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Postnatal Growth Rates Covary Weakly with Embryonic Development Rates and Do Not Explain Adult Mortality Probability among Songbirds on Four Continents

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Abstract: Growth and development rates may result from genetic programming of intrinsic processes that yield correlated rates between life stages. These intrinsic rates are thought to affect adult mortality probability and longevity. However, if proximate extrinsic factors (e.g., temperature, food) influence development rates differently between stages and yield low covariance between stages, then development rates may not explain adult mortality probability. We examined these issues based on study of 90 songbird species on four continents to capture the diverse life-history strategies observed across geographic space. The length of the embryonic period explained little variation (ca. 13%) in nestling periods and growth rates among species. This low covariance suggests that the relative importance of intrinsic and extrinsic influences on growth and development rates differs between stages. Consequently, nestling period durations and nestling growth rates were not related to annual adult mortality probability among diverse songbird species within or among sites. The absence of a clear effect of faster growth on adult mortality when examined in an evolutionary framework across species indicates that species that evolve faster growth rates may also evolve mechanisms for ameliorating physiological costs on mortality and longevity, such as shifts in fatty acid composition of mitochondrial membranes or cellular repair mechanisms (e.g., Hulbert et al. 2007). Such effects could obviate a relationship between growth rates and adult mortality among species. Instead, adult mortality rates of species in the wild may be determined more strongly by extrinsic environmental causes.

Keywords: life history, adult mortality, nest predation, nestling growth rates, incubation periods.

Introduction

Growth and development rates are critical elements of life-history strategies and theory (Roff 2002). Faster growth and development can arise from various physiological mechanisms, such as faster metabolism (Arendt 1997; West et al. 2001) or trade-offs among developing physiological systems (reviewed in Arendt 1997). These mechanisms and trade-offs can create physiological and phenotypic costs that may compromise adult survival and longevity (McCay 1933; Olsson and Shine 2002; Rollo 2002; Metcalfe and Monaghan 2003). Across various taxa, studies within species have provided experimental support for this intrinsic processes hypothesis, where faster growth rates yielded higher adult mortality (e.g., Olsson and Shine 2002; Rollo 2002; Lee et al. 2013). Yet, experimental manipulations of growth rates within species are tests of proximate responses. Species that evolve faster growth rates may also evolve mechanisms for ameliorating physiological costs on mortality and longevity, such as shifts in fatty acid composition of mitochondrial membranes or cellular repair mechanisms (e.g., Hulbert et al. 2007). Such effects could obviate a relationship between growth rates and adult mortality among species. Moreover, adult mortality may be influenced more strongly by environmental influences such as predation, migration, or non-breeding season stressors (e.g., Rowley and Russell 1991; Sillett and Holmes 2002; Evans et al. 2006; Turbill et al. 2011) than by physiological costs of growth and development. Thus, the influence of variation in growth rates on adult mortality among species in the wild is unclear and deserves a broad test because of the implications for life-history theory.

The influence of intrinsic processes on growth and development rates is thought to be genetically based (Arendt 1997; West et al. 2001). As such, growth and development rates of differing life stages might be expected to positively covary when intrinsic processes are the dominant cause of growth and development rates. In other words, species with long embryonic periods might be expected to also exhibit long postnatal periods and slow growth rates. On the other hand, growth and development rates also may be influenced by extrinsic factors, such as temperature or...
food (Arendt 1997; Badyaev and Martin 2000; Gillooly et al. 2002; Martin et al. 2007, 2013; Remes 2007). Such proximate extrinsic effects may not create the intrinsic costs that can affect adult mortality as expected when development rates are determined primarily by intrinsic processes (Martin and Schwabl 2008; Martin et al. 2013). If proximate extrinsic factors play a strong but differential role between stages, then we might expect low covariance between life stages.

Songbirds provide an interesting system to test these issues. First, lengths of embryonic and nestling periods are thought to be strongly correlated (Moreau and Moreau 1940; Lack 1968; Skutch 1976), potentially reflecting a strong role of intrinsic processes. However, broad phylogenetic tests are lacking. Second, lengths of embryonic periods are negatively correlated with adult mortality probability across species both within and among geographic regions (Martin 2002; Ricklefs 2006; Remes 2007), which may support the intrinsic processes hypothesis. Yet, this correlation may be an indirect result of adult mortality acting on parental effort to influence embryonic development time (Martin 2002). As a result, the importance of physiological costs from faster growth and development for adult mortality remain unclear, at least in the embryonic stage.

Third, the nestling stage may provide a clearer test of the intrinsic processes hypothesis. Variation in embryonic development rates among species is strongly influenced by the proximate extrinsic effects of temperature (Martin 2002; Martin et al. 2007). Temperature is less important in the nestling period as young develop the ability to thermoregulate, and food delivery does not explain variation in nestling growth and development rates (Martin et al. 2011). As a result, intrinsic effects may be more clearly expressed in the nestling stage, as suggested by correlations between growth rates and metabolic rates (Drent and Klaassen 1989). These differing influences of intrinsic and extrinsic effects between the two stages predict low covariance of development rates between the two stages.

Fourth, increased nest predation risk can exert selection to favor evolution of faster growth rates among diverse species (Remes and Martin 2002; Martin et al. 2011). This faster growth appears to be achieved in part through physiological trade-offs (Cheng and Martin 2012). Such physiological trade-offs, together with metabolic influences on growth rates (Drent and Klaassen 1989), provide a reasonable basis for the intrinsic processes hypothesis. An initial test among songbird species of North America found that adult mortality probability was weakly related to nestling growth rates but not to length of nestling periods (Remes 2007). Yet, nestling growth rates and periods in this test were corrected for nest predation effects (Remes 2007), and nest predation can underlie intrinsic trade-offs (Cheng and Martin 2012) that are thought to influence adult mortality. The low amount of variance explained for growth rates, the inconsistent results between metrics (growth rates vs. nestling period duration), and the removal of potential intrinsic effects due to nest predation suggest that further tests are needed. Moreover, songbirds exhibit much greater variation in these life-history traits when examined across geographic space than within North America alone (Martin et al. 2000, 2007, 2011; Sandercock et al. 2000; Ghalambor and Martin 2001; Lloyd et al. 2014). Thus, comparisons across latitudes are needed to allow examination of potential relationships across relatively large geographic shifts in life-history strategies.

Here, we report tests based on field studies of 90 species on four continents. We examined whether lengths of nestling periods and growth rates covaried with lengths of embryonic periods, to explore whether species-specific intrinsic processes underlie the development rates of these two life stages. We further tested the hypothesis that nest predation exerts selection on nestling periods and growth rates that are potentially achieved through physiological trade-offs that explain variation in adult mortality probability. Finally, we tested the ability of nestling period durations and growth rates to explain variation in adult mortality probability.

Material and Methods

Study Areas

We studied 90 passerine species on four continents (fig. A1, available online). We were able to obtain exact observations of embryonic and nestling development times (see below) for these 90 species. We measured growth rates on 80 of these species and were able to estimate annual adult mortality probability for 66 species. The species were studied in north temperate Arizona (34°N), tropical Venezuela (9°N), tropical Malaysia (6°N), and south temperate South Africa (34°S), representing a broad phylogenetic range of songbirds (fig. A1). Nests were studied in north-central Arizona for 26 years (1987–2012) and adult mortality for 21 years (1993–2013) at about 2,350-m elevation in mixed deciduous and coniferous forest (Martin et al. 2007). Nests and adult mortality were studied in the tropics for 7 years (2002–2008) in primary forest in Yacambu National Park, Venezuela, at elevations of 1,400–2,000 m (Martin et al. 2007) and for 5 years (2009–2013) in Kinabalu Park, Malaysia, at 1,450–1,950-m elevation (Martin et al. 2013). Nests were studied for 5 years (2000–2004), and adult mortality for 7–8 years (2001–2007), in south temperate coastal dwarf shrubland near Cape Town, South Africa, at sea level (Martin et al. 2007).
Nest Predation Rates, Development Times, and Nestling Growth Rates

For species examined here, large numbers of nests were monitored following long-term protocols (Martin et al. 2007, 2011). The embryonic period was quantified as the number of days between last egg laid and last egg hatched (Martin et al. 2007). The nestling period was quantified as the number of days between the last egg hatch and the last nestling to leave the nest (Martin et al. 2011). Nests were checked every 2–4 days to determine status and predation events but were checked daily or twice daily during egg laying, near hatching, and near fledging to obtain exact period durations. Nest predation was assumed to be constant, sex specific, or a transient model based on the first year of capture versus all subsequent years. Thus, the global model was $\Phi$ (sex + transient) $p$ (sex + transient). Sex in tropical sites included an unknown category because many species do not exhibit sexual dimorphism and many individuals are not in breeding condition when captured. Parameter estimates were based on averaging across all 16 models based on model weights (Burnham and Anderson 2002) for all species except those in the South Africa site, where we used previous estimates (Lloyd et al. 2014).

We first examined the covariance of embryonic and nestling development times based on the mean estimates for each species using a linear mixed model based on LME4 (Bates et al. 2014). We examined effects of within- and among-site variances using the approach described by van de Pol and Wright (2009) and included site as a random effect in the model. We also tested the same model with nestling growth rate as the dependent variable. We initially included log-transformed mass as a covariate because of potential allometric effects (e.g., Rahn and Ar 1974; Calder 1984), but mass was never a significant influence on these traits, as previously found for songbirds (e.g., Martin et al. 2007, 2011). As a result, we dropped mass from analyses.

We next examined the ability of nestling period length and nestling growth rate to explain variation in adult mortality probability. Again, we initially included body mass as a covariate because both adult mortality and nest predation may be influenced by adult size (Roff 2002; Bianucci and Martin 2010), but mass was not significant and was dropped from analyses. Adult mortality probability was the dependent variable, with nestling period or nestling growth rates as the covariate. We again examined effects of within- and among-site variances using the approach described by van de Pol and Wright (2009) and included site as a random effect in the model.

We followed this analysis with one where we tested the importance of adult and offspring (i.e., nest predation) mortality to variation in nestling period and nestling growth rates. Nestling period or growth rate was the dependent variable in separate analyses with adult mortality probability and nest predation rates as within- and among-

Adult Mortality

In Arizona, Venezuela, and Malaysia, nets were deployed in stations of 10 or 12 nests as subplots within and across all nest-searching plots. These netting subplots were deployed three times per breeding season, with 20–25 days before subplots were revisited. Nets at a station were deployed for 6 h starting at dawn. Netting methods for the South Africa site are detailed in Lloyd et al. (2014). All birds that were captured were banded with numbered metal bands and unique combinations of three color bands (two bands per leg), unless it was a recapture. Color bands were used for resighting by nest searchers who visited each nest plot daily or every other day throughout the season. Resighting and recaptures were used in RMARK (Laake 2013) to estimate annual adult survival probabilities (see “Statistical Analyses”).
Comparisons of a few pairs of related species provide examples that lengths of embryonic and nestling periods can vary independently of each other (fig. 1). First, comparisons of three pairs of related species between the new world tropics (Venezuela) and north temperate zone (Arizona) show large differences in embryonic periods but no difference in nestling periods between latitudes in three different avian families (fig. 1). Similarly, related species within a site demonstrated the same kinds of divergence between stages. For example, two related species in the family Cettidae in Malaysian Borneo showed large differences in opposing directions between stages; *Urosphena*

Figure 1: Comparisons of embryonic period (days; A) and nestling period (days; B) between related species of three avian families in the north temperate Arizona site versus the tropical Venezuela site and in the Cettid family within the Malaysian Borneo site. Troglodytidae includes *Troglodytes aedon* in Arizona and *Hemicorhina leucophrys* in Venezuela. Emberizidae includes *Junco hyemalis* in Arizona and *Arremon bruneinucha* in Venezuela. Parulidae includes *Setophaga virginiae* in Arizona and *Basileuterus tristriatus* in Venezuela. Cettidae includes *Cettia vulcania* first and *Urosphena whiteheadi* second, both in Malaysian Borneo.

Analyses were first made on raw data. Phylogenetically independent contrasts were also calculated and analyzed to control for possible phylogenetic effects (Felsenstein 1985) using the recent comprehensive phylogeny provided by Jetz et al. (2012). Phylogenetic trees were obtained from http://www.birdtree.org (Jetz et al. 2012) using the Hackett et al. (2008) backbone and imported into Mesquite (Madison and Maddison 2011) to construct a majority-rule consensus tree based on 500 trees (fig. A1). The trees were time calibrated, such that branch lengths were included in analyses. Contrasts were calculated using the PDAP module (Midford et al. 2002) and imported into IBM SPSS (ver. 22) for linear regressions through the origin. Results using independent contrasts were reported along with analyses of raw data.

Results

Comparisons of a few pairs of related species provide examples that lengths of embryonic and nestling periods can vary independently of each other (fig. 1). First, comparisons of three pairs of related species between the new world tropics (Venezuela) and north temperate zone (Arizona) show large differences in embryonic periods but no difference in nestling periods between latitudes in three different avian families (fig. 1). Similarly, related species within a site demonstrated the same kinds of divergence between stages. For example, two related species in the family Cettidae in Malaysian Borneo showed large differences in opposing directions between stages; *Urosphena*

Figure 2: Covariance of nestling period (90 species; A) and nestling growth rates (80 species; B) with embryonic development time among four sites spanning north temperate, tropical, and south temperate latitudes. The red ellipses encompass the majority of species from the north temperate Arizona site to demonstrate the narrow range of embryonic development time but large variation in nestling development time and growth rates.
Table 1: Tests of the covariance of nestling period and nestling growth rate with embryonic period based on mixed-model analyses of raw data with site as a random effect and partitioning within- versus among-site variance and a regression analysis of phylogenetic independent contrasts for songbird species from Arizona, Venezuela, Malaysia, and South Africa

<table>
<thead>
<tr>
<th>Variable</th>
<th>B (SE)</th>
<th>P</th>
<th>$r^2_p$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed model:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nestling period (days) as dependent variable ($n = 90$ spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Embryonic period (among sites)</td>
<td>.208 (.238)</td>
<td>.45</td>
<td>...</td>
</tr>
<tr>
<td>Embryonic period (within sites)</td>
<td>.477 (.139)</td>
<td>&lt;.001</td>
<td>...</td>
</tr>
<tr>
<td>Nestling growth rate ($k$) as dependent variable ($n = 80$ spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Embryonic period (among sites)</td>
<td>-.012 (.004)</td>
<td>.004</td>
<td>...</td>
</tr>
<tr>
<td>Embryonic period (within sites)</td>
<td>-.010 (.003)</td>
<td>.001</td>
<td>...</td>
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<tr>
<td>Independent contrasts:</td>
<td></td>
<td></td>
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<tr>
<td>Nestling period (days) as dependent variable ($n = 90$ spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Site</td>
<td></td>
<td>.44</td>
<td>.031</td>
</tr>
<tr>
<td>Embryonic period</td>
<td></td>
<td>.440 (.122)</td>
<td>.001</td>
</tr>
<tr>
<td>Nestling growth rate ($k$) as dependent variable ($n = 80$ spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Site</td>
<td></td>
<td>.021</td>
<td>.121</td>
</tr>
<tr>
<td>Embryonic period</td>
<td></td>
<td>-.008 (.002)</td>
<td>.002</td>
</tr>
</tbody>
</table>

whiteheadi has a much longer embryonic period but a shorter nestling period compared to Cettia vulcania (fig. 1). Many other similar examples exist. Thus, even related species can exhibit strong independent variation in durations of these two developmental stages.

A more general test among 90 species for the covariance of development time between the two stages indicated that embryonic periods were related to nestling period lengths within sites but not among sites (fig. 2A; table 1). However, the independent contrast analyses showed that embryonic development time explained only 13% of the variation in nestling period lengths and nestling periods did not differ among sites (fig. 2A; table 1). Indeed, the red ellipse includes most of the Arizona species, which show a relatively narrow variation in duration of the embryonic period but large variation in the nestling period (fig. 2A), demonstrating that nestling period durations can vary independently of embryonic period durations across species. The lack of significant site effects reflects that variation in nestling period durations in Arizona encompassed the range of variation observed in the other three geographic locations. In contrast, few of the tropical species from Venezuela and Malaysia exhibit embryonic development times within the ellipse representing most of Arizona. Instead, the tropical species generally have longer and more variable embryonic periods than north temperate species.

Table 2: Tests of the ability of nestling period length (days) and nestling growth rate $k$ to predict annual adult mortality probability based on mixed-model analyses of raw data with site as a random effect and partitioning within- versus among-site variance and a regression analysis of phylogenetic independent contrasts for songbird species from Arizona, Venezuela, Malaysia, and South Africa

<table>
<thead>
<tr>
<th>Variable</th>
<th>B (SE)</th>
<th>P</th>
<th>$r^2_p$</th>
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</thead>
<tbody>
<tr>
<td>Mixed model:</td>
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</tr>
<tr>
<td>Adult mortality probability as dependent variable ($n = 66$ spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nestling period (among sites)</td>
<td>-.066 (.099)</td>
<td>.57</td>
<td>...</td>
</tr>
<tr>
<td>Nestling period (within sites)</td>
<td>.005 (.003)</td>
<td>.20</td>
<td>...</td>
</tr>
<tr>
<td>Adult mortality probability as dependent variable ($n = 62$ spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Growth rate $k$ (among sites)</td>
<td>2.432 (1.360)</td>
<td>.33</td>
<td>...</td>
</tr>
<tr>
<td>Growth rate $k$ (within sites)</td>
<td>-.026 (.212)</td>
<td>.92</td>
<td>...</td>
</tr>
<tr>
<td>Independent contrasts:</td>
<td></td>
<td></td>
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<tr>
<td>Adult mortality probability as dependent variable ($n = 66$ spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Site</td>
<td></td>
<td>&lt;.001</td>
<td>.449</td>
</tr>
<tr>
<td>Embryonic period</td>
<td></td>
<td>.004 (.004)</td>
<td>.33</td>
</tr>
<tr>
<td>Adult mortality probability as dependent variable ($n = 62$ spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Site</td>
<td></td>
<td>&lt;.001</td>
<td>.342</td>
</tr>
<tr>
<td>Embryonic period</td>
<td></td>
<td>.138 (.240)</td>
<td>.57</td>
</tr>
</tbody>
</table>
These longer embryonic periods were often observed for species that had nestling periods similar to relatives in north temperate Arizona (figs. 1, 2A), demonstrating why embryonic development time explained little variation in nestling development time.

Embryonic development time also was related to nestling growth rate ($k$) within sites and among sites (fig. 2B; table 1). The significant among-sites effect reflects that average growth rates for a site were correlated with average embryonic development times of sites (fig. 2B; table 1). Nonetheless, embryonic development time still explained only 13% of the variation in nestling growth rates within and among sites based on the independent contrast analyses (table 1). Indeed, comparisons of temperate species in the red ellipse versus tropical species show similar independence of growth rates between stages as seen for nestling period durations. In particular, tropical species show large differences in embryonic development time from temperate species with similar nestling growth rates (fig. 2B).

A test of the intrinsic processes hypothesis that nestling periods or nestling growth rates explained variation in adult mortality probability across species within and among sites found no hint of support (table 2; fig. 3A, 3B). The significant site effects in the independent contrast analyses indicated that adult mortality probability differed among sites, being lower on average in tropical sites (table 2). However, this among-site difference was not correlated with nestling development time, as shown by the nonsignificant among-site test in the mixed models (table 2).

A test of the effects of age-specific mortality on nestling period and nestling growth rates similarly showed that adult mortality probability was not important but nest predation was correlated with both measures of nestling development rates (table 3; fig. 3C, 3D). The independent contrast analyses showed that nest predation explained 34% of the variation in nestling period lengths and 27% of the variation in nestling growth rates (table 3; fig. 3C, 3D).

Figure 3: Plots of adult mortality probability as a function of nestling period length (days; A) and nestling growth rate ($k$; B), plus partial regression plots of nestling period length (days; C) and nestling growth rates ($k$) as a function of nestling predation rates while accounting for adult mortality probability. We did not correct for site differences to allow illustration of any differences.
Table 3: Nestling period length (days) and nestling growth rate $k$ relative to daily nest predation rate during the nestling period and annual adult mortality probability based on mixed-model analyses of raw data with site as a random effect and partitioning within- versus among-site variance and a regression analysis of phylogenetic independent contrasts for songbird species from Arizona, Venezuela, Malaysia, and South Africa.

<table>
<thead>
<tr>
<th>Variable</th>
<th>B (SE)</th>
<th>P</th>
<th>$r_p^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed model:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nesting period as dependent variable (n = 64 spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nest predation rate (among sites)</td>
<td>-167.7 (232.0)</td>
<td>.60</td>
<td>...</td>
</tr>
<tr>
<td>Nest predation rate (within sites)</td>
<td>-154.7 (18.42)</td>
<td>&lt;.001</td>
<td>...</td>
</tr>
<tr>
<td>Adult mortality (among sites)</td>
<td>-17.08 (23.54)</td>
<td>.60</td>
<td>...</td>
</tr>
<tr>
<td>Adult mortality (within sites)</td>
<td>1.932 (3.211)</td>
<td>.55</td>
<td>...</td>
</tr>
<tr>
<td>Nestling growth rate $k$ as dependent variable (n = 61 spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nest predation rate (among sites)</td>
<td>5.116 (5.804)</td>
<td>.54</td>
<td>...</td>
</tr>
<tr>
<td>Nest predation rate (within sites)</td>
<td>2.427 (.392)</td>
<td>&lt;.001</td>
<td>...</td>
</tr>
<tr>
<td>Adult mortality (among sites)</td>
<td>.715 (.590)</td>
<td>.44</td>
<td>...</td>
</tr>
<tr>
<td>Adult mortality (within sites)</td>
<td>.052 (.067)</td>
<td>.44</td>
<td>...</td>
</tr>
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</table>

Independent contrasts:

<table>
<thead>
<tr>
<th>Variable</th>
<th>B (SE)</th>
<th>P</th>
<th>$r_p^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nesting period as dependent variable (n = 64 spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Site</td>
<td>.010</td>
<td>.177</td>
<td></td>
</tr>
<tr>
<td>Nest predation rate</td>
<td>-109.2 (19.92)</td>
<td>&lt;.001</td>
<td>.341</td>
</tr>
<tr>
<td>Adult mortality</td>
<td>.874 (3.227)</td>
<td>.79</td>
<td>.001</td>
</tr>
<tr>
<td>Nestling growth rate $k$ as dependent variable (n = 61 spp.):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Site</td>
<td>.001</td>
<td>.274</td>
<td></td>
</tr>
<tr>
<td>Nest predation rate</td>
<td>1.856 (.418)</td>
<td>&lt;.001</td>
<td>.267</td>
</tr>
<tr>
<td>Adult mortality</td>
<td>.089 (.069)</td>
<td>.17</td>
<td>.034</td>
</tr>
</tbody>
</table>

Discussion

The inability of nestling periods and growth rates to explain any variation in adult mortality probability (fig. 3A, 3B; table 2) across a diverse suite of songbird species is counter to long-standing life-history expectations (McCay 1933; Arendt 1997; Metcalfe and Monaghan 2003; Lee et al. 2013). Moreover, this result differs from the relationship between embryonic development time and adult mortality probability observed in passerine species (Martin 2002; Ricklefs 2006; Remesˇ 2007). These differences between life stages in their correlated variation with adult mortality probability are consistent with the results that nestling development times and growth rates varied largely independently of embryonic development times (figs. 1, 2; table 1).

The independent variation in development rates between stages likely reflects differential effects of extrinsic and intrinsic inputs. The association of adult mortality probability with embryonic development time may result largely through selection on parental effort in warming embryos and influencing development time (Martin 2002). In contrast, temperature is less important to development during the nestling stage of passerine birds because nestlings develop endothermy (Ricklefs 1973; Cheng and Martin 2012). As a result, related species that differ in embryonic periods (fig. 1A) because of parentally induced temperatures (see Martin et al. 2007; Martin and Schwabl 2008) do not necessarily differ in lengths of the nestling period (fig. 1B). Of course, nestling growth rates may be influenced by other extrinsic influences of parental effort, such as feeding rates. Yet, adult mortality should exert selection on parental effort (Williams 1966; Michod 1979; Reznick et al. 1990; Charlesworth 1994; Ghalambor and Martin 2001), and if parental effort (i.e., feeding rates) has a strong influence on growth rates, then growth rates should be positively correlated with adult mortality. Yet, variation in parental feeding rates did not explain variation in growth rates among diverse species (Martin et al. 2011). Moreover, the absence of any relationships between nestling growth rates and adult mortality (fig. 3A, 3B) further suggests that selection by adult mortality on parental effort is of minimal importance to variation in growth rates.

The results here are for a single order of birds (Passeriformes), although it encompasses nearly 50% of the bird species of the world. The applicability of our results to other taxa is unclear. The covariance of embryonic and postnatal development times and rates has not been tested across other taxa with external ectothermic embryos. Yet, substantial independence of the development rates of embryonic and postnatal stages, as we found for songbirds, also might be expected in other taxa. The extrinsic influence of temperature on embryonic development time is...
well documented across diverse ectothermic taxa (Gillooly et al. 2002), in addition to songbirds (Martin 2002; Martin et al. 2007). Yet, in contrast, postnatal growth rates of these taxa may be more strongly influenced by physiological influences (West et al. 2001). One difference between birds and other taxa with external ectothermic embryos is that birds can more strongly alter the temperature environment of the embryo through parental effort in warming eggs (Martin et al. 2007). In contrast, the embryos of other ectothermic species are often simply left exposed to environmental temperatures of the site where they place eggs. Of course, ectothermic species can alter the temperature environment through nest site selection and even parental care in some species, but such temperature modulation is still limited and can have large effects on embryo development rate (Deeming and Ferguson 1991; Madsen and Shine 1999; Angilletta et al. 2009). In contrast to altricial songbirds, temperature effects on development rate may be reduced for postnatal young of ectothermic species because they have the ability to move and behaviorally regulate temperature. Thus, embryonic and postnatal development times of other taxa may also show large divergences due to differing extrinsic and intrinsic inputs between stages, and these possibilities deserve study.

As thought to be true of diverse taxa (i.e., West et al. 2001), variation in postnatal growth rates of birds may be more strongly influenced by intrinsic processes than extrinsic parental effects compared with the embryonic period (Martin et al. 2007; Cheng and Martin 2012). Faster postnatal growth seems to reflect the role of faster metabolism, a key intrinsic process (Drent and Klaassen 1989; West et al. 2001). Moreover, increased nest predation plays an important role in favoring evolution of faster nestling growth rates among bird species (fig. 3B; also Remes and Martin 2002; Martin et al. 2011) and may underlie physiological trade-offs of birds and other taxa (Arendt 1997; Cheng and Martin 2012). These physiological processes and trade-offs underlying growth rate variation can create the physiological costs that are thought to compromise adult survival and longevity among diverse organisms (McGy 1933; Arendt 1997; Metcalfe and Monaghan 2003; Lee et al. 2013). Yet, the lack of covariance of nestling growth rates with adult mortality suggests that these intrinsic costs are of minor importance to the broader range of variation in adult mortality among wild species of differing geographic regions (fig. 3A, 3B; table 2).

While faster growth rates may incur physiological costs, species that evolve faster growth may also evolve physiological mechanisms to offset these costs and their effects on longevity, such as variation in the fatty acid composition of mitochondrial membranes or cellular repair mechanisms (Hulbert et al. 2007). Indeed, the fact that nonflying organisms have greater longevity despite higher metabolic rates for the same body size as nonflying organisms (Holmes and Austad 1995) demonstrates that physiological costs, such as those produced by metabolism, can be ameliorated. Instead, external sources of mortality imposed by predators, migration, or stressors during reproduction or lean seasons (Rowley and Russell 1991; Sillett and Holmes 2002; Leyer et al. 2013) may be more important in driving the majority of adult mortality and longevity for species in the wild (also see Reznick et al. 2004).

In conclusion, development times and rates may vary independently between life stages because proximate external influences, such as temperature, can differ between stages and mask underlying physiological programs that might be similar between stages (e.g., Martin et al. 2013). Physiological costs associated with evolved differences in growth rates among species may not manifest in effects on adult mortality and longevity because species may also evolve mechanisms to ameliorate such costs. Instead, external sources of adult mortality may have a more important role in explaining broad variation in adult mortality among diverse species in the wild. Assumptions that growth rates influence adult mortality and longevity need to be viewed with more caution in an evolutionary framework.

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