Ecology predicts life history evolution in birds

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Abstract:
Although a number of studies describe current evolutionary patterns concerning life-history evolution, how historical changes in the way organisms interact with their environment have shaped life-history evolution still remains unresolved. In this study, I integrate prospective and retrospective comparative approaches to ask what ecological factors have driven current variation in lifespan of bird passerines. An analysis of >500 species suggest that lifespan is higher in cooperative breeders and in species that build the nest in more secure sites, consistent with the age-specific theory of life history evolution. A retrospective analysis further indicated that these two traits likely evolved through an Ornstein-Uhlenbeck evolutionary model with different optima for each selective regime. Specifically, transitions to cooperative breeding behavior and to nesting in less exposed sites resulted in changes towards longer lifespan optima. These results are the first evidence that ecological and behavioral changes produced in life history strategies in the past, and provide new insights to understand and predict current and future life-history evolutionary patterns.

Key words:
Life-history, fast-slow continuum, Passerines, Ornstein-Uhlenbeck
INTRODUCTION

Evolution has led to a great diversity of life cycles. Such variety in life history strategies is reflected in how all the living forms differ in the investment of energy to grow, reproduce or survive. As different life history traits affect reproduction and survival, and therefore fitness, the study of life history remains crucial to understand evolutionary processes. However, how such diversity of life histories has evolved remains a major unresolved question in biology.

There is consensus that the enormous diversity of life-history strategies results from environmental factors influencing life history traits and tradeoffs that act as evolutionary constrains (Stearns 2000). One of the major axis of life-history variation is explained by the tradeoff between survival and fecundity, which defines the fast-slow continuum (Stearns 1983, Bielby et al. 2007). Thus, at the fast extreme there are species that prioritize reproduction over survival through an earlier reproductive age, a greater reproductive effort and a lower survival, whereas the slow extreme is represented by species prioritizing survival over reproduction.

Life history theory regards age-specific mortality as a major agent of selection that drives the fast-slow continuum evolution. This is based on demographic models and predicts life history evolution depending on which age groups are selected. When adult mortality is high, then earlier maturation age and higher reproductive effort should be selected, while when adult mortality is low, selection should favor a later maturation age and a longer lifespan because there is a lower risk to die before reproducing. On the contrary, increased juvenile mortality rates should favor a slow-lived strategy, while increased adult mortality rates should select for a fast-lived strategy (Charlesworth 1980, Reznick et al. 2002).

The age-specific hypothesis is supported by several empirical studies in different taxa. First, there is evidence that in wild populations high mortality rates lead to an earlier maturation age and higher reproductive effort, whereas at low mortality rates selection drives life-history evolution to the slow extreme (Reznick et al. 1990, Crowl and Covich 1990). Second, several comparative studies also support that species living in less risky environments are associated with slow-lived strategies and viceversa (Shattuck and Williams 2000, Holmes and Austad 1994, Martin 2011).

Although there are a number of prospective studies concerning the evolution of the fast-slow continuum, there has been little effort to use retrospective analyses. However, documenting current patterns of variation is insufficient to study evolutionary processes because the processes we currently observe are not necessarily the same that occurred in the past. Thus, some authors have highlighted the need to study adaptations with an integrative approach that combines prospective and retrospective perspectives (Losos 1994). Admittedly, a major challenge of retrospective analyses is the need to infer evolutionary changes that occurred millions of years ago. Yet, the current availability of life
history and ecological information, together with the development of new methods to reconstruct past changes in a phylogeny (Revell 2012, Beaulieu et al. 2012), provide a unique opportunity to address the evolution of life histories within a historical framework.

Here, I use these advances to ask how changes in ecology have shaped the fast-slow continuum evolution in passerines. My focus is on maximum lifespan, defined as the longest period any individual of a species has been recorded to survive, which is considered a major component of the fast-slow continuum. According to theory, several ecological and behavioral factors are predicted to affect lifespan. First, nesting behavior has been related with juvenile and adult survival in several ways. For example, it has been suggested that open nests or those located on the ground suffer higher predation rates than those located in cavities or in tall trees or cliffs (Martin and Li 1992, Martin 1995, Shattuck and Williams 2010). Second, foraging behavior may also affect lifespan, increasing with height where foraging activity is performed. Indeed, previous work in birds and mammals suggest that both flight and arboreality are related with longer lifespan probably due to a reduced exposure to terrestrial predators (Shattuck and Williams 2010, Pomeroy 1990). Third, social factors such as cooperative breeding and colonial behavior are also hypothesized to be related with lifespan, predicting longer lifespan for both cooperative breeders and colonial species due to a reduction of mortality (Arnold and Owens 1998, Varela et al. 2007). Fourth, relations between lifespan and both parental care and mating system have also been reported; although in this case the underlying mechanisms remain unclear, these could be related to the costs of the increment in mating competition and parental care (Liker and Székely 2005). Fifth, classical theories predict fewer species at equilibrium on islands rather than in the mainland and, therefore, islands should support fewer predators (MacArthur and Wilson 1967, Blumstein 2002). Thus, insularity should be related with a longer lifespan through a reduction in predation risk. Finally, it has been suggested that migrant species should exhibit lower survival rates than residents because of the costs of travelling long distances over unfamiliar regions (Sillett and Holmes 2002).

My initial goal is to use a prospective approach to identify which of the above ecological and behavioral factors are related to lifespan. To this purpose, I use phylogenetic least square regressions within a model selection framework as a way to assess the relative importance of each factor.

Next, I reconstruct the most relevant factors in a phylogeny to test whether past evolutionary changes have occurred several times independently across the phylogeny, and if so, use new developed Brownian and Ornstein-Uhlenbeck evolutionary models to directly assess whether and how changes in these factors bring associated changes in lifespan, and, therefore, in the fast-slow continuum. Brownian motion models assume that changes in life history randomly accumulate over time whereas Ornstein-Uhlenbeck models instead assume that there is a single optimum value of life history or different optima for each selective regime (Beaulieu et al. 2012). Integrating prospective and
retrospective approaches, I pretend to address how the variation of ecological and behavioral patterns influenced evolutionary changes towards fast lived or slow lived strategies in the past.

MATERIAL AND METHODS

Birds have played an important role in the study of life-history evolution because ecological, behavioral, life history and phylogenetic information are easily available. For all these reasons this study is focused on birds and in particular, on Passerines in order to reduce the error resulting from the variation that exists in some traits, such as the mode of development, between different orders.

Data collection

Records of lifespan, ecological, behavioral and confounding variables were obtained from published literature for 555 species of passerines (see source references in Appendix S1). However, because information of these variables was lacking for some species, the initial dataset was reduced to 327 species in analyses where all the variables were tested simultaneously. Phylogenies for the species analyzed were downloaded from the complete phylogeny of extant birds, which comprises 9,993 species, compiled in a Bayesian framework (Jetz et al. 2012). Subsets of 100 phylogenetic trees were obtained from the complete phylogeny for both Hackett and Ericson pseudo-posterior distributions, to ensure that the obtained results were not affected by phylogenetic uncertainties.

Description of variables

Lifespan records, obtained from published material (see sources in Appendix S1), included both wild and captivity maximum longevity data. Previous work has shown that there is a good correlation between wild and captivity maximum longevity data (Ricklefs 2000), which justifies their use in comparative analyses. When both wild and captivity records were available, then the maximum lifespan was used as it is a closer estimate of the maximum potential lifespan (Shattuck and Williams 2010).

The ecological and behavioral variables hypothesized to affect either juvenile or adult extrinsic mortality were: i. nest site (mean height where the nest is placed, categorized as ground= 0m, shrub= 0-3m, canopy= >3m, cliff); ii. nest protection (open, dommed, non-excavator= hole nesting); iii. parental care (biparental, female care); iv. cooperative breeding (non-cooperative, occasional, cooperative breeder); v. foraging behavior (main foraging substrate classified as: ground (0m), low vegetation (0-3m), arboreal (>3m), aerial, generalist); vi. coloniality (colonial, loosecolonies, facultative, solitary); vii. mating system (monogamous, monogamous/polygynous, polygynous); viii. insularity (mainland, islands); ix. migration (migrant, resident). When one species could not be unambiguously assigned
to a unique category, I decided to assign it to the category where predation risk was lower, assuming that these sites would be preferred when predation risk is high.

Body size, biogeographical region and diet were included as confounding factors. Body size is related to lifespan in birds and mammals through extrinsic mortality effects (Calder 1983, Ricklefs 2000) or intrinsic mortality effects associated with basal metabolic rates (Speakman 2005, Hulbert et al. 2007). The biogeographic region (coded as Africa, Australia, Indomalayan, Nearctic, Neotropical, Palearctic, Multiregion) can also affect life history traits, with tropical species tending toward the slow extreme of the fast-slow continuum compared with non-tropical species (Ghalambor and Martin 2001, Martin 2004). Finally, diet (carnivorous, herbivorous, omnivorous) can also affect lifespan; the underlying mechanisms still remain unclear, but these could be related with indirect effects on body size and/or somatic maintenance effects (Munshi-South and Wilkinson 2006, Wasser and Sherman 2010). The sources for all these variables are presented in Appendix S1.

Data analysis

PHILOGENETIC LEAST SQUARE REGRESSION

To identify what ecological and behavioral factors affect lifespan, I conducted a Phylogenetic Least Square Regression Analysis (PGLS, hereafter) using the R Caper package (Orme 2012). Lifespan and body size were log-transformed to improve the linearity of the relationship. To assess what factors mostly affected lifespan, I first conducted univariate analyses using the maximum sample size for each variable. To ensure that the results were not affected by phylogenetic uncertainties, each univariate analysis was repeated 10 times with different randomly selected trees of the posterior distribution of both Hackett and Ericson phylogenies. Then, using the MuMIn R package (Barton 2012), I validated all the possible combination of variables performing a model selection based on AICc values. Because several “best models” models were selected (ΔAICc < 4), I calculated the sum of the AICw values over all the models where each variable was included to infer the relative importance of the variable.

STOCHASTIC CHARACTER MAPPING AND EVOLUTIONARY MODELS

The factors selected in the PGLS analyses were reconstructed on the phylogenies using a Stochastic Character Mapping procedure (Nielsen 2002, Huelsenbeck 2003). This method uses a Bayesian approach to model character changes following a continuous-time Markov process (Nielsen 2002, Huelsenbeck et al. 2003). In order to reconstruct the potential trait changes on the phylogeny, the R package Phytools (Revell 2012) was used to obtain 300 stochastic character maps for each factor by running five simulations per each phylogenetic tree obtained from subsets of the complete phylogeny of Hackett and Ericson distributions. To test whether there are independent evolutionary changes across the phylogeny, the
number and type of character transitions were calculated for each factor from the obtained character stochastic maps.

The stochastic character maps were then analyzed using the R package OUwie to assess what evolutionary model best explains the evolution of lifespan under the different selective regimes (Beaulieu et al. 2012). I considered a variety of both Brownian motion (BM), and Ornstein-Uhlenbeck (OU) models (Beaulieu et al. 2012). The OU models fitted were: 1) a simple OU model with a single optimum ($\theta$) for all the species (“OU1” model), 2) an “OUM” model with different optimum means and a single strength of selection ($\alpha$) and rate of stochastic motion around the optima ($\sigma^2$), 3) OU models with different optimum means and multiple $\sigma^2$ (“OUMV” model) or $\alpha$ (“OUMA” model) across the selective regimes, and 4) an “OUMVA” model that allows $\theta$, $\sigma^2$ and $\alpha$ variation.

Moreover, two different BM models were also fitted: a single rate “BM1” model and a “BMS” model with different rate parameters for each state or phylogeny. Brownian motion models can describe drift, drift-mutation balance and stabilizing selection toward a moving optimum (Beaulieu et al. 2012). Although I hypothesize that the studied ecological changes should lead to changes in the rate of lifespan change (BMS model) or to different optima for each selection regime (OUM models), models assuming that different factors do not affect differently lifespan changes (BM1 and OU1 models) were also fitted as control. Furthermore, the rejection of BMS model would indicate that lifespan evolution has not followed random processes.

The performance of evolutionary models was evaluated with 80 randomly selected stochastic character maps obtained for both Hackett and Ericson phylogenies. To find the best model supported by the data, a model selection based on Akaike weights (AICw) was conducted through the calculation of the relative likelihood of each model in each phylogeny, and then averaging the AICw of each model overall phylogenetic trees (Burnham and Anderson 2002). Then, the parameter estimates of the models selected were averaged to obtain their mean and their 2.5 and 97.5% quantiles.

RESULTS

PGLS

Univariate PGLS models revealed an association between lifespan and two of the studied ecological factors: cooperative breeding and nest site. According to the models, cooperative breeders exhibit a longer maximum life than non-cooperative breeders whereas ground nesters showed shorter lives than canopy nesters (Table 1A). These results are consistent regardless of the phylogenetic hypothesis used (Table 1A). A third variable, diet, also seem to be associated with lifespan, yet in this case evidence is less clear. Thus,
although some models suggest significant differences between carnivores and omnivores, the overall model is non-significant and only the 50% of the sampled phylogenies are significant (Table 1B). Finally, body size is positively associated with lifespan, being the overall model consistent in all the phylogenetic hypotheses used (Table 1B).

Model selection analyses indicate that there are several models that best explain the relation between lifespan and all the ecological and behavioral variables (Appendix S2). Cooperative breeding and nest site have a consistent importance across the models (relative importance = 0.754 and 0.347, respectively), despite the importance of the confounding effects of body size and diet (relative importance = 0.999 and 0.8789, respectively) and the notable reduction in sample size (N = 327) due to missing values (Figure 1). Again, the results are consistent regardless of the phylogenetic hypothesis used (Figure 1).

STOCHASTIC CHARACTER MAPPING AND EVOLUTIONARY MODELS

The 500 stochastic character maps generated for cooperative breeding and nest site (see Fig 2 for an example) shows that several independent changes have occurred along the phylogenies (Appendix S3). In the case of cooperative breeding, most transitions are from no cooperation to occasional or frequent cooperation (Appendix S3A), whereas in the case of nest site transitions are more evenly distributed (Appendix S3B).

When trying to fit the evolutionary models, it turned out that some of the OU models were far more complex than the information contained in our data, and as a result some of the parameters were poorly estimated. For this reason, the OUMA and OUMVA models could not be fitted. The other simpler models were fitted but their eigenvalues were examined in order to detect and remove those cases containing non accurate parameter estimates.

Based on AICw (Table 2), the best model for both cooperative breeding behavior and nest site is the OUM model, with all the alternative models receiving little support (AICw<0.1226). Indeed, there are striking differences between OU and BM models, with OU models receiving far more support than BM models (Table 2).

The model-averaged estimates of the parameter for both cooperative breeding and nest site show different lifespan optima for each selective regime (Table 3 and 4).

Specifically, the inferred optimum values suggest that cooperative and cliff nester species evolved to longer lifespan, that occasional cooperative breeders and canopy and shrub nesters evolved to intermediate longevity values (with canopy nesters having a longer lifespan than shrub nesters) and non-cooperative breeders, and that ground nesters evolved toward shorter lifespan.
DISCUSSION

The results presented here represent the first evidence to date that historical changes in the ecology and behavior of animals have brought associated changes in their lifespan. Specifically, changes in cooperative breeding and in nesting behavior appear to have been associated with important lifespan adjustments in the direction predicted by life history theory. Below I discuss the results and their implications for life history evolution.

OU models received more support that BM models, indicating that lifespan evolution has not followed random processes. Moreover, OUM support shows that different optima for each selective regime have driven lifespan evolution in the past.

Changes from non-cooperative breeding to occasional cooperative breeding, and from occasional cooperative breeding to cooperative breeding seem to have resulted in an increment of lifespan whereas changes from lower nests sites to higher or inaccessible nests sites have also resulted in changes to an increment of lifespan.

The findings that cooperative breeding behavior brought associated changes in lifespan are in agreement with Arnold and Owens (1998), who showed that cooperative breeding is related with low adult mortality, and therefore, long lifespan. In winter helpers contribute in sentinel behavior to detect predators, so that pairs can decrease their sentinel behavior and increase foraging time, and during the breeding season pairs with helpers are benefited through a greater nest protection than lone pairs (Hailman et al. 1994). Thus, cooperative breeding seems to imply a reduction of predation, and, therefore, extrinsic mortality. Although this finding was supported by posterior studies (Wasser and Sherman 2010), others found no relationship between cooperative breeding and lifespan (Blumstein and Moller 2008). My results not only show that cooperative breeding is associated with lifespan, but also yield evidence that past evolutionary changes from non-cooperative to cooperative breeding are associated with changes towards longer lifespan.

Taken together, the results indicate that historical changes in cooperative breeding brought associated changes in the fast-slow continuum, favoring long-lived strategies presumably through mortality effects.

The results concerning lifespan changes associated with the different nest site selective regimes also support previous predictions that adult survival is associated to nest sites (Martin 1995). The results are consistent with predictions that arboreality is related with longer lifespan due to a reduction of terrestrial predators while species living on the ground are associated with shorter lifespan as they suffer lower survival due to higher predation rates (Shattuck and Williams 2010). Moreover, songbirds inhabiting in cliffs exhibited longer lifespan than canopy nesters, suggesting that cliffs are probably a more secure nest site than canopies as they are more inaccessible to predators. Thus, as changes in nest site brought associated changes in lifespan through extrinsic mortality effects,
results show how changes from nest sites exposed to high predation risk to nest sites with lower predation risk resulted in changes towards long lived strategies in the past. My results however show some discrepancies with Martin 1995, who found that due to reduced predation effects, ground nesters exhibited greater adult survival than canopy and shrub nesters, and that survival of canopy nesters was higher than shrub nesters. Although the present results also show greater lifespan for canopy nesters respect shrub nesters, ground nesters exhibit the lowest lifespan, and, therefore the lowest survival.

Inferring which factors have affected past evolutionary changes in the fast-slow continuum is of great importance to understand current evolutionary patterns of life history variation or to predict evolutionary patterns in the future. The integration of prospective and retrospective perspectives has allowed me to show how past changes in ecological and behavioral variables brought associated changes towards fast or slow strategies, and, therefore can contribute to the understanding of current patterns associated with life history evolution and to the prediction of future evolutionary patterns.

However, the unbalanced data resulting from focusing the study on passerines have not allowed the possibility to fit complex evolutionary models. Thus, it would be interesting to analyze in the future how variation in the strength of selection and in the rate of stochastic motion has explained transitions towards the different selective regimes optimums. Further research is also needed to understand how ecology and behavior have shaped life history evolution through both adult and juvenile mortality. While there is currently abundant information on adult survival, how changes in ecological and behavioral patterns influenced past changes in life-history evolution through juvenile mortality still remains unresolved due to the paucity of mortality information for this age-stage. Filling this gap can represent an important avenue for future research.

**ACKNOWLEDGEMENTS**

I’m grateful to Sol’s lab for their helpful support and discussions. Moreover, I would like to thank all the previous studies in which this study is based, as without the important work of experimental studies, comparative approaches would not be possible.
REFERENCES


Barton, K., 2012. Model selection and model averaging based on information criteria (AICc and alike). R package version 1.9.5


Table 1A. PGLS univariate analyses integrating 20 sampled phylogenies of both Hackett and Ericson phylogenies. Min. and max. values of the estimates, st. error, factor and model sign., as well as the proportion of the significant variables among the 10 phylogenies for the different distributions is also given.

| Factors                  | N     | Estimates   | Est. error | Pr(|t|)          | Proportion significant factors | Model p-value | Proportion significant models |
|--------------------------|-------|-------------|------------|-----------------|-------------------------------|---------------|-------------------------------|
|                          |       |             |            |                 | Ericson                      | Hackett       | Ericson                      | Hackett |
| **Foraging**             |       |             |            |                 |                               |               |                               |         |
| Aerial*                  | 444   | 0.0000      | —          | —               | —                             |               |                               |         |
| High veg.                | 444   | -0.0629/-0.0564 | 0.1342 / 0.1377 | 0.6117 / 0.7496 | 0%                            | 0%            | 0.4453 / 0.5556               | 0%      |
| Ground                   | 444   | 0.0095 / 0.0193 | 0.1370 / 0.1405 | 0.8476 / 0.9971 | 0%                            | 0%            | 0%                            | 0%      |
| Generalists              | 444   | 0.0366 / 0.0419 | 0.1244 / 0.1285 | 0.6773 / 0.8165 | 0%                            | 0%            | 0%                            | 0%      |
| Low veg.                 | 444   | -0.1158/-0.1127 | 0.1559 / 0.1596 | 0.4140 / 0.5424 | 0%                            | 0%            | 0%                            | 0%      |
| **Colonial behavior**    |       |             |            |                 |                               |               |                               |         |
| Colonial*                | 553   | 0.0000      | —          | —               | —                             |               |                               |         |
| Facultative              | 553   | -0.1122/-0.1108 | 0.1211 / 0.1223 | 0.3239 / 0.3983 | 0%                            | 0%            | 0%                            | 0%      |
| Loose colonies           | 553   | -0.0720/-0.0712 | 0.1278 / 0.1292 | 0.5384 / 0.6450 | 0%                            | 0%            | 0%                            | 0%      |
| Solitary                 | 553   | -0.0785/-0.0729 | 0.0996 / 0.1014 | 0.4044 / 0.5310 | 0%                            | 0%            | 0.8520 / 0.9124               | 0%      |
| **Coop. breeding**       |       |             |            |                 |                               |               |                               |         |
| Non-cooperative*         | 547   | 0.0000      | —          | —               | —                             |               |                               |         |
| Occasional               | 547   | 0.1559 / 0.246 | 0.0708 / 0.0716 | **0.0251 / 0.0362** | 100%                         | 100%          |                               | 0%     |
| Cooperative              | 547   | 0.1633 / 0.1742 | 0.0778 / 0.0787 | **0.0266 / 0.0418** | 100%                         | 100%          | **0.0047 / 0.0078**          | 100%    |
| **Mating system**        |       |             |            |                 |                               |               |                               |         |
| Monogamous*              | 436   | 0.0000      | —          | —               | —                             |               |                               |         |
| Polygynyous              | 436   | 0.1098 / 0.1193 | 0.1129 / 0.1157 | 0.2710 / 0.3952 | 0%                            | 0%            | 0.4203 / 0.6023               | 0%      |
| Monog/Polyg.             | 436   | 0.0649 / 0.0738 | 0.0842 / 0.0856 | 0.3548 / 0.4463 | 0%                            | 0%            | 0%                            | 0%      |
| **Nest protection**      |       |             |            |                 |                               |               |                               |         |
| Dommed*                  | 522   | 0.0000      | —          | —               | —                             |               |                               |         |
| Non-exavator             | 522   | 0.0058 / 0.0086 | 0.1206 / 0.1242 | 0.8588 / 0.9920 | 0%                            | 0%            | 0%                            | 0%      |
| Open                     | 522   | -0.1153/-0.1044 | 0.1111 / 0.1164 | 0.3025 / 0.3848 | 0%                            | 0%            | 0.1832 / 0.3043               | 0%      |

* These categories have been set to zero and have been used as baseline for comparison.
Table 1B. PGLS univariate analyses integrating 20 sampled phylogenies of both Hackett and Ericson phylogenies. Min. and max. values of the estimates, st. error, factor and model sign., as well as the proportion of the significant variables among the 10 phylogenies for the different distributions is also given.

| Factors             | N    | Estimates          | Est. error          | Pr(>|t|) | Proportion significant factors | Model p-value | Proportion significant models |
|---------------------|------|--------------------|---------------------|---------|--------------------------------|----------------|-----------------------------|
| Nest Site           |      |                    |                     |         | Ericson                        | Hackett        | Ericson                     | Hackett |
| Canopy*             | 553  | 0.0000             | —                   | —       | —                              |                |                             |         |
| Clift               | 553  | 0.0326 / 0.0385    | 0.1042 / 0.1069     | 0.6847 / 0.8008 | 0% | 0%                            |                |                             |         |
| Ground              | 553  | -0.2907/-0.2849    | 0.0784 / 0.0805     | <0.001 / <0.001 | 100% | 100%                         |                |                             |         |
| Shrub               | 553  | -0.0135/-0.0051    | 0.0499 / 0.05036    | 0.7444 / 0.9459 | 0% | 0%                            | <0.001 / 0.0014 | 100% | 100%                         |
| Parental Care       |      |                    |                     |         |                                |                |                             |         |
| Biparental*         | 529  | 0.0000             | —                   | —       | —                              |                |                             |         |
| Female only         | 529  | -0.1428/-0.1282    | 0.1217 / 0.1233     | 0.2355 / 0.3052 | 0% | 0%                            | 0.2449 / 0.3495 | 0% | 0%                         |
| Insularity          |      |                    |                     |         |                                |                |                             |         |
| Island*             | 552  | 0.0000             | —                   | —       | —                              |                |                             |         |
| Mainland            | 552  | -0.0214/-0.0071    | 0.0848 / 0.0865     | 0.8037 / 0.9325 | 0% | 0%                            | 0.9400 / 0.9928 | 0% | 0%                         |
| Migration           |      |                    |                     |         |                                |                |                             |         |
| Migrant*            | 543  | 0.0000             | —                   | —       | —                              |                |                             |         |
| Resident            | 543  | 0.0113 / 0.0229    | 0.0502 / 0.0508     | 0.6494 / 0.8237 | 0% | 0%                            | 0.8132 / 0.9515 | 0% | 0%                         |
| Diet                |      |                    |                     |         |                                |                |                             |         |
| Carnivores*         | 520  | 0.0000             | —                   | —       | —                              |                |                             |         |
| Herbivores          | 520  | 0.1242 / 0.1270    | 0.0739 / 0.0755     | 0.0602 / 0.1290 | 0% | 0%                            |                |                             |         |
| Omnivores           | 520  | 0.1128 / 0.1162    | 0.0547 / 0.0556     | **0.0196** / 0.0570 | 50% | 60%                          | **0.0244** / 0.1005 | 0% | 10%                         |
| Biogeogr. Region    |      |                    |                     |         |                                |                |                             |         |
| Africa              | 552  | -0.3007/-0.2575    | 0.4843 / 0.4953     | 0.5368 / 0.5999 | 0% | 0%                            |                |                             |         |
| Australia           | 552  | -0.2165/-0.1744    | 0.4823 / 0.4934     | 0.6560 / 0.7214 | 0% | 0%                            |                |                             |         |
| Nearctic            | 552  | -0.0316/0.00260    | 0.4749 / 0.4859     | 0.9472 / 0.9984 | 0% | 0%                            |                |                             |         |
| Palearctic          | 552  | -0.2557/-0.2274    | 0.4755 / 0.4868     | 0.5929 / 0.6373 | 0% | 0%                            |                |                             |         |
| Indomalayan         | 552  | -0.3341/-0.2876    | 0.4827 / 0.4939     | 0.4913 / 0.5569 | 0% | 0%                            |                |                             |         |
| Neotropical         | 552  | -0.1372/-0.1001    | 0.4776 / 0.4885     | 0.7751 / 0.8363 | 0% | 0%                            |                |                             |         |
| Multiregion         | 552  | -0.0293/-0.0008    | 0.4849 / 0.4955     | 0.9521 / 0.9987 | 0% | 0%                            | 0.0136 / 0.0366 | 0% | 0%                         |
| Body size (log)     | 552  | 0.1765 / 0.1874    | 0.0225 / 0.0287     | **<0.001** / **<0.001** | 100% | 100%                        | **<0.001** / **<0.001** | 100% | 100%                     |

* These categories have been set to zero and have been used as baseline for comparison.
Table 2. Model selection based on AICw of lifespan evolution under different ecological and behavioral factors.

<table>
<thead>
<tr>
<th></th>
<th>BM1</th>
<th>BMS</th>
<th>OU1</th>
<th>OUM</th>
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<tbody>
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<td>Coop breeding</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
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<td>0.8774</td>
</tr>
<tr>
<td>Nest Site</td>
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<td>&lt;0.001</td>
<td>0.0015</td>
<td>0.9985</td>
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Table 3. Model-averaged parameter estimates and their associated 2.5 and 97.5% quantiles of OUM models selected for lifespan evolution under different intensities of cooperative breeding.

<table>
<thead>
<tr>
<th>Coop br. OUM</th>
<th>Non-cooperative</th>
<th>Occasional</th>
<th>Cooperative</th>
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Table 4. Model-averaged parameter estimates and their associated 2.5 and 97.5% quantiles of OUM models selected for lifespan evolution for different nesting behaviors.

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**Fig 1.** Relative importance of each variable based on AICw of the model selection performed for 327 species of passerines.

**Fig 2.** Sampled stochastic character map showing cooperative breeding behavior changes on an Ericson phylogeny. Colored branches indicate cooperative breeding behaviors estimated in each branch (blue = non-cooperative breeding; green = occasional cooperative breeding; red = cooperative breeding).
APPENDIX

Appendix S1. List of sources used to construct the database:

Lifespan data


Body size data


Behavioral and ecological data:


Birdlife International. World Bird Database (www.birdlife.org/datazone)


Appendix S2. Model selection based on AICc values to find the best model defining the relation between lifespan and ecological and behavioral factors (N=327).

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Appendix S3. Number and types of transitions for each phylogenetic tree between the different cooperative breeding (a) and nest site strategies (b) for Hackett and Ericson phylogenies, resulting from 300 stochastic character maps for each factor and phylogeny. In cooperative breeding behavior no= non-cooperative breeding.

Appendix S4. Code used for the different analyses.

#CODE (PGLS, STOCHASTIC CHARACTER MAPING & OUwie)

#1.PGLS####

setwd("C:/Users/Mar/Desktop/LifespanTOT")
library(ape)
library(caper)
library(MuMIn)

#PGLS with 1 phylogenetic tree:
tree<-'read.nexus("TotesBS.214.Ericson.tre")  #100 phylogenies
PGLSpasser<-'read.csv("PGLS.N552.csv",sep=";")
attach(PGLSpasser)
Num <- row.names(PGLSpasser)
dades <- cbind(Num, PGLSpasser)

passeriformes <- comparative.data(phy=tree[[16]],data=dades,names.col=SpeciesPhylo,vcv=TRUE, na.omit = F, warn.dropped = TRUE, vcv.dim=3)
model<-p gls(log(lifespanMax)~log_bodySize+DietGroup+F2+C1+CB1+NP2+NS2+MS2+PC1+Insularity +Migration+Biogeography,data=passeriformes,lambd a="ML")

#Model Selection and relative importance calculations:
```r
model.selection <- dredge(pglS(log(lifespanMax) ~
  log_bodySize + DietGroup + F2 + C1 + CB1 + NP2 + NS2 + MS2 + PC1 + Insularity + Migration + Biogeography,
data = passeriformes, lambda = "ML"))
imp.modSel <- importance(model.selection)

#2. STOCHASTIC CHARACTER MAPPING#

setwd("C:/Users/Mar/Desktop/LifespanTOT/StochCharMap")

library(phytools)
library(OUwie)

trees100 <- read.nexus("CB1.N547.Ericson.tre") # 100 phylogenies
tree_sample <- sample(trees100, 5) # 5 phylogenies

passer <- read.csv("CB1.N547.csv", sep = ",")
dades <- data.frame(passer[, 2:3])
rownames(dades) <- passer[, 1]
attach(dades)
names(CB1) <- rownames(dades)
names(log_lifespanMax) <- rownames(dades)

# 10 simulations for 5 phylogenies:
CB1Simmap50 <- make.simmap(tree_sample, CB1, model = "ER", nsim = 10, message = FALSE)
CB1_colors <- c("blue", "green", "red"); names(CB1_colors) <- c("1", "2", "3")
plotSimmap(CB1Simmap50[[1]], CB1_colors, pts = FALSE, lwd = 5)

# Calculation of transitions:
describeCB1 <- describe.simmap(CB1Simmap50)

#3. EVOLUTIONARY MODELS#

setwd("C:/Users/Mar/Desktop/LifespanTOT/StochCharMap")

library(phytools)
library(OUwie)
trees <- read.simmap("CB1Simmap50.trees", format = "phylip")
tree_sample <- sample(trees, 10) # sample with 5 phylogenetic trees
CB1 <- read.csv("CB1.N547.csv", sep = ",")
```
#OUwie models with one phylogeny:

BM1Output = OUwie(trees[[1]], CB1, model = "BM1", simmap.tree = TRUE)
BMSOutput = OUwie(trees[[1]], CB1, model = "BMS", simmap.tree = TRUE)
OU1Output = OUwie(trees[[2]], CB1, model = "OU1", simmap.tree = TRUE)
OUMVOutput = OUwie(trees[[1]], CB1, model = "OUMV", simmap.tree = TRUE)
OUMAOutput = OUwie(trees[[1]], CB1, model = "OUMA", simmap.tree = TRUE)
OUMVOutput = OUwie(trees[[1]], CB1, model = "OUMVA", simmap.tree = TRUE)

#MultiOUwie with several phylogenies:

require (OUwie)

multiOUwie <- function(phy, data, model, nregimes, simmap.trees){
  if(model=="BMS"){
    num.cols <- (nregimes*2) + (nregimes+1)
  }else{
    num.cols <- (nregimes*3)
  }
  #res< matrix(nrow=length(phy),ncol=3+num.cols)
  res<-matrix(nrow=length(phy),ncol=7+num.cols)
  for(i in 1:length(phy)){
    print(i)
    tmp<-OUwie(phy[[i]], data, model=model, simmap.tree=TRUE)
    rownames(tmp$theta)<-colnames(tmp$solution)
    res[i,]<-c(i, tmp$eigval[1:4], tmp$loglik, tmp$AICc,
    tmp$solution[1,order(colnames(tmp$solution))], tmp$solution[2,order(colnames(tmp$solution))],
    tmp$theta[order(rownames(tmp$theta))],1))
  }
  return(res)
}

Multi.OUM <- multiOUwie(phy=tree_sample, data=CB1, model=c("OUM"), nregimes=3, simmap.trees=TRUE)