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1	Reproductive costs in terrestrial male vertebrates: what can we
2	learn from birds?
3	
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20 Abstract

Reproduction requires resources that cannot be allocated to other functions resulting in direct 21 reproductive costs, i.e. trade-offs between current reproduction and subsequent 22 23 survival/reproduction. In wild vertebrates, direct reproductive costs have been widely 24 described in females, but their occurrence in males remains to be explored. To fill this gap, we gathered 53 studies on 48 species testing direct reproductive costs in male vertebrates. We 25 26 found a trade-off between current reproduction and subsequent performances in 29% of the 27 species and in every taxa. As 73% of the studied species are birds, we focused on that class to investigate whether such trade-offs are associated with i) levels of paternal care, ii) polygyny 28 29 or iii) pace of life. More precisely for this third question, it is expected that fast species (i.e. short lifespan, early maturity, high fecundity) pay a cost in terms of survival whereas slow 30 species (with opposite characteristics) in terms of fecundity. Our findings tend to support this 31 hypothesis. Finally, we pointed out the potential confounding effects that should be accounted 32 for when investigating reproductive costs in males and strongly encourage the investigation of 33 34 such costs in more taxa to understand to what extent our results are relevant for other 35 vertebrates.

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Keywords: Costs of reproduction, Generation time, Life history, Paternal care, Polygyny,
Trade-off, Vertebrates

40 1. Introduction

Trade-offs, resulting from constraints on the evolution of linked traits, play a central 41 role in life history theory [1]. According to the principle of allocation [2], individuals allocate 42 their limited amount of resources to a function at the cost of other ones. Therefore, the 43 maximum fitness an individual can reach is limited by trade-offs among fitness components. 44 45 The theory about reproductive trade-offs, first formalized by Williams [3], suggested that, as reproduction is energy-demanding, individuals should trade current reproduction versus future 46 reproduction via reduced future fecundity (i.e., fecundity costs of reproduction) and/or 47 reduced future survival (i.e., survival costs of reproduction) [1,4]. However, as formulated by 48 the van Noordwijk and de Jong's model [5], even if two traits compete for the same resource 49 at the individual level, such a negative trade-off can remain undetected at the population level, 50 because individuals can differ in both resource acquisition and resource allocation to each trait 51 depending on the quality of their habitat or on their own quality. 52

53 Nearly 50 years after Williams's publication [3], studies investigating costs of reproduction have flourished. Stearns reviewed studies investigating the effects of 54 reproduction on growth, parental survival, late fecundity and longevity in a wide range of taxa 55 and with different methods, including laboratory experiments, unmanipulated and 56 manipulated field populations [1]. More recently, studies investigating specifically direct 57 58 reproductive costs in wild mammals, i.e. the co-variations between current reproduction (at time t) and subsequent reproduction and/or survival (at time t+1), have been reviewed [6]. 59 Interestingly, most of the studies gathered in this review focused on females [6]. There have 60 in fact been considerably fewer attempts to assess trade-offs among fitness components in 61 wild males. This bias may be in part explained by methodological problems, i.e. it is often 62 more difficult to assess reproductive effort in wild males than females [7]. Also, while it has 63

become an accepted notion that reproduction is costly for males as well [8–10], studies 64 65 dealing with costs of reproduction in males are often rooted in the theory of sexual selection and therefore refer to the cost of producing or maintaining sexual traits on future survival [8] 66 (see [11] for a recent review on the relationship between strength of sexual selection and age-67 specific survival patterns across vertebrates). Investigating reproductive costs in males, 68 specifically through the co-variations between life history traits (also called direct fitness 69 70 traits sensu Roff [4], i.e. fecundity and survival) can be important because such costs that are expressed at the individual level may also have implications at the population level [12]. 71 Thus, in the perspective of the development of more realistic population models that include 72 73 both females and males, a better understanding of constraints shaping fitness traits in males appears important. 74

In a broad evolutionary context, theory predicts that i) the intensity of mating 75 76 competition in polygynous species should translate into higher costs of reproduction for males compared to socially monogamous species [13]. At the same time, ii) species characterized 77 with high level of paternal care are expected to be the ones with the highest reproductive 78 costs, as parental care is energy-demanding [3,14,15]. While a high level of polygyny or 79 paternal care is expected to be costly for males, such costs of reproduction can also strongly 80 81 depend upon the position of the species on the fast-slow continuum of life history variation [16,17]. Indeed, the recent extension of the van Noordwijk and de Jong's model [5] predicts 82 that reproductive costs are closely linked to the position of a given species on the fast-slow 83 84 continuum [6], that contrasts fast species with early maturity, high fecundity, and short lifespan, to slow species with opposite characteristics (see [18] for a recent review). Briefly, 85 iii) fast species characterized with high variance in survival and low variance in reproduction 86 should exhibit survival costs of reproduction, whereas slow species should rather suffer from 87

fecundity costs. While this evolutionary model has been first supported in female mammals[6], its validity for male vertebrates remains to be investigated.

In this review, we summarize the empirical tests of reproductive trade-off between two 90 breeding seasons in males in unmanipulated wild terrestrial and seasonally reproducing birds, 91 mammals, squamates and amphibians (figure S1 and ESM1). We choose to focus on these 92 taxa because most of the long-term individual based field studies, essential in this context, are 93 94 done on these species [19]. Experimental approaches have been useful tools to study tradeoffs and to show relationship of causality [20]. Thus, numerous studies have investigated 95 costs of reproduction in males from experimental field populations, by assessing for instance 96 97 the effect of brood size manipulation in birds (i.e., reduced or enlarged) on subsequent survival and/or reproduction (e.g. [21,22] for reviews of experimental studies). However, as 98 we aim here to report tests of direct reproductive costs with potential demographic 99 100 consequences, we focus our literature survey on males of unmanipulated wild populations. Then, we compare the occurrence of direct costs of reproduction between taxa and species. As 101 102 most of the gathered studies in our review use birds as case studies, we focus on that taxon to question the link between direct costs of reproduction and i) level of polygyny, ii) level of 103 paternal care, and iii) pace of life (i.e. the position on the fast-slow continuum). Moreover, we 104 105 discuss possible biases of correlative studies in the wild with regard to identifying reproductive costs in males. Finally, we discuss the implication of our results for the other 106 taxa of terrestrial vertebrates and suggest some future lines of research. In particular, we 107 encourage the inclusion of trade-offs between fitness-related traits in males into demographic 108 models of population growth to better predict the fate of wild vertebrate populations. 109 110

111 2. Tests of direct costs of reproduction in males

We gathered 53 studies on 48 vertebrate species investigating the co-variations 112 113 between current (t) reproduction and subsequent (t+1) reproduction/survival (search protocol in ESM2, results of the literature search in table S1). It is noteworthy that the most 114 represented class is the bird one with 35 species for which such co-variations have been 115 investigated. We found only 7 mammalian species, 4 squamate species and 2 amphibian 116 species. In total, we reported 116 co-variations (including one non-informative co-variation) 117 118 between reproductive trait at t and reproduction/survival at t+1 (tables S1 and S2, figure 1). Because of high variability of traits considered at t in the reviewed studies (table S1), 119 we summarized them into 6 types of traits namely "number of young" (e.g. when clutch size 120 121 has been considered as a reproductive trait at t), "breeding status" (e.g. comparison of subsequent performances of breeders vs. non breeders), "mating" (e.g. number of matings), 122 "paternal care" (e.g. a measure of feeding rate), "timing" (e.g. comparison of early vs. late 123 124 breeders) and "number of breeding" (e.g. number of broods produced). In 74% of the covariations (i.e. 85/115), the number of young produced or breeding status was used to assess 125 reproductive effort at time t (table S2). The use of breeding status as a trait at t means that 126 direct costs of reproduction are investigated through the comparison of subsequent 127 performances of breeders (or successful breeders) with subsequent performances of non-128 129 breeders (or failed breeders) (table S1). Thus, negative co-variations between current reproduction and subsequent performances, i.e. when non-breeders at t (or failed breeders at t) 130 outperformed breeders at t (or successful breeders at t) the subsequent breeding season (t+1), 131 132 suggest the existence of direct costs of reproduction. Reproductive traits linked to the level of paternal care, the timing of breeding, the number of matings or the number of breeding may 133 be difficult to measure in the field and are specific of the ecology/reproductive cycle of each 134 species. That can explain why these traits are less often used in the reviewed studies (table 135 S2). For example, in the black grouse (*Tetrao tetrix*), a lekking species for which the amount 136

137 of paternal care is absent, the authors used the number of matings as a reproductive trait at 138 time t [23].

Negative co-variations between reproduction at t and survival or reproduction at time 139 t+1 were detected in 29% of the reported species (table S1). Despite the low number of 140 studies on some taxa, it seems that males of all the taxa have equal probability to exhibit 141 direct costs of reproduction (figure 1, table S3). More precisely, these costs correspond to 16 142 143 negative co-variations between life history traits among the 115 co-variations reported (tables S1 and S2). Interestingly, even if the traits "breeding status" and "matings" have a tendency to 144 be more frequently involved in such negative co-variations, there is no significant effect of the 145 146 types of trait considered at t on the probability to detect reproductive costs (tables S2 and S4). Therefore, breeding (or successfully breeding) in a given breeding season may negatively 147 affect male survival to the next breeding season (this is the case for 3 species out of 21) and 148 149 may also affect reproduction (for 4 out of 15 species) (table S2). Similarly, for some species, a high number of young, of paternal care, of matings or of breeding reduces male survival or 150 151 reproduction the next breeding season (table 1). Moreover, the probability to find a negative co-variation in a study does not depend on the number of co-variations tested (table S5) or the 152 number of types of traits tested neither (table S6), suggesting that the differences in sample 153 154 sizes between studies do not bias our results.

Regarding traits considered at t+1, 57% of the studies investigated only survival costs, 34% both survival and fecundity costs, and only 9% investigated fecundity costs alone (table S1). Interestingly, no study investigating both types of costs provided support for both. For instance, in Laysan albatross (*Phoebastria immutabilis*) or Nazca booby (*Sula granti*), male breeders (or successful breeders) can pay a cost in terms of survival to the next breeding season but not in terms of fecundity (i.e. probability to reproduce given survival) [24,25]. In king penguin (*Aptenodytes patagonicus*), Southern giant petrel (*Macronectes giganteus*) and

spotted owl (*Strix occidentalis*), the effect was opposite: breeding affected future breeding
probability but not survival [26–28]. Similarly, in great reed warbler (*Acrocephalus arundinaceus*), there is a cost of paternal care on future reproduction (timing of settlement)
but no detectable cost on survival [29]. This stresses the importance to study both fecundity
costs of reproduction and survival costs of reproduction.

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168 3. Reproductive costs in male birds and their link with paternal care,

169 polygyny and pace of life

For some species, negative co-variation between reproduction at time t and subsequent 170 reproductive performance occurred, whereas for others species, the co-variation was null or 171 positive (figure 1). These differences among species may be explained by differences in 172 mating systems, levels of paternal care or pace of life resulting in different life history 173 174 strategies. Indeed, we expect higher costs for males in polygynous mating systems, or when males invest more in paternal care. Moreover, as highlighted for female mammals, a close 175 relationship between pace of life and the type of direct reproductive costs detected may be 176 177 expected, i.e. fast species should be affected by survival costs of reproduction whereas slow species by fecundity costs of reproduction [6]. Generation time provides a relevant measure to 178 179 rank the species on the fast-slow continuum [17]. Therefore, we expect an increase of the probability to find a fecundity cost of reproduction compared to the probability to find a 180 181 survival cost of reproduction with longer generation times.

We chose to test these predictions only for male birds because 73% of the species in the reviewed studies are birds. Moreover, the different bird species gathered in this review exhibit a wide range of avian life histories along the fast-slow continuum, diverse mating systems and levels of paternal care which is essential to test these three hypotheses. More

precisely, 11 studies revealed direct costs of reproduction in males (7 found survival costs and 186 187 4 fecundity costs of reproduction) among the 35 different bird species we gathered (table S1). For most species, a score of paternal care and polygyny was available from the literature [30] 188 (see table 1 for details on score calculation) and, for each species, the generation time (in 189 years) was retrieved from species specific demographic studies (table 1). We found that the 190 probability to observe a cost of reproduction in one species did not depend on the level of 191 192 polygyny (table S7.A) or the level of paternal care (table S7.B). Thus, with the current 193 dataset, there is no evidence that species suffering from direct reproductive costs are the ones with the highest levels of paternal care or polygyny. However, as expected, species suffering 194 195 from fecundity costs of reproduction have longer generation times than species suffering from survival costs of reproduction, except for one outlier, the Laysan albatross [25] (figure 2). 196 197 This result tends to support a relationship between the pace of life and reproductive costs [6]. 198 It remains to see from studies in other taxa whether this is a general pattern in vertebrates. The results of these statistical comparisons should be considered with some caution. 199 200 First, the power of our tests may be low due to the number of studies that include data on costs of reproduction. Second, the levels of polygyny, of paternal care and the generation 201 times were calculated at the species level. However, it is true that some variation in these 202 203 variables can exist among populations of the same species [31-33] and even among individuals within a population [e.g. for paternal care 34]. As a consequence, investigating the 204 relationship between reproductive trade-offs and levels of paternal care, polygyny or pace of 205 206 life measured at different levels of biological organization (such as at the population level) may be really interesting. It is noteworthy that in this review, we reported opposite results in 207 208 terms of reproductive costs in two populations of the same species, the willow tit (*Parus montanus*). In one of them, survival costs of reproduction have been detected [35] but not in 209 the other one [36]. Whether these two populations differ in their level of paternal care, 210

polygyny or pace of life and whether these differences are translated into different costs ofreproduction remain to be explored.

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4. Negative, positive or null co-variations between current reproduction and subsequent performances and the detection of reproductive costs

We reported 14% of negative co-variations between current reproduction and
subsequent performances and also 58% of null and 28% of positive co-variations (figure 1).
The high proportions of null and positive co-variations may partly explain why studies
dealing with reproductive costs are often oriented towards females in the literature.

221 A negative co-variation between current reproduction and subsequent performances 222 may indicate that males trade their current reproduction versus their subsequent performances and thus that males suffer from direct reproductive costs [3]. However, this pattern may be 223 224 more complex. In mating systems where males and females interact, the investment of one sex in a current reproductive event may depend on the characteristics of the other sex. For 225 instance, the theory of differential allocation predicts that a mate may invest more or less in 226 227 reproduction when paired with their preferred mate (or with high quality mate) [positive or negative differential allocation: 37]. This pattern can create positive or negative correlations 228 between life history traits for males that are driven by females' investment in reproduction 229 and that are not linked to male reproductive costs. However, because positive differential 230 allocation seems more frequent than negative differential allocation (at least in birds [38]), it 231 should be rare to find a negative co-variation between male life history traits in absence of 232 real reproductive costs for males. 233

A null or a positive co-variation between current reproduction and subsequent 234 235 performances may indicate the absence of direct reproductive costs in males. Again, interaction between the sexes may drive this pattern, for example if the absence of 236 237 reproductive costs in males is correlated to the presence of such costs in females. Such correlations may depend on how strong the sexual conflict over parental investment is and 238 239 how it is resolved in each species [30,39]. Therefore, it would be particularly interesting to 240 compare both males and females in the same study. Moreover, it is noteworthy that males that do not exhibit direct reproductive costs may pay a cost later in life [40], or may suffer from 241 other types of costs, such as inter-generational or cumulative costs of reproduction (see 242 243 ESM1).

Remarkably, null or positive co-variations between current reproduction and 244 subsequent performances may be found even if direct reproductive costs are present. First, in 245 246 certain cases, individuals that try to reproduce but do not succeed to sire offspring may pay quite similar fitness costs than successful breeders. This could be more important in males 247 248 than females because typically males can invest large amount of energy prior to mating (e.g. actively searching for mates, trying to defend a territory, injuries during combat) without 249 managing to successfully sire offspring [7]. Also, brood loss may happen late in the season or, 250 251 at least, after that most of the energy allocated for reproduction has already been invested in the current reproductive event. Therefore, in these cases, one will fail to detect any difference 252 in terms of future survival and/or reproduction between breeders (or successful breeders) and 253 non-breeders (or failed breeders) even if direct costs of reproduction occur. Second, the 254 detection of reproductive costs may be masked by phenotypic differences in individual 255 quality. Indeed, in case of high variance in resources acquisition, individuals able to acquire 256 more resources (i.e. high quality individuals) are also able to allocate more resources than 257 other individuals to both current reproduction and future survival and/or reproduction, which 258

can prevent the detection of costs of reproduction at the population level [5]. These limitations 259 260 due to individual heterogeneity explain in part why experimental studies have been widely used to study trade-offs [e.g. 22]. However, studies of unmanipulated wild populations are 261 262 still essential to understand the ecological consequences of life history variations and to obtain realistic estimates of demographic parameters [19]. Thanks to the development of appropriate 263 statistical tools, accounting for individual heterogeneity is possible, making the correlative 264 265 studies more powerful [6,41]. Yet, it is likely that among the high proportion of studies that reported no negative co-variation between current reproduction and subsequent performances, 266 some of them actually concluded to the absence of direct costs of reproduction even if they 267 268 occurred in the considered species.

More generally, it is also possible that some studies may fail to detect negative co-269 variation between reproduction at time t and survival/fecundity at time t+1, even if there are 270 271 some real costs of reproduction, because appropriate co-factors are not taken into account. For example, differences in individual quality may be more pronounced in years with harsher 272 273 environmental conditions when resources are more limited, resulting in annual variations in trade-off detection [e.g. 24,42]. Accounting for age-effects may also be recommended while 274 investigating direct reproductive costs. Indeed, reproductive performances may be age-275 276 specific, with for example lower reproductive output at old ages compared to younger ages due to senescence [43,44], or at the opposite lower reproductive output at young ages due to 277 inexperience [45], possibly preventing the detection of reproductive costs. Carefully 278 disentangling the age effects from the costs of previous reproduction appears crucial. 279 Moreover, it is also important to keep in mind that the age of the individuals can mediate the 280 trade-offs between current reproduction and subsequent reproduction and/or survival. In other 281 words, reproductive costs themselves can be dependent on the age. For example, reproduction 282 can be more costly in young individuals [46] or on the contrary more costly in old individuals 283

[47], or appear more costly due to terminal investment [48]. Therefore, studies combining the
information of age effects and life history trade-offs should be developed to strengthen
comparative studies and to improve our general understanding of such patterns.

Finally, in many of the studies included in the present review, it is assumed that the 287 social and genetic father is the same. In birds for instance, the number of eggs/chicks present 288 in the nests is used as an estimator of male reproductive success. However, thanks to 289 290 molecular genetic tools, it is now accepted that, even in socially monogamous species, individuals can engage in extra-pair copulations [14,49]. In particular, males involved in such 291 extra-pair copulations can increase their reproductive success without increasing their amount 292 293 of paternal care. This means that some traits measured at time t may be more or less correlated to the reproductive success and the paternal investment of the males. For example, 294 certainty of paternity has been shown to covary with paternal care in birds [50]. Thus, even if 295 296 a male has a large clutch, its investment may be low if some chicks are sired by a different male. This is why the quantification of extra-pair parternity can allow more precise 297 measurement of reproductive effort, which may allow highlighting different relationships 298 between reproduction at time t and fitness-related trait at t+1. 299

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5. Conclusion and perspectives

In this review, we gathered studies exploring the co-variations between current reproduction and subsequent reproduction and/or survival of wild unmanipulated terrestrial male birds, mammals, squamates and amphibians. It is noteworthy that our review reports some studies highlighting positive co-variations between life history traits, suggesting that the individual quality hypothesis is often supported in male vertebrates. But we also found empirical evidence of direct reproductive costs in several species, belonging to all taxa, even with the inherent difficulties of correlative studies. It is thus obvious that direct reproductivecosts concern both males and females in wild populations.

We showed that the occurrence of reproductive costs in males is not correlated to 310 polygyny and levels of paternal care but is associated with pace of life, in birds at least. 311 Unfortunately, the small number of studies in the other taxa did not allow us testing our 312 evolutionary hypotheses in other terrestrial vertebrates. However, we are confident that our 313 314 results, drawn from birds, may also be relevant to other taxa. Indeed, after some evidence of a link between pace of life and costs of reproduction in female mammals [6], our review 315 provides support for that life-history model in male birds. Yet, exploring to what extent such a 316 317 model can be generalized to all terrestrial vertebrates, and, in particular, unravelling the factors that may explain variations among taxa remain an exciting challenge. For example, in 318 line with a comparative study that has shown that birds have a slower life-history than 319 320 mammals for the same body mass [51], one could expect different relationships between pace of life and costs of reproduction in males in these two taxa. Another important difference 321 322 within vertebrates is the mode of temperature regulation. Indeed, ectotherms can store energy more efficiently than endotherms like birds, and thus rely more often on stored resources to 323 fuel reproduction (capital vs. income breeders) [52]. Such different reproductive strategies 324 may induce different reproductive costs. Thus, we strongly encourage further studies in more 325 taxa with diverse mating systems and life history strategies to be able to broaden these results. 326 Correlations between life history traits and in particular, reproductive trade-offs, can 327 328 have demographic consequences and can influence population dynamics [53]. It is true that most models in population dynamics are female-based and neglect males [54] but there is now 329 growing evidence that males may markedly influence population dynamics as well (e.g. 330 [55,56]). While methodological developments now provide the tools to integrate costs of 331 reproduction into population models [53,57], a promising avenue of research could be to take 332

333	into account reproductive costs in males as well as females into population dynamic models.
334	Even if all models are approximations, capturing the fluctuation of demographic parameters
335	and accounting for it into population models is the best way to provide sustainable and
336	relevant management and conservation scenarios.
337	
338	Data accessibility
339	The datasets supporting this article are in tables 1 and S1.
340	Competing interests
341	We have no competing interests.
342	Authors' contributions
343	JB and MG proposed the study and performed the literature search; JB, MG and BES wrote
344	the paper.
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353	
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567	Table 1. Scores of paternal care, polygyny and pace of life. Details on the bird species for which direct reproductive costs have been tested in
568	males. We present their order and family, and their associated scores of paternal care and polygyny. These scores were calculated by Olson et al.
569	[30]. Briefly, the method to calculate the score of paternal care consists in scoring paternal investment in 5 different activities: nest building,
570	incubation, brooding, chick feeding, and chick defence. For each activity, the participation of males was scored on a 5-points scale: 0 (no male
571	care),1 (1–33% male care), 2 (34–66% male care), 3 (67–99% male care) or 4 (100% male care). Thus the maximum score for each species is 20.
572	The score of polygyny represents the percentage of males exhibiting polygyny, with: 0 (no polygyny or less than 0.1% of individuals), 1 (rare
573	polygyny: 0.1–1%), 2 (uncommon polygyny: 1–5%), 3 (moderate polygyny: 5–20%) and 4 (common polygyny, greater than 20%). Generation
574	time (in years), a measure ranking the species on the fast-slow continuum, was extracted from Birdlife International database
575	(http://www.birdlife.org/datazone/home). The column "Negative co-variation?" indicates whether at least one negative co-variation between
576	reproduction at t and fecundity ("F") and/or survival ("S") at $t+1$ was found for the considered species or not ("N").

Species	Order	Family	Paternal care	Polygyny	Generation time	Negative co-variation?	Ref.
Barn swallow (Hirundo rustica)	Passeriformes	Hirundinidae	7	1	3.9	Ν	[58]
Barnacle goose (Branta leucopsis)	Anseriformes	Anatidae	6	0	10.5	Ν	[59]
Black grouse (Tetrao tetrix)	Galliformes	Phasianidae	0	4	6.4	Ν	[23]
Blue tit (Cyanistes caeruleus)	Passeriformes	Paridae	4	3	4.4	Ν	[60]
Brown Thornbill (Acanthiza pusilla)	Passeriformes	Acanthizidae	4**	NA	5.7	Ν	[61]
Cliff swallow (Petrochelidon pyrrhonota)	Passeriformes	Hirundinidae	10	0	4.3	S	[62]

Collared Flycatcher (Ficedula albicollis)	Passeriformes	Muscicapidae	4	3	3.9	Ν	[63]
Crested tit (Parus cristatus)	Passeriformes	Paridae	2	0	4	Ν	[35]
Great reed warbler (Acrocephalus arundinaceus)	Passeriformes	Sylviidae	6	2	4.2	F	[29]
Great tit (Parus major)	Passeriformes	Paridae	4	1	4.3	Ν	[64,65]
Greater prairie-chicken (Tympanuchus cupido)	Galliformes	Phasianidae	0	4	5.5	Ν	[66]
Green-rumped Parrotlet (Forpus passerinus)	Psittaciformes	Psittacidae	2	0	4.1	Ν	[67]
Hawai'i 'Elepaio (Chasiempis sandwichensis sandwichensis)	Passeriformes	Monarchidae	NA	NA	5.9	Ν	[68]
House martin (Delichon urbica)	Passeriformes	Hirundinidae	10	0	4.3	Ν	[69]
Indigo bunting (Passerina cyanea)	Passeriformes	Cardinalidae	3	3	4.1	Ν	[70]
Jackdaw (Corvus monedula)	Passeriformes	Corvidae	NA	NA	7.4	Ν	[71]
King penguin (Aptenodytes patagonicus)	Ciconiiformes	Spheniscidae	6	0	12.7	F	[26]
Kittiwake gull (Rissa tridactyla)	Ciconiiformes	Laridae	10	0	12.9	Ν	[72,73]
Laysan albatross (Phoebastria immutabilis)	Procellariiformes	Diomedeidae	NA	NA	28.5	S	[25]
Long-tailed tit (Aegithalos caudatus)	Passeriformes	Aegithalidae	NA	NA	4.2	Ν	[74,75]
Marsh tit (Parus palustris)	Passeriformes	Paridae	NA	NA	4.2	Ν	[76,77]
Monteiro's storm-Petrel (Oceanodroma monteiroi)	Procellariiformes	Hydrobatidae	NA	NA	16.5	Ν	[78]
Mountain white-crowned sparrow (Zonotrichia	Desserifermes	Eringillidaa	2	2	1.2	S	[70]
leucophrys oriantha)	Fassemonnes	rnngnndae	2	2	4.5	5	[/9]
Nazca booby (Sula granti)	Suliformes	Sulidae	NA	NA	10	S	[24]
Northern giant petrel (Macronectes halli)	Procellariiformes	Procellariidae	NA	NA	17	Ν	[27]

Oystercatcher (Haematopus ostralegus)	Charadriiformes	Haematopodidae	9	1	13.7	Ν	[80]
Savannah sparrow (Passerculus sandwichensis)	Passeriformes	Fringillidae	4	4	3.4	S	[81]
						Ν	[82]
Snowy Plover (Charadrius nivosus)	Ciconiiformes	Charadriidae	NA	NA	5	Ν	[83]
Southern giant petrel (Macronectes giganteus)	Procellariiformes	Procellariidae	NA	NA	21.3	F	[27]
Spotted owl (Strix occidentalis)	Strigiformes	Strigidae	4	0	10.1	F	[28]
Tengmalm's owl (Aegolius funereus)	Strigiformes	Strigidae	3	4	5.8	Ν	[84]
Tree swallow (Tachycineta bicolor)	Passeriformes	Hirundinidae	3	2	4	S	[42]
Wheatear (Oenanthe oenanthe)	Passeriformes	Muscicapidae	4**	NA	4.1	Ν	[85,86]
Willow ptarmigan (Lagopus lagopus)	Galliformes	Phasianidae	3	3	4.2	Ν	[87]
Willow tit (Parus montanus)	Passeriformes	Paridae	3	0	4.6	S	[36]
						Ν	[35]

⁵⁷⁷ ** Personal communication from Andras Liker and Tamas Szekely

579 Figure captions

580

- 581 Figure 1. Distribution of the number of tested co-variations between current
- 582 reproduction and subsequent performances in male terrestrial vertebrates. Number of
- 583 co-variations between reproduction at time t and survival/reproduction at time t+1 collected
- in the literature in amphibians, birds, mammals and squamates.

585

586 Figure 2. Types of direct costs of reproduction in male birds and pace of life.

- 587 Differences in generation times (in years) between the bird species for which fecundity costs
- 588 of reproduction (in blue) and survival costs of reproduction (in red) have been reported for at
- least one reproductive trait. Dots represent the data points (see table 1).

591 Figure 1









