Bright moonlight triggers natal dispersal departures

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ABSTRACT

Upon leaving their natal area, dispersers are confronted with unknown terrains. Species-specific perceptual ranges (i.e., the maximum distance from which an individual can perceive landscape features) play a crucial role in spatial movement decisions during such wanderings. In nocturnal animals that rely on vision, perceptual range is dramatically enhanced during moonlight, compared to moonless conditions. This increase of the perceptual range is an overlooked element that may be responsible for the successful crossing of unfamiliar areas during dispersal. The information gathered from 143 radio-tagged eagle owl *Bubo bubo* juveniles in Spain, Finland and Switzerland shows that, although the decision to initiate dispersal is mainly an endogenous phenomenon determined by the attainment of a given age (~6 months), dispersers leave their birthplace primarily under the best light conditions at night, i.e. when most of the lunar disc is illuminated. This sheds new light into the mechanisms that may trigger dispersal from parental territory.

Keywords: dispersal age, lunar cycle, moon phase, perceptual range, timing of dispersal, visual cues, *Bubo bubo*

Word count: 3808
The start of natal dispersal represents a risky stage in the life of animals, which entails multiple costs (Bonte et al. 2012). Consequently, when the time comes to leave the birthplace, individuals are expected to start their dispersal under the best conditions.

Visual cues are crucial for animals moving at night; for instance, many migratory birds refer to moonlight position to establish and maintain orientation during their nocturnal journey (Martin 1990). The lunar cycle (full moon to full moon, 29.5 days) determines important environmental changes that influence and can be perceived by animals. Night-time light levels are far from uniform: around full moon, the increased light levels at night may determine the occurrence of occasional nocturnal activities (like night foraging and migration) in otherwise strictly diurnal birds (Pienkowski 1982; Martin 1990; Bulyuk et al. 2009). Nocturnal birds also benefit from the additional light provided by a completely, or nearly, full moon (Martin 1990; Brigham and Barclay 1992): night-time movements and feeding appear to be enhanced by moonlight, independent of the visual sensitivity of owls and other nocturnal birds. Thus, although there are certainly many more favourable conditions than those provided by moonlight, the lunar light has the potential to represent an important factor affecting nocturnal movement across unknown landscapes.

The perceptual range of animals, i.e. the maximum distance from which an individual can perceive landscape features and elements, plays an important role in spatial processes (Zollner and Lima 1997). The perceptual range affects movement strategies and behaviours during dispersal, as well as the probability of successfully reaching suitable habitats (Lima and Zollner 1996; Zollner and Lima 1999). For nocturnal species, the perceptual range is strongly affected by the ambient light (Zollner and Lima 1997; Zollner...
and Lima 1999): their perceptual abilities increase as ambient illumination increases, even though they have considerable visual acuity (Zollner and Lima 1999). It remains poorly understood to which extent moonlight phases may interfere with spatial movement decision such as natal dispersal in nocturnal species.

Here, we show that a nocturnal avian predator, the eagle owl *Bubo bubo*, starts dispersal primarily when juveniles are ~6 months old and the ambient light is the most favourable for perceiving and exploring unfamiliar lands. This phenomenon occurs within its whole distribution range, as demonstrated here by eagle owl natal dispersal in three geographically and ecologically diverse European areas spread over a wide latitudinal gradient.

**STUDY AREA AND METHODS**

*Data collection*

From 2003 to 2012 we marked 95 juveniles (from 23 nests) in Spain with conventional units (Biotrack, http://www.biotrack.co.uk; for details on study area and radiotracking procedure see Delgado et al. 2010, Penteriani and Delgado 2011) and 24 juveniles (from 18 nests) in Finland with satellites units (Microwave Telemetry, http://www.microwavetelemetry.com). 24 juveniles (from 17 nests) were marked in Switzerland (for details on the study area and radiotracking procedure see Aebischer et al. 2010, Schaub et al. 2010) with: (i) 30 g battery-powered satellite transmitters supplied by North Star (http://www.northstarst.com); and (ii) 15 g VHF tags (Holohil, http://www.holohil.com). The weight of the transmitters was between the 3 and the 3.5 % of the weight of the smallest individual at the time of tagging (Spain: 850 g,
mean ± SD = 1267 ± 226.4 g; Finland: 1370 g, mean ± SD = 1861.2 ± 283.8 g;
Switzerland: 1000 g, mean ± SD = 1506 ± 260.9 g; see also Penteriani et al. 2011). Signals
from the satellite transmitters were recorded by the ARGOS satellite system (www.cls.fr).
To make comparable the data acquired by different radiotracking equipments and
methods, for each individual we estimated the start of dispersal in the same way (Delgado
and Penteriani 2008): we defined dispersal as starting when the distance of each location
from the nest became larger than the global mean distance travelled to the nest, that is,
when the distance of each location from the nest started progressively increasing rather than
fluctuating around a low value. Owls were sexed by molecular procedures using DNA
extracted from blood samples (Delgado et al. 2010).

Moon phases
Over the study years, daily variations of the moon phases were obtained from the Naval
Oceanography Portal (http://aa.usno.navy.mil/data/docs/RS_OneYear.php) and expressed
in terms of the fraction of moon disk illuminated and whether the moon was waxing or
waning. Following the periodic regression approach suggested by deBruyn and Meeuwig
(2001) and applied elsewhere (e.g. Kuparinen et al. 2010; Penteriani et al. 2011, 2013), the
fraction of moon disk illuminated was converted into radians (θ), with one lunar cycle
corresponding to a gradual change from 0 to 2π radians (0 and 2π radians correspond to
the full moon, and π radians corresponds to the new moon). Cos(θ), sin(θ), cos(2θ) and
sin(2θ) transformations were included in the statistical model as explanatory variables, to
investigate possible lunar effects on eagle owl behaviour throughout the lunar cycle (see
deBruyn and Meeuwig 2001, for details). Given that the probability to have a cloudy night
is equally distributed over the study period and, consequently, among all moon phases, we
considered such variation to lead to additional noise, which is likely to weaken the signal strength in our results rather than to create systematic biases (Penteriani et al. 2013).

Statistical analyses

To test the effects of moon phase, age (days after hatching) and sex of individuals, as well as the interactions between age and the different moon phases on the decision to start dispersal (as Julian date), we used linear mixed-effects models. We first selected the optimal structure of the random component, which was the one containing three levels of random effects, represented by birthplace nested in country nested in year (ESM1). The response variables were scaled, i.e. normalised to zero mean and unit variance. Model simplification was performed by backward selection of variables from the full model, and models were compared using likelihood ratio tests until a minimal adequate model was obtained (Crawley 2007). An ANOVA test compared age of dispersal among the three countries. All statistical analyses were performed in R 2.10.1 statistical software (R Development Core Team 2009), nlme (Pinheiro et al. 2009) package.

RESULTS

Mean age of dispersal (±SD) was 163.6 ± 20.1 days, (range = 116 – 222 days), not differing significantly (F2,140 = 2.40, P = 0.10) between Spain (mean = 166.8 ± 20.1 days, range = 128 – 222 days), Finland (mean = 150.0 ± 18.6 days, range = 116 – 185 days), and Switzerland (mean = 164.1 ± 16.6 days, range = 140 – 209 days).

The linear mixed-effects model (Table 1 and ESM1) demonstrated that the decision to start dispersal was influenced by: (i) age of individuals, i.e. dispersal mainly started when juveniles were ~6 months old; (ii) sex, with males (age of dispersal = 161.3 ±
22.1 days, range = 116 – 222 days) showing more variation in their departure age than females (age of dispersal = 166.6 ± 16.9 days, range = 127 – 206 days); (iii) the moon phase, with dispersal departures mostly occurring during a full moon (Fig. 1); and (iv) the interaction between age and moon phase: the moonlight effect was the highest when owls dispersed at the average dispersal age of this species.

**DISCUSSION**

Dispersal age appears to be a constant trait of eagle owl life-history: most departures occurred in a relatively narrow time window, despite the diverse ecological conditions and wide latitudinal gradient in this study (from Mediterranean to boreal habitats). Yet, most juveniles approaching dispersal age preferred to initiate dispersal when most of the lunar disc was illuminated. However, when individuals are not able to disperse within that favourable temporal window, they cannot pay much attention to the environmental conditions under which to initiate dispersal. Despite the importance of age specific dispersal, this latter has been ignored in most of the evolutionary theory of dispersal (Johst and Brandl 2000). In an evolutionary context, the evolution of age specific dispersal strategies may be due to an increase in the efficiency of dispersal at certain age classes (Johst and Brandl 2000). Actually, within the life cycle, timing of dispersal may influence the evolution of dispersal strategies and can be important in predicting the favoured dispersal strategies (Johst and Brandl 1997). One common advantage driving all juveniles of a same species to disperse as soon as they are ready (i.e., at the same age) is the increased access to higher quality breeding sites: the rapid acquisition of an area where to settle is likely to be especially important in resident, territorial birds, principally if residence is determined by
order of arrival (Ellsworth and Belthoff 1999). Though the proximate factors that stimulate juveniles to initiate dispersal are not well understood, either exogenous or endogenous factors may influence the timing of dispersal (Belthoff and Dufty 1998). Several behaviours have been suggested to appear as the post-fledging period progresses, e.g. parental aggression towards young (Bunn et al. 1982; Wiggett and Boag 1993), aggression of young toward each other (De Laet 1985; Strickland 1991) and decreasing food availability within the natal area (Veltman 1989; Kenward et al. 1993). Despite evidence that exogenous factors may drive dispersal, in some birds neither diminishing food supplies nor parental aggression elicited dispersal, dispersal being mostly initiated by endogenous mechanisms (Nilsson 1990; Belthoff and Dufty 1998). For example, body condition or social status have been considered to play a role in triggering dispersal (Nilsson and Smith 1985; Ellsworth and Belthoff 1999). In two owl species, the western *Otus kennicottii* and the eastern *Otus asio* screech-owls, the action of a specific hormone, the adrenal glucocorticoid corticosterone, seems to be responsible of increasing activity levels and changes in body condition (see also Silverin 1997; Wingfield and Ramenofsky 1997), which ultimately trigger juvenile dispersal (Ritchison et al. 1992; Belthoff and Dufty 1995, 1998). That is, exogenous stimuli related to the termination of parental feeding, increased aggression among siblings or other factors may not be required for dispersal departure to start, a hypothesis which is supported by the evidence that eagle owls under different exogenous stimuli started dispersal at the same age. However, the interaction between endogenous and external stimuli at the origin of dispersal initiation demonstrates that the dispersal of eagle owls provides a good example of an ecological process modulated by the combination of
internal and environmentally determined behaviours, which have evolved to fit within the
natural geophysical periods (Brown 1972).

Because light and dark cycles have existed throughout evolutionary time, many
organisms have evolved behaviours that are cued by moon cycles. Indeed, this is not the
first time that animal movements have shown an association with moon phases. Increased
lunar illumination can increase: (a) swimming depth of pelagic sharks and seals in response
to changes in the vertical distribution of their prey related to the aforementioned
illumination (e.g. Trillmich and Mohren 1981; Saunders et al. 2011); (b) fish migrations
given that they may utilize moonlight to navigate and relocate themselves more easily (e.g.,
Leatherland 1992; Hasegawa 2012); (c) dispersal and migration movements in some insects
and crustaceans, which may use the moon for vision, orientation and navigation (e.g.
Danthanarayana 1986; Scapini et al. 1997); (d) perceptual abilities of dispersing rodents
(Zollner and Lima 1999); (e) locomotor activity in owl monkeys of the genus *Aotus*
(Fernández-Duque et al. 2010); and (f) nocturnal bird migrations (e.g., Richardson 1978;
James 2000). In the case of eagle owls, we consider plausible the possibility that starting
dispersal during the brightest nights increases the perceptual range of dispersers (Zollner
and Lima 1997), helping inexperienced juveniles deal with unfamiliar habitats and find
prey. Actually, the potential for owls to detect prey increases with increasing light levels
(e.g. Clarke 1983; Kotler et al. 1991), which could be crucial for juvenile survival in the
days immediately following departure from the natal area. Furthermore, moonlight
intensity (due to moon phases) and the time during which the moon is visible in the night
sky are correlated (i.e. the nights are brighter for longer), which might represent an
additional advantage for individuals to disperse during full moon. Lunar light can thus be
considered both an environmental condition that is as important as diurnal light and a
resource, similar to time, space and temperature (Gerrish et al. 2009).

Finally, it is well known that predator-prey interactions are subject to monthly
changes owing to the lunar cycle, with predators adjusting their activity rhythms and
strategies in response to the increased concealment of prey during the brightest nights
(Mukherjee et al. 2009; Kotler et al. 2010; Penteriani et al. 2011). In view of the current
results, the lunar cycle might also have the potential to influence predator-prey
relationships on a seasonal basis. If, as we may expect, other nocturnal predators also take
advantage of the lunar light to start dispersal (Zollner and Lima 1997), we might observe
an increased number of predators (adults + dispersers) actively moving around the full
moon phase after reproductions. This phenomenon could in turn increase overall predation
risk and pressure, exerting a temporally short but profound influence on predator-prey
dynamics.

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Ethical standards. We manipulated and marked owls under: (i) Spanish Junta de
Andalucía-Consejería de Medio Ambiente authorizations No. SCFFS-AFR/GGG RS-

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**Table 1** The linear mixed-effects model showing the effect of moon phase, age and sex of individuals on the decision to start dispersal

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**Figure Legends**

**Fig. 1** Dispersal departures of eagle owls at various moon phases. When individuals reach their dispersal age (see Results), they mainly leave the natal area during bright nights.
Figure 1