Ecological and life-history drivers of avian skull evolution

Eloise S.E. Hunt^{1,2,1}, Ryan N. Felice^{2,3}, Joseph A. Tobias^{4,10}, Anjali Goswami^{2,5,10}

¹Department of Life Sciences and Grantham Institute, Imperial College London, London, United Kingdom

²Department of Life Sciences, The Natural History Museum, London, United Kingdom

³Centre for Integrative Anatomy, Department of Cell and Developmental Biology, University College London, London, United Kingdom ⁴Department of Life Sciences, Imperial College London, Ascot, United Kingdom

Department of Life Sciences, imperial Coneye London, Ascol, Onited Kingdon

⁵Department of Genetics, Evolution, and Environment, University College London, London, United Kingdom

Corresponding author: Department of Life Sciences, Natural History Museum, Cromwell Road, London SW7 5BD, United Kingdom. Email: e.hunt@nhm.ac.uk

Abstract

One of the most famous examples of adaptive radiation is that of the Galápagos finches, where skull morphology, particularly the beak, varies with feeding ecology. Yet increasingly studies are questioning the strength of this correlation between feeding ecology and morphology in relation to the entire neornithine radiation, suggesting that other factors also significantly affect skull evolution. Here, we broaden this debate to assess the influence of a range of ecological and life-history factors, specifically habitat density, migration, and developmental mode, in shaping avian skull evolution. Using 3D geometric morphometric data to robustly quantify skull shape for 354 extant species spanning avian diversity, we fitted flexible phylogenetic regressions and estimated evolutionary rates for each of these factors across the full data set. The results support a highly significant relationship between skull shape and both habitat density and migration, and developmental mode. We further found heterogenous rates of evolution between different character states within habitat density, migration, and developmental mode, with rapid skull evolution in species that occupy dense habitats, are migratory, or are precocial. These patterns demonstrate that diverse factors affect the tempo and mode of avian phenotypic evolution and that skull evolution in birds is not simply a reflection of feeding ecology.

Teaser Text

Almost 200 years ago, Darwin found that Galápagos finches' beaks were different shapes in birds with different diets. Today, it is well established that phylogeny, allometry, and ecology can also be key factors in shaping skulls. Yet, the influence of specific aspects of ecology, as well as life history, on morphological evolution remains poorly constrained. Here, we examined whether three novel factors also influence the shape of bird skulls and rates of evolution: habitat density, migration, or developmental mode. To do so, we combine high-resolution 3D quantification of skull shape with dense taxonomic sampling across living birds. Our analyses revealed that skull shape varies in birds based on vegetation density in their habitats and the extent to which they migrate, but not their developmental mode. Despite these differences, habitat density, migration, and life history all influence the rate at which bird skulls evolve. Birds evolved fastest if they live in densely vegetated habitats, migrate long distances, or are precocial. This adds to the growing body of evidence that avian skull evolution is affected by a diverse range of factors and suggests that habitat density, migration, and life history should be considered in future analyses on drivers of phenotypic evolution.

Keywords: macroevolution, morphological evolution, life-history evolution

Background

The Galápagos finches are a classic "textbook" example of avian adaptive radiations where beak morphology is considered an adaptation to diet (Grant & Grant, 1989). In the last 5 years, there have been significant efforts to robustly quantify this interaction of cranial and beak shape and various ecological and developmental factors, particularly feeding ecology (Bright et al., 2016; Coonev et al., 2017; Felice & Goswami, 2018; Felice et al., 2019; Navalón et al., 2019; Pigot et al., 2020, Natale & Slater, 2022), which have demonstrated that this relationship is highly complex and differs across scales and across lineages. Diet has been found to strongly correlate with beak shape in waterfowl (Anseriformes; Olsen, 2017), and corvids (Corvidae; Kulemeyer et al., 2009), as well as brain shape in kingfishers (Alcedinidae; Eliason et al., 2021) and skull shape in shorebirds and relatives (Charadriiformes; Natale & Slater, 2022). Conversely, beak and braincase morphology is largely controlled by size in raptors (Bright et al., 2016), and diet only predicts 2.4% of skull shape variation in parrots and cockatoos (Psittaciformes; Bright et al., 2019). Large-scale studies across Neornithes have also yielded variable results: Diet can be predicted from linear measurements (Pigot et al., 2020), but there is only a weak correlation between diet and cranial morphology (Felice et al., 2019) or beak morphology (Navalón et al., 2019) when using geometric morphometrics. Recently, Crouch and Tobias (2022) found no association between bursts of morphological evolution and rates of dietary evolution at a global scale.

It is well established that diverse aspects of ecology can be key factors in determining both skull morphology (Bardua et al., 2021; Dumont et al., 2016; da Silva et al., 2018; Vidal-García & Scott Keogh, 2017) and rates of shape evolution (Collar et al., 2010; Millien, 2006). Phenotypic convergence occurs when different lineages adapt to similar habitats

which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.

Received February 3, 2023; revisions received April 4, 2023; accepted April 26, 2023

[©] The Author(s) 2023. Published by Oxford University Press on behalf of The Society for the Study of Evolution (SSE).

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/),

(McGhee, 2011). A range of aspects of ecology has been associated with bursts in morphological evolution, such as transitions to a new ecological niche (Price et al., 2011; Sherratt et al., 2017), ecological opportunity (Losos, 2010), habitat stability (Crouch & Tobias, 2022), and competition (Rosenzweig, 1978). Given that diet, as currently measured, is an incomplete predictor of skull shape variation and evolutionary tempo across birds, alternative aspects of life history or ecology warrant investigation. Chira et al. (2018) found low support for an association between rates of beak evolution and generation length, temperature, UVB levels, range size, proportion living on islands, or competition, but 80% of variation in species-level evolutionary rates remained unexplained. Across Neornithes, there are correlations between ecological traits and morphology, for instance, down feather morphology is adapted to habitats (Pap et al., 2020), and there is widespread convergence linking cranial and postcranial linear measurements to trophic niches (Pigot et al., 2020). Within passerines, there is evidence of correlations between body form and foraging mode (Fitzpatrick, 1985); correlations between the lengths of the tarsus and midtoe and substrate utilization (Miles & Ricklefs, 1984), as well as a correspondence between tangers bill morphology and the filling of ecomorphospace (Vinciguerra & Burns, 2021). So, there is evidently a robust correlation between ecology and avian morphology, but it is not clear which components of ecology are shaping avian skull evolution.

Additionally, phylogeny (Brusaferro & Insom, 2009; Degrange & Picasso, 2010), ontogeny (Navalón et al., 2021), allometry (Bright et al., 2016; Tokita et al., 2017; Yamasaki et al., 2018), phenotypic integration (Felice & Goswami, 2018; Navalón et al., 2020; Shatkovska & Ghazali, 2020), and encephalization (Marugán-Lobón et al., 2021) are all intrinsic factors, which have been found to significantly influence skull morphology within various avian lineages, but most have not been assessed across the breadth of avian diversity. Collectively, this research calls into question the primacy of the relationship between diet and avian skull shape.

Here, we interrogate the relationship between cranial morphology and three key ecological/life-history traits: habitat density, migration behavior, and developmental mode. We chose to investigate habitat density as one of our ecological traits due to evidence that habitat openness influences kingfisher brain shape evolution, with forest dwellers undergoing more rapid rates of brain shape evolution (Eliason et al., 2021). This study did not find any single brain shape associated with forest living and instead suggested that brain shape in the forest dwellers was diverging stochastically, possibly in response to genetic drift in fragmented habitats. Given that the skull roof tracks the brain in birds (Fabbri et al., 2017), factors that drive shifts in brain shape may also result in changes in skull shape. However, the impact of the density of habitats on the tempo and mode of avian phenotypic evolution on a broad macroevolutionary scale has not been investigated until now.

Migration is widespread in seasonal environments, with approximately 40% of all birds migrating (El-Sayed, 2019), and it has well-established adaptive value (Hedenström, 2008; Lack, 1968). It has been proposed that the genes for migratory behavior are ancestral in all birds (Pulido, 2007) and that seasonal migration is heritable and can rapidly change in response to selection (Berthold et al., 1992). Thus, transitions between migratory and non-migratory behavior do not

require repeated innovation, but merely selection driving a pre-existing genetic program (Alerstam et al., 2003; Salewski & Bruderer, 2007; Winger et al., 2012; Zink, 2002), which may explain the dynamic fluctuations in migration across extant birds (Piersma et al., 2005; Winger et al., 2012; Zink, 2002). Despite the rate at which avian migration can evolve, the degree to which this affects evolutionary rates has not been assessed. Migratory birds have evolved a suite of adaptations to minimize weight, such as organs reducing size before migration (Battley et al., 2000) and hearts being relatively smaller in migrants (Vágási et al., 2016). Additionally, a negative correlation has been identified between migration distance and brain size (Sol et al., 2010; Vincze, 2016). As there are strong correlations between the shapes and sizes of brains and endocasts in birds (Watanabe et al., 2019), and differences in endocranial anatomy are correlated with craniofacial differences in birds (Iwaniuk & Nelson, 2002; Marugán-Lobón & Buscalioni, 2009; Marugán-Lobón et al., 2021), it is possible that migratory birds have also evolved weight-saving adaptations to cranial anatomy.

Finally, we integrate a fundamental aspect of life history that varies widely across birds: the altricial-precocial spectrum. Precocial developmental mode, where juveniles are relatively mature at birth or hatching, is more common than altricial development among vertebrates. This strategy was proposed to be an adaptation to high rates of predation on juveniles (Arnold & Wassersug, 1978; Wassersug & Sperry, 1977). By contrast, altricial developmental mode is associated with more extensive parental care, which promotes rapid growth rates that can average four times that of similarly sized precocial species (Case, 1978; Ricklefs, 1979), as well as poor locomotor performance, and short developmental periods. This variation in life history creates different selective pressures acting on juveniles that fall into different character states along the altricial-precocial spectrum, so it has been suggested that selection on the juvenile morphology could act more strongly than selection on adult morphology for precocial species (Carrier, 1996; Dial & Carrier, 2012).

Furthermore, there is a correlation between degree of precociality and smaller relative brain sizes across birds (Griesser et al., 2023; Hardie & Cooney, 2022), providing evidence for the altricial–precocial spectrum driving morphological differences. However, the influence of developmental mode on avian cranial shape evolution has yet to be investigated across crown birds.

We used 3D geometric morphometric data from 354 species across Neornithes and a phylogenetic comparative framework to address two key questions about the relationship between avian skull shape and ecological and life-history traits. First, we assessed whether avian skull shape covaries with size, habitat density, migration, and developmental mode. Second, we tested whether evolutionary rates differ between different character states within habitat density, migration, and developmental mode.

Methods

Morphological data

Our analyses use a previously published three-dimensional geometric morphometric data set of 354 adult species, representing 159 families of extant birds (Supplementary Table S1, Felice & Goswami, 2018). One skull was used per species, and species were selected based on the availability of complete or almost complete adult skulls. These were subjected to the previously published procedure of landmarking using IDAV Landmark (Felice & Goswami, 2018; Wiley et al., 2005) to place anatomical landmarks and curve semi-landmarks on digital three-dimensional skull models formed from CT and surface scans. We then used the R package "Morpho" v2.5.1 (Schlager, 2017) to project surface semi-landmarks onto each specimen from a template. A total of 757 landmarks were used to quantify three-dimensional cranial morphology, divided into the rostrum, cranial vault, sphenoid region, palate, pterygoid/quadrate, naris, and occipital, as in Felice and Goswami (2018) (Figure 1). The effects of size, position, and rotation were removed with a generalized Procrustes analysis using the R package "geomorph" v3.0.6 (Adams & Otárola-Castillo, 2013). We extracted log centroid size of the cranium during the Procrustes superimposition and used this as a proxy for size in further analyses. Following the finding by Natale and Slater (2022) that some shorebirds followed different scaling patterns thus body mass was a more appropriate size measure for the skull, we assessed the correlation between log body mass and log centroid size of the cranium and found that they are

Α

highly correlated for our sample ($r^2 = 0.885$, Supplementary Figure S1).

Phylogenetic hypothesis

A previously published composite phylogenetic tree was utilized for the phylogenetic comparative analyses (Felice et al., 2019). This composite topology was generated by following the procedure described by Cooney et al. (2017). This tree incorporates the backbone of relationships among major clades from (Prum et al., 2015), with the fine-scale species relationships from a maximum clade credibility tree generated by Jetz et al. (2012). The tree of 9,993 species was then pruned down to the 354 species in our data set.

Ecological and life-history trait data

Habitat density, migration, and developmental mode of birds were all classified using three character states (Figure 1). Habitat density was categorized as "dense" (n = 120), "semi-open" (n = 91), or "open" (n = 143) following Tobias et al. (2016), sourced from Tobias et al. (2022). Dense habitats are those where species primarily occupy dense thickets, shrubland, or forest (except species perching habitually



Figure 1. (A) The ecological and life-history trait states of every species in our sample mapped onto the phylogenetic tree used in analyses. (B) The landmarking scheme used in our analyses, presented in lateral view. The landmarks are colored as follows: golden, rostrum; pale blue, cranial vault; green, sphenoid region; yellow, palate; navy, pterygoid/quadrate; orange, naris; and pink, occipital (Felice & Goswami, 2018).

on top of the canopy). Semi-open habitats include primarily living in open shrubland scattered bushes or deciduous forest. Open habitats are where species primarily live in desert, grassland, open water, seashores, cities, or habitually perching on top of forest canopy. Migration behavior was classed as "non-migratory" (n = 218), "partially migratory" (n = 63), or "migratory" (n = 73) following Tobias and Pigot (2019; Tobias et al. 2022). Whereas the migratory class is comprised of species where most of the population embark on long-distance (typically intercontinental) migrations, partially migratory species travel much shorter distances on average. This includes short-distance elevational movements, nomadism, and cases in which a minority of the population migrates relatively short (intracontinental) distances. Finally, non-migratory birds do not migrate and tend to be either sedentary or undergo only local movements. Developmental mode was categorized as "precocial" (n = 60), "semi-precocial" (n = 80), and "altricial" (n = 214; Cooney et al., 2020; Hoyo et al., 1992; Starck, 1993). Where data were not available in an existing database (Cooney et al., 2020), we classified species using Hoyo et al. (1992) and Botelho et al. (2015). Where information was not available at species level, the developmental mode was inferred by information on other species within the genus or family, as previous studies have suggested that there is little intrafamily variation in position on the altricial-precocial spectrum (Ducatez & Field, 2021).

Data analyses

We ran preliminary phylogenetic ANOVAs using the "procD. pgls" function in the geomorph R package (Adams et al., 2022) to assess whether there are any interactions between our three traits (habitat density, migration, and life history) and the previously examined or potentially related traits of diet, habitat, and primary lifestyle, sourced from Tobias et al. (2022) using "trophic niche" as a measure for diet. We found no significant interactions between diet, habitat, or primary lifestyle and our factors at the p < .01 level except a marginally significant interaction between diet and migration (Supplementary Table S2). For this reason, we did not incorporate diet into our further analyses given the heavy focus on this factor in other analyses using npMANOVAs to assess the strength of covariation between diet and shape, as well as estimating rates of evolution for the same morphological data as we use in this study (Felice et al., 2019). We then used type II phylogenetic MANOVAs (phylogenetic regressions) to assess the significance of habitat density, migration, and developmental mode for avian skull shape. We fit these models using the full geometric morphometric data set, with log centroid size, habitat density, migration, and developmental mode as predictors for the "mvgls" and "manova. gls" functions in the R package mvMORPH 1.1.4 (Clavel et al., 2015). We used the "mvgls" function to fit multivariate phylogenetic linear models with Pagel's lambda by penalized likelihood (Clavel et al., 2015). We employed the "manova. gls" function to assess the significance of the four predictors via type II MANOVA tests with Pillai's statistic over 1,000 permutations (Clavel et al., 2019). Principle component analvsis was used to visualize the main axes of variation for the whole skull. Morphospaces were plotted in ggplot2 v.3.3.6 (Wickham, 2016), with convex hulls plotted for the different character states of our three traits. The primary axes of shape variation are shown by extreme shapes along the first two principal component (PC) axes.

We further estimated the evolutionary rates for each habitat density, migration, and developmental mode character state following the protocol in Bardua et al. (2021). First, we utilized the "ace" function in ape v5.3 (Paradis & Schliep, 2019) to calculate the ancestral states for habitat density, migration, and developmental mode. We used the "make. simmap" function in the "phytools" package v.1.2-0 (Revell, 2012) to reconstruct the evolutionary history of these factors by stochastic character mapping, which we then used to fit flexible BMM (Brownian motion with multiple rates matrix) models. We conducted model fitting using the "mvgls" function in mvMORPH with the "error = TRUE" setting. We additionally ran our evolutionary rates analyses using this protocol for each the seven anatomical modules of the bird skull (Felice & Goswami, 2018).

Results

PC 1 explains 45.3% of the total variance and mainly describes skull elongation (Figure 2). PC 2 explains 10.2% of variance and represents the dorsoventral beak curvature as well as the mediolateral expansion of the palatine bones (see Supplementary Figure S2 for additional morphospaces). The region of morphospace associated with moderate PC1 and PC2 scores contains the greatest density of taxa and corresponds to passerines and species with passerine-like cranial proportions. Both migration and habitat density states have heavily overlapping convex hulls with broad morphospace occupation, indicating that there are a number of viable phenotypes within each ecological trait state. Non-migratory birds occupy a region of morphospace with higher PC 2 values (Figure 2B), associated with high beak curvature in a convex direction compared with migratory birds which occupy a region of morphospace with lower PC 2 scores. Partially migratory birds overlap with migratory and non-migratory species, but also exhibit both the highest and lowest PC 2 scores of our sample. Whereas altricial species explore a region of morphospace defined by low PC 1 scores and higher PC 2 values, precocial taxa are restricted to a smaller region of morphospace relative to semi-precocial or altricial taxa.

Significant relationships were observed between shape and size, habitat density, and migration categories (P < 0.01), but there was not a statistically significant relationship between shape and developmental mode (p = .096; Table 1). Additionally, there are significant interactions between size and habitat density (p = .001), among size, habitat density, and developmental mode (p = .001), and size and developmental mode (p = .002). There are also significant interactions between size, habitat, and migration (p = .037).

We further identified significant differences in evolutionary rates ($\sigma_{\rm mult}$) among the character states of the three traits (Figure 3). Birds living in dense or semi-open habitats evolve approximately three times more rapidly (1.97×10^{-7} and 1.50×10^{-7} , respectively) than those in open habitats (5.85×10^{-8}). Migratory birds have a faster rate of skull evolution (1.64×10^{-7}) than non-migratory or partially migratory birds (7.07×10^{-8} and 1.06×10^{-7} , respectively). Precocial birds have a rate of cranial evolution approximately three times faster (3.03×10^{-7}) than semi-precocial birds (9.63×10^{-8}) and approximately four times faster than altricial birds (7.48×10^{-8}).







Figure 2. Principal component analyses of whole-skull shape. PC 1 describes 45.3% and PC 2 represents 10.2% of the overall shape variation, as illustrated by the landmark configurations along the PC axes in (A). Convex hulls indicate variation of states for the following ecological and life-history traits: (A) Habitat density; (B) migration; (C) developmental mode.

Discussion and conclusion

Our analyses demonstrate two additional factors, habitat density and migration, are significantly associated with avian

skull shape. Furthermore, both ecological and life-history traits affect rates of cranial shape evolution across a globally distributed and speciose sample of birds. These results add

	Pillai's test statistics	SES (effect sizes)	<i>p</i> -values
Size	0.977	7.48	.001**
Habitat density	1.77	3.35	.001**
Migration	1.79	3.82	.001**
Developmental mode	1.73	1.23	.096
Size:habitat density	1.82	3.67	.001**
Size:migration	1.74	0.749	.248
Habitat density:migration	3.49	1.07	.151
Size:developmental mode	1.79	2.55	.002**
Habitat density:developmental mode	3.50	1.13	.127
Migration:developmental mode	3.44	-0.181	.585
Size:habitat density:migration	3.57	1.69	.037*
Size:habitat density:developmental mode	3.64	2.77	.001**
Size:migration:developmental mode	3.50	0.224	.451
Habitat density:migration:developmental mode	4.36	-0.256	.637
Size:habitat density:migration:developmental mode	2.58	-0.671	.766

Table 1. Results of Type II phylogenetic non-parametric MANOVA of skull shape against whole-skull centroid size, habitat density, migration, and developmental mode.

Results for MANOVA, including effect sizes, for interactions between our three traits and size are listed with a colon denoting an interaction between the listed traits. Significances of Pillai's test statistics are based on permutations (n = 1,000) with *p*-values significant at the following alpha levels: * \leq .05, ** \leq .01.



Figure 3. Estimated evolutionary rates (σ_{mult}) for the three different character states of habitat density, migration, and developmental mode.

to the growing body of research suggesting that there is a complex interplay of intrinsic (Bright et al., 2016; Marugán-Lobón et al., 2021; Navalón et al., 2020) and extrinsic factors (Natale & Slater, 2022; Pigot et al., 2020) contributing to avian skull shape evolution.

Our discovery of a significant relationship between skull shape and migration is consistent with previous studies reporting smaller brain sizes in migratory birds (Vincze, 2016), as well as smaller forebrains of migratory "warblers" compared with non-migratory species (Burish et al., 2004). These patterns may be explained by skull size being under strong selection to be lightweight for aerodynamics, driving weight-reducing adaptations in cranial anatomy. Furthermore, brain size may be developmentally or energetically constrained in migrants because of the metabolic costs of migration (Winkler et al., 2004; McGuire & Ratcliffe, 2011) and high-energy use of the brain (Isler & van Schaik, 2009). Alternately, birds with small brains may migrate to compensate for low behavioral flexibility (Winkler et al., 2004). Additionally, the majority of brain size variation is often found superficially in the nidopallium and hyperstriatum regions of the forebrain (Nicolakakis et al., 2003; Rehkämper et al., 1991; Winkler et al., 2004). It is therefore possible that this forebrain region is also responsible for the skull shape covariation with migration which we uncovered.

Analysis of evolutionary rates across character states demonstrated that migrants' skulls evolve faster than those of non-migratory birds. We found that migratory birds evolved faster than partially migratory birds which, in turn, evolved faster than non-migratory birds. Similarly, Winkler et al. (2004) also found the effect of migration on brain size was stronger in long-distance migrants. We propose that these rapid rates of evolution are associated with migratory syndrome, that is., the adaptations of behavior and morphology for migration (e.g. Dingle, 1996; Piersma et al., 2005). In this case, the rapid rates of skull evolution in migrants may be associated with smaller forebrains and dorsoventrally lower skull vaults relative to non-migratory species. Focusing on skull regions, the vault in particular, and to a lesser extent the rostrum, evolves faster in migratory birds compared to non-migratory species (Supplementary Table S3). This result lends further support to the notion that the rapid rates of evolution in migrants is associated with migratory syndrome. Taken as a whole, our results suggest migration exerts a significant selective pressure on brain development, which results in the rapid evolution of different vault morphologies.

Beyond migration, habitat density also affects both avian skull shape and rates of skull evolution across birds. Habitat density covaries with overall skull shape, corroborating work by Kennedy et al. (2020), which found that habitat and strata differentiate corvoid passerine morphology. We discovered heterogenous rates of evolution among birds inhabiting more or less dense habitats, with birds in dense habitats evolving most rapidly. Birds in semi-open habitats evolve more rapidly than those in open habitats which corroborates one of the findings of Eliason et al. (2021) that kingfishers living in forests experience faster brain shape evolution than those in more open habitats. Faster evolutionary rates in dense habitats may be explained by birds in forest habitats adapting to microhabitats, which are not captured by our broad habitat density categories. In addition, birds in open habitats must be highly adapted to extreme environments, which may act as a constraint on cranial morphological evolution; for instance, penguins are adapted to extreme Antarctic conditions and have the slowest evolutionary rates detected in birds (Cole et al., 2022).

In contrast to the results for the ecological traits, developmental mode is not significantly associated with cranial shape variation. The difference in association between ecological and developmental traits may reflect the fact that the two ecological traits are associated with lifelong resource acquisition (Pigot et al., 2016; Ricklefs, 2005; Winkler & Leisler, 1985), while developmental mode may not affect selective pressures experienced by adult birds. Whereas this sample was comprised of adult specimens, an avenue for future research may be investigating whether juvenile bird skull shape or ontogenetic trajectory covary with developmental mode.

Nonetheless, precocial birds have a significantly higher rate of evolution than semi-precocial or altricial species, similar to patterns observed in placental mammals (Goswami et al., 2022). Rates of evolution are fastest in the vault module, particularly for precocial birds (Supplementary Table S3). We hypothesize that these differences are due to precocial hatchlings independently living and interacting with their environment at an earlier age than do altricial hatchlings, including all passerines, which are fed by parents. This earlier independence also drives more rapid neurocranial morphological evolution in precocial birds than in semi-precocial birds such as gulls, which are fed by parents despite being capable of leaving the nest soon after hatching. This study aimed to comprehensively investigate the role of ecological and life-history traits in the accumulation of phenotypic diversity in a major global radiation. Our results demonstrate that whereas developmental mode only influences evolutionary rates, habitat density and migration shape both the tempo and mode of avian phenotypic evolution. This highlights the importance of investigating a range of factors that may influence evolution, as opposed to presuming a form-function relationship focused on solely one function, particularly for complex, multifunctional structures such as the skull. Skull evolution in birds is not simply a reflection of feeding ecology, but also a product of complex interactions between morphology, life-history, and ecological traits.

Supplementary material

Supplementary material is available online at *Evolution* (https://academic.oup.com/evolut/qpad079).

Data availability

Three-dimensional surface models scans are freely available at www.phenome10k.org. The raw data and code are archived at www.github.com/EloiseSEHunt/Hunt_et_al_2023_ Bird_skulls.

Author contributions

R.N.F., J.A.T., and E.S.E.H. collected the data. A.G., R.N.F., and E.S.E.H. conceived the study and designed the analyses. All authors prepared the manuscript.

Conflict of interest: The authors declare no conflict of interest.

Acknowledgments

We thank Judith White, Chris Milensky, Christine Lefevre, Steve Rogers, Ben Marks, Janet Hinshaw, Paul Sweet, Lydia Garetano, Kristof Zyskowski, and Greg Watkins-Colwell for facilitating specimen access for morphometric data collection. E.S.E.H. received funding from a Natural Environment Research Council studentship (grant no. NE/S007415/1). Data collection was supported by European Research Council grant STG-2014–637171 (to A.G.), Natural Environment Research Council grant no. NE/I028068/1 (to J.A.T.), and SYNTHESYS grant no. FR-TAF-5635 (to R.N.F.).

References

- Adams, D. C., & Otárola-Castillo, E. (2013). Geomorph: An R package for the collection and analysis of geometric morphometric shape data. *Methods in Ecology and Evolution*, 4(4), 393–399. https:// doi.org/10.1111/2041-210x.12035
- Adams, D. C., Collyer, M., Kaliontzopoulou, A. & Baken, E. (2022). Geomorph: Software for geometric morphometric analyses. R package version 4.0.4. https://cran.r-project.org/package=geomorph.
- Alerstam, T., Hedenström, A., & Åkesson, S. (2003). Long-distance migration: Evolution and determinants. Oikos, 103(2), 247–260. https://doi.org/10.1034/j.1600-0706.2003.12559.x
- Arnold, S. J., & Wassersug, R. J. (1978). Differential predation on metamorphic anurans by garter snakes (Thamnophis): Social behavior

as a possible defense. *Ecology*, 59(5), 1014–1022. https://doi. org/10.2307/1938553

- Bardua, C., Fabre, A-C., Clavel, J., Bon, M., Das, K., & Stanley, E. L., Blackburn, D. C., Goswami, A. (2021). Size, microhabitat, and loss of larval feeding drive cranial diversification in frogs. *Nature Communications*, 2503, 1–13.
- Battley, P. F., Piersma, T., Dietz, M. W., Tang, S., Dekinga, A., & Hulsman, K. (2000). Empirical evidence for differential organ reductions during trans-oceanic bird flight. *Proceedings Biological Sciences*, 267(1439), 191–195. https://doi.org/10.1098/ rspb.2000.0986
- Berthold, P., Helbig, A. J., Mohr, G., & Querner, U. (1992). Rapid microevolution of migratory behaviour in a wild bird species. *Nature*, 360(6405), 668–670. https://doi.org/10.1038/360668a0
- Botelho, J. F., Smith-Paredes, D., & Vargas, A. O. (2015). Altriciality and the evolution of toe orientation in birds. *Evolutionary Biology*, 42(4), 502–510. https://doi.org/10.1007/s11692-015-9334-7
- Bright, J. A., Marugán-Lobón, J., Cobb, S. N., & Rayfield, E. J. (2016). The shapes of bird beaks are highly controlled by nondietary factors. *Proceedings of the National Academy of Sciences of the United States of America*, 113(19), 5352–5357. https://doi.org/10.1073/ pnas.1602683113
- Bright, J. A., Marugán-Lobón, J., Rayfield, E. J., & Cobb, S. N. (2019). The multifactorial nature of beak and skull shape evolution in parrots and cockatoos (Psittaciformes). BMC Evolutionary Biology, 19, 31101003.
- Brusaferro, A., & Insom, E. (2009). Morphometric analysis of the kingfisher cranium (Aves). *Italian Journal of Zoology*, 76, 53–63.
- Burish, M. J., Kueh, H. Y., & Wang, S. S. H. (2004). Brain architecture and social complexity in modern and ancient birds. *Brain Behavior* and Evolution, 63(2), 107–124. https://doi.org/10.1159/000075674
- Carrier, D. R. (1996). Ontogenetic limits on locomotor performance. *Physiological Zoology*, 69(3), 467–488. https://doi.org/10.1086/ physzool.69.3.30164211
- Case, T. J. (1978). On the evolution and adaptive significance of postnatal growth rates in the terrestrial vertebrates. *Quarterly Review of Biology*, 53(3), 243–282. https://doi.org/10.1086/410622
- Chira, A. M., Cooney, C. R., Bright, J. A., Capp, E. J., Hughes, E. C., & Moody, C. J., Nouri, L. O., Varley, Z. K., Thomas, G. H. (2018). Correlates of rate heterogeneity in avian ecomorphological traits. *Ecology Letters*, 21, 1505–1514.
- Clavel, J., Aristide, L., & Morlon, H. (2019). A penalized likelihood framework for high-dimensional phylogenetic comparative methods and an application to new-world monkeys brain evolution. *Systematic Biology*, 68(1), 93–116. https://doi.org/10.1093/sysbio/ syy045
- Clavel, J., Escarguel, G., & Merceron, G. (2015). MvMORPH: An R package for fitting multivariate evolutionary models to morphometric data. *Methods in Ecology and Evolution*, 6(11), 1311–1319. https://doi.org/10.1111/2041-210x.12420
- Cole, T. L., Zhou, C., Fang, M., Pan, H., Ksepka, D. T., Fiddaman, S. R., Emerling, C. A., Thomas, D. B., Bi, X., Fang, Q., Ellegaard, M. R., Feng, S., Smith, A. L., Heath, T. A., Tennyson, A. J. D., Borboroglu, P. G., Wood, J. R., Hadden, P. W., Grosser, S., ... Zhang, G. (2022). Genomic insights into the secondary aquatic transition of penguins. *Nature Communications*, 13, 1–13.
- Collar, D. C., Schulte, J. A., O'Meara, B. C., & Losos, J. B. (2010). Habitat use affects morphological diversification in dragon lizards. *Journal of Evolutionary Biology*, 23(5), 1033–1049. https://doi. org/10.1111/j.1420-9101.2010.01971.x
- Cooney, C. R., Bright, J. A., Capp, E. J., Chira, A. M., Hughes, E. C., & Moody, C. J., Nouri L. O., Varley, Z. K., Thomas, G. H. (2017). Mega-evolutionary dynamics of the adaptive radiation of birds. *Nature*, 542, 344–347.
- Cooney, C. R., Sheard, C., Clark, A. D., Healy, S. D., Liker, A., Street, S. E., Troisi, C. A., Thomas, G. H., Székely, T., Hemmings, N., & Wright, A. E. (2020). Ecology and allometry predict the evolution of avian developmental durations. *Nature Communications*, 11, 1–9.

- Crouch, N. M. A., & Tobias, J. A. (2022). The causes and ecological context of rapid morphological evolution in birds. *Ecology Letters* 25(3), 611–623. https://doi.org/10.1111/ele.13962
- Da Silva, F. O., Fabre, A. -C., Savriama, Y., Ollonen, J., Mahlow, K., Herrel, A., Müller, J., & Di-Poï, N. (2018). The ecological origins of snakes as revealed by skull evolution. *Nature Communications*, 9(1), 376. https://doi.org/10.1038/s41467-017-02788-3
- Degrange, F. J., & Picasso, M. B. J. (2010). Geometric morphometrics of the skull of Tinamidae (Aves, Palaeognathae). Zoology, 113(6), 334–338.
- Dial, T. R., & Carrier, D. R. (2012). Precocial hindlimbs and altricial forelimbs: Partitioning ontogenetic strategies in Mallard ducks (*Anas platyrbynchos*). Journal of Experimental Biology, 215(Pt 21), 3703–3710. https://doi.org/10.1242/jeb.057380
- Dingle, H. (1996). *Migration. The biology of life on the move*. Oxford University Press.
- Ducatez, S., & Field, D. J. (2021). Disentangling the avian altricial-precocial spectrum: Quantitative assessment of developmental mode, phylogenetic signal, and dimensionality. *Evolution*, 75(11), 2717– 2735. https://doi.org/10.1111/evo.14365
- Dumont, M., Wall, C. E., Botton-Divet, L., Goswami, A., Peigné, S., & Fabre, A-C. (2016). Do functional demands associated with locomotor habitat, diet, and activity pattern drive skull shape evolution in musteloid carnivorans? *Biological Journal of the Linnean Soci*ety, 117, 858–878.
- Eliason, C. M., McCullough, J. M., Andersen, M. J., & Hackett, S. J. (2021). Accelerated brain shape evolution is associated with rapid diversification in an avian radiation. *American Naturalist*, 197(5), 576–591. https://doi.org/10.1086/713664
- El-Sayed, A. (2019). Bird migration. In A. F. El-Sayed (Ed.), *Bird Strike in Aviation* (pp. 183–184). John Wiley & Sons Ltd.
- Fabbri, M., Mongiardino Koch, N., Pritchard, A. C., Hanson, M., Hoffman, E., Bever, G. S., Balanoff, A. M., Morris, Z. S., Field, D. J., Camacho, J., Rowe, T. B., Norell, M. A., Smith, R. M., Abzhanov, A., & Bhullar, B. S. (2017). The skull roof tracks the brain during the evolution and development of reptiles including birds. *Nature Ecology and Evolution*, 1(10), 1543–1550. https://doi. org/10.1038/s41559-017-0288-2
- Felice, R. N., & Goswami, A. (2018). Developmental origins of mosaic evolution in the avian cranium. Proceedings of the National Academy of Sciences of the United States of America, 115(3), 555–560. https://doi.org/10.1073/pnas.1716437115
- Felice, R. N., Tobias, J. A., Pigot, A. L., & Goswami, A. (2019). Dietary niche and the evolution of cranial morphology in birds. *Proceedings of the Royal Society B: Biological Sciences*, 286, 1–9.
- Fitzpatrick, J. W. (1985). Form, foraging behaviour, and adaptive radiation in the Tyrannidae. Ornithological Monographs, 36, 447–470.
- Goswami, A., Noirault, E., Coombs, E. J., Clavel, J., Fabre, A-C., & Halliday, T. J., Churchill, M., Curtis, A., Watanabe, A., Simmons, N. B., Beatty, B. L., Geisler, J. H., Fox, D. L., Felice, R. N. (2022). Attenuated evolution of mammals through the Cenozoic. *Science*, 378, 377–383.
- Grant, B. R., & Grant, P. R. (1989). Natural selection in a population of Darwin's finches. American Naturalist, 133, 377–393.
- Griesser, M., Drobniak, S. M., Graber, S. M., & van Schaik, C. P. (2023). Parental provisioning drives brain size in birds. Proceedings of the National Academy of Sciences of the United States of America, 120, 1–10.
- Hardie, J. L., & Cooney, C. R. (2022). Sociality, ecology and developmental constraints predict variation in brain size across birds. *Journal of Evolutionary Biology*, 36, 144–155.
- Hedenström, A. (2008). Adaptations to migration in birds: Behavioural strategies, morphology and scaling effects. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 363(1490), 287– 299. https://doi.org/10.1098/rstb.2007.2140
- del Hoyo, J., Cabot, J. & Baptista, L. F. (1992). The handbook of the birds of the world. Lynx Edicions.
- Isler, K., & van Schaik, C. P. (2009). The Expensive Brain: A framework for explaining evolutionary changes in brain size. *Journal of*

Human Evolution, 57(4), 392–400. https://doi.org/10.1016/j.jhevol.2009.04.009

- Iwaniuk, A. N., & Nelson, J. E. (2002). Can endocranial volume be used as an estimate of brain size in birds?. *Canadian Journal of* Zoology, 80, 16–23.
- Jetz, W., Thomas, G. H., Joy, J. B., Hartmann, K., & Mooers, A. O. (2012). The global diversity of birds in space and time. *Nature*, 491(7424), 444–448. https://doi.org/10.1038/nature11631
- Kennedy, J. D., Marki, P. Z., Fjeldså, J., & Rahbek, C. (2020). The association between morphological and ecological characters across a global passerine radiation. *Journal of Animal Ecology*, 89(4), 1094–1108. https://doi.org/10.1111/1365-2656.13169
- Kulemeyer, C., Asbahr, K., Gunz, P., Frahnert, S., & Bairlein, F. (2009). Functional morphology and integration of corvid skulls—A 3D geometric morphometric approach. *Frontiers in Zoology*, 6, 1–14.
- Lack, D. (1968). Bird Migration and Natural Selection. Oikos, 19(1), 1–9. https://doi.org/10.2307/3564725
- Losos, J. B. (2010). Adaptive radiation, ecological opportunity, and evolutionary determinism. *American Naturalist*, 175(6), 623–639. https://doi.org/10.1086/652433
- Marugán-Lobón, J., & Buscalioni, A. D. (2009). New insight on the anatomy and architecture of the avian neurocranium. *Anatomical Record*, 292, 364–370.
- Marugán-Lobón, J., Nebreda, S. M., Navalón, G., & Benson, R. B. J. (2021). Beyond the beak: Brain size and allometry in avian craniofacial evolution. *Journal of Anatomy*, 240, 197–209.
- McGhee, G. (2011). Convergent evolution: Limited forms most beautiful. The MIT Press.
- McGuire, L. P., & Ratcliffe, J. M. (2011). Light enough to travel: Migratory bats have smaller brains, but not larger hippocampi, than sedentary species. *Biology Letters*, 7(2), 233–236. https://doi. org/10.1098/rsbl.2010.0744
- Miles, D. B., & Ricklefs, R. E. (1984). The correlation between ecology and morphology in deciduous forest passerine birds. *Ecology*, 65(5), 1629–1640. https://doi.org/10.2307/1939141
- Millien, V. (2006). Correction: Morphological evolution is accelerated among island mammals. *PLoS Biology*, 4, e384384. https://doi. org/10.1371/journal.pbio.0040384
- Natale, R., & Slater, G. J. (2022). The effects of foraging ecology and allometry on avian skull shape vary across levels of phylogeny. *American Naturalist*, 200, 174–188.
- Navalón, G., Bright, J. A., Marugán-Lobón, J., & Rayfield, E. J. (2019). The evolutionary relationship among beak shape, mechanical advantage, and feeding ecology in modern birds. *Evolution*, 73, 422–435.
- Navalón, G., Marugán-Lobón, J., Bright, J. A., Cooney, C. R., & Rayfield, E. J. (2020). The consequences of craniofacial integration for the adaptive radiations of Darwin's finches and Hawaiian honeycreepers. *Nature Ecology & Evolution*, 4, 270–278.
- Navalón, G., Nebreda, S. M., Bright, J. A., Fabbri, M., Benson, R. B., & Bhullar, B-A., Marugán-Lobón, J., & Rayfield, E. J. (2021). Craniofacial development illuminates the evolution of nightbirds (Strisores). *Proceedings of the Royal Society B: Biological Sciences*, 288, 1–10.
- Nicolakakis, N., Sol, D., & Lefebvre, L. (2003). Behavioural flexibility predicts species richness in birds, but not extinction risk. *Animal Behaviour*, 65, 445–452.
- Olsen, A. M. (2017). Feeding ecology is the primary driver of beak shape diversification in waterfowl. *Functional Ecology*, 31(10), 1985–1995. https://doi.org/10.1111/1365-2435.12890
- Pap, P. L., Osváth, G., Daubner, T., Nord, A., & Vincze, O. (2020). Down feather morphology reflects adaptation to habitat and thermal conditions across the avian phylogeny. *Evolution*, 74(10), 2365–2376. https://doi.org/10.1111/evo.14075
- Paradis, E., & Schliep, K. (2019). ape 5.0: An environment for modern phylogenetics and evolutionary analyses in R. *Bioinformatics*, 35(3), 526–528. https://doi.org/10.1093/bioinformatics/bty633
- Piersma, T., Pérez-Tris, J., Mouritsen, H., Bauchinger, U., & Bairlein, F. (2005). Is there a "migratory syndrome" common to all migrant birds? *Annals of the New York Academy of Sciences*, 282, 293.

- Pigot, A. L., Sheard, C., Miller, E. T., Bregman, T. P., Freeman, B. G., Roll, U., Seddon, N., Trisos, C. H., Weeks, B. C, & Tobias, J. A. (2020). Macroevolutionary convergence connects morphological form to ecological function in birds. *Nature Ecology and Evolution*, 4(2), 230–239. https://doi.org/10.1038/s41559-019-1070-4
- Pigot, A. L., Trisos, C. H., & Tobias, J. A. (2016). Functional traits reveal the expansion and packing of ecological niche space underlying an elevational diversity gradient in passerine birds. *Proceedings* of the Royal Society B: Biological Sciences, 283, 2013–2015.
- Price, S. A., Holzman, R., Near, T. J., & Wainwright, P. C (2011). Coral reefs promote the evolution of morphological diversity and ecological novelty in labrid fishes. *Ecology Letters*, 14(5), 462–469. https://doi.org/10.1111/j.1461-0248.2011.01607.x
- Prum, R. O., Berv, J. S., Dornburg, A., Field, D. J., Townsend, J. P., Lemmon, E. M., & Lemmon, A. R. (2015). A comprehensive phylogeny of birds (Aves) using targeted next-generation DNA sequencing. *Nature*, 526(7574), 569–573. https://doi.org/10.1038/nature15697
- Pulido, F. (2007). The genetics and evolution of avian migration. Bio-Science, 57(2), 165–174. https://doi.org/10.1641/b570211
- Rehkämper, G., Frahm, H. D., & Zilles, K. (1991). Quantitative development of brain and brain structures in birds (galliformes and passeriformes) compared to that in mammals (insectivores and primates). *Brain Behav. Ecol.*, 37, 125–134.
- Revell, L. J. (2012). phytools: An R package for phylogenetic comparative biology (and other things). *Methods in Ecology and Evolution*, 3, 217–223.
- Ricklefs, R. E. (1979). Adaptation, constraint, and compromise in avian postnatal development. *Biological Review*, 54(3), 269–290. https:// doi.org/10.1111/j.1469-185x.1979.tb01013.x
- Ricklefs, R. E. (2005). Small clades at the periphery of passerine morphological space. *American Naturalist*, 165(6), 651–659. https://doi.org/10.1086/429676
- Rosenzweig, M. (1978). Competitive speciation. *Biological Journal of the Linnean Society*, 10, 275–289.
- Salewski, V., & Bruderer, B. (2007). The evolution of bird migration—A synthesis. *Naturwissenschaften*, 94(4), 268–279. https:// doi.org/10.1007/s00114-006-0186-y
- Schlager, S. (2017). Morpho and Rvcg—Shape analysis. In R. G. Zheng, S. Li, & G. J. Szekely (Eds.), *Statistical shape and deformation analysis* (pp. 217–256). Academic Press.
- Shatkovska, O., & Ghazali, M. (2020). Integration of skeletal traits in some passerines: Impact (or the lack thereof) of body mass, phylogeny, diet and habitat. *Journal of Anatomy*, 236(2), 274–287. https:// doi.org/10.1111/joa.13095
- Sherratt, E., Serb, J. M., & Adams, D. C. (2017). Rates of morphological evolution, asymmetry and morphological integration of shell shape in scallops. BMC Evolutionary Biology, 17(1), 248. https:// doi.org/10.1186/s12862-017-1098-5
- Sol, D., Garcia, N., Iwaniuk, A., Davis, K., Meade, A., Boyle, W. A., & Székely, T. (2010). Evolutionary divergence in brain size between migratory and resident birds. *PLoS One*, 5(3), e9617. https://doi. org/10.1371/journal.pone.0009617
- Starck, J. M. (1993). Evolution of Avian Ontogenies, in. Current Ornithology, 10, 275–383.
- Tobias, J. A., & Pigot, A. L. (2019). Integrating behaviour and ecology into global biodiversity conservation strategies. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 374(1781), 20190012. https://doi.org/10.1098/rstb.2019.0012
- Tobias, J. A., Sheard, C., Seddon, N., Meade, A., Cotton, A. J., & Nakagawa, S. (2016). Territoriality, social bonds, and the evolution of communal signaling in birds. *Frontiers in Ecology and Evolution*, 4, 103389.
- Tobias, J. A., Sheard, C., Pigot, A. L., Devenish, A. J. M., Yang, J., Sayol, F., Neate-Clegg, M. H. C, Alioravainen, N., Weeks, T. L, Barber, R. A., Walkden, P. A., MacGregor, H. E. A., Jones, S. E. I, Vincent, C., Phillips, A. G., Marples, N. M., Montaño-Centellas, F. A., Leandro-Silva, V., Claramunt, S., ... Schleuning, M. (2022). AVONET: Morphological, ecological and geographical data for all birds. *Ecology Letters*, 25(3), 581–597. https://doi.org/10.1111/ ele.13898

- Tokita, M., Yano, W., James, H. F., & Abzhanov, A. (2017). Cranial shape evolution in adaptive radiations of birds: Comparative morphometrics of Darwin's finches and Hawaiian honeycreepers. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 372(1713), 20150481. https://doi.org/10.1098/rstb.2015.0481
- Vágási, C. I., Pap, P. L., Vincze, O., Osváth, G., Erritzøe, J., & Møller, A. P. (2016). Morphological adaptations to migration in birds. *Evolutionary Biology*, 43, 48–59.
- Vidal-García, M., & Scott Keogh, J. (2017). Phylogenetic conservatism in skulls and evolutionary lability in limbs—Morphological evolution across an ancient frog radiation is shaped by diet, locomotion and burrowing. BMC Evolutionary Biology, 17(1), 165. https:// doi.org/10.1186/s12862-017-0993-0
- Vinciguerra, N. T., & Burns, K. J. (2021). Species diversification and ecomorphological evolution in the radiation of tanagers (Passeriformes: Thraupidae). *Biological Journal of the Linnean Society*, 133, 920–930.
- Vincze, O. (2016). Light enough to travel or wise enough to stay? Brain size evolution and migratory behavior in birds. *Evolution*, 70(9), 2123–2133. https://doi.org/10.1111/evo.13012
- Wassersug, R. J., & Sperry, D. G. (1977). The relationships of locomotion to differential predation on *Pseudacris triseriata* (Anura: Hylidae). *Ecology*, 58(4), 830–839. https://doi.org/10.2307/1936218
- Watanabe, A., Gignac, P. M., Balanoff, A. M., Green, T. L., Kley, N. J., & Norell, M. A. (2019). Are endocasts good proxies for brain size

- Wickham, H. (2016). *Ggplot2: Elegant graphics for data analysis* (2nd edn.). Springer International Publishing.
- Wiley, D. F., Amenta, N., Alcantara, D. A., Ghosh, D., Kil, Y. J., Delson, E. Harcourt-Smith, W., Rohlf, F. J., St John, K., & Hamann, B. (2005). Evolutionary morphing. In *Proceedings of IEEE visualization 2005* (pp. 431–438). IEEE.
- Winger, B. M., Lovette, I. J., & Winkler, D. W. (2012). Ancestry and evolution of seasonal migration in the Parulidae. *Proceedings of the Royal Society B: Biological Sciences*, 279, 610–618.
- Winkler, H. & Leisler, B. (1985). Morphological aspects of habitat selection in birds. In M. Cody (Ed.), *Habitat selection in birds* (pp. 415–434). Academic Press.
- Winkler, H., Leisler, B., & Bernroider, G. (2004). Ecological constraints on the evolution of avian brains. *Journal fuer Ornithologie*, 145, 238–244.
- Yamasaki, T., Aoki, S., & Towkita, M. (2018). Allometry and integration do not strongly constrain beak shape evolution in large-billed (*Corvus macrorhynchos*) and carrion crows (*Corvus corone*). Ecology and Evolution, 8, 10057–10066.
- Zink, R. M. (2002). Towards a framework for understanding the evolution of avian migration. *Journal of Avian Biology*, 33(4), 433–436. https://doi.org/10.1034/j.1600-048x.2002.03081.x