. J. Anim. Ecol. 67, 78-90.
- Davies, N. B. & Lundberg, A. 1985 The influence of food on time budgets and timing of breeding of the dunnock Prunella modularis. Ibis 127, 100-110.
- Houston, A. I. & McNamara, J. M. 1993 A theoretical investigation of the fat reserves and mortality levels of small birds in winter. Ornis. Scand. 24, 205-219.

- Houston, A. I., McNamara, J. M. & Hutchinson, J. M. C. 1997 General results concerning the trade-off between gaining energy and avoiding predation. Phil. Trans. R. Soc. Lond. B 341, 375 - 397.
- Hutto, R. L. 1990 Measuring the availability of food resources. Studies Avian Biol. 13, 20-28.
- Kullberg, C., Fransson, T. & Jakobsson, S. 1996 Impaired predator evasion in fat black caps (Sylvia atricappilla). Proc. R. Soc. Lond. B 263, 1671-1675.
- Lilliendahl, K., Carlson, A., Welander, J. & Ekman, J. B. 1996 Behavioural control of daily fattening in great tits (Parus major). Can. J. Zool. 74, 1612-1616.
- Lima, S. L. 1986 Predation risk and unpredictable feeding conditions: determinants of body mass in birds. Ecology 67, 377 - 385.
- McNamara, J. M. & Houston, A. I. 1987 Starvation and predation as factors limiting population size. Ecology 68, 1515-1519.
- McNamara, J. M. & Houston, A. I. 1990 The value of fat reserves and the tradeoff between starvation and predation. Acta Biother. 38, 37-61.
- McNamara, J. M., Houston, A. I. & Lima, S. L. 1994 Foraging routines of small birds in winter: a theoretical investigation. 7. Avian Biol. 25, 287-302.
- Metcalfe, N. B. & Ure, S. E. 1995 Diurnal variation in flight performance and hence potential predation risk in small birds. Proc. R. Soc. Lond. B 261, 395-400.
- Monaghan, P. & Nager, R. C. 1997 Why don't birds lay more eggs? Trends Ecol. Evol. 12, 270-274.
- Olsson, O. 1998 Through the eyes of a woodpecker: understanding habitat selection, territory quality and reproductive success from individual behaviour. PhD thesis, Lund University, Sweden.
- Olsson, O. & Holmgren, N. M. A. 1998 The survival-rate-maximizing policy for Bayesian foragers: wait for good news. Behav. Ecol. 9, 345-353.
- Olsson, O. & Holmgren, N. M. A. 1999 Gaining ecological information about Bayesian foragers through their behaviour. I. Models with predictions. Oikos 87, 251–263.
- Olsson, O., Wiktander, U., Holmgren, N. M. A. & Nilsson, S. G. 1999 Gaining ecological information about Bayesian foragers through their behaviour. II. A field test with woodpeckers. Oikos 87, 264-276.
- Poulin, B. & Lefebvre, G. 1997 Estimation of arthropods available to birds: effects of trapping technique, prey distribution, and bird diet. J. Field Ornithol. 68, 426-442.
- Rogers, C. M. & Smith, J. N. M. 1993 Life-history theory in the nonbreeding period-trade-offs in avian fat reserves. Ecology 74, 419-426.
- Werner, E. E. & Anholt, B. R. 1993 Ecological consequences of the trade-off between growth and mortality-rates mediated by foraging. Am. Nat. 142, 242–272.
- Wiktander, U. 1998 Reproduction and survival in the lesser spotted woodpecker: effects of life history, mating system and age. PhD thesis, Lund University, Sweden.
- Wiktander, U., Olsson, O. & Nilsson, S. G. 2000 Annual and seasonal reproductive trends in the lesser spotted woodpecker Dendrocopos minor. Ibis. (In the press.)
- Witter, M. S. & Cuthill, I. C. 1993 The ecological costs of avian fat storage. Phil. Trans. R. Soc. Lond. B 340, 73-92.