Influence of different weather aspects on breeding performance, food supply and nest-space use in hoopoe offspring

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Abstract
In this study, we investigated the influence of different weather aspects on breeding performance, food supply and nest-space use in hoopoe offspring (Upupa epops). Camera recordings of 88 nests were used to examine how ambient environmental conditions influence food supply, offspring nest-space use and the trade-off nestlings face regarding the two mainly used locations in the nest. Therefore, we provide a comprehensive analysis involving different factors including weather parameters together with food provisioned to nestlings on different temporal scales to identify the factors having the most influence on nest-space use. We found that different breeding conditions significantly influenced how nestlings used the nest. During excessively humid weather, nestlings spent more time under the entrance hole when small food was delivered. However, nestlings supplied with large prey more often remained hidden in the distant area, despite the adverse weather situation. In all three aspects and temporal scales, our analysis confirmed that prey was the most important factor influencing offspring nest-space use, suggesting a crucial role of large insects for hoopoes. Finally, we found that long-term effects of weather affect overall food provisioned to nestlings and thus offspring behaviour. We provide evidence that parental feeding location and prey size, which are in turn influenced by weather conditions, are the most influential factors for nest-space use. This study expands our knowledge of parent–offspring communication and how environmental factors may lead to differential nest-space use, which may be regarded as the earliest form of habitat preference in birds.

Significance statement
Nests are usually constrained in space but designed to protect offspring from the environment while giving them limited possibilities to express behavioural diversity. This is particularly true for bird nests, where nestlings are usually packed in close contact with one another and without much space for movement, except begging. Here we demonstrate that nest features, such as available nest space together with environmental conditions surrounding a nest, influence nesting strategies and behaviours, including social interactions between nest mates, which further leads to fitness consequences. Our results seem to be of great importance for habitat selection theory in birds, in particular regarding the early development of habitat preferences (imprinting) and use. On the other hand, the result may also have important implications for conservation issues given that nestling behaviour may be used as a determinant of environmental quality.

Keywords Altricial nestlings · Weather conditions · Food supply · Nest-space use · Hoopoe · Upupa epops

Introduction
The ability to choose a proper habitat is an important component of an individual, influencing its life expectancy, performance and fitness (Morris 2011; Uboni et al. 2017; Maresh Nelson et al. 2020). Habitat choice can concern a variety of aspects of life, including the choice of a proper foraging or roosting site, in relation to successful mate attraction or in providing the necessary requirements for reproduction (Gibson et al. 2016; Schwemmer et al. 2016; Evens et al. 2016).
The ability to select the right habitat could be innate, learned or achieved by imitating the choice behaviour of conspecifics, as suggested by the social attraction (Piper 2011; Nielsen et al. 2013; Dixson et al. 2014) or commodity hypothesis (Danchin and Wagner 1997).

Differential habitat use may be important throughout the whole lifetime of an individual, but it can occur even at a very early stage of life. For instance, juvenile fish or anuran larvae actively select a specific habitat during their development (Alford 1986; Armstrong and Nislow 2006). For birds, not much information is available regarding early habitat use, since parent birds usually provide sophisticated rearing environments (nests) for offspring development but limited space to move (Collias and Collias 1985). Thus, for altricial nestlings, the need for differential nest-space use is limited, and whether it is even developed at such an early age is largely unknown.

Weather and predation risk are important environmental factors that may be difficult to forecast, at least in specific landscapes and climatic regions (Lima 2009; Chalfoun and Martin 2010; Rivrud et al. 2010; Reichert et al. 2012; Sauve et al. 2021). Thus, when choosing a habitat, animals have to cope with more or less unpredictable environmental conditions throughout their whole life and adjust their life history traits to changes in the environment (Tököli et al. 2012). Weather conditions during the early stages of life can affect offspring both directly and indirectly, e.g. in terms of thermal conditions (Sauve et al. 2021), food availability (Arlettaz et al. 2010, 2017), parental care, e.g. the provisioning effort (Dawson and Bortolotti 2000; Öberg et al. 2015), or changes in parental allocation rules and decisions (Chalfoun and Martin 2010; Ryser et al. 2016). Several studies have been conducted on the effect of seasonal variation in weather conditions on breeding performance (e.g. Tobolka et al. 2015; Wann et al. 2016; Coppes et al. 2021), but less is known about short-term, less predictable changes in the weather, which may require nestlings to respond immediately to, e.g. thermal stress or reduced food availability during rainy periods (Jovani and Tella 2004; Pipoly et al. 2013; França et al. 2020). As a result, offspring may develop strategies to cope with unpredictable situations (Tököli et al. 2012), for instance by selecting the most profitable areas in terms of food acquisition, in particular when food is scarce (Budden and Wright 2005). In this context, habitat selection represents a complex decision-making behaviour, where individuals are faced with choices that differ in terms of habitat quality and which can influence fitness outcomes via the costs (e.g. predation risk) or benefits (e.g. food acquisition) of particular habitat use. Consequently, individuals are often assumed to use the habitat where fitness is the highest (Jones 2001; Morris 2003; Piper 2011; Chalfoun and Schmidt 2012).

In a previous study on cavity-nesting European hoopoes (Upupa epops), we demonstrated that when provided with sufficient nest space, nestlings reveal differential nest-space use and even express certain behaviours at specific nest locations (Nuhlíčková et al. 2021). The use of these locations is likely to be affected by parental feeding location, namely whether parents feed inside a nest or from the outside through an entrance hole, but also by nestling age and brood size, which seem to be determinants for nesting nest-space use (Nuhlíčková et al. 2021). For instance, it has been shown that older siblings move more towards the entrance hole, suggesting better locomotor skills with increasing age of hoopoes. Nestlings fed from outside the nest box spent more time under the entrance hole, waiting for parents with food. Furthermore, there is an indication that nestlings face a trade-off regarding nest-space use (Nuhlíčková et al. 2021). If they stay in front of the entrance hole, they receive more food but may suffer higher predation risk and less favourable thermal conditions. On the other hand, if they stay in the dark corner, they may spend more time resting, better protected from predators or avoiding unfavourable ambient thermal conditions but are less likely fed (Wiebe 2001; Wesołowski 2002, 2017; Dawson et al. 2005; Rhodes et al. 2009; Wesołowski and Maziarz 2012; Maziarz and Wesołowski 2013).

Therefore, the main question of this study is whether local environmental (e.g. weather) conditions surrounding a nest influence food provisioned to nestlings, offspring nest-space use and consequently the trade-off nestlings face regarding these two main locations in the nest? Hoopoes are an insectivorous bird species that strongly depend on large prey items and react very sensitively to varying weather conditions. This could be due to changes in prey activity and thus food availability (Arlettaz et al. 2010; Nuhlíčková et al. 2016), altered foraging patterns (Dawson and Bortolotti 2000; Radford et al. 2001) or increased energy demands (Naef-Daenzer 2000; Grieco 2002; Tremblay et al. 2005; Arlettaz et al. 2010), often leading to starvation and the death of subordinate chicks (Martin-Vivaldi et al. 1999; Krištín 2001; Arlettaz et al. 2010). For this reason, we provide a comprehensive analysis involving different weather parameters together with food provisioned to nestlings on different time scales to examine the use of the two most important nest-space locations dedicated to either the food acquisition or resting (trade-off) of hoopoe nestlings (Nuhlíčková et al. 2021).

In the first step, we compare offspring nest-space use in two study years representing different environmental conditions and try to determine which of the following socio-ecological factors may act as important influential parameters for nest-space use: (1) parental feeding strategy (feeding from inside or outside), (2) prey size (in cm of prey items delivered by parents), (3) feeding rate, including feeding

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visits of both parents, (4) brood size and (5) nestling age. In the second step, we investigate the influence of weather parameters (temperature, humidity and size of prey items delivered) on two different temporal scales: (i) the weather conditions over a longer period — i.e. the nestling period from day of hatching to the day of the behavioural recording (the long-term effect), and (ii) the weather conditions just prior to the observed response, namely the weather on the day of the camera recording up to the moment when the recording was finished (the short-term effect).

We hypothesize that the two study years, which differed in the overall breeding conditions, will be reflected in the distinct parent–offspring behaviour at the nest, changing the importance of the two most frequently used locations and hence the trade-off in their use. We predict that hoopoe nestlings will use the frontal area of the nest box more often in the year with less favourable conditions, for example to be closer to parents with food (Ostreiher 2001; Kolliker and Richner 2004; Dickens et al. 2008) when less energetic food is delivered (see Arlettaz et al. 2010). We assume that parental control over the location where food is delivered will be one of the most important determinants of nest-space use (Nuhlicková et al. 2021). In this context, offspring may be fed more from the entrance hole even at a younger age, suggesting a change in parental provisioning behaviour (Dawson and Bortolotti 2000; Radford et al. 2001) in the season with less favourable conditions.

Depending on the temporal scale, we also expect that the influence of ambient weather conditions will differ. Weather parameters for example could indirectly but also directly influence, how much time nestlings would need to stay at a specific location. For example, an indirect influence of weather on nest-space use could happen via food availability. Long-term effects of weather e.g. long periods of cold and rainy weather, could affect overall food availability representative for the whole season and hence affect parental food supply. In this context, food availability depends not only the time but also the extent of adverse weather events, which might be important for insect availability, e.g. in terms of fluctuations in prey populations (either seasonal or inter-annual), on prey detectability (partly determined by the prey activity pattern), or on prey accessibility (reviewed in Arlettaz et al. 2010, 2017). For instance, rain and cold periods during emergence and the mating flight, can cause shorter life span of adult scarabaeid beetles and a prolonged maturation of the eggs in the body. The older larvae are able to avoid extreme high temperatures and drought by moving to deeper soil (Huiting et al. 2006), making them inaccessible for hoopoes. In this context, adverse weather events can be of different importance. Thus, we expect that weather conditions may operate indirectly, namely through the food supply and for longer temporal scales, shaping the overall parent–offspring behaviour and thus the importance of the two most frequently used locations. As a result, hoopoe nestlings may be forced to spend more time under the entrance hole, in periods with less favourable conditions (Dawson and Bortolotti 2000; Fournier and Arlettaz 2001; Arlettaz et al. 2010; Tagmann-Ioset et al. 2012; Öberg et al. 2015). In line with this, we would not necessarily expect that nestlings immediately respond to each change in weather conditions, especially if the subsequent weather event (situation) does not last for long. For instance, short occasions of rainy weather may not necessarily elicit a nestling response in terms of an adjustment of the position in the nest. Offspring may, despite changing weather, remain either hidden in the corner of the nest box or stay under the entrance hole. Therefore, we predict that short-term weather changes may have a weak impact on nest-space use. However, if weather also operates indirectly, e.g. through changes in food availability, even short weather events could, to some degree, influence nest-space use (prey size). Hence, we predict that differential nest-space use is more likely obvious on a longer temporal scales, i.e. weather conditions predominant and characteristic for the whole nesting or even breeding period.

Material and methods

General methods

The study was conducted from May to July 2010 and 2011 in the region of Wagram, Austria (119°3 km², 230–350 m, 48°45’–48°47’ N 15°66’–15°73’ E). The entire hoopoe population (approximately 100 breeding pairs) bred in nest boxes of size 50 cm (length) × 35 cm (width) × 25 cm (height) (Nuhlicková et al. 2016, 2021). The nest-box size was based on the naturally occupied cavities in the upper size range (see Kubík 1960; Krštin 2001; Hudec and Šťastný 2005). The spacious nest sites allowed nestlings to show a variety of behaviours and to distinguish them in relation to nest-space use (for further details, see Nuhlicková et al. 2021). Nests were inspected twice a week to determine clutch and brood size, hatching date and hence nestling age to determine when they are old enough to start actively exploring the nest. To facilitate this, only broods in the mean age of 8.3–25.7 days were selected from the 88 nests (average age: 16.5 days). Furthermore, at this age offspring also start to engage in sibling competition, including gaping, neck stretching, standing and wing-flapping (see Kilner 2002), and are able to actively move around in the nest box.

In the Wagram vineyards, soil-inhabiting scarabaeid larvae have been determined to be the most dominant prey of hoopoes, making up 73.4% of identified prey items (Nuhlicková et al. 2016). The Common Kestrel (Falco tinnunculus) represents one of the most frequent predators in our study population, grabbing hoopoe nestlings through the
entrance hole or hunting them at or shortly after fledgling. For further specifications about the study area and species, see the detailed description in Krištofík et al. (2013) and Nuhlíčková et al. (2016).

**Field study**

Two infrared cameras were used. To determine the nest-space use of nestlings, camera 1 was installed inside the nest box. To collect data on nestling diet and parental feeding ecology, camera 2 was installed outside the nest box. For each nest, 2 h of recordings were used either in the morning (6:00–9:00 am) or in the afternoon (5:00–8:00 pm), since parents were most active in terms of food provisioning during these periods (Nuhlíčková et al. 2016). The recording time (120 min) was automatically controlled via the software. Altogether, camera recordings were obtained from 88 nests, whereby 40 and 48 nests were obtained in 2010 and 2011, respectively. Tarsus length and weight of nestlings measured at the age of approx. 18–19 days were used as supplementary information to compare chick condition in the two different breeding periods. In total, body measurements from 39 different nests were included in this study (19 and 20 nests were included in 2010 and 2011, respectively, and in total, 84 and 83 chicks were measured with the average ± SE age of 19.7 ± 2.6 and 18.8 ± 2.12 days in 2010 and 2011) (see Supplementary material 1). For further specification related to the field study, see Nuhlíčková et al. (2021).

**Determination of nest-space use**

Overall nest-space use was previously determined by Nuhlíčková et al. (2021), in which the relative frequency (%) of nestlings occurring in six same-sized areas (16.6×17.5 cm) within the overall nest space was observed (Fig. 1). In this study, nestlings seemed to preferentially use specific areas, namely the frontal area (FH) under the entrance hole and the distant area (FC) in the front corner of the cavity (see Fig. 1). Vigorous begging of chicks (e.g. calling, wing-flapping and moving towards the parent (see Kilner 2002), usually starts from the distant (FC) or frontal area (FH) immediately after parental arrival with food. Thus, with the help of recordings from camera 1, nest-space use was determined by stopping the footage for exactly one minute prior to a feeding event (parental arrival with food). At that moment, the position of each nestling was noted. When a bird was standing on the border of two areas, the nestling position was assigned to the area where the larger part of the body was located.

In this study, 6584 observations of nestlings’ positions were determined from 88 nests during two successive years (40 and 48 nests with 2174 and 4410 positions in 2010 and 2011, respectively) (see Supplementary material 2). Similar to the previous study, we found that offspring do not use the nest space evenly but show a strong preference for two locations on the grid. Thus, 79.64% (n = 5244/6584) of all positions were recorded in the frontal (FH) and the distant area (FC) of the cavity (see Fig. 1), which were subsequently included in this analysis. The nest-space use was analysed for all nestlings within a nest as one cohort, and individual nestlings were not considered separately.

**Determination of socio-ecological conditions**

The socio-ecological conditions in the study area were expressed as (1) prey size delivered to the nestlings and (2) parental provisioning effort expressed as feeding rate, calculated as the number of parental visits with food per hour. For each feeding event, the time of parental arrival with food, prey type and prey size were determined from the recordings of camera 2. Hoopoes are single-prey loaders (they usually bring only one prey item at a time to the nest) (Orians and Pearson 1979), which clearly facilitated prey identification (Fournier and Arlettaz 2001; Arlettaz et al. 2010; Nuhlíčková et al. 2016). Thus, the size of each item was possible to distinguish according to the following categories:

1. Small prey: the food item was smaller than half of a nestling’s beak size (e.g. spiders and flies up to 10 mm in body length);
(2) Medium-sized prey: the food item was larger than half of a nestling’s beak size (e.g. adult beetles up to 20 mm in body length);

(3) Large-sized prey: the food item was larger than a nestling’s beak size (e.g. larvae of beetles and butterflies, up to 40 mm in body length) (Nuhlíčková et al. 2016).

To determine the effect of social interactions on the nestlings’ nest-space use, nestling age, brood size and parental feeding strategy were included in the analysis, since they are known to influence intra-brood competition (Mock and Parker 1997; Roulin and Dreiss 2012) and hence potentially also nest-space use (see Nuhlíčková et al. 2021). In terms of parental feeding strategies, hoopoe parents enter the nest box and feed their chicks from inside the nest chamber or parents stay outside and feed them through the entrance hole (see Arlettaz et al. 2010; Ryser et al. 2016).

Determination of the weather situation

Data on weather variability were received by the nearest meteorological station in Langenlois; (48°28′N, 15°41′E; see www.zamg.at), which is located in the centre of the study area (approx. 3 to 6 km away from the nearest and farthest nest-box, respectively). To describe the annual differences in breeding conditions, four meteorological parameters were measured during the 2010 and 2011 breeding periods and were used as a supplementary information (see Table 1). A comparison of meteorological data from May to August of both 2010 and 2011 shows that during the two most important months, when the majority of camera recordings were made, May and June 2010 had almost double the rainfall compared to those two months in 2011 (see Table 1; Fig. 2), which was also reflected in higher proportion of overall humidity in 2010 than in 2011.

To determine to what extent nest-space use is affected by weather conditions, daily temperature (°C) and humidity (%) measurements were among the most important predictors used for insect activity and thus food availability for hoopoes (Arlettaz et al. 2010; Jaworski and Hilszczański 2014). To determine the importance of weather on nestling nest-space use, weather parameters were analysed in two ways: (1) an average value calculated from daily measurements, which included the period (days) from birth when the oldest sibling hatched to the day the camera recording was made (long-term effect), and (2) as the current weather variables measured on the same day when the nest-space use of nestlings was recorded (short-term effect).

Statistical analysis

It was not possible to record data blind because our study involved focal animals in the field. However to minimize observer bias, blinded methods were used for data analyses.

Annual differences in breeding conditions and nest-space use

Based on previous results and our own predictions, only the two most frequently used areas (FH and FC) were included in the analyses. A generalized linear model (GLM) was used to compare year differences in (1) feeding rate, (2) prey size, (3) chick residual body mass and (4) average offspring age, when parents started to feed them from outside a cavity. Regarding feeding rate, total feeding rate per hour for both parents was used as the dependent variable, year as the independent parameter, and nestling number as the covariate. Regarding prey size, average length (cm) of the prey size categories provisioned by both parents per nest and recording (2 h) was used as the dependent variable, year as the independent variable, and the oldest sibling hatched to the day the camera recording was made (long-term effect), and (2) as the current weather variables measured on the same day when the nest-space use of nestlings was recorded (short-term effect).

| Parameter          | May       | June      | July      | August    |
|--------------------|-----------|-----------|-----------|-----------|
|                    | 2010-2011 | 2010-2011 | 2010      | 2011      | 2010-2011 | 2010-2011 |
| Sunshine duration [h] | 99        | 291       | 212       | 230       | 249       | 167       | 217       | 255       |
| Temperature [°C]    | 14.6      | 15.3      | 18.8      | 19.7      | 22.1      | 19.5      | 19.0      | 20.9      |
| Humidity [%]        | 76        | 62        | 70        | 63        | 68        | 67        | 75        | 68        |
| Rainfall [mm]       | 113       | 63        | 117       | 50        | 83        | 50        | 162       | 31        |

Fig. 2 Monthly precipitation (mm) in two nestling seasons, 2010 and 2011. Separately shown for May (black), June (light grey) and July (dark grey). Source: www.zamg.ac.at
factor and nestling number as the covariate. To determine the year difference in body condition, residual condition, based on the relationship between body size (tarsus length in mm) and body mass in g ($r=0.45$, $F=41.5$, $p<0.0001$, $df=1.165$), was used as the dependent variable and year as an independent factor. To control for brood size, nestling number was used as the covariate. To assess the differences in offspring age between the two study years, (when parents began to be feed from outside the nest box) the average age (in days) of all chicks observed per nest was used as the dependent variable and the year as the independent factor. A multivariate analysis of variance (MANOVA) was used to determine whether there were annual differences in nest-space use, using the time that nestlings spent in the frontal area (FH) under the entrance hole and in the distant area (FC) of the nest box as dependent variables and year and brood size as independent factors. IBM SPSS 20.0 software was used for these statistical analyses.

**Importance of socio-ecological conditions**

To determine the importance of socio-ecological conditions on nest-space use, five variables were included as explanatory variables: (1) parental feeding strategy (feeding from inside or outside a nest box), (2) prey size (the size in cm of prey items delivered by parents), (3) feeding rate per one hour, including feeding visits of both parents, (4) brood size and (5) nestling age. The percentage of the time that nestlings spent in front of the cavity entrance was used as the dependent variable, and the year as the covariate. An information-theoretic approach (Burnham and Anderson 2002) was employed in this study to find the best model for describing the relationship between the five major variables and nest-space use, specifically based on individual discrete choice models. The Akaike information criterion corrected for small sample sizes ($\text{AIC}_c$) was used to evaluate and compare these different models. To achieve the best model, the methods of model averaging and multi-model inference were applied (Burnham and Anderson 2002). This method allows inference over all the models considered, but it was weighted according to the model as supported by the data. These calculations were conducted in R using the MuMIn package (Barton 2013). Akaike weights were used to quantify the relative variable importance (RVI) within the model.

**Influence of the weather situation**

A generalized linear model (Nelder and Wedderburn 1972) was constructed to explain the variability of nest-space use by chicks in relation to weather conditions, specifically in terms of temperature (°C), humidity (%) and food quality expressed by the size of prey items delivered (in cm). The dependent variable was represented by a matrix with two vectors (together capturing the ratio of time nestlings spent in the frontal area). Such a variable is firmly limited by the interval 0 to 1 (i.e. 0 to 100%); thus, we used a binomial regression procedure with logit link. The initial models were compiled as the most complex possible and their terms subsequently tested by analysis of deviance. In the next step, we simplified them by step-by-step removal of their insignificant terms, while respecting the rule of marginality — it is not possible to remove an insignificant term which is part of a significant higher-order interaction. The final model hence represents a further irreducible — minimal adequate model. Based on values of null and residual deviance, we calculated pseudo-$R^2$, which captures the proportion of the variation in the dependent variable that is predictable from the independent variable(s). Models were visualized using sigmoid curves with 95% confidence intervals derived from fitted values $\pm 2 \times \text{SE}$ on the link series, whereas the inverse of the link function was used to map the fitted values and the upper and lower limits of the interval back on to the response series. The modelling and creation of plots were done in the R 3.4.4 environment for statistical computing (R Core Team 2018).

**Results**

**Annual differences in breeding conditions and nest-space use**

We found a significant year-on-year difference in (i) average feeding rate (GLM: $F=4.71$, $p=0.033$; Fig. 3a), while controlling for nestling number ($F=0.07$, $p>0.792$), (ii) size of prey items provided by the parents (GLM: $F=29.02$, $p<0.0001$; Fig. 3b) and (iii) chick body mass not explained by body size (GLM: $F=14.47$, $p<0.0001$, $df=1$; Fig. 3c), while controlling for brood size ($F=1.09$, $p>0.3$) and average offspring age when parents started to feed them from outside the cavity (GLM: $F=5.045$, $p=0.031$, $df=1.37$; Fig. 3d), when controlling for brood size ($F=1.565$, $p>0.219$). As a result, feeding rate, prey size and nestling body condition were reduced in the adverse breeding season of 2010 in comparison to the 2011 season, which had more regular conditions (see Fig. 3a–c). In addition, in the adverse season 2010, parents started to feed chicks at a younger average age from outside than in the season with more regular conditions (see Fig. 3d).

Regarding the nest-space use compared in two nestling seasons, we found a significant effect on the use of the two nest positions investigated (MANOVA: Wilks’ lambda $=0.89$; $F=4.94$; $df=2.8$; $p=0.009$), but no effect of brood size (MANOVA: Wilks’ lambda $=0.96$; $F=1.58$; $df=2.8$; $p>0.21$). In 2011, nestlings spent less time in the frontal area (FH) but more time in the distant area (FC) of
the nest box than in the adverse 2010 season (Fig. 4). The results further suggest a stronger impact on nest-space use in the distant (FC) ($F = 9.78$, $p = 0.002$) than in the frontal area (FH) ($F = 4.33$, $p = 0.04$) (see Fig. 4).

**Importance of socio-ecological conditions**

Analysing the importance of the five parameters to explain nest-space use, but independent of year-to-year differences, an information-theoretic approach revealed the highest relative variable importance (RVI) for parental feeding location. The RVI value was 1, which suggests parent feeding location to be definitely in the model. The positive parameter estimate indicates that the time nestlings spend in the frontal area (FH) increases with parents starting to feed from outside. According to the RVI value (0.83), prey size is also very likely in the best model to explain the time spent in the frontal area (FH) under the entrance hole. The negative parameter estimates, however, suggests a negative relationship between the size of delivered prey by the parents and time nestlings spent in front of the entrance hole (FH). Finally, the relatively high RVI value for nestling age (0.78) also indicates that nestling age is likely to be in the best model, suggesting that nestling time spent in the frontal area (FH) of the nest box increases with their age. Interestingly, brood size, which could be seen as an indicator of competition over food resources, seems highly unlikely to even be in the model (0.29), and the feeding rate does not seem to be important, given the lower RVI value (0.65), which indicates that the probability being in the model is mere chance (for the overall model output based on particular RVI values, see Table 2 and Supplementary material 3).

**Influence of the weather situation: long-term effect**

The model examining the influence of the weather situation on offspring that faced such influence throughout the
whole nesting period, explains 31.1% of the nest-space use variability (see Table 3). When small prey was delivered, the model exhibits a positive relationship between humidity (%), temperature (°C) and the time nestlings spent in the frontal area (FH) of the nest box. This pattern was also positively related to an ambient temperature of approximately 20 °C. Nestlings spent more than half of their time under the entrance hole (FH) when humidity was excessively high at the times when small prey was delivered (Fig. 5). However, the model suggests a completely different pattern when large prey was supplied. At lower to moderate temperatures (~10 °C and ~15, respectively), we found only a slightly positive relationship between humidity (%) and the time hoopoes spent under the entrance hole (FH). Despite increasing humidity outside the cavity, hoopoes supplied with large prey still spent the majority of their time hidden in the distant corner (FC) of the cavity. Hoopoes tended to move more often towards the entrance hole (FH) only when the ambient temperature was higher (~20 °C).

### Influence of the current weather situation: short-term effect

Examining the immediate response of nestlings in relation to the current weather situation, the model exhibits only a very weak relationship between humidity (%), temperature (°C) and the time nestlings spent in the frontal area (FH) of the nest box, as less than 1% of the variability was explained (see Table 4). However, examining the effect of the current weather situation in combination with prey size, the

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**Table 2** Model output specifically based on individual discrete choice models. Given are model-averaged parameter estimates (Estimate), standard error (SE), adjusted standard error (Adjust. SE), z-value, probability (P) and relative variable importance (RVI) for the relationship between the different parameters and the time spent in the frontal area (FH) of the nest box.

| Parameter            | Estimate | SE   | Adjust. SE | z       | P(>|z|)   | RVI |
|----------------------|----------|------|------------|---------|----------|-----|
| Feeding location     | 0.4059   | 0.1062 | 0.1076     | 3.773   | 0.000161 *** | 1.00 |
| Prey size            | –20.7556 | 7.6239 | 7.7293     | 2.685   | 0.007246 ** | 0.83 |
| Nestling age         | 15.2393  | 6.8297 | 6.9320     | 2.198   | 0.027921 *  | 0.78 |
| Feeding rate         | 0.5040   | 0.2729 | 0.2761     | 1.825   | 0.067942  | 0.65 |
| Brood size           | –1.7614  | 2.6246 | 2.6597     | 0.662   | 0.507812  | 0.29 |
| (Intercept)          | 10.1979  | 32.0197| 32.2448    | 0.316   | 0.751801  | 0.00 |

Null deviance = 3731.6; residual deviance = 2572.9; pseudo-$R^2$ = 0.311

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**Table 3** A generalized linear model, its main factors and their second-order interactions affecting the proportion of time (%) that hoopoe nestlings spent in the frontal area (FH) of the nest box. Influence of the long-term weather situation.

| Parameter            | Estimate | Std. error | z       | p       |
|----------------------|----------|------------|---------|---------|
| Intercept            | –2.1514  | 1.7668     | –1.22   | 0.223   |
| Humidity             | 0.1196   | 0.0067     | 17.97   | <0.001 *** |
| Temperature          | –0.2163  | 0.0864     | –2.51   | 0.012 *  |
| Average prey size    | –5.2575  | 0.8956     | –5.87   | <0.001 *** |
| Temperature x average prey size | 0.2250  | 0.0480     | 4.69    | <0.001 *** |

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**Fig. 5** Visualization of a generalized linear model which captures the influence of weather conditions (temperature, humidity, size of prey items delivered) on the proportion of time hoopoe nestlings spent in the frontal area (FH) of the nest box. Influence of the long-term weather situation.
The model explains 22.4% of the variability. The strong effect of prey size (Table 4) was reflected in the negative relationship between prey size and the time nestlings spent in the frontal area (FH) of the nest box. When small food was delivered, hoopoes spent the majority of their time under the entrance hole (see Fig. 6). In contrast, when large food was delivered, nestlings spent most of their time in the distant corner of the nest box (see Fig. 6). We found that most of the variability was explained exclusively by prey size and the current weather conditions had almost no effect on nest-space use.

Discussion

Our results suggest differential habitat use in hoopoes already at a very early stage of life (Nuhlíčková et al. 2021). In contrast to other taxa (Alford 1986; Armstrong and Nislow 2006), not much information is available on the possibility of early habitat use in birds. Given that parents provide nests which are specifically designed to meet their offspring’s needs during their early development and, on the other hand, offspring mobility is usually restricted due to space constraints (Collias and Collias 1985), early habitat use is unlikely to play a significant role. In this context, hoopoes are slightly different, because hoopoe parents do not build nests; they rather search for more or less suitable cavities for raising their offspring (Kubík 1960; Krištín 2001; Hudec and Šťastný 2005). More importantly, we can show that local environmental (e.g. weather) conditions surrounding the nest influence offspring nest-space use. Specifically, two important environmental factors, weather and predation risk (Lima 2009; Chalfoun and Martin 2010; Rivrud et al. 2010; Reichert et al. 2012; Sauve et al. 2021), seem to affect the trade-off nestlings face when choosing between the two preferred localities in the nest. Given the relatively long developmental period of hoopoe nestlings (Krištín 2001; Hudec and Šťastný 2005) and the fact that they are still immature and clumsy when the recordings were made, it is surprising to observe nestlings actively responding to environmental changes with differential nest-space use. So, the question arises whether differential habitat use of nestlings is an independent decision and innate (Nielsen et al. 2013; Dixson et al. 2014) or simply driven by physiological needs, e.g. nestlings following the state of hunger (Dickens et al. 2008; Marques et al. 2011) or thermal requirements (Gilbert et al. 2010). When provided with sufficient space, nestlings differentially used their nest space, namely they showed a preference for two locations. To receive food, nestlings apparently prefer to move towards the entrance hole to await the arriving parent, a behaviour that has also been observed in other bird species (e.g. Ostreiher 2001; Köllicher and Richner 2004; Dickens et al. 2008). On the other hand, the distant corner of the cavity was used for other behaviours, especially those related to resting or huddling activities (Nuhlíčková et al. 2021), suggesting that this is a site safer from predators (Wesołowski and Maziarz 2012; Wesołowski 2017) or a more stable site regarding thermal

Table 4 A generalized linear model, its main factors and their second-order interactions affecting the proportion of time (%) that hoopoe nestlings spent in the frontal area (FH) of the nest box. Influence of the current weather situation

| Estimate | Std. error | z     | p     |
|----------|------------|-------|-------|
| Intercept| 17.5459    | 1.2349| 14.21 | <0.001***|
| Humidity | −0.0089    | 0.0030| −3.01 | 0.003**  |
| Temperature| −0.7312    | 0.0619| −11.80| <0.001***|
| average prey size | −9.9386 | 0.6840| −14.53| <0.001***|
| temperature x average prey size | 0.4144 | 0.0345| 12.00 | <0.001***|

Null deviance = 3731.6; residual deviance = 2895.3; pseudo- \( R^2 \) = 0.224

Fig. 6 Visualization of a generalized linear model which captures the influence of weather conditions (temperature, humidity, size of prey items delivered) on the proportion of time hoopoe nestlings spent in the nest frontal area (FH). Influence of the current weather situation
conditions (Amat-Valero et al. 2014; Larson et al. 2015). Thus, if possible, nestlings appear to actively prospect different nest areas for different purposes. This suggests that nestlings are already able to follow some, probably innate rules when selecting their whereabouts.

Beside the importance of environmental conditions, we could also confirm the role of parental feeding strategy as already demonstrated in an earlier study (Nuhlíčková et al. 2021). Based on the high relative variable importance (RVI) value derived using the information-theoretic approach, parental feeding strategy is also in this study the most important determinant for explaining nest-space use. Similarly to Nuhlíčková et al. (2021), we could show that when parents feed from outside, nestlings spend more of their time under the entrance hole compared to the resting patch in the distant corner of the cavity (Nuhlíčková et al. 2021). This indicates the ability of nestlings to predict parental arrival through the entrance hole, given that the entrance hole represents the only access of parents to feed offspring. The predictability of the feeding location further means that hoopoe nestlings are forced to beg from this patch to successfully receive food of their arriving parents. These results are also in accordance with previous studies on other bird species (Bonabeau et al. 1998; Parker et al. 2002; Royle et al. 2002; Kolliker and Richner 2004; Budden and Wright 2005; Porkert and Špinka 2006). However, when comparing the parental feeding strategy in relation to variation in breeding conditions, namely the two different nestling seasons, we also found that during the adverse season parents started to feed their chicks already in an earlier age from outside the nest box compared to the season with more regular conditions. Changing the feeding location from inside the cavity to the entrance hole may indicate a parental strategy to avoid time-consuming decision-making on which offspring to feed at any given feeding event (see Tanner et al. 2007; Ryser et al. 2016). In this context, parents may increase their search time at the cost of time they spend with their offspring, which may be especially beneficial when food is restricted (Naef-Daenzer 2000; Grieco 2002; Tremblay et al. 2005). This further implies, that when the need for investment in gathering food is increased (Grieco 2002), e.g. when food availability is low (Arlettaz et al. 2010; Öberg et al. 2015), hoopoe parents started feed from outside earlier, with the consequences that only chicks in good condition may manage to get enough food and to fledge (Ryser et al. 2016). Our results thus indicate that the choice of the feeding location as well as the timing of the switch between them is under parental control (Nuhlíčková et al. 2021) and, furthermore, is influenced by food availability, e.g. reflected by the different conditions of the two study years.

Given the importance of parents to determine the feeding location and consequently also nestling nest-space use, the question of whether nestling have any independent decision options arises. One parameter identified in this context seems to be prey size, which could directly influence nestling condition (e.g. their state of hunger) and hence affect their nest-space use independent of parental control. The high relative variable importance (RVI) suggests prey size to play a role for nest-space use. The negative parameter estimate in fact implies that the time in front of the entrance hole decreases with increasing prey size. This indicates that prey size and not feeding rate is better reflecting foraging conditions during a breeding season. Thus, when examining the role of ambient weather on nestling nest-space use, prey size has to be taken into account.

Our results from examining weather parameters together with food delivered to offspring indicate that prey size was the most important factor influencing offspring nest-space use, suggesting a crucial role of large insects for hoopoe nestlings (Martin-Vivaldi et al. 1999; Fournier and Arlettaz 2001; Arlettaz et al. 2010), which was supported by both models applied in this study. When parents fed with smaller prey, nestlings spent more time under the entrance hole, but when supplied with large prey, they remained hidden in the distant area of the cavity and started to beg from this patch. These results were additionally supported by comparing the overall breeding situation in two study years. Smaller prey, lower feeding rates, parental provisioning done almost exclusively through the entrance hole and lower chick body mass were found in the adverse year of 2010. Further support is provided by the negative effect of high humidity or rainy weather on food availability, which has been shown to be important for (i) offspring survival (Jovani and Tella 2004; Golawski 2006; Geiser et al. 2008; Arlettaz et al. 2010; Öberg et al. 2015), (ii) parental care in relation to a change of food allocation rules (Caro et al. 2016; Ryser et al. 2016), (iii) provisioning effort (Davis et al. 1999; Radford et al. 2001; Hoset et al. 2004) and finally (iv) offspring response in terms of scramble competition to occupy the most promising areas in front of arriving parents (Ostreicher 2001; Kolliker and Richner 2004; Budden and Wright 2005; Tanner et al. 2007).

Identifying parental feeding strategy and prey size as the most important determinants of nest-space use also raises questions about how much prey supplied to hoopoe chicks depends on weather conditions. Weather conditions are in general very important for species relying on poikilothermic insects, which reduce their overall activity under adverse weather conditions and thereby becoming unavailable to predators (Arlettaz et al. 2010, 2017). The importance of weather for insect availability may be particularly complex in hoopoes, with regard to the timing and extent of adverse weather events (de Zwaan et al. 2020; Sauve et al. 2021). We found that the effects of weather parameters were stronger on the long-term temporal scales. If the weather was humid for a longer period, our results indicate that nestlings were
provided with smaller prey and spent more time under the entrance hole. Depending to a great extent on soil insects (Fournier and Arlettaz 2001; Tagmann-Ioset et al. 2012; Nuhlíčková et al. 2016), hoopoes may have difficulties reaching their prey, even with their long-curved bills (Krištín 2001). This also indicates that adverse weather conditions operating over a longer period may result in an overall decline of the most profitable prey, namely large scarabaeid larvae (Nuhlíčková et al. 2016). In this context, adverse weather events can be of different importance, which in turn influence parent–offspring behaviour in the nest. Our results indicate that hoopoe parents switch to other prey types, such as large larvae, as already shown in Fournier and Arlettaz (2001) and Arlettaz et al. (2010). Thus, these results suggest that excessively humid weather operating on longer temporal scales indirectly affects food provisioned to nestlings and subsequently parental provisioning behaviour (Geiser et al. 2008; Arlettaz et al. 2010; Caro et al. 2016) and offspring response, for instance in terms of competitive positioning (Kölliker and Richner 2004; Budden and Wright 2005; Tanner et al. 2007). Thus, under scrambled competition to be closer to the feeding parents, nestlings attempt to occupy the most promising locations, which is the frontal area of the nest.

On the other hand, if short-term weather parameters operate directly on offspring behaviour, our model showed that the current weather conditions had almost no effect on nest-space use. This could be partly due to the fact that in our population nestlings develop in nest boxes and therefore might be less exposed to ambient weather. This may consequently mean that nest box conditions may moderate short-term weather effects on offspring nest-space use and the influence of ambient weather conditions likely depends on the temporal scale. The shorter a weather event, the less likely nestlings may respond with adjusting the nest position. However, by including also food size into the model, we found that also short weather events may elicit a nestling response; nevertheless, in this case the effects seem to be weaker. Thus, it is actually not weather but food availability and food delivered to nestlings, which has the more profound impact on nestling behaviour. This backs up other studies suggesting that prey size is an important determinant of food availability and a major driving force influencing the breeding success of insectivorous bird species relying on large prey (Martín-Vivaldi et al. 1999; Fournier and Arlettaz 2001; Arlettaz et al. 2010, 2017).

In conclusion, our results show that the way nestlings make use of their post-natal environment is not trivial. Further research would be necessary to better understand the role of the interactions among nestlings in relation to nest-space use. Given the hatching asynchrony in hoopoe broods (Hildebrandt and Schaub 2018), it would be interesting to determine individual variation in nest-space use in relation to body size asymmetry and age differences. Furthermore, it would be interesting to consider the role of parental control and sibling competition on nest-space use. For instance, conditional aspects like state of hunger or competitive abilities could be crucial to occupy the best position already before parents arrive with food.

Taken together, nest features, such as available nest space, together with ecological conditions surrounding the nest, influence nestling strategies and provide evidence for an ability in habitat selection, which may even result in fitness consequences. Due to different foraging conditions in the two study years, we found that hoopoe offspring differ in the use of the two most important areas in the nest. To explain nest-space use, parental feeding strategy and prey size, which are in turn influenced by weather conditions, are the most influential factors for nest-space use. This indicates on the one hand parental control over food allocation (Davis et al. 1999; Forbes 2007; Roulin and Dress 2012; Ryser et al. 2016; Smith et al. 2017) and on the other hand the crucial role of prey size being sensitive to weather conditions for insectivorous birds (Arlettaz et al. 2010). Thus, parental feeding behaviour, together with decision-making processes of altricial nestlings, may explain the post-natal nest-space use of cavity-nesting birds. Therefore, this study expands our knowledge on parent–offspring interactions and how environmental factors may lead to differential nest-space use, which might be regarded as the earliest form of habitat preferences in birds known in habitat selection theory.

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Analysed the data: FK, HH, SN
Wrote the manuscript: SN, HH
Originally formulated the idea: HH, SN, FK
Developed the methodology: SN, HH, ME, JS
Conducted the fieldwork: SN, HH, ME, JS
Collaborated on the imaging analysis: SN, JS

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Data availability All data generated or analysed during this study are included in this published article and its supplementary information files.
Code availability Not applicable

Declarations

Ethics approval Our research adhered to the requirements of the Austrian welfare law, and the necessary ethical approval and licences were obtained under the institutional guidelines of the Austrian Academy of Sciences. Since this is not an animal experiment in a strict sense, no license number was allocated.

Consent for publication All the authors have approved the submitted version and have agreed to be personally accountable for their own contributions and regarding questions related to the accuracy or integrity of any part of the work.

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