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A 'slow pace of life' in Australian old-endemic passerine birds is not accompanied by low basal metabolic rates

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Abstract

Life history theory suggests that species experiencing high extrinsic mortality rates allocate more resources toward reproduction relative to self-maintenance and reach maturity earlier ('fast pace of life') than those having greater life expectancy and reproducing at a lower rate ('slow pace of life'). Among birds, many studies have shown that tropical species have a slower pace of life than temperate-breeding species. The pace of life has been hypothesized to affect metabolism and, as predicted, tropical birds have lower basal metabolic rates (BMR) than temperate-breeding birds. However, many temperatebreeding Australian passerines belong to lineages that evolved in Australia and share 'slow' life-history traits that are typical of tropical birds. We obtained BMR from 30 of these 'old-endemics' and ten sympatric species of more recently arrived passerine lineages (derived from Afro-Asian origins or introduced by Europeans) with 'faster' life histories. The BMR of 'slow' temperate-breeding old-endemics was indistinguishable from that of new-arrivals and was not lower than the BMR of 'fast' temperatebreeding non-Australian passerines. Old-endemics had substantially smaller clutches and longer maximal life spans in the wild than new arrivals, but neither clutch size nor maximum life span was correlated with BMR. Our results suggest that low BMR in tropical birds is not functionally linked to their 'slow pace of life' and instead may be a consequence of differences in annual thermal conditions experienced by tropical versus temperate species.

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A 'slow pace of life' in Australian old-endemic passerine birds is not accompanied by low

basal metabolic rates

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Abstract

Life history theory suggests that species experiencing high extrinsic mortality rates allocate more resources toward reproduction relative to self-maintenance and reach maturity earlier ('fast pace of life') than those having greater life expectancy and reproducing at a lower rate ('slow pace of life'). Among birds, many studies have shown that tropical species have a slower pace of life than temperate-breeding species. The pace of life has been hypothesized to affect metabolism and, as predicted, tropical birds have lower basal metabolic rates (BMR) than temperate-breeding birds. However, many temperate-breeding Australian passerines belong to lineages that evolved in Australia and share 'slow' life-history traits that are typical of tropical birds. We obtained BMR from 30 of these 'old-endemics' and 10 sympatric species of more recently arrived passerine lineages (derived from Afro-Asian origins or introduced by Europeans) with 'faster' life histories. The BMR of 'slow' temperate-breeding old-endemics was indistinguishable from that of new-arrivals and was not lower than the BMR of 'fast' temperatebreeding non-Australian passerines. Old-endemics had substantially smaller clutches and longer maximal life spans in the wild than new arrivals, but neither clutch size nor maximum life span was correlated with BMR. Our results suggest that low BMR in tropical birds is not functionally linked to their 'slow pace of life' and instead may be a consequence of differences in annual thermal conditions experienced by tropical versus temperate species.

Keywords Australian passerines; basal metabolic rate; pace-of-life, life history, tropical vs. temperate

Introduction

Biologists interpret life history traits as evolved solutions to ecological challenges that allow organisms to maximize their reproductive potential (Stearns 2000). Life history is shaped by intrinsic organismal properties (genetics, physiology, morphology, etc.) as well as by extrinsic ecological factors, both biotic (e.g., predation, competition) and abiotic (e.g., temperature, rainfall). Together with phylogenetic history, this complex selective matrix generates a strikingly diverse assemblage of phenotypes, but there are common features among animals that occupy particular regions of the life history continuum. For example, species with high rates of extrinsic mortality are selected to emphasize current reproduction over survival: they mature rapidly, reproduce early in life, show high fecundity, have low survival rates and reduced maximum lifespan potential (MLSP), and are said to live a 'fast pace of life' (Ricklefs & Wikelski 2002). By contrast, species with low rates of extrinsic mortality are selected to favor survival over current reproduction to maximize lifetime reproductive success, have slow maturation, low fecundity, high survival rates, and higher MLSP. Species with these life histories are said to have a 'slow pace of life' (Ricklefs & Wikelski 2002, Magrath et al. 2000).

The different emphasis on investing in fecundity versus survival as a function of the urgency of reproduction forms the basis of the physiology/life history nexus proposed by Ricklefs and Wikelski. They suggested that long-lived species should invest more in immune processes and other survival-promoting functions than species with short life spans, and some evidence indicates that immune responses do differ among birds and mammals with varying MSLP (Lee 2006, Prevital et al. 2012). They also predicted that long-lived and slow-paced animals should have relatively low basal metabolic rates. Basal metabolic rate (BMR) - the minimum energy cost of homeostatic existence in an endothermic animal - is probably rarely under direct selection (as by definition it is the cost of performing no growth, activity, or reproduction). However, it is mechanistically linked to traits that are likely to be 'seen' by selection, such as exercise or thermogenic capacity and field metabolic rate (the daily flux of energy through the animal; White

& Seymour 2004; Hulbert & Else 2004). The idea that BMR should be lower in slow-paced animals was supported by a common-garden experiment with stonechats (Saxicola torquata and S. rubicola) obtained as nestlings from populations with contrasting life histories (Wikelski et al. 2003). Adults from a sedentary tropical population had lower metabolic rates than birds from a migratory north-temperate population. Similarly, two extensive comparative analyses showed that birds from Neotropical forests in Panama and Peru have lower BMR (Wiersma et al. 2007b; Londoño et al. 2015) and, in the Panamanian birds, lower maximal metabolic rates than species from temperate latitudes (Wiersma et al. 2007a). Given that tropical birds have smaller clutches, longer parental care, and higher juvenile and adult survival than their north-temperate counterparts [Moreau 1944; Skutch 1985; Martin 1996; Cardillo 2002; Russell et al. 2004)], these studies support the notion that a slow 'pace of life' engenders a slow 'pace' of energy metabolism (e.g., Pearl 1928). However, it is unclear if low BMR in the tropics is a functional necessity or a byproduct of 'slow' life history, or instead is simply an outcome of living in warm, stable habitats with low requirements for thermoregulation, migration, or other energetically demanding activities (Wiersma et al. 2007a). One way to resolve this is to examine the metabolic rates of temperate-breeding birds sharing the same 'slow' life-history traits as tropical passerines, and compare them with sympatric passerines with 'fast' life-histories.

The Australian avifauna offers a good opportunity for such a test. Passerine species that breed in Australia include numerous 'old-endemic' species that evolved in relative isolation in Australia and New Guinea. A smaller number of 'new invaders' of Afro-Eurasian origin is thought to have colonized Australia much later. Additionally, a few north-temperate passerines were introduced by European settlers during the last two centuries (Rowley & Russell 1991; Keast 1981a). Of 338 passerine species that breed in Australia, 258 are old-endemics while 64 and 16 are 'new invaders' and introduced species, respectively (Keast 1981b). For simplicity we refer to the latter two groups as 'new-arrivals'.

Old-endemics and new-arrivals occur sympatrically in a variety of habitats, most of which are climatically temperate (warm summers, cool winters, large seasonal changes in rainfall and daylength). However, old-endemics have a 'slow pace of life,' with the small clutches, prolonged incubation and fledging care, and high survival rates that typify tropical birds (Magrath et al. 2000; Russell et al. 2004; Rowley & Russell 1991; Woinarski 1985; Yom-Tov 1987; Geffen & Yom-Tov 2000; Russell 2000). In contrast, new-arrivals have 'fast' life histories like those of temperate breeders from the Northern hemisphere. For example, clutch sizes average 2.7 in old-endemics and 4.7 in new-arrivals (Yom-Tov 1987). While the ecophysiology of Australian birds has received much attention (Astheimer & Buttemer 2002), no study has examined the energetics of Australian Passerines in the context of the 'slow-fast' contrast in their life histories. To determine if the BMR of old-endemics is indeed lower that that of new-arrivals, we measured BMR for 33 species from temperate climates, obtained BMR for several others from the literature, and used the combined dataset to compare BMR between 'slow' versus 'fast' life histories. To verify the postulated 'slow-fast' contrast in life history of our study species, we used published information on clutch size and maximum lifespan. To place our findings in a broader context, we compared our results to a global database of almost 500 temperate and tropical bird species (Londoño et al. 2015).

Material and methods

Bird capture and handling

We obtained most of the birds measured for this study near Wollongong, NSW (34.25°S, 153.53°E) or at the Fowlers Gap Research Station, NSW (31.09°S, 141.70°E). Wollongong-area birds were captured between October 1998 and August 1999 in urban areas, gardens, parks, and coastal eucalyptus woodland and temperate rainforest. Birds at Fowlers Gap were captured between September and November 2007 in eucalyptus and saltbush habitats. White-browed

Babblers were caught at Back Yamma State Forest, NSW (33.33°S, 148.21°E) in May 1998 and held in large flight cages at the University of Wollongong. Birds were mist-netted between 15:00 and 17:00 local time and transported to the laboratory in individual cloth bags for metabolic measurements that night. Prior to the start of measurements, birds were caged individually with access to water but not food. Birds were released the following morning close to the site of capture (white-browed babblers were returned to flight cages). Each individual was used only once for determination of BMR.

We obtained maximum life span data for free-living birds from the Australian Bird and Bat Banding Scheme database (www.environment.gov.au/science/bird-and-bat-banding; October 2015) and clutch sizes from published sources (Yom-Tov 1987; Higgins et al. 2001, 2006; Higgins and Peter 2002).

Basal metabolism

We measured metabolism as rates of oxygen uptake ($\dot{V}O_2$). BMR trials started at approximately 1800 (local time) and continued for 12-14 h. Up to four birds were tested concurrently in individual chambers constructed from paint cans (2 or 5 L) equipped with air flow ports, a thermocouple, and a perch. Chambers were placed in an environmental cabinet maintained at 31.0 \pm 1.0 °C, which we assumed to be within thermoneutrality for all species (in some species this was confirmed by measuring metabolism at several ambient temperatures). Separate mass flow controllers (Tylan; Bedford, Massachusetts, USA) maintained a constant flow of dry air to each chamber (500-2500 ml min⁻¹ STP depending on bird size). Mass flow controllers were calibrated by timing the rate of soap bubble travel in a precision burette, corrected to STP conditions. Excurrent air was subsampled at about 100 ml min⁻¹, scrubbed of CO₂ and water vapor, and routed to a two-channel O₂ analyzer (S3A-2; Applied Electrochemistry, Sunnyvale, California, USA) through a computer-controlled air stream

selector (Respirometry Multiplexer V2.0, Sable Systems, Las Vegas, Nevada, USA), such that two birds were measured simultaneously (one by each S3A channel). Birds were monitored for 42 minutes, followed by a 3-min reference reading, before switching to another individual. Accordingly, each bird was monitored approximately 47% of the time. The reference interval was sufficient for O_2 concentrations to reach stable, low values. Data were recorded on a Macintosh computer interfaced to a DataTaker 500 A/D converter (Data Electronics, Rowville, Victoria, Australia) with LabHelper software (www.warthog.ucr.edu). We computed $\dot{V}O_2$ as:

$$\dot{V}O_2$$
 (ml O₂ min⁻¹) = F * (FiO₂ – FeO₂)/(1 – FeO₂)

where F is flow rate (ml min⁻¹ STPD), FiO₂ is incurrent fractional O₂ concentration and FeO₂ is excurrent fractional O₂ concentration. BMR was calculated as the lowest $\dot{V}O_2$ averaged over a continuous 10-minute interval when $\dot{V}O_2$ was low and stable. We converted $\dot{V}O_2$ to Watts using an oxycaloric value of 19.8 kJ LO₂⁻¹. Birds were weighed just prior to being placed in the respirometer chamber in the evening and again in the morning upon retrieval from the chamber, with the mean of the two values used as body mass (M_b).

Literature and unpublished data

We measured BMR in 33 free-living Australian passerine species (24 old-endemics, four new invaders and five introduced species). In addition, we included published BMR for seven old-endemics and two new-arrivals. We used only data that conformed to requirements for true BMR (Mckecknie & Wolf 2005) and excluded data from long-term captives (white-browed babblers were held in large flight cages and were recent captures). When multiple measurements were available we used mean values, for a total of 40 species (Table 1).

BMR for additional species were obtained from Londoño et al. (2015). To avoid confounding analyses, we excluded the Australian species from that study that we had measured ourselves (N=9).

Statistics

Analyses are based on species means. Both M_b (in grams) and BMR (in Watts) were log_{10} -transformed to account for allometric scaling. Where necessary, a further Box-Cox transformation was applied to achieve normal distributions.

For 12 of the Australian species, data were available for only one or two individuals. Given these low sample sizes (N), we performed two sets of analyses, one including only species for which N > 2 (28 species) and the other including all 40 Australian species. There were very few difference between the two analyses. Accordingly, we show results derived from the complete dataset, except when there were qualitative differences in significance; these instances are noted in the Results section.

Phylogenetic signal (a measure of the statistical dependence among species' trait values due to their phylogenetic relationships; Blomberg et al. 2003) was estimated for BMR, M_b, and residuals of regressions between BMR and M_b (Revell 2010) with Pagel's lambda (Pagel 1999) using 'phytools' (Revell 2002). Since significant phylogenetic signal was present (see below), we made between-group comparisons with phylogenetic analyses performed in R (R Core Team 2013), based on trees derived from the avian 'tree of life' (Londoño et al. 2015;Jetz et al. 2012; Hackett et al. 2008; Fig. 1; Supplemental Fig. 1). Phylogenetic Independent Contrasts (PIC; Felsenstein 1985) were used to estimate correlations between BMR and M_b ('ape' package; Paradis et al. 2004). To compare BMR and M_b across origins (old-endemic and new-arrivals), orders, and geographic locations, we performed Phylogenetic Generalized Least Squares (PGLS) with 'nlme' (Pinheiro et al. 2013). We evaluated five models of evolution for the correlation structure (Londoño et al. 2015; Revell 2010; Blomberg et al. 2003; Grafen 1989; Martins & Hansen 1997). Akaike Information Criterion tests ('qpcR' [39]) indicated that Pagel's model was the best fit for Australian passerines and for temperate-tropical and passerine/non-passerine

comparisons. The Grafen model was the best fit for comparisons between Australian and non-Australian passerines. For life span comparisons between old-endemic and new-arrival Australian passerines, the Ornstein-Unlenbeck model was the best fit.

Since the use of phylogenetically-aware statistical models remains controversial (McNab 2015) we also performed conventional analyses (ordinary least squares; OLS), using JMP Pro 11 (SAS institute, Cary, North Carolina, USA).

Unless stated otherwise, all data are reported as mean \pm standard error.

Results

Tests revealed significant phylogenetic signal in M_b , BMR, and their residuals (P < 0.006 for all groups), and there were significant effects of M_b on BMR in all groups (Tables 2A and 2B).

Maximum life span

OLS ANCOVA of maximum free-living life span (body mass as covariate) revealed that the old-endemic species in our dataset have maximum life spans (MLS) 62% longer than those of new arrivals (13.8 \pm 0.8 years versus 8.5 \pm 1.5 years, respectively; $F_{1,37}$ = 10.0, P = 0.0031; Supplemental Fig. 1). Maximum life span increased with M_b in old endemics (t = 2.76, P = 0.010), but the relationship was not statistically significant in the smaller sample of new arrivals (P = 0.95). Results were similar if data for the species with the shortest MLS were discarded (*Eremiornis carteri*; 1 year, possibly an artifact of low sample size): old endemics live longer than new arrivals (13.8 \pm 0.8 years versus 9.4 \pm 1.4 years, respectively; $F_{1,36}$ = 6.9, P = 0.0125).

Phylogenetic analysis (PGLS) yielded qualitatively similar results: old endemics have maximum life spans longer than those of new arrivals ($F_{1,37}=6.66$, P=0.0139), and larger species have longer MLS than small birds ($F_{1,37}=4.90$, P=0.033).

Seasonal effects on BMR

Seasonal changes in avian BMR are well known (McKechnie & Swanson 2010) and could bias comparisons. Most (but not all) of our data were obtained in the spring and summer (September – March; Supplemental Table 1), as were the BMR data for northern hemisphere passerines (April-July; Wiersma et al. 2007). Capture dates (± 30 days for literature sources) were available for 37 of the 40 Australian species in our study. The remaining 3 species were sampled in the spring and summer (September – February; Vitali et al. 1999). For most species, all individuals were captured and tested within intervals of 60 days; for these we computed a mean capture date (day of the year). Seven species were captured in two discrete periods separated by more than 60 days; for these we computed mean values and capture dates for each of the two periods. Since capture date could not be normalized, we used nonparametric tests. For all 37 species for which we had dates, there was no significant influence of capture date on BMR (P = 0.119, Kendall's τ ; P = 0.135, Spearman's ρ). Similarly, in the seven species sampled within two distinct periods, there was no significant effect of capture date (P = 0.154, Kendall's τ ; P = 0.253, Spearman's ρ).

Australian passerine BMR

An OLS ANCOVA of the BMR of 40 Australian passerines including M_b , origin (old-endemic versus new arrival) and clutch size showed strong effects of M_b ($F_{1,38} = 262$, P < 0.0001) and origin ($F_{1,38} = 6.5$, P = 0.0155;), with old-endemics having about 12% higher BMR than new arrivals. There was also an origin X M_b interaction (F = 4.7, P = 0.0376), but no effect of clutch size on BMR. Removal of clutch size from the model did not change the qualitative results. Clutch size was independent of M_b ($F_{1,38} = 0.05$, P = .82) but was significantly higher in new arrivals than in old endemics ($3.62 \pm .21$ and $2.68 \pm .12$, respectively; F = 15.3, P = 0.0004).

There was no effect of maximum life span on BMR, after accounting for M_b and origin ($F_{3,36} = 0.3$, P = 0.58).

Results from phylogenetically-aware analyses were similar. Regressions describing PIC of M_b and PIC of BMR were not affected by origin ($F_{1,38} = 0.15$, P = 0.70; Group-PIC M_b interaction = 0.324 in a GLM; Fig. 2). PGLS revealed a strong dependence of BMR on M_b ($F_{1,38} = 448.3$, P < 0.0001), but no M_b X origin interaction ($F_{1,38} = 0.15$, P = 0.70). Clutch size was independent of M_b and BMR (P > .75 for both) but was significantly higher in new-arrivals than in old-endemics (3.61 ± 0.20 and 2.68 ± 0.11 respectively, P = 0.0011). Maximum life span did not significantly affect BMR after accounting for M_b and origin (PGLS; $F_{1,36} = 0.091$, P = 0.76).

Comparison of Australian to Northern Hemisphere Temperate-breeding Passerines

In an OLS ANCOVA of 129 temperate-breeding passerines (40 Australian plus 89 Northern Hemisphere species), M_b was strongly correlated to BMR ($F_{2,126}$ = 444, P < 0.0001), origin was also significant ($F_{2,126}$ = 5.10, P = 0.0074) and there was a significant origin X M_b interaction ($F_{2,126}$ = 5.23, P = 0.0066). Post-hoc comparisons (Tukey-Kramer) showed no difference between new arrivals and either north-temperate breeders (P = .77) or old-endemics (P = 0.064), but 11% higher BMR in old-endemics than in north-temperate breeders (P = 0.0095).

In phylogenetically-aware tests, there was no M_b X origin interaction (PGLS; $F_{2,126}$ = 1.66, P = .20). As expected, BMR and M_b were strongly correlated ($F_{2,126}$ = 624, P < 0.0001), with a slope (mass exponent) of 0.71 ± 0.04 (SE). However, BMR differed between Australian and north-temperate passerines ($F_{2,126}$ = 7.73, P < 0.0001), with BMR of Australian species averaging 7.7% lower. Most of the difference stemmed from new-arrivals, which had BMR 21.5 % less than that of north-temperate passerines, whereas the BMR of old-endemics was only 3.1 % lower. When analysis was restricted to the 28 Australian species for which sample size

was greater than 2 individuals, there was no effect of origin (Australian vs. north-temperate) on BMR ($F_{2.114} = 2.67$, P = 0.105).

Temperate/tropical and passerine/non-passerine comparisons

In 519 bird species (40 Australian passerines plus data from 5 datasets in Londoño et al. 2015), OLS ANCOVA found significantly lower BMR in tropical than in temperate species $(F_{1,517}=9.36;\ P=0.0023)$, significantly higher BMR in passerines than in non-passerines, a significant effect of dataset $(F_{5,513}=9.68,\ P<0.0001)$, a significant dataset X M_b interaction $(F_{5,513}=6.90,\ P=0.0001)$, but no M_b X latitude or M_b X order interactions $(F_{1,517}=1.60,\ P=0.21$ and $F=1.29,\ P=0.26$, respectively). As expected, BMR was strongly correlated with M_b $(F_{1,517}=2049,\ P<0.0001)$ with a slope (mass exponent) of 0.678 ± 0.015 (SE). On average, the BMR of temperate species was 12% higher than that of tropical species and the BMR of passerine birds averaged 19.5% higher than that of non-passerines.

Phylogenetically-aware tests (PGLS) also indicated that tropical birds have lower BMR than temperate birds ($F_{1,517} = 17.53$, P < 0.0001) and that Passeriformes have higher BMR than non-Passeriformes ($F_{1,517} = 86.02$, P < 0.0001) with a strongly positive overall correlation between M_b and BMR ($F_{1,517} = 2368$, P < 0.0001). There were significant interactions between M_b and taxonomic order ($F_{1,517} = 8.54$, P = 0.0038), latitude ($F_{1,517} = 15.58$, P = 0.0001) and dataset ($F_{5,513} = 3.69$, P = 0.0013). Restricting the PGLS analysis to the 28 Australian species for which N > 2 resulted in similar overall findings, although there was no Mb X taxonomic order interaction ($F_{1,505} = 0.02$, P = 0.88).

Discussion

The addition of our findings for Australian species does not change the conclusions of analyses showing lower BMR in tropical-breeding birds than in temperate-breeding species (Wiersma et al. 2007b; Londoño et al. 2015) and higher BMR in passerines than in non-passerine

orders (Londoño et al. 2015). However, in the context of metabolism and life history, our most interesting findings are that the BMR of temperate-zone, old-endemic Australian passerines is indistinguishable from that of new-arrivals inhabiting the same region, and that BMR is not affected by clutch size or by maximum life span. This lack of differentiation was unexpected, as both old-endemics and tropical forest passerines have a 'slow pace of life' but new-arrivals are 'faster.' In the Australian species we sampled, clutch size was 35% higher in new-arrivals than in old endemics, and for a much broader sampling of Australian passerines it was >70% higher in new-arrivals (Yom-Tov 1987). Maximum life span was 62% longer in the old endemics in our sample, also consistent with this group having a 'slow' life history. If life history and energy metabolism were coupled, then BMR and clutch size should be positively correlated whereas BMR and MLS should be negatively correlated after accounting for body mass, and, in general, old-endemics should have low BMR compared to new-arrivals. None of these predictions was supported by our data.

Our comparison of Australian passerines to north-temperate passerines also did not reveal low BMR in the old-endemics (in fact, it may be higher than that of 'fast' north-temperate passerines, as suggested by OLS post-hoc tests). Moreover, the new-arrivals have maintained similar clutch sizes in Australia as their conspecifics in Europe (Yom-Tov 1987), again showing lack of correlation between BMR and reproductive 'pace of life'.

Seasonal variation in some bird species, particularly those from high latitudes, is well documented (e.g., Mckechnie & Swanson 2010), so if our comparisons of old endemics, new arrivals, and north-temperate passerines were based on data obtained in contrasting seasons, results could be biased. Many of the north-temperate species were sampled in spring and summer (April –July; Wiersma et al. 2007b), and most of the Australian samples were obtained in the same season (September –February). The Australia data also contained some samples obtained outside that period, but we found no significant effect of sampling date on BMR, either

for all Australian species combined or within a subset of species with data spanning several months. A previous study of superb fairy-wrens (Lill et al. 2006) also found no effect of season on BMR. Therefore we do not believe our intergroup comparisons were confounded by sampling in contrasting seasons.

Of the old-endemics we sampled, only the Pomatostomidae (babblers), which are relatively weak flyers with clutch sizes averaging 3 eggs, had consistently low BMR (Table 1). The two old-endemic families with the highest BMR included the one with the lowest average clutch size (honeyeaters; Meliphagidae; 2.3 eggs) and the Pardalotidae (pardalotes, scrubwrens, thornbills), which averaged 3 eggs per clutch (Table 1). While clutch size is only one minor component of total reproductive effort in birds, it is a fundamental life history trait and the basis of other, larger investments in fecundity (e.g., incubation and especially parental care). The independence of clutch size and BMR in Australian passerines suggest that the difference in BMR between temperate and Neotropical birds found by Wiersma et al. (2007b) and Londoño et al. (2015) may not result from a linkage between the reproductive 'pace of life' and metabolism. Instead, it may derive from latitudinal contrasts in temperature or other factors separate from life history *per se* (Wiersma et al. 2007a).

At variance with that suggestion is the lack of effect of a 2.6 km altitude gradient on BMR in Andean birds in Peru, despite a 12 °C difference in environmental temperature between lowland and highland habitats (annual means 23 °C and 11 °C, respectively; Londoño et al. 2015). Such temperature differences should strongly affect rates of energy use for thermoregulation, especially for small birds, which typically have lower critical temperatures of 25-28 °C (unpublished data). However, some recent analyses indicate that variation in BMR across bird species is best explained not only by average ambient temperature, but also by its annual variation (White et al. 2007; Jetz et al. 2008). Hence, seasonal stability, despite differing average temperatures, may explain the across-altitude similarity of BMR among Peruvian species

(Londoño et al. 2015); in contrast, the large seasonal variation in temperate latitudes may engender high BMR. Birds expend more energy thermoregulating in cool habitats than in warmer regions and hence may have greater demand on muscular thermogenesis and digestive functions. Wiersma and colleagues found that Panamanian birds have lower maximal metabolic rates than temperate birds during forced exercise and shivering thermogenesis, as well as lower BMR (Wiersma et al. 2007a). They suggested that the reduced aerobic capacity might derive from lower demands for flight performance in tropical forests, where most birds are non-migratory and many are relatively sedentary; this, as well as low BMR (Vezina et al. 2006; Daan et al. 1990; Chappell et al. 1999) is consistent with data in tropical species showing relatively small organ masses (pectoral muscle, heart, lung, liver, kidney, liver, pancreas; Wiersma et al. 2012).

Overall, we found that BMR and two indices of the 'pace of life' -- reproductive investment (indicated by clutch size) and maximum life span -- are not linked in Australian passerines. Our findings are more consistent with the idea that low BMR in tropical birds is a consequence of living in a benign and relatively aseasonal climate rather than to having a 'slow' set of life-history traits, as predicted by the 'physiology-life history nexus' concept (Ricklefs and Wikelski 2002). To further test this interpretation, characterization of BMR in Australian oldendemic species living in temperate versus tropical regions is an obvious opportunity.

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Figures

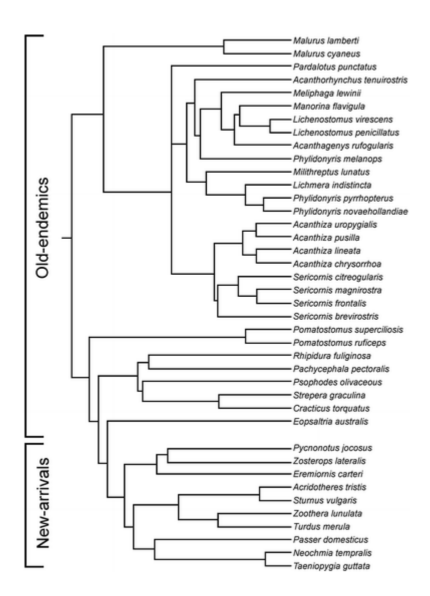


Figure 1. Phylogenetic tree for Australian passerines measured in this study. See Supplemental Figure 1 for a tree including all 519 species.

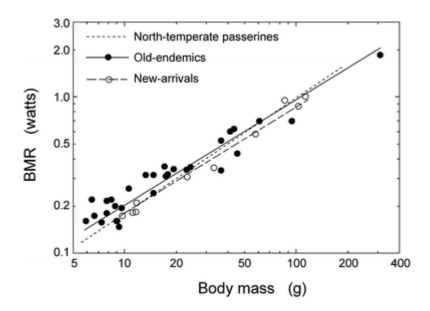


Figure 2. Phylogenetically independent contrast (PIC) of BMR against PIC of body mass for 40 species of Australian passerines (old-endemics and new-arrivals) and 89 species of temperate-breeding passerines from the Northern Hemisphere (individual data points not shown for clarity). The latter was based on data in Wiersma et al. (2007b) and Londoño et al. (2015). See text for statistics.

Table 1Body mass and BMR (in ml $O_2 min^{-1}$) in Australian old and new endemic passerine birds (n = number of individuals)

Species	Common name	Group ^a	n	Mass (g)	BMR	MLS	Clutch	Referencesb
Maluridae		•		•	-		•	
Malurus cyaneus	Superb fairy- wren	Old	10	8.7 ± 0.5	0.53 ± 0.08	10.4	3.1	This study (Woll)
Malurus cyaneus	Superb fairy- wren	Old		9.2	0.45			Lill et al. (<u>2006</u>) ^c
Malurus lamberti	Variegated fairy-wren	Old	7	7.8 ± 0.4	0.55 ± 0.03	11	2.8	This study (FG)
Meliphagidae	-	•		•	-		•	
Acanthagenys rufogularis	Spiny- cheeked honeyeater	Old	7	43.4 ± 4.4	1.90 ± 0.21	13.6	2.2	This study (FG)
Manorina flavigula	Yellow- throated miner	Old	5	41.1 ± 3.7	1.83 ± 0.22	12.1	3.2	This study (FG)
Meliphaga lewinii	Lewin's honeyeater	Old	7	36.2 ± 3.0	1.60 ± 0.10	15.9	2.2	This study (Woll)
Lichenostomus virescens	Singing honeyeater	Old	12	25.0	1.03	12	2.2	Collins et al. (1980)
Lichenostomus virescens	Singing honeyeater	Old	4	20.7 ± 0.8	1.06 ± 0.05			This study (FG)
Lichenostomus penicillatus	White- plumed honeyeater	Old	9	17.7 ± 1.3	0.97 ± 0.06	15.9	2.3	Buttemer and Astheimer (2000)
Melithreptus lunatus	White-naped honeyeater	Old	11	14.3 ± 0.6	0.74 ± 0.01	12.3	2.7	Vitali et al. (<u>1999</u>)
Lichmera indistincta	Brown honeyeater	Old	15	9.0	0.55	13.2	1.9	Collins et al. (1980)
Lichmera indistincta	Brown honeyeater	Old	6	10.1 ± 0.7	0.63 ± 0.03		2	Vitali et al. (1999)
Phylidonyris pyrrhoptera	Crescent honeyeater	Old		14.6	0.96	10.9	2.1	Weathers et al. (1996)
Phylidonyris novaehollandiae	New Holland honeyeater	Old		17.3	0.95	14.4	2.2	Weathers et al. (<u>1996</u>)
Phylidonyris melanops	Tawny- crowned honeyeater	Old	13	18.8 ± 0.7	0.87 ± 0.03	4.9	2.0	Vitali et al. (<u>1999</u>)
Acanthorhynchus tenuirostris	Eastern spinebill	Old	7	11.3 ± 0.9	0.83 ± 0.07	15.5	2.2	This study (Woll)
Acanthorhynchus tenuirostris	Eastern spinebill	Old		9.7	0.74			Weathers et al. (<u>1996</u>)
Pardalotidae								

Species	Common name	Group	n	Mass (g)	BMR	MLS	Clutch	Referencesb
Pardalotus punctatus	Spotted pardalote	Old	2	7.8	0.66	4	3–4	This study (Woll)
Sericornis citreogularis	Yellow- throated scrubwren	Old	3	16.9 ± 0.5	1.09 ± 0.08	15.6	2.5	This study (Woll)
Sericornis frontalis	White- browed scrubwren	Old	11	13.2 ± 0.8	0.96 ± 0.10	17.6	2.8	This study (Woll)
Sericornis magnirostris	Large-billed scrubwren	Old	1	8.7	0.61	15.3	3	This study (Woll)
Smicrornis brevirostris	Weebill	Old	1	5.9	0.49	7.4	2.6	This study (FG)
Acanthiza pusilla	Brown thornbill	Old	6	6.4 ± 0.7	0.67 ± 0.07	17.7	2.9	This study (Woll)
Acanthiza uropygialis	Chestnut- rumped thornbill	Old	2	6.6	0.53	12.7	3.3	This study (FG)
Acanthiza chrysorrhoa	Yellow- rumped thornbill	Old	8	8.3 ± 0.6	0.67 ± 0.07	13.1	3.2	This study (FG)
Acanthiza lineate	Striated thornbill	Old	1	7.3	0.48	23.5	2.9	This study (Woll)
Pomatostomidae								
Pomastostomus superciliosus	White- browed babbler	Old	4	36.1 ± 0.7	1.03 ± 0.06	14.4	2.6	This study (Woll)
Pomastostomus ruficeps	Chestnut- crowned babbler	Old	8	45.4 ± 2.5	1.32 ± 0.11	8.3	ca. 4	This study (FG)
Petroicidae								
Eopsaltria australis	Eastern yellow robin	Old	7	19.2 ± 2.3	1.05 ± 0.10	14.6	2.3	This study (Woll)
Cinclostomatidae								
Psophodes olivaceus	Eastern whipbird	Old	1	61.0	2.12	12.3	2–3	This study (Woll)
Pachycephalidae								
Pachycephala pectoralis	Golden whistler	Old	4	24.0 ± 0.3	1.08 ± 0.06	18	2.2	This study (Woll)
Dicruridae								
Rhipidura fuliginosa	Grey fantail	Old	1	8.9	0.49	9.7	2.7	This study (Woll)
Artamidae								
Cracticus torquatus	Grey butcherbird	Old	2	93.7	2.13	20.2	3.2	This study (Woll)
Strepara graculina	Pied currawong	Old	1	306.7	5.65	24.1	3.1	This study (Woll)

Species	Common name	Group ^a	n	Mass (g)	BMR	MLS	Clutch	References ^b
Passeridae				•	•		•	
Passer domesticus	House sparrow	New	35	23.0 ± 2.0	0.94 ± 0.11	7.5	4.1	This study (Woll) ^d
Taeniopygia guttata	Zebra finch	New	8	11.7 ± 0.6	0.64 ± 0.05	4.2	4.4	This study (FG)
Neochmia temporalis	Red-browed firetail	New	2	9.6	0.53	7.4	5	This study (Woll)
Pycnonotidae								
Pygnonotus jacosus	Red- whiskered bulbul	New	2	32.9	1.07	11.1	3.2	This study (Woll)
Sylviidae								
Eremiornis carteri	Spinifexbird	New	23	11.6 ± 0.2	0.56 ± 0.02	1	2–3	Ambrose et al. (<u>1996</u>)
Zosteropidae								
Zosterops lateralis	Silvereye	New	6	10.4 ± 0.4	0.63 ± 0.04	18.7	2.8	This study (Woll)
Zosterops lateralis	Silvereye	New	9	11.8 ± 1.5	0.48 ± 0.08			Maddocks and Geiser (1997)
Muscicapidae								
Zoothera lunulata	Bassian thrush	New	2	113.2	3.07	3.4	2–2.5	This study (Woll)
Turdidae								
Turdus merula	Eurasian blackbird	New	3	85.1 ± 2.74	2.88 ± 0.03	9.4	3.5	This study (Woll)
Sturnidae								
Sturnus vulgaris	Common starling	New	8	57.5 ± 3.1	1.76 ± 0.23	14.1	4.6	This study (Woll)
Acridotheres tristis	Common myna	New	4	102.3 ± 2.9	2.65 ± 0.25	12.2	3.7	This study (Woll)

Clutch sizes (egg numbers) are averages or ranges (Yom-Tov 1987; Higgins et al. 2001, 2006; Higgins and Peter 2002). Maximum life span (MLS; in years) is from Australian Bird and Bat Banding Scheme data (http://www.environment.gov.au/science/bird-and-bat-banding)

^a'Old' indicates old-endemics, while 'new' includes both new invaders and introduced species

^bWoll = Wollongong, FG = Fowlers Gap

^cMean values for three seasons (winter, spring and summer; no seasonal differences were found)

^dIn part published in Chappell et al. (<u>1999</u>)

Table 2

Allometric equations for BMR in several groups of birds, based on ordinary least squares regression, for the relationship between body mass and BMR, where BMR (Watts) = $a \times M_b^b$ (in g)

Group	N	а	b	r^2
Old-endemic	30	0.0547	0.602 ± 0.033	0.92
New-arrivals	10	0.0327	0.722 ± 0.030	0.98
North-temperate passerines	89	0.0329	0.739 ± 0.036	0.83
Temperate breeders ^a	203	0.0425	0.644 ± 0.012	0.94
Tropical breeders ^a	316	0.0482	0.592 ± 0.014	0.85
Passerines ^a	376	0.0438	0.637 ± 0.017	0.79
Non-passerine orders ^a	143	0.0351	0.659 ± 0.015	0.93

Slope (b) is shown \pm SE. In all cases, F > 200 and P < 0.0001

^aData from Londoño et al. (2015)

Table 3 Regression of phylogenetic independent contrasts (PIC) of body mass versus PIC of BMR, where PIC BMR = $a + b \times PIC$ mass

Group	N	a	b	r^2
Old-endemics	30	0.0007	0.678 ± 0.07	0.82
New-arrivals	10	-0.004	0.757 ± 0.04	0.98
North-temperate passerines ^a	89	-0.002	0.737 ± 0.05	0.69
Temperate breeders ^a	203	-0.002	1.033 ± 0.04	0.75
Tropical breeders ^a	316	0.0005	0.981 ± 0.03	0.58
Passerines ^a	376	0.0002	0.876 ± 0.04	0.52
Non-passerine orders ^a	143	-0.0008	1.095 ± 0.05	0.76

Slope (b) is shown \pm SE. In all cases, F > 190 and P < 0.0001

^aData from Londoño et al. (<u>2015</u>)