

## Risky foraging leads to cost-free mate guarding in male teal *Anas crecca*

Matthieu Guillemain · Céline Arzel · Pierre Legagneux · Johan Elmberg ·  
Hervé Fritz · Michel Lepley · Christophe Pin · Antoine Arnaud ·  
Grégoire Massez

Received: 11 July 2006 / Revised: 20 October 2006 / Accepted: 21 October 2006 / Published online: 12 December 2006  
© Dt. Ornithologen-Gesellschaft e.V. 2006

**Abstract** Mate guarding by males is common in species with long-lasting pair bonds. We tested if the need to guard females affected foraging depth in male teal (*Anas crecca*), and if they were more vigilant than females when foraging with submerged eyes (preventing monitoring of competing males and predators). These predictions were not supported, suggesting that foraging depth selection is primarily driven by other factors, presumably food related. A likely reason why deeply foraging males did not increase vigilance is that 37.5% of the foraging time was already dedicated to it. The

apparent lack of guarding costs in foraging male teal may explain why such small ducks can maintain pair bonds for up to 7 months.

**Keywords** Circumannual · Foraging behaviour · Mate guarding · Teal · Vigilance

### Introduction

Many bird species form annual pair bonds long before nesting (e.g., Black 1996). Mate guarding by males is a common phenomenon in these species. This may affect daily time budgets for up to 7 or 8 months per year (e.g., Paulus 1983), not to mention species with lifelong monogamy, in which males may have to guard their mate throughout life (e.g. geese; Black et al. 1996). Consequently, mate guarding may have positive effects on fitness in both sexes in terms of natural as well as sexual selection. For males, this is often seen as a trade-off between vigilance, which reduces foraging efficiency, and enhanced breeding success, by sustaining the pair bond and preventing extra-pair copulations (e.g., Birkhead and Parker 1997).

Pairing in fall or early winter is considered to be adaptive in dabbling ducks *Anas* spp. (Hepp and Hair 1983), as mated birds get a dominant status within flocks that confers better access to food (e.g., Hepp 1986; Black 2005). This in turn allows paired females to improve body condition earlier than unpaired birds, which is likely to later translate into higher breeding success (due to the negative relationship between breeding date and success in *Anatidae*; Bowler 2005; Elmberg et al. 2005). However, precocial pairing also has drawbacks, especially for males; due to male-

---

Communicated by F. Bairlein.

---

M. Guillemain (✉) · C. Arzel  
Office National de la Chasse et de la Faune Sauvage,  
CNERA Avifaune Migratrice, La Tour du Valat,  
Le Sambuc, 13200 Arles, France  
e-mail: m.guillemain@oncfs.gouv.fr

C. Arzel · J. Elmberg  
Department of Mathematics and Natural Sciences,  
Kristianstad University, 291 88 Kristianstad, Sweden

C. Arzel  
Laboratoire d'Ecologie des Hydrosystèmes,  
Université Paul Sabatier, 118 route de Narbonne,  
31062 Toulouse, France

P. Legagneux · H. Fritz  
Centre d'Etudes Biologiques de Chizé,  
CNRS UPR 1934, 79360 Beauvoir sur Niort, France

M. Lepley · C. Pin · A. Arnaud  
Station Biologique de la Tour du Valat,  
Le Sambuc, 13200 Arles, France

G. Massez  
Les Marais du Vigueirat, Mas Thibert,  
13200 Arles, France

biased sex-ratios in duck populations (Baldassarre and Bolen 1994), males have to devote much time to guarding their female (e.g., Davis 2002; Guillemain et al. 2003). Male vigilance thus offers the female safer efficient foraging conditions, but is costly to males, if nothing else through reduced foraging efficiency.

In an earlier flyway-level study (Guillemain et al. 2006, in press), we observed that teal (*Anas crecca*) gradually increase foraging depth from September to August. This change is apparently due to predation risk restricting the behavioural repertoire in winter to safer shallow feeding methods, i.e., those with the eyes above the water surface (see also Guillemain et al. 2001). Conversely, the lower frequency of fly-overs by potential predators as the annual cycle progresses allows teal to use a wider variety of methods, including riskier deeper foraging when prey availability and energy requirements make it necessary. However, deep foraging with the eyes submerged not only limits the ability to detect predators, for a male it also prevents monitoring other males. Given (1) the costs of mate guarding, (2) that teal are paired for several months before breeding (e.g. Johnson and Rohwer 1998), and (3) the theoretically high relative energy requirements due to small body size compared to other dabbling ducks (ca. 300 g on average), we hypothesized that there should be intersexual differences in teal foraging behaviour.

Our aims were: (1) to determine if both sexes switch to deeper foraging methods over the annual cycle and, if so, if they do it at the same rate; alternatively, males may maintain vigilance by foraging shallower, e.g. by feeding on different food types than females, and (2) to test whether males and females partition foraging time into feeding bouts and interruptions in the same way when they rely on deep foraging. As deep foraging prevents monitoring of competing males, we hypothesized that deep foraging males should have shorter feeding bouts and/or longer foraging interruptions.

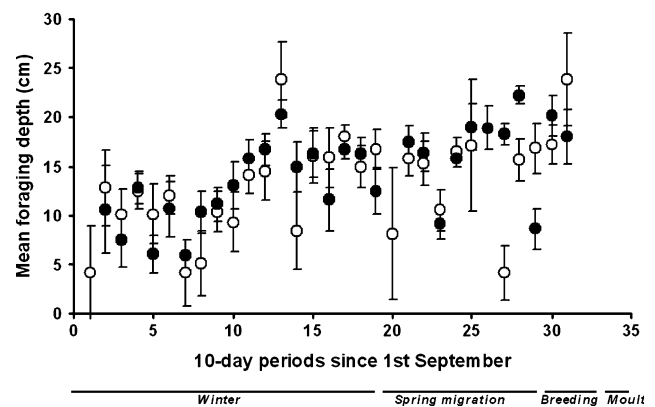
Study sites and methods are described in detail in Guillemain et al. (2006, in press). In brief, we used focal individual observations (Altman 1974) of foraging teal, chosen at random within flocks during daylight hours at 25 different sites in seven geographical areas within its flyway in western Europe (from Camargue in southern France to Västerbotten in north-central Sweden), covering the whole annual cycle except autumn migration stopovers. In total, 658 focal observations, i.e., the duration of ten successive foraging bouts using the same foraging method, plus the duration of the ten associated foraging interruptions, were recorded (244 females and 414 males). For each bird, we calculated an average value of bout length for foraging as well as for interruptions. Each foraging posture was

associated with a feeding depth based on body measurements in Thomas (1982), e.g., bill length for dabbling, head + neck length for dipping, etc. Foraging depth does not necessarily depend on the depth of lakes, since ducks may also seek floating seeds and invertebrates in the water column. High turnover rates in teal assemblages (Pradel et al. 1997) made double recording of the same individual unlikely, with the possible exception of breeding lakes where density is low. Because each focal individual relied on one foraging method, associated with a certain depth, hence a discontinuous variable, means of behavioural parameters were computed for all birds per 10-day period (foraging depth and duration of foraging and interruption bouts).

We have earlier demonstrated that increased use of deep foraging methods from September to August is associated with a relaxation of predation risk (Guillemain et al. 2006, in press). Here, we do not address why birds change behavior, but rather use ANCOVAs to compare sex-specific patterns of mean foraging depth and duration of feeding bouts and interruptions over the year.

The model exploring mean foraging depth in relation to date and sex was significant ( $F_{3,55} = 7.49$ ,  $r^2 = 0.29$ ,  $P = 0.0003$ ). However, neither sex nor sex  $\times$  date had a significant effect (sex:  $F = 0.03$ ,  $P = 0.8539$ ; sex  $\times$  date:  $F = 0.08$ ,  $P = 0.7792$ ). Instead, all variance was explained by date, i.e. foraging depth increased seasonally (10-day periods:  $F = 21.37$ ,  $P < 0.0001$ ; Fig. 1) at the same rate in both sexes, which on average foraged at the same depth.

For individuals foraging with submerged eyes, the model including sex, date, and sex  $\times$  date did not fit with variation in average length of feeding bouts



**Fig. 1** Annual variation in foraging depth of male (dots) and female (circles) teal (*Anas crecca*) (mean  $\pm$  SE), expressed per 10-day period starting 1–10 September. The main phases of the annual cycle are indicated under the x-axis. See main text for statistics

( $F_{3,52} = 0.36$ ,  $r^2 = 0.02$ ,  $P = 0.7848$ ), nor feeding interruptions ( $F_{3,52} = 1.35$ ,  $r^2 = 0.07$ ,  $P = 0.2682$ ). The partitioning of behaviour into foraging and vigilance thus remained unchanged over time, and did not differ between sexes when foraging in a posture preventing vigilance. Deep foraging bouts lasted 1.88 s on average ( $\pm 0.09$  SE,  $n = 56$ ), feeding interruptions lasted 1.20 s ( $\pm 0.06$  SE,  $n = 56$ ).

Among dabbling ducks, mate guarding has been shown to strongly influence male behavior in wigeon (*A. penelope*) (Mayhew 1987; Guillemain et al. 2003) and gadwall (*A. strepera*) (Dwyer 1975). Deeply foraging male teal cannot detect competing males, which is why we hypothesized that they should use this behaviour less than females do. However, this is not what we observed. Both sexes had similar foraging depths, and switched from shallow to deep foraging at the same rate over time. In a previous study, we hypothesized that relaxing predation risk over time would allow teal in general to increase foraging depth in response to higher energy requirements and/or changing food availability (Guillemain et al. 2006, in press). The latter factors may thus be more important than intrasexual competition to foraging depth selection in males or, alternatively, deep foraging is not as costly as hitherto assumed in terms of decreased ability to monitor competing males.

Our second prediction, i.e. that deeply foraging males unlike females alter their partitioning of behaviors by increasing vigilance, was not supported either. Average foraging bout length was similar (<2 s) between sexes, and interruptions lasted 1.2 s in both males and females. We suggest that deeply foraging males already have such high anti-predator vigilance (in terms of frequency of bouts as well as proportion of time) that they do not have to increase it further to guard their mate. As a comparison, male wigeon increased vigilance from 3.32 to 14.36% of foraging time after they paired (Guillemain et al. 2003). The corresponding percentage here is 37.5% for deeply foraging teal. This implies that, as opposed to earlier studies documenting high costs of mate-guarding, male teal do not pay a significant price for guarding their mate, at least when foraging. This may explain how such relatively small birds, with high relative energy requirements, can afford to form and defend pairs for up to 7 months a year (Johnson and Rohwer 1998).

## Zusammenfassung

Riskante Nahrungssuche erlaubt männlichen Krickenten *Anas crecca* ein kostenfreies Bewacherverhalten

Bei Arten mit langer Paarbindung findet man bei den Männchen häufig ein Bewacherverhalten. Wir haben untersucht, ob die Notwendigkeit, die Weibchen zu bewachen männliche Krickenten (*Anas crecca*) in der Intensität der Nahrungsaufnahme beeinträchtigte, und ob sie bei der Nahrungssuche mit den Augen unter Wasser wachsamer waren als die Weibchen (um einer Überwachung konkurrierender Mänchen und Prädatoren vorzubeugen). Diese Hypothesen bestätigten sich nicht, was nahelegt, dass der Selektionsdruck auf die Intensität bei der Nahrungssuche primär von anderen, vermutlich an die Nahrung gekoppelten Faktoren angetrieben wird. Ein Grund, wieso intensiv nach Nahrung suchende Männchen ihre Wachsamkeit nicht erhöhten, liegt wahrscheinlich darin, dass 37.5% der Zeit auf Nahrungssuche bereits auf Wachsamkeitsverhalten verwendet wurde. Dass es bei Krickentmännchen anscheinend keine Bewachungskosten gibt, könnte erklären, wieso diese kleine Entenart eine bis zu sieben monatige Paarbindung aufrecht erhalten kann.

**Acknowledgments** We are most grateful to the managers of the study sites, especially the Marais du Vigueirat for the Camargue, the Ligue pour la Protection des Oiseaux and the Fédération Départementale des Chasseurs (FDC) de Charente-Maritime for the Atlantic sites, the Réserve Naturelle de Chérine and the FDC de l'Indre for the Brenne lakes, the FDC de l'Eure and R. Legay for the Grand'Mare reserve and hunting ponds, and the county administration of Västerbotten for permission to work within the Brånsjön reserve. The staff at the Biosphere Reserve 'Kristianstads Vattenrike' is acknowledged for valuable help in coordinating landowner contacts. We also thank those who helped in the field, especially N. Guillon, F. Latraube and L. Tison, and two referees for their comments on an earlier version of the manuscript. M. Guillemain received a doctoral grant from the Conseil Général Poitou-Charentes when working at the Atlantic sites. C. Arzel was supported by a PhD Grant from ONCFS and grants V-124-01 and V-98-04 from the Swedish Environmental Protection Agency. P. Legagneux received a grant from the university Louis Pasteur of Strasbourg, France.

## References

- Altman J (1974) Observational study of behaviour: sampling methods. *Behaviour* 49:227–267
- Baldassarre GA, Bolen EG (1994) Waterfowl ecology and management. Wiley, New York
- Birkhead TR, Parker GA (1997) Sperm competition and mating systems. In: Krebs JR, Davies NB (eds) Behavioural ecology—an evolutionary approach. Blackwell, Oxford, pp 121–145
- Black JM (1996) Introduction: pair bonds and partnerships. In: Black JM (ed) Partnerships in birds—the study of monogamy. Oxford University Press, Oxford, pp 3–20
- Black JM (2005) Ecology of social behaviour. In: Kear J (ed) Ducks, geese and swans. Oxford University Press, Oxford, pp 57–67

- Black JM, Choudhury S, Owen M (1996) Do barnacle geese benefit from lifelong monogamy? In: Black JM (ed) Partnerships in birds—the study of monogamy. Oxford University Press, Oxford, pp 91–117
- Bowler J (2005) Breeding strategies and biology. In: Kear J (ed) Ducks, geese and swans. Oxford University Press, Oxford, pp 68–111
- Davis ES (2002) Female choice and the benefits of mate guarding by male mallards. *Anim Behav* 64:619–628
- Dwyer TJ (1975) Time budget of breeding gadwalls. *Wilson Bull* 87:335–343
- Elmberg J, Nummi P, Pöysä H, Gunnarsson G, Sjöberg K (2005) Early breeding Teal *Anas crecca* use the best lakes and have the highest reproductive success. *Ann Zool Fenn* 42:37–43
- Guillemain M, Duncan P, Fritz H (2001) Switching to a feeding method that obstructs vision increases head-up vigilance in dabbling ducks. *J Avian Biol* 32:345–350
- Guillemain M, Caldow RWG, Hodder KH, Goss-Custard JD (2003) Increased vigilance of paired males in sexually dimorphic species: distinguishing between alternative explanations in wintering Eurasian Wigeon. *Behav Ecol* 14:130–135
- Guillemain M, Arzel C, Legagneux P, Elmberg J, Fritz H, Lepley M, Pin C, Arnaud A, Massez G (2006) Teal (*Anas crecca*) adjust foraging depth to predation risk: a flyway-level circum-annual approach. *Anim Behav* (in press)
- Hepp GR (1986) Effects of body weight and age on the time of pairing of American black ducks. *Auk* 103:477–484
- Hepp GR, Hair JD (1983) Reproductive behaviour and pairing chronology in wintering dabbling ducks. *Wilson Bull* 95:675–682
- Johnson WP, Rohwer FC (1998) Pairing chronology and agonistic behaviors of wintering green-winged Teal and Mallards. *Wilson Bull* 110:311–315
- Mayhew PW (1987) Vigilance levels in European wigeon—sexual differences. *Wildfowl* 38:77–81
- Paulus SL (1983) Dominance relations, resource use, and pairing chronology of Gadwalls in winter. *Auk* 100:947–952
- Pradel R, Rioux N, Tamisier A, Lebreton JD (1997) Individual turnover among wintering teal in Camargue: a mark-recapture study. *J Wildl Manage* 61:816–821
- Thomas GJ (1982) Autumn and winter feeding ecology of waterfowl at the Ouse washes, England. *J Zool* 197:131–172