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Journal of Ornithology

ISSN 2193-7192 Volume 161 Number 1

J Ornithol (2020) 161:89-101 DOI 10.1007/s10336-019-01717-z



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ORIGINAL ARTICLE



Stopover ecology of autumn-migrating Bluethroats (*Luscinia svecica*) in a highly anthropogenic river basin

Philippe Fontanilles¹ · Iván De la Hera^{1,2} · Kevin Sourdrille¹ · Florent Lacoste¹ · Christian Kerbiriou^{3,4}

Received: 14 March 2019 / Revised: 27 August 2019 / Accepted: 15 September 2019 / Published online: 28 September 2019 © Deutsche Ornithologen-Gesellschaft e.V. 2019

Abstract

Wetlands are highly productive habitats used by many avian species as stopover sites during their migrations. However, these habitats are highly threatened by anthropogenic activities, such as land-use changes, the introduction of exotic species, and global warming. Further understanding on the spatiotemporal use of wetlands and their surrounding areas by migrating birds is essential to predict how these changes might affect avian en route ecology. We selected a habitat-generalist passerine, the Bluethroat Luscinia svecica, as model of how migratory birds exploit a highly anthropogenic river basin in southwestern France (i.e. Barthes de la Nive) during autumn migration. We captured and radiotracked 29 young Bluethroats in this region to shed light on different aspects of their stopover ecology and behavior such as stopover duration, habitat selection, and home-range size. We also characterized Bluethroat diet and arthropod availability in different habitats. Bluethroats positively selected pure or mixed reed beds (associated with sedge), hydrophilous tall grasslands, and corn crops. Birds staying more than one day, 8.4 days on average, used preferably corn crops. Home-range sizes were on average 5.8 ha (fixed kernels K95) and high-occupancy area (K50) was 1.36 ha with large individual variation. Bluethroats stopping over with low fuel loads tended to have larger home ranges and used preferentially corn crops, wet, or mesotrophic grasslands and rural paths. Reed beds were typically used as roosting habitat for the majority of birds, being on average 397 m apart from their daytime core areas. Short-staying birds tended to show higher fuel loads and restricted their activities to a smaller home range (1 ha) in pure and mixed reed beds. The diet of Bluethroats was dominated by ants, spiders, and beetles that were particularly abundant in corn crops. The use of corn crops by autumn-migrating Bluethroats in our study site seems to be a reasonable solution in a highly altered environment. Reducing the use of insecticides in these crops and delaying the harvesting time after mid-October are two supplemental measures that, together with a good management of the remaining wetland patches, could greatly favor Bluethroats and other migratory species in this region.

Keywords Luscinia svecica · Home range · Fuel load · Habitat selection · Corn crop · Diet · Trophic resources

Zusammenfassung

Rastplatzökologie im Herbst ziehender Blaukehlchen (*Luscinia svecica*) in einem stark anthropogenen Flussgebiet Feuchtbiotope sind ausgesprochen produktive Gebiete und werden von vielen Vogelarten als Rasthabitate während ihres Zuges genutzt. Diese Biotope sind aber auch besonders gefährdet durch anthropogene Einflüsse wie z.B. eine veränderte

Communicated by N. Chernetsov.

Electronic supplementary material The online version of this article (https://doi.org/10.1007/s10336-019-01717-z) contains supplementary material, which is available to authorized users.

Philippe Fontanilles fontanilles.oiso@laposte.net

- ¹ Observatoire d'Intérêt Scientifique Ornithologique (OISO), Cami deth Sailhetou, 65400 Lau Balagnas, France
- ² School of Biological, Earth and Environmental Sciences, University College Cork, Cork T23XA50, Ireland
- ³ Centre d'Ecologie et des Sciences de la Conservation, UMR7204 MNHN-CNRS-UPMC-Sorbonne Université, 55 Rue Buffon, 75005 Paris, France
- ⁴ Station de Biologie Marine, Place de la croix, 29900 Concarneau, France

Landnutzung, die Ansiedlung exotischer Arten und die Klimaerwärmung. Ein tieferes Verständnis der zeitlichen und räumlichen Nutzung der Feuchtgebiete und der sie umgebenden Gebiete durch Zugvögel ist wesentlich, um einschätzen zu können, wie diese anthropogenen Veränderungen die Zugökologie beeinflussen mögen. Wir wählten einen Habitat-Generalisten, das Blaukehlchen (Luscinia svecica), als Modellorganismus dafür, wie Zugvögel ein hochgradig anthropogenes Flussgebiet in Südwest-Frankreich (z.B. Barthes de la Nive) während des Herbstzugs ausnutzen. Hierfür fingen wir in der Region 29 junge Blaukehlchen und verfolgten sie per Radiotelemetrie, um unterschiedliche Aspekte ihrer Rastplatzökologie und ihres Verhaltens wie z.B. die Dauer des Zwischenstopps, die Wahl des Habitats und ihren Aktionsradius dort aufzuklären. Für die unterschiedlichen Habitate bestimmten wir außerdem die Nahrung der Blaukehlchen sowie generell die Menge der verfügbaren Arthropoden. Die Blaukehlchen suchten gezielt reine oder gemischte Schilfgebiete (mit Abwässern), am Wasser gelegene hochbewachsene Grünflächen und Maisfelder auf. Tiere, die länger als nur einen Tag - im Schnitt 8.4 Tage - blieben, bevorzugten Maisfelder. Der Aktionsradius betrug im Schnitt 5,8 ha (fixed kernels K95), wobei die Hochdichtefläche (K50) bei 1,36 ha mit großen individuellen Unterschieden lag. Blaukehlchen, die den Zwischenstopp mit geringen Energievorräten erreichten, tendierten zu größeren Aktionsradien und bevorzugten Maisfelder und sehr feuchte oder mesotrophische Wiesen und Feldwege. Röhrichte wurden von der Mehrheit der Vögel üblicherweise als Schlafplätze genutzt und lagen im Schnitt 397 m von ihren Aufenthaltsgebieten tagsüber entfernt. Vögel, die nur kurz blieben, besaßen in der Regel größere Energievorräte und begrenzten ihre Aktivitäten auf einen kleineren Aktionsradius (1 ha) innerhalb der Schilfgebiete. Die Nahrung der Blaukehlchen bestand hauptsächlich aus vor allem in Mais häufig vorkommenden Ameisen, Spinnen und Käfern. Die Verwendung von Mais schien für die Blaukehlchen in unserer Untersuchung auf ihrem Herbstzug eine sinnvolle Ausnutzung der stark veränderten Umwelt zu sein. Weniger Insektizide im Mais einzusetzen und die Maisernte auf die Zeit nach Mitte Oktober zu verschieben wären zwei Maßnahmen, die zusätzlich zu einem guten Management der verbliebenen Feuchtgebiete den Blaukehlchen und anderen Zugvögeln in dieser Region sehr nutzen würden.

Introduction

The global conservation of migratory birds not only relies on the protection of their breeding and wintering habitats, but also on the presence of suitable stopover sites along their migratory routes (Hutto 2000; Yong et al. 1998; Newton 2008). Coastal marshes, estuaries, and other types of wetlands are highly productive habitats largely used during migration by waterbirds and other avian species, constituting key conservation areas (Czech and Parsons 2002). Wetland habitats experienced a dramatic reduction and an increased fragmentation during the 20th century, mainly as a consequence of the pressure of human activities such as urbanization and the drainage for agricultural use (von Behren 2007; Whited et al. 2000; Czech and Parsons 2002). As a result, wetlands usually remain now embedded in a complex and highly altered human matrix, where every year migrating birds rest and search for food to undertake their next flight bout.

The wetland remnants that many birds use as stopover sites are normally small and face some common problems that, in some cases, can be mitigated by human intervention. For example, many wetland managers implement measures to avoid clogging, which could lead to bush encroachment and the progressive disappearance of marsh vegetation (Clark and Wilson 2001). The control of invasive and exotic species, that can alter wetland biodiversity and functioning, is also a common practice (Fontanilles et al. 2014; Arizaga et al. 2013). These measures would increase the quality and carrying capacity of the wetlands, which would be very beneficial for migrating birds (e.g. fuel deposition; Delingat and Dierschke 2000). Another aspect that has hitherto received little attention is the potential management and planning of the areas surrounding the wetland. These adjacent areas might be intensively used by migrants given the restricted size of wetland patches and the potential high intra and inter-specific competition with other migratory and resident individual birds. This situation could be particularly exacerbated during the migration peaks, when large number of individuals can co-occur in a single location (Newton 2004). Such circumstance predicts that many migrants will be forced to expand their home ranges and use alternative habitats out of the wetland. In this context, identifying which alternative habitats are positively selected by migrating birds during their brief stopovers would be essential to design buffer areas that best meet the requirements of these migrants en route. However, this knowledge is limited in many key stopover areas.

In this study, we analyzed the stopover ecology of migrating Bluethroats *Luscinia svecica* in Barthes de la Nive (France) during autumn migration. Barthes de la Nive is a mosaic of wetlands, riparian woodlands, and farmlands located near the Adour River mouth in the southwestern Atlantic coast of France. The scattered wetlands in this region attract a large diversity of both aquatic and nonaquatic birds during migration periods (Fontanilles et al. 2012). The only radiotracking study carried out in this area so far showed that the globally endangered Aquatic Warbler *Acrocephalus paludicola* had a strong preference for reed beds but, to a lesser extent, it also made use of some farmlands (Fontanilles et al. 2014). This highlights the relevance that the surrounding matrix might have even for a wetland specialist as the Aquatic Warbler. Conversely, Bluethroats are migratory birds that can use a large range of habitats, mainly reed beds and marshlands (Arizaga et al. 2006; Musseau et al. 2017), but also bushlands, woodland, and farming habitats (Cramp 1988; Cornulier et al. 1997; Chiron 2017; Berndt and Hölzel 2012), making it an interesting model to study its stopover ecology and habitat selection during migration. Arizaga et al. (2011) radiotracked 20 autumn-migrating Bluethroats in Txingudi (North Spain), a wetland located less than 30 km away from Barthes de la Nive. Their results showed a strong selection of reed beds, low-halophytic vegetation, and tidal flats. However, Txingudi is located in a more urbanized environment that probably restricts bird movements and its proximity to the river mouth allows the presence of habitats that are lacking in Barthes de la Nive (e.g. halophytic vegetation). On the other hand, bird-ringing information obtained from both wetlands during autumn migration showed dramatic differences in the number of within-season recoveries, which are much more usual in Txingudi than in Barthes de la Nive (Arizaga et al. 2011; Fontanilles unpublished data). These differences could be a consequence of longer stopover duration of Bluethroats in Txingudi than in Barthes de la Nive, but they could also be caused by the existence of larger home ranges in Barthes de la Nive, something that would be possible in its much less urbanized surrounding. All these aspects suggest a different stopover ecology and behavior of Bluethroats between both wetlands in spite of their geographic proximity that remains still unresolved.

To better understand the stopover ecology of the Bluethroat in Barthes de la Nive, we performed a radiotracking study during autumn migration that aims to shed light on (1) the stopover duration and the proximate factors that affect it, (2) habitat selection and home-range sizes, and (3) we complemented the study with an analysis of Bluethroat diet and the availability of food resources. Although Bluethroats are not globally threatened, some populations have been considered to be under high risk of extinction (Huntley et al. 2007). Consequently, we expect that the spatial information generated in this study could be valuable for the competent authorities to design management practices contributing to the conservation of Bluethroats and other birds during their migrations.

Materials and methods

Study area and vegetation map

Barthes de la Nive (43°27'N; 01°28'W) is a 442-ha mosaic of natural and anthropogenic habitats (Fig. A1), whose

remaining wetlands have been maintained safe from complete drainage due to its role in reducing the risk of flooding in Bayonne city. The wetlands are connected to the Adour River by channels so that its hydrology is influenced by the flow of the river and the tidal regimes. However, its natural influence and ecologic functions in Adour estuary were drastically affected to urbanize the city, to develop port facilities and dykes, which virtually destroyed the original mudflats and intertidal areas. Our study site is now a protected area included in the European Natura 2000 network (FR7200786). The non-urban areas near the river have been widely used for pasture, as hay meadows and other agricultural purposes, being corn plantations the primary crop nowadays. Farming in the region uses a reasonably low quantity of fertilizers and no insecticide. Field abandonment and lack of management is increasing gradually bush and tree encroachment, which is changing dramatically the landscape. A vegetation map was obtained from photo interpretation and field validation for summer 2014 and 2015 (see Fig. A1 in Supplementary Material, Table 1). This approach showed the relative importance of each habitat in the region (see Table 1).

Bluethroat trapping and body measurements

Mist-netting sessions were performed from mid-August to late September, when migrating Bluethroats stop over in this region during their post-breeding migratory period. Note that Bluethroats do not breed in Barthes de la Nive or use it for moulting (Fontanilles et al. 2012). Ringing sessions took place in five different locations across the study area: four wetlands and one corn crop (see details in Supplementary

 Table 1
 Abbreviations for the different habitats available in Barthes

 de la Nive with its corresponding percentage of cover in the study
 area

Code	Area (ha)	Habitats
WAT	32.91	Open water
PRB	19.35	Pure reed bed <i>Phragmites australis</i> > 75%
MRB	11.89	Reed bed mixture <i>Phragmites australis</i> (> 25%) + <i>Carex sp</i>
WOR	14.63	Wooded reed bed (wood $< 50\%$)
WGR	21.37	Wet grassland
MGR	64.04	Mown grassland
CGR	10.79	Mesophilic grassland
CRO	69.4	Crop field
FER	0.84	Fern
TAL	3.79	Hydrophilous tall herb
WOL	171.5	Wood land
PAT	12.05	Natural path
HOU	7.64	House
RAW	1.54	Railway

Material Fig. A1, Ap. 3, Table 2). These five sites were reasonably distant to each other (see Fig. A1) to reduce potential geographic bias and, according to previous studies (Fontanilles et al. 2012; Fontanilles 2014), they represent the two main habitats (i.e. reed bed, corn crops), where Bluethroats occur during their stopover at Barthes de la Nive.

To maximize the number of captures, we used one male song playback from 30 min before dawn (when mist-nets were open) until a tagged bird was released (when mist-nets were furled; de la Hera et al. 2017). Using the tape lures only 30 min before sunrise, we reduced the possibilities of forcing the landing of migrating Bluethroats and we expected to capture only birds that decided voluntarily to stop over in our study area (Schaub et al. 1999; Arizaga et al. 2015). During our ringing sessions, we captured 58 Bluethroats (28 in 2014 and 30 in 2015).

Several morphological measurements were taken from each bird to obtain a composite index of bird body size (Freeman and Jackson 1990; Tellería et al. 2013, see Supplementary Material). We also recorded body mass (± 0.1 g) and standard scores of fat and muscle (Busse 2000). Body size-corrected measurements of body mass were used to estimate individual fuel loads. This was done using a linear regression of body mass on body size (see Supplementary Material Ap. 2, Fig. A2), where individuals with higher fuel loads showed more positive residuals (Salewski et al. 2009).

Radiotracking information

Out of the 58 Bluethroats captured, 29 juveniles were equipped with a radio-transmitter, with a minimum of four

radio-tagged individuals in each of the five ringing sites (25 birds were trapped in the reed beds and four in the corn crops; see details in Supplementary Material, Ap. 3, Table A2). Life expectancy of the emitters is typically more than 17 days and the average detection distance is 80–300 meters. We only tagged first-year birds, because the study site is mainly used by this age group, and because juveniles face their first autumn migration and, consequently, their habitat choice cannot be influenced by previous experience (Piper 2011).

On the day of capture, monitoring of tagged Bluethroats started at least 1 h after the bird was released to avoid bias linked to potential stress behavior. For birds that stayed in the area the following days after capture (see below), they were normally monitored 8 h per day during, normally, 3–4 days in long-staying individuals. After this period, we checked their presence every day, but the intensity of the radiotracking typically got reduced and depended on other duties associated with the project, particularly, the trapping and monitoring of other individuals (see Supplementary Material Ap.3 for more details).

The positions of the birds were obtained normally by triangulation (three vectors taken consecutively within less than 10 min), but we used biangulation in those cases in which landscape barriers (like canals, rivers or dense vegetation) impeded taking more than two informative vectors, or just one vector plus an estimation of the actual distance to the bird when this was observed. We used Sika receivers (Biotrack Ltd) and Yagi antennas to find the birds and vectors were delimited using a GPS (to determine observer position; Garmin Ltd.) and a compass (to obtain magnetic

Home range	50% kernel			95% kernel			
	$\beta_{\text{cond.}}$	SE _{adj.}	P value	$\beta_{\text{cond.}}$	SE _{adj.}	P value	
Global home rai	nge						
CGR	0.485	0.151	0.001	_	-	-	
PRB	- 0.362	0.208	0.081	- 0.876	0.250	< 0.001	
WGR	0.404	0.163	0.008	0.683	0.278	0.014	
WOL	0.325	0.162	0.045	0.694	0.224	0.002	
FUEL	_	-	_	- 0.664	0.210	0.002	
SOD	0.337	0.182	0.061	0.895	0.289	< 0.001	
Daily home rang	ge						
CRO	_	-	_	- 1.532	0.278	< 0.001	
MGR	0.8355	0.2645	0.002	1.509	0.286	< 0.001	
PAT	_	-	_	- 1.883	0.365	< 0.001	
PRB	- 1.3473	0.3181	< 0.001	- 2.027	0.585	< 0.001	
WAT	0.6355	0.2279	0.005	_	-	-	
WGR	-	-	_	0.683	0.269	0.014	
WOL	0.6010	0.2229	0.007	_	-	-	

CRO crop field, *CGR* mesophilic grassland, *MGR* mown grassland, *PAT* natural path, *PRB* pure reed bed, *WAT* open water, *WGR* wet grassland, *WOL* wood land, *SOD* stopover duration, *FUEL* fuel loads

Table 2 Conditional average of estimates ($\beta_{cond.}$), adjusted SE (SE_{adj.}) and *P* value for the variables included in the model set for daily and global home ranges (K50 & K95), (–) indicated a variable unselected

Azimuth). All this information was computerized using the software Cartoexploreur (Bayo Ltd.) and we considered the centroid of the triangle determined by the three corresponding vectors as the most likely geographic position of the bird or the intersections between lines, when two vectors were only available. Prior to the start of radiotracking, observers were trained in the study area and the accuracy of the triangulation method was assessed. The average error found in the estimation of the positions of transmitters hidden in the study area was $14.5 \pm 1 \text{ m} (n=69)$, which can be considered enough given the purpose of studying main habitat selection.

Stopover duration and its determinants

We used the number of days spent by each Bluethroat after being tagged (assessed by the radiotracking survey) as a proxy of stopover duration. This approach is expected to provide a better idea than ringing recaptures on how long Bluethroats stopover in Barthes de la Nive, since ringrecovery data are spatially restricted, have a lower detection probability and strongly depend on a good sampling strategy (Chernetsov 2012). According to the non-Gaussian distribution of stopover durations (see Fig. A3), we performed nonparametric tests (Wilcoxon tests, Kruskal–Wallis tests, and Spearman correlations) to evaluate which factors (year, sex, site, date of capture, muscle development, fat accumulation, and body condition) better correlated with the observed variation in stopover duration.

Habitat selection

Initially, we tested whether Bluethroat locations were randomly distributed across habitats. We applied the quadrats method (Zanimetti 2005) by defining a grid of *K* squares 100×100 m enveloping all the radiotracking points. The theoretical distribution $K_{\text{th}}(n)$ was obtained following the Poisson's law: $K_{n\text{th}} = [K \times (D^n/n!)] \times \exp(-D)$, where n is the number of points per mesh, K(n) the number of stitches having *n* points, *N* is the total number of locations, and *D* is the average density of locations by cell that is defined as *N* divided by *K*. The existence of significant differences between the two distributions (i.e., K(n) and $K_{\text{th}}(n)$) was assessed using a Kolmogorov–Smirnov test.

Secondly, for assessing habitat selection, we calculated for each habitat, the electivity index of Jacobs (1974) $I_h = U$ $-D/(U+D-2U \times D)$, where U is the proportion of locations in habitat h, and D the proportion of this habitat in the study area. The index ranges between -1 (strong rejection of a particular habitat) and 1 (strong selection). We used the proportion of locations instead of the proportion of areas in core areas to take into account the edge effects and to be more precise with the actual habitat used. We excluded habitats poorly represented showing less than five locations.

Home range estimates

The overall home-range size of each individual Bluethroat was estimated by the model of 95% kernel (K95; Worton 1989). The K95 approach is usually used in this type of studies, which would allow between study comparisons (Börger et al. 2006). Areas of high occupancy were also estimated using the 50% kernel (K50), considered as a good estimator of core areas (Börger et al. 2006). We calculated the overall home range with all the locations for each bird and also for each day, when there was a minimum of ten positions. Home-range sizes were processed using the Ranges 8v2.10 software (Anatrack ltd).

Home range analysis

We explored whether overall home-range size varied in relation to the proportion of habitats contained within it (10 habitat variables, see Table 1), several metrics of heterogeneity (mean patch size, number of habitat, or Shannon index perform of patch size distribution) and individual-associated variables (body condition, sex, and stopover duration). For this purpose, we performed General Linear Models (GLM) using the home-range size (K50 and K95) as a response variable, while habitat cover and individual-associated variables were included as explanatory variables. According to the distribution of home-range size (positive long tail distribution), we applied a negative binomial error distribution to the GLM to minimize issues related to the over-dispersion ratio in the models (i.e., as close as possible to 1; Zuur et al. 2009). Following a multi-model inference (Burnham et al. 2011; Grueber et al. 2011), we generated a set of candidate models containing all possible variable combinations and ranked them by corrected Akaike information criterion (AICc) using the dredge function (R package MuMIn; Barton 2019). We only integrated the models complying with the following conditions: (1) models do not include simultaneously correlated covariates $(R^2 > 0.7)$ and (2) models do not include more than three variables to avoid over-parameterization due to the limited data set. Since the simultaneous inclusion of habitat cover and measures of habitat heterogeneity generated important multi-collinearity problems, we decided to perform two separate modeling: one with habitat cover within the home range and individual-associated variables and a second one, with measures of habitat heterogeneity and individual-associated variables (results of this second modeling are shown in Appendix 5). We restricted this set of models using a cut-off of 2 AICc. The modeling with habitat cover resulted in four and three top models for K50 and K95, respectively (see Supplementary Material Ap. 5, Table A3). The modeling with measures of habitat heterogeneity resulted in five and two top models for K50 and K95, respectively (see Supplementary Material Ap. 5 Table B2).

Following the same approach, we explored whether daily home-range size (with a minimum of 10 positions) varied in relation to habitat cover within home range and individualassociated variables. According to the hierarchical structure of this data set (radio-tagged individuals survey each day), we treated the variable "individual identity" as a random effect, while considering the other explanatory variables (home range and individual-associated variables) as fixed effects (Zuur et al. 2009). Following a multi-model inference using General Linear Mixed Models with a negative binomial error distribution, the daily home-range analyses resulted in a total of two and three top models for the K50 and K95, respectively (see Supplementary Material Ap. 5, Table A4). The modeling with measures of habitat heterogeneity resulted in two top models for K50 and K95 (see Supplementary Material Ap. 5, Table B3). We also used a secondary method based on PCA analysis detailed in Supplementary Material Ap. 6.

Roosting habitat selection

To provide information on roosting behavior, we identified the habitats used between sunset and sunrise (roosting locations) and also estimated their distance to the daytime areas, where birds were potentially foraging. We identified the habitat of all those locations on the vegetation map. We compared the proportions of roosting locations occurring in each habitat to identify the most used by a posteriori Wilcoxon test. For birds flying to a roosting location far away from the area, used during daytime hours, we calculated the distance between the centroid of the home range exploited during daytime hours and the core of roosting locations. We then compared the proportions of habitats between these two areas (home range during daytime hours vs. roosting locations core) by a posteriori Wilcoxon test.

Diet of Bluethroat

To elucidate why birds tend to select specific habitats, we analyzed their diet and the invertebrate availability. We collected 105 fecal samples during ringing operations for all sites in 2015 (n=52) and 2014 (n=30) and only in Urdains in 2012 (n=23). All remains identified in feces were used to estimate the minimum number of prey items and the occurrence of each taxonomic group within each sample. While some bias in diet analyses was possible, because small or soft-bodied preys are less easily detected, a strong correlation has been found between prey remains in droppings and the actual composition of the diet (Davies 1977). We also assessed prey biomass using predictive models based on the relationship between body length and mass of terrestrial arthropods (Hódar 1996; see Supplementary Material Ap. 7 for additional details). We explored which factors

influenced prey abundance by testing explicitly the effects of age (young vs. adult birds), sex, and day using a Generalized Linear Model (GLM) with a Poisson error distribution. P values were corrected for over dispersion. Only taxa representing more than 2% of the prey abundance were considered.

Invertebrate availability between habitats

We estimated the variation in the abundance of (near) ground invertebrates (the main feeding substrate of Bluethroats) between the two main habitats occupied by Bluethroats (corn crops and reed beds). Given the large extension of the study area, we decided to focus our invertebrate sampling on three of the five trapping areas: the corn crop area, and Villefranque and Urdains reed beds (Fig. A1). The corn crops in the other bank of the Nive River had similar management practices so they are expected to have a similar invertebrate availability. We used three standardized colored (yellow, white, and blue) bowl traps and a pitfall glass per station (for a total of 12 stations, six in the corn crops, three in each reed bed). All stations were sampled simultaneously at three temporal stages in August 2015. Traps were deployed for 4 days (For more details, see Supplementary Material Ap. 8). For subsequent analyses, we pooled together the data of the three bowl traps of each station. We focused the analyses on those taxa, whose remains had been found in feces of trapped Bluethroats. We compared invertebrate availability, abundance, and biomass between the two reed beds and the corn crop using General Linear Models with a Poisson error distribution.

Unless specified, mean values are given \pm SE (standard error).

Results

Stopover duration and body condition effect

Tagged Bluethroats stayed on average 5 ± 1 (range 1–20) days upon capture in Barthes de la Nive, but showing clearly a non-Gaussian distribution, where up to 13 individuals left the study area, the next night after being captured (Fig. A3). The remaining 16 Bluethroats stayed on average 8.4 ± 1.3 days (see Table A1, Fig. A3).

Observed variation in stopover duration did not differ significantly between years (W = 91, p = 0.536), sites (Kruskal–Wallis H(3, 29) = 1.11, p = 0.774), sex (W = 133, p = 0.189), date of capture (r Spearman = -0.11, $t_{27} = -0.59$, p = 0.558), or muscle development (r Spearman = -0.17, $t_{27} = -0.92$, p = 0.365). The fat score was significantly associated with stopover duration (r Spearman = -0.40, $t_{27} = -2.24$, p = 0.034). However, the

individual trait that better explained stopover duration was fuel load, which was estimated from the residuals of body mass on body size (effects of body size on body mass: $\beta = 0.888$, $F_{1,27} = 100.9$, p < 0.001). Thus, Bluethroats with larger size-corrected fuel loads left the Nive basin, earlier on average (r Spearman = -0.59, $t_{27} = -3.78$, p < 0.001; Fig. 1) than birds with smaller fuel stores. Additionally, we analyzed whether stopover duration differed between habitats used. Irrespective of where they were trapped, birds that used corn crops remained significantly (W = 23, p < 0.001) longer (10 days ± 1.1 SE, n = 10) in the area than those not using this habitat type (2.7 days ± 0.6 SE, n = 19; see Fig. 2).

Habitat selection

The 29 radiotracked juvenile Bluethroats provided 1718 positions during their stay. Bluethroat locations were not randomly distributed among habitats. Their distributions differed significantly from the theoretical null distributions (Kolmogorov–Smirnov test D = 0.896, p < 0.0001). The main habitats used by all birds were corn crops and reed beds, either pure or mixed (Fig. 3). The distribution differed between birds staying 1 day, which preferred pure reed bed (W=55.5, p=0.034), and the remaining, which positively selected the corn crops (W = 162, p = 0.013). Birds stopping over for just 1 day made use of less habitat types (2.2 ± 0.3) [1-4] habitats) than birds staying longer $(4.7 \pm 0.5 [2-10])$; W = 180.5, p < 0.001). However, during the first day of stay, there were no differences between them in the preferred habitat and number of them (tests Wilcoxon, p > 0.05, Fig. A4). Bluethroats positively selected pure and mixed reed beds, corn crops, and hydrophilous tall grasslands (Fig. 4). They moderately selected paths, like wooded reed beds and mesophilic grasslands, while they avoided water,



Fig. 1 Relationship between size-corrected fuel loads and stopover duration in Barthes de la Nive, n = 29



Fig. 2 Variation in stopover duration between bluethroats that used corn crops and birds that did not in Barthes de la Nive, mean \pm SE

railways, ferns, wet grasslands, houses, woodlands, and hay meadows.

Home range analysis

The average number of locations per individual was 59.2 ± 9 (22.5 ± 1.6 for birds staying 1 day, and 89 ± 12 for birds staying longer). The average overall K95 home range was 5.8 ± 1.8 ha with a large variation between individuals (range 0.016-46.5). The core area (i.e., K50) was on average 1.36 ± 0.35 ha (range 0.004-7.6). K95 areas were larger in birds staying more than one day (9.72 ± 3 ha) than in birds stopping over just 1 day (1.07 ± 0.46 ha; W=23, p < 0.001). Similar results were obtained for the K50 core area: 0.28 ± 0.12 ha for long-staying birds and 2.24 ± 0.53 ha for birds staying only 1 day (W=18, p < 0.001).

Bluethroats with greater overall home range included more wet grassland, woodland, and mesophilic grassland



Fig. 3 Proportion (mean \pm SE) of positions occurring in each habitat type of Barthes de la Nive for all Bluethroats, those staying 1 day and those staying more than 1 day. Abbreviations as in Table 1. The number of individual home ranges in which each habitat type was found is shown between brackets



Fig. 4 Jacobs index values for each habitat in the Bluethroats radiotracked in Barthes de la Nive. Values below 0 represent habitat rejection, while values above 0 represent positive selection

within their range and also exhibited a longer stopover duration. On the other hand, Bluethroats with smaller overall home range exhibited larger quantities of body reserves and occupied areas with a higher proportion of reed bed (Table 2). Analyses performed for daily home ranges highlighted that Bluethroats with greater home range included more hay grasslands, open water, and woodlands; while Bluethroat with smaller daily home range occupied areas with a higher proportion of crop fields, natural paths, and pure reed beds (Table 2). In addition, analyses performed with measures of habitat heterogeneity showed that Bluethroats with greater overall home range exhibited a more diversified home range (Appendix 5). According to AICc, measures of habitat heterogeneity performed better than habitat cover variables for K95, while it was the opposite for K50 (Appendix 5). A second analytical method based on PCA analysis gave similar results (see Supplementary Material Ap. 6).

Roosting habitat selection

We identified the overnight habitat (n = 103 nights) in 26 Bluethroats. Pure reed bed was the main roosting habitat, with 44.2% of the nights (Fig. 5). Mixed or wooded reed beds were also well-represented within the roosting habitats, representing 17.5% and 16.8%, respectively. So, reed beds (pure, mixed, and wooden reed bed) were the preferred habitats for roosting ($80.1 \pm 8\%$ of nights) in 23 birds that stayed for at least one night. Corn crops were also used by five birds, but less frequently than reed beds: $13.1 \pm 6\%$ of nights.

83% of the birds that spent the night in reed beds (pure, mixed, and wooden reed bed) occupied the same reed bed during the day. This took place in $59 \pm 9\%$ of the nights





Fig. 5 Proportion (mean \pm SE) of overnight (roosting) positions occurring in each habitat for 26 Bluethroats and 103 nights at Barthes de la Nive. See abbreviations in Table 1. The number of individuals/ number of nights is also shown at the top. Letters group habitats for which no significant differences were detected with an a posteriori Wilcoxon test

(Fig. 6). This particularly concerned the 13 Bluethroats, which stopped over for just one day. Individuals staying for at least 2 days showed relatively predictable movements and selected repeatedly the same areas during the night. 27% of the birds (7) spent the night in a reed bed (Urdains or Ville-franque) and normally moved during the day to a corn crop. This was noticed for $12.5 \pm 5\%$ of the nights. We calculated the distance between roosting core areas and daytime (foraging) core areas. Birds flew on average 397 ± 33 m to reach its roosting location (n = 10, range 80–692 m). Five birds also used corn crops to stay during the night, although using it during the daytime (19 nights).



Fig. 6 Proportion (mean \pm SE) of different combinations of daytime (foraging) and overnight (roosting) habitats for 26 Bluethroats and 103 identified cases at Barthes de la Nive. Abbreviations as in Table 1, with the exception of RB that lumped PRB, MRB, and WOR together. The number of individuals/number of nights is shown at the top. Letters group habitats for which no significant differences were detected with an a posteriori Wilcoxon test

Diet analysis and habitat-specific invertebrate availability

We obtained 105 fecal samples: 84 fecal samples from firstyear birds and 21 from adults, 55 from males and 47 from females. A total of 431 prey items were identified; 4.1 ± 0.8 on average per dropping. We found 14 orders of invertebrates $(2.2 \pm 0.3 \text{ on average per sample})$. The diet of Bluethroats was dominated by ants (Hymenoptera Formicidae), representing 45.5% of prey items and occurring in 54% of the samples (Table 3). Spiders (Araneidae) were the second most abundant group (20.4%) with a similar occurrence to ants (53%); and beetles (Coleoptera) were the third most common group (10.2% of all prey items and 31% of the samples). These two last taxa were the main contributors to the consumed biomass, representing 64.7% and 14.9%, respectively. We did not detect any significant effects of Bluethroat's age and sex on prey abundance (Table 3). We did also observe that leafhoppers (Cicadellidae) were less consumed over time (Table 3).

Corn crops hosted significantly more invertebrates and biomass than reed beds for beetles, flies (Diptera), and spiders in pitfalls and for leafhoppers in bowls (Table 4). Corn crops also had higher availability of non-Formicidae hymenopterans than Urdains in bowls, but less than Villefranque. Finally, ants were more abundant in Urdains than in the other two sites.

Discussion

Variation in stopover duration and home-range size

Bluethroats showed large variation in their stopover duration and home-range size in Barthes de la Nive. The observed mean differed from that obtained by Arizaga et al. (2013). Thus, stopover duration was shorter and home range was larger in Barthes de la Nive (mean stopover duration of 5 days and mean home range size of 5.8 ha) than in Txingudi (9.6 days and 2 ha, respectively). Differences in stopover duration between these two sites were mediated by the fact that 45% of the radiotracked Bluethroats in Barthes de la Nive apparently departed the following night after capture, a circumstance that never took place in Txingudi (Fig. A3). If these short-staying individuals were excluded from the calculations, the stopover duration in Barthes de la Nive (i.e., 8.4 days) would conform better to the values obtained in Txingudi. After the signal of a bird disappeared from the place it was last detected, we carefully explored the whole study area by car the following day in its search and sporadically during subsequent days. This makes us very confident that the stopover duration estimates obtained in Barthes de la Nive are reliable. Likewise, the fact that only one of the 29 radiotracked Bluethroats stayed longer than the life expectancy of the transmitter would not alter these betweensite differences or the general conclusions of our study (see below). The number of Bluethroats whose stay reached the

Table 3 Percentage of each arthropod group in the fecal samples of 105 Bluethroats, occurrence frequency, proportion of total biomass and factors influencing the relative prey abundance (age: yearlings *vs* adults; sex and day)

Order	Family	Abundance%	Occurrence%	Biomass%	Age effec	ct	Sex effect		Day effect	
					z	р	z	р	z	р
Hymenoptera	Formicidae	45.48 (196)	54.29	8.34	0.998	0.318	- 1.218	0.223	0.868	0.386
Arachnida Araneae	Araneidae	20.42 (88)	53.33	14.92	- 0.986	0.324	0.373	0.709	- 1.326	0.185
Coleoptera		10.21 (44)	31.43	64.70	- 0.466	0.641	0.385	0.700	- 0.614	0.539
Diptera		7.42 (36)	28.57	2.39	- 0.010	0.992	0.462	0.644	0.281	0.779
Hymenoptera	Non Formicidae	4.41 (19)	15.24	4.73	- 0.193	0.847	0.803	0.422	-0.705	0.481
Hemiptera	Cicadellidae	3.25 (14)	13.33	2.45	- 1.388	0.165	0.549	0.583	- 1.993	0.046
Crustacea		3.32 (10)	9.52	2.48	0.008	0.993	- 0.872	0.383	- 0.772	0.440
Gastropoda		1.62 (7)	5.71	-						
Lepidoptera		1.62 (7)	6.67	-						
Neuroptera	Myrmeleontidae	1.16 (5)	3.81	-						
Arachnida Acarina		0.93 (4)	3.81	-						
Heteroptera		0.7 (3)	2.86	-						
Malacostraca Amphipoda		0.23 (1)	0.95	-						
Odonata		0.23 (1)	0.95	-						

The number of prey items is given between brackets. "-" Indicates no data available. We tested the effects of age (young vs adult birds), sex, and day using a Generalized Linear Model (GLM) with a Poisson error distribution. Pvalues were corrected for over dispersion. Only taxa representing more than 2% of the prey abundance were considered

Order	Family	Trap	Abundance			Biomass (mg)			
			Crop	Ru	Rv	Сгор	Ru	Rv	
Hymenoptera	Formicidae	Pitfall	0.2 ± 0.1	1.1±0.3*	0.2 ± 0.1	0.04 ± 0.02	0.6±0.2/Crop*	0.2 ± 0.1	
		Bowl	0.3 ± 0.1	-	0.6 ± 0.2	0.03 ± 0.02	0 ± 0	0.2 ± 0.1	
Arachnida	Araneidae	Pitfall	$3.5 \pm 1^{**}$	0.8 ± 0.3	1.3 ± 0.4	$7.5 \pm 2^*$	2.5 ± 1	4.8 ± 2.3	
		Bowl	1.9 ± 0.4	1.2 ± 0.5	2 ± 0.6	3.4 ± 0.8	5.9 ± 2.9	9.4±3.7**	
Coleoptera		Pitfall	$52.8 \pm 12.2^{**}$	0.1 ± 0.1	2.1 ± 0.5	$2006.9 \pm 449.1^{***}$	1.9 ± 1.8	12.4 ± 4.2	
		Bowl	$5.5 \pm 1.4^{***}$	0.7 ± 0.3	$3.8 \pm 1.1^{***}$	$49.2 \pm 40.7 ^{***}$	0.9 ± 0.6	8.2 ± 3.7	
Diptera		Pitfall	$10.3 \pm 5.1^{***}$	0.3 ± 0.2	1.1 ± 0.3	$17.9 \pm 7.1^{***}$	0.8 ± 0.8	1.1 ± 0.5	
		Bowl	$24.5\pm6^{***}$	1.4 ± 0.5	12.3 ± 2.5	$42.2 \pm 8.8^{***}$	1.8 ± 0.7	24.1 ± 5	
Hymenoptera	Non-Formicidae	Pitfall	0.9 ± 0.3	0.2 ± 0.1	1.4 ± 0.5/Ru*	0.7 ± 0.3	0.1 ± 0.1	2.7 ± 1.3	
		Bowl	4.5 ± 1	0.6 ± 0.4	$8.9 \pm 1.6^{***}$	14.4 ± 4.5	1.2 ± 1.1	43.6 ± 8.2	
Hemiptera	Cicadellidae	Pitfall	0.3 ± 0.1	-	0.1 ± 0.1	0.2 ± 0.2	0 ± 0	0 ± 0	
		Bowl	$2.7\pm0.7^{**}$	0.1 ± 0.1	0.7 ± 0.3	4 ± 1.9/Ru***	0 ± 0	0.8 ± 0.4	

Table 4 Abundance and biomass (mean \pm SE) of the most consumed invertebrates by Bluethroats in corn crops (Crop), Villefranque reed bed (Rv) and Urdains reed bed (Ru), which were trapped in pitfall and colored bowls

We compared abundance and biomass between sites using General Linear Models with a Poisson error distribution. Numbers in bold indicate the site for which the abundance or biomass was significantly (***p < 0.001, **p < 0.01, *p < 0.05) higher than one of the other sites (site shown after a slash symbol) or the other two sites. Two hyphens represent no data available

expected functional life of the transmitters in Txingudi was slightly higher (3 out of 20) than in Barthes de la Nive, supporting the longer stopover duration of Bluethroats in the former site.

It could be argued that the higher number of short-staying Bluethroats in Barthes de la Nive was caused by the use of playbacks during the trapping sessions, which were not implemented in Txingudi. Although we cannot rule out completely this confounding factor as an alternative explanation to observed results, current evidence does not support this idea. Thus, the use of playbacks typically attracts Bluethroats with lower body reserves (Arizaga et al. 2015) and, as our results show, birds with less fuel loads would tend to have longer stopover durations (Alerstam and Lindström 1990; Salewski and Schaub 2007). This impact of playback on fuel loads of autumn-migrating Bluethroats was consistent in the three wetlands on the Bay of Biscay (Gironde, Txingudi, Urdaibai) analyzed by Arizaga et al. (2015) that are geographically very close to Barthes de la Nive. If Bluethroats do show transient vs. non-transient divergent behaviors during migration as have been described for other migratory species (Rappole and Warner 1976), playbacks might have also biased stopover duration estimates if they do preferentially attract transient individuals. However, a study of stopover duration using tape lures did not detect these transient Bluethroats during autumn migration (Arizaga et al. 2010). Likewise, neither the results obtained by Arizaga et al. (2015) would support this possibility, since playbacks do not seem to affect the number of Bluethroat recaptures and this would be an expected outcome if playbacks promoted the capture of more transient individuals that would have less recapture prospects. Likewise, and contrary to what would be expected for their transient condition, these short-staying individuals did not move as much as would be expected after being trapped in Barthes de la Nive, and their first-day home range did not differ significantly from that observed for long-staying conspecifics.

Observed differences in stopover duration between Barthes de la Nive and Txingudi might explain the relatively low number of recaptures obtained in Barthes de la Nive, when compared to Txingudi and also other nearby sites for which there is no radiotracking information so far (i.e., Gironde and Urdaibai; Arizaga et al. 2015). Additionally, these differences in recapture rates could be also promoted by the existence of overall home ranges that are nearly twice to thrice larger in Barthes de la Nive than in Txingudi (Arizaga et al. 2013) and in Gironde (Musseau et al. 2017). These contrasting differences between Txingudi and Barthes de la Nive are potentially mediated by the strong differences in the degree of urbanization and habitat composition that seem to have profound consequences in the stopover behavior of Bluethroats between two relatively close locations.

Habitat and roosting site selection

Migrating Bluethroats in Barthes de la Nive strongly selected reed beds (pure and mixed) and other hydrophilic natural grasslands, which concurs with the favourite habitat that Arizaga et al. (2013) and Musseau et al. (2017) found for migrating and moulting conspecifics, respectively. Much less known was the preference of migrating Bluethroats for corn crops, although one study had detected this species in this crop type during migration (Gottschalk and Cover 2016) and some populations of Bluethroats seem to thrive and even breed successfully in some agricultural fields such as oilseed rape crops (Cornulier et al. 1997; Berndt and Hölzel 2012, Chiron 2017). We did not find the preference of Bluethroats for tidal mudflats or low-halophytic vegetation described in other studies (Arizaga et al. 2013; Godet et al. 2015; Musseau et al. 2017), but the representation of these habitats in Barthes de la Nive is relatively limited, which could explain this lack of use.

Our results showed that home-range size is smaller when birds occupied reed beds (for overall home ranges) and corn crops (for daily home ranges). A potential explanation for this result is that these habitats might offer more trophic resources, which would prevent birds from making long foraging trips (Bibby and Green 1980; Chernetsov et al. 2004). Another complementary hypothesis for the higher occurrence of Bluethroats in these two habitat is that, given their dense structure, they could also provide a suitable shelter, not only for roosting during the night, but also to rest during daytime between migration flight bouts. The use of reed beds as roosting habitat has been described before in Bluethroats (Eybert et al. 2004; Harmange et al. 2016) and our study confirmed this preference and suggested that corn crops might occasionally play a similar role. In contrast, long-staying Bluethroats seem to expand their home range probably, because they are in high demand of food for refueling and, perhaps, because of the competition with conspecifics and other species (Chernetsov and Titov 2001; Fransson et al. 2008).

If home-range size is considered a proxy of habitat quality, inland reed beds could be interpreted as a refuge or a more optimal habitat than corn crops, which might represent the most commonly available substitutionary habitat (Godet et al. 2018), where birds needing to refuel would move due to competition and the limited availability of reed beds.

In our site, Bluethroats encounter also some potentially unsuitable habitats such as woodlands, open water, mown grasslands, and infrastructures (house, road, and railway) that would explain why they had to fly away a relatively long distance from their preferred roosts (Harmange et al. 2016), expanding their home-range size. Similar patterns have been shown in Briere for breeding Bluethroats (Godet et al. 2015).

We did not detect any difference between males and females in home-range size, flight distance, or stopover duration. Typically, small-sized, young and female Bluethroats are subordinated to large, adult and male conspecifics, respectively (Lindström et al. 1990; Moore et al. 2003). Our results suggest that Barthes de la Nive was occupied, not under an ideal despotic distribution, but under an ideal free distribution, where home-range size would be determined by its habitat composition.

Diet of birds and invertebrate availability

The diet description for our study site fitted well with the trophic characterization made in other Bluethroat studies (Allano et al. 1988; Orłowski et al. 2014; Musseau et al. 2017). In decreasing order of importance, our study showed the relevance of ants (40.6% of prey; 8.7% of biomass), flies (resp. 14% and 6.5%), beetles (13.7%, 40%) and spiders (9.4%, 22.5%). Bluethroats typically forage on the ground (Orlowski et al. 2014; Allano et al. 1988; Cramp 1988), where ants are particularly abundant. As in other agricultural farms (Cornulier et al. 1997), corn crops showed a relatively high insect abundance and biomass, which could explain why Bluethroats positively selected this habitat. Again and as a consequence of the relative scarcity of intertidal habitats, arthropods such as amphipods (Malacostraca Amphipoda) were absent from the diet of Bluethroat in Barthes de la Nive, although they have been found in other studies (Musseau et al. 2017; Allano et al. 1988).

Habitat management implications

Our study confirmed the importance of wetlands and their associated reed beds in Barthes de la Nive for migrating Bluethroats, as it was also shown for other bird species stopping over in the region (Fontanilles et al. 2014). This reaffirms the need to conserve and potentially expand these valuable habitats, whose largest remnants in southwestern France persist in the Nive/Adour river basin. These wetlands are mainly threatened by clogging and bush encroachment, particularly by invasive exotic tree species (i.e. *Acer negundo*, *Baccharis halimifolia*), so regular intervention is needed to maintain their characteristic aquatic vegetation.

Likewise, our results unexpectedly revealed a positive selection of corn crops by Bluethroats. The tall and dense arrangements of corn plants resemble the habitat structure of reed beds. However, corn crops do not seem to constitute an ecological trap, but a substitutionary habitat (Godet et al. 2018), a human-induced opportunity for migrating birds. Anyway, further research would be required to further understand this selection process and whether it also takes place in other sites. Our preliminary results suggest that corn crops would provide plenty of food resources and a suitable shelter for birds stopping over in Barthes de la Nive. Our study using the Bluethroat as model species raises the question of how other bird species respond to the presence of corn crops during their stopovers. The corn crops of Barthes de la Nive are used by a large group of migratory species (Fontanilles not published) so that measures to maintain or increase the suitability of this common crop for birds around the limited wetland remnants should be encouraged. Among them, promoting organic farming and postponing the harvesting time after mid-October, when most insectivorous migrants are in or near their southern wintering quarters, would be relatively easy to implement and might have immediate payoff (Dänhardt et al. 2010).

Acknowledgements This work was supported by the Conseil Départemental des Pyrénées Atlantiques, Syndicat Nive Maritime, Villefranque and Bayonne councils. We are particularly grateful to F. Esnault, S. Valles, R. Damestoy and R. Dufourcq. We are extremely thankful to L. Delalande, F. Delfour, A. Glad, M. Janczyszyn-Le Goff, T. Sarraude, for their commitment in the collection of the radiotracking data, to L. Julien for invertebrate sampling, and members of OISO association who made this research possible, especially F. Lartigau. National Museum of Natural History authorized the bird-ringing and radiotracking scheme. The experiments complied with the current laws of the country in which they were performed. We also thank two anonymous reviewers who gave insightful comments that improved an early version of the manuscript.

Author's contribution All of the authors contributed to data collection. KS prepared the habitat map, FL and CK gathered data on bird diet and arthropod availability. PF, IH, and CK analyzed the data, designed the methods, and conducted the research. All of the authors took part in the formulation of the questions and hypotheses. PF wrote the paper in cooperation with IH and CK.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflicts of interest.

Ethical statement All data and information from external sources used for the analyses and preparation of the manuscript are properly cited. The conclusions are entirely based on our own results. All applicable international, national, and/or institutional guidelines for the care and use of animals were followed.

References

- Alerstam T, Lindström C (1990) Optimal bird migration: the relative importance of time, energy and safety. In: Gwiner E (ed) Bird migration: the physiology and ecophysiology. Springer, Heidelberg, pp 331–351
- Allano L, Bonnet P, Constant P, Eybert M (1988) Premières données sur le régime alimentaire des jeunes gorgebleues (Luscinia svecica namnetum, Mayaud) au nid dans un marais salant exploité (Guérande, Loire-Atlantique). C R Acad Sci Paris 306:369–374
- Arizaga J, Alonso D, Campos F, Unamuno JM, Monteagudo A, Fernandez G, Carregal XM, Barba E (2006) Do subspecies of bluethroat *Luscinia svecica* show a geographic segregation during the autumn migration period in Spain? Ardeola 53:285–291
- Arizaga J, Barba E, Alonso D, Vilches A (2010) Stopover of Bluethroats (*Luscinia svecica cyanecula*) in northern Iberia during the autumn migration period. Ardeola 57:69–85
- Arizaga J, Mendiburu A, Alonso D, Cuadrado JF, Jauregi JI, Sánchez JM (2011) A comparison of stopover behavior of two subspecies of the Bluethroat (*Luscinia svecica*) in Northern Iberia during the autumn migration period. Ardeola 58:251–265
- Arizaga J, Andueza M, Tamayo I (2013) Spatial behaviour and habitat use of first-year Bluethroats *Luscinia svecica* stopping over at coastal marshes during the autumn migration period. Acta Ornithol 48:17–25

- Arizaga J, Musseau R, Laso M, Esparza X, Unamuno E, Azkona A, Fontanilles P (2015) Biases associated with the use of a playback in stopover ecology studies in small passerine birds. Bird Study 62:280–284
- Barton K (2019) MuMIn package: Multi-Model Inference. https:// cran.r-project.org/web/packages/MuMIn/MuMIn.pdf
- Berndt AM, Hölzel N (2012) Energy crops as a new bird habitat utilization of oilseed rape fields by the rare Bluethroat (*Luscinia svecica*). Biodivers Conserv 21:527–541
- Bibby CJ, Green RE (1980) Foraging behaviour of migrant pied flycatchers, *Ficedula hypoleuca*, on temporary territories. J Anim Ecol 49:507–521
- Börger L, Francon N, De Michele G, Gantz A, Meschi F, Manica A, Coulson Lovari S (2006) Effects of sampling regime on the mean and variance of home range size estimates. J Anim Ecol 75:1393–1405
- Burnham KP, Anderson DR, Huyvaert KP (2011) AIC model selection and multimodel inference in behavioral ecology: some background, observations, and comparisons. Behav Ecol Sociobiol 65:23–35
- Busse P (2000) Bird station manual. SE European Bird Migration Network, Gdansk
- Chernetsov N (2012) Passerines migration: stopovers and flight. Springer, Berlin
- Chernetsov N, Titov NV (2001) Movement patterns of European reed warblers Acrocephalus scirpaceus and sedge warblers A. schoenobaenus before and during autumn migration. Ardea 89:509–515
- Chernetsov N, Mukhin A, Ktitorov P (2004) Contrasting spatial behaviour of two long-distance passerine migrants at spring stopovers. Avian Ecol Behav 12:53–61
- Chiron D (2017) Range extension of the Bluethroat *Luscinia svecica namnetum* towards intensive farming habitats in Central western France. Alauda 85:81–91
- Clark DL, Wilson MV (2001) Fire, mowing, and hand-removal of woody species in restoring a native wetland prairie in the Willamette valley of Oregon. Wetlands 21:135–144
- Cornulier Th, Bernard R, De Arroyo B, Bretagnolle V (1997) Extension géographique et écologique de la Gorgebleue à miroir *Luscinia svecica* dans le Centre-Ouest de la France. Alauda 65:1–6
- Cramp S (1988) The birds of the Western Palearctic. Oxford University Press, Oxford
- Czech HA, Parsons CK (2002) Agricultural Wetlands and Waterbirds: a Review. Waterbirds 25:56–65
- Dänhardt J, Green M, Lindström Å, Rundlöf M, Smith HG (2010) Farmland as stopover habitat for migrating birds effects of organic farming and landscape structure. Oikos 119:1114–1125
- Davies NB (1977) Prey selection and the search strategy of the Spotted Flycatcher *Muscicapa striata*, a field study on optimal foraging. Animal Behav 25:1016–1033
- De la Hera I, Fontanilles P, Delalande L, Glad A, Sarraude T (2017) Attraction of other species by bluethroat *Lusscinia svecica* song playback during autumn migration: an experimental test using bird-ringing data. Ardeola 64:91–99
- Delingat J, Dierschke V (2000) Habitat utilization by Northern Wheatears (*Oenanthe oenanthe*) stopping over on an offshore island during migration. Vogelwarte 40:271–278
- Eybert MC, Bonnet P, Geslin T, Questiau S (2004). La Gorgebleue. Belin, p 71
- Fontanilles P (2014) Migration et régime alimentaire de la Gorgebleue à miroir dans les Barthes de l'Adour, Actes du séminaire "Conservation des espèces et gestion des zones humides", Bayonne
- Fontanilles P, Chanchus B, Chauby X, De Franceschi C, Dufresne L, Fourcade JM, Lapios JM, Lartigau F, Legay P, Thouy P, Tillo S, Urbina-Tobias P, Short R, Williams H (2012) Bilan du baguage des passereaux migrateurs et nicheurs sur les barthes de la Nive (Pyrénées-Atlantiques) en 2011. Le Casseur d'os 12:80–84

- Fontanilles P, Laval B, Diribarne M (2014) Sélection des habitats et occupation spatiale du Phragmite aquatique *Acrocephalus paludicola* sur une halte migratoire du sud-ouest de la France, mise en place d'une gestion intégrée. Alauda 82:343–351
- Fransson T, Barboutis C, Mellroth R, Akriotis T (2008) When and where to refuel before crossing the Sahara desert extended stopover and migratory fuelling in first-year garden warblers *Sylvia borin*. J Avian Biol 39:133–138
- Freeman S, Jackson WM (1990) Univariate metrics are not adequate to measure avian body size. Auk 107:69–74
- Godet L, Marquet M, Eybert MC, Gregoire E, Monnet S, Fournier J (2015) Bluethroats *Luscinia svecica namnetum* offset landscape constraints by expanding their home range. J Ornithol 156:591–600
- Godet L, Harmange C, Marquet M, Joyeux E, Fournier J (2018) Differences in home-range sizes of a bird species in its original, refuge and substitution habitats: challenges to conservation in anthropogenic habitats. Biodivers Conserv 27:719–732
- Gottschalk TK, Cover L (2016) Migratory and visiting birds in a maize field near Giessen, Germany in summer and autumn. Vogelwarte 54:1–14
- Grueber CE, Nakagawa S, Laws RJ, Jamieson IG (2011) Multimodel inference in ecology and evolution: challenges and solutions. J Evol Biol 24:699–711
- Harmange C, Godet L, Marquet M, Dietrich J, Monnet S, Grégoire E, Eybert MC, Fournier J (2016) [Unexpected movements of male Bluethroats Luscinia svecica namnetum during the breeding period] Déplacements inattendus chez des mâles de Gorgebleue à miroir blanc Luscinia svecica namnetum en période de nidification. Alauda 84:195–202
- Hódar JA (1996) The use of regression equation for estimations of arthropod biomass in ecological studies. Acta Oecol 17:421-433
- Huntley B, Green RE, Collingham YC, Willis SG (2007) A climatic atlas of European breeding birds. Lynx Edicions. Barcelona
- Hutto RL (2000) On the importance of en route periods to the conservation of migratory landbirds. Stud Avian Biol 20:109–114
- Jacobs J (1974) Quantitative measurement of food selection: a modification of the forage ratio and Ivlevs' selectivity index. Oecologia 14:413–417
- Lindström C, Hasselquist D, Bensch S, Grahn M (1990) Asymmetric contests over resources for survival and migration. A field experiment with Bluethroats. Animal Behav 40:453–461
- Moore FR, Mabey S, Woodrey M (2003) Priority access to food in migratory birds: age, sex and motivational asymmetries. In: Berthold P, Gwinner E, Sonnenschein E (eds) Avian migration. Springer, Heidelberg, pp 281–291
- Musseau R, Beslic S, Kerbiriou C (2017) Importance of intertidal wetlands for the French coastal endemic Bluethroat (*Cyanecula*

svecica namnetum) and conservation implications in the context of global changes. Ardeola 64:325–345

- Newton I (2004) Population limitation in migrants. Ibis 146:197–226 Newton I (2008) The migration ecology of birds. Academic Press, London
- Orłowski G, Rusiecki S, Karg J (2014) Partial dietary segregation between adult and nestling Bluethroats Luscinia svecica. Acta Ornithol. 49:107–118
- Piper WH (2011) Making habitat selection more "familiar": a review. Behav Ecol Sociobiol 65:1329–1351
- Rappole JH, Warner DW (1976) Relationships between behavior, physiology and weather in avian transients at a migration stopover site. Oecologia 26:193–212
- Salewski V, Schaub M (2007) Stopover duration of Palearctic passerine migrants in the Western Sahara; independent of fat stores? Ibis 149:223–236
- Salewski V, Kéry M, Herremans M, Liechti F, Jenni L (2009) Estimating fat and protein fuel from fat and muscle scores in passerines. Ibis 151:640–653
- Schaub M, Schwilch R, Jenni L (1999) Does tape-luring of migrating Eurasian Reed-Warblers increase number of recruits or capture probability? Auk 116:1047–1053
- Tellería JL, de la Hera I, Pérez-Tris J (2013) Morphological variation as a tool for monitoring bird populations: a review. Ardeola 60:191–224
- von Behren CR (2007) Impact of agriculture and urban development on the community structure of wetland birds in East Central Minnesota. Honors Projects, Paper, p 5
- Whited D, Galatowitsch SM, Tester JR, Schik K, Lehtinen R, Husvethc J (2000) The importance of local and regional factors in predicting effective conservation Planning strategies for wetland bird communities in agricultural and urban landscapes. Landscape Urban Plan 49:49–65
- Worton BJ (1989) Kernel methods for estimating the utilisation distribution in home range studies. Ecology 70:164–168
- Yong W, Finch DM, Moore FR, Kelly JF (1998) Stopover ecology and habitat use of migratory Wilson's Warblers. Auk 115:829–842
- Zanimetti JM (2005). Statistique spatiale. Ed Lavoisier
- Zuur AF, Ieno EN, Walker N, Saveliev AA, Smith GM (2009) Mixed effects models and extensions in ecology with R Statistics for Biology and Health. Springer, New York

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