Avian thermoregulation in the heat: efficient evaporative cooling allows for extreme heat tolerance in four southern Hemisphere columbids

Running title
Evaporative cooling in columbids

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Summary statement
Arid zone pigeons and doves dissipate heat primary through cutaneous evaporative pathways, which has minimal metabolic costs, is highly efficient, and allows for the defense of relatively a low T_b under conditions of extreme heat.
Abstract

Birds show phylogenetic variation in the relative importance of respiratory versus cutaneous evaporative cooling capacity. The consequences for heat tolerance and evaporative cooling capacity remain unclear. We measured evaporative water loss (EWL), resting metabolic rate (RMR) and body temperature ($T_b$) in four arid-zone columbids from southern Africa [Namaqua dove ($Oena capensis$, ~37 g), laughing dove ($Spilopelia senegalensis$, ~89 g) and Cape turtle dove ($Streptopelia capicola$, ~148 g)] and Australia [crested pigeon ($Ocyphaps lophotes$, ~186 g)] at air temperatures ($T_a$) of up to 62 °C. There was no clear relationship between body mass and maximum $T_a$ tolerated during acute heat exposure. Maximum $T_b$ at very highest $T_a$ was 43.1 ± 1.0 °C, 43.7 ± 0.8 °C, 44.7 ± 0.3 °C and 44.3 ± 0.8 °C in Namaqua doves, laughing doves, Cape turtle doves and crested pigeons, respectively. In all four species, RMR increased significantly at $T_a$ above thermoneutrality, but the increases were relatively modest with RMR at $T_a = 56 °C$, 32 %, 60 %, 99 % and 11 % higher, respectively, than at $T_a = 35 °C$. At the highest $T_a$ values reached, evaporative heat loss was equivalent to 466 %, 227 %, 230 % and 275 % of metabolic heat production. The maximum ratio of evaporative heat loss to metabolic production observed in Namaqua doves, 4.66, exceeds by a substantial margin previous values reported for birds. Our results support the notion that cutaneous evaporation provides a highly efficient mechanism of heat dissipation and an enhanced ability to tolerate extremely high $T_a$.

Introduction

The defense of a body temperature ($T_b$) set point below environmental temperature is possible only via evaporative heat dissipation, and animals that regularly experience extremely hot conditions rely heavily on this avenue of heat exchange to avoid lethal hyperthermia. The capacity for evaporative cooling is particularly pronounced in birds, reflecting their predominantly diurnal activity and limited use of thermally buffered underground microsites, even among species occupying the hottest regions on the planet (Dawson and Bartholomew, 1968; Dawson and Fisher, 1969; Dawson and Schmidt-Nielsen, 1964; Grant, 1982; Williams and Tieleman, 2005), but see also (Williams and Tieleman, 2005). A number of studies have shown that birds can maintain $T_b$ 10-20 °C below air temperature ($T_a$) during acute (Dawson and Fisher, 1969; Tieleman et al., 2002b; Whitfield et al., 2015; Wolf and Walsberg, 1996) and
chronic (Marder and Arieli, 1988) heat exposure, via rapid increases in the rate of evaporative water loss (EWL) when $T_a$ exceeds normothermic $T_b$. Under such conditions, a trade-off arises between the avoidance of hyperthermic $T_b$ and the avoidance of dehydration resulting from rapid depletion of body water; at $T_a$s approaching 50 °C, rates of EWL in very small birds may be equivalent to ~7 % of body mass per hour (Wolf and Walsberg, 1996).

There are two major physiological mechanisms whereby birds increase rates of evaporative heat loss during acute heat exposure. The first involves accelerating heat dissipation across the surfaces of the respiratory tract via increases in respiration frequency and decreases in tidal volume (panting) and/or rapid vibration of gular membranes (gular flutter; Dawson, 1982). The second involves increases in the rate of trans-cutaneous evaporation (Marder and Arieli, 1988; Webster and Bernstein, 1987), a process regulated over short time scales by adjustments to peripheral microcirculation (Ophir et al., 2002), and over longer time scales by changes in epidermal lipid composition (Haugen et al., 2003; Menon et al., 1989; Menon et al., 1988; Muñoz-Garcia et al., 2008). The relative contributions of respiratory and cutaneous evaporative water loss (REWL and CEWL, respectively) to evaporative heat dissipation at high $T_a$ vary phylogenetically, with the data currently available indicating that REWL predominates in the Passeriformes (Tieleman and Williams, 2002; Wolf and Walsberg, 1996) and Galliformes (Bouverot et al., 1974; Richards, 1976). In contrast, CEWL represents the major avenue of heat dissipation at high $T_a$ in Columbiformes (Hoffman and Walsberg, 1999; Marder and Arieli, 1988; McKechnie and Wolf, 2004; Smith and Suthers, 1969; Webster and Bernstein, 1987; Withers and Williams, 1990). The partitioning of total evaporative water loss into CEWL and REWL in columbids shows some phenotypic flexibility, with CEWL/REW ratios typically higher in individuals acclimated to hot conditions (Marder and Arieli, 1988; McKechnie and Wolf, 2004; Ophir et al., 2003).

The available data also suggest that birds rely on increases in either REWL or CEWL, but generally not both, as the primary avenue of evaporative heat loss at $T_a$ above normothermic $T_b$ (Wolf and Walsberg, 1996). However, the ecological and evolutionary significance of these different modes of heat dissipation remains unclear. Respiratory evaporation appears to be a less energetically efficient mode of heat dissipation compared to cutaneous evaporation, on account of the muscle activity required for panting and consequent metabolic heat production. In dune larks
Mirafra erythrocephalus), for example, resting metabolic rate (RMR) increases by 100% between $T_a = 35$ and 48 °C (Williams, 1999). Experimental evidence for the notion that CEWL is more energetically efficient is provided by heat-acclimated western white-winged doves (Zenaida asiatica mearnsii) which increased CEWL compared to conspecifics acclimated to cooler conditions; both $T_h$ and resting metabolic rate (RMR) at $T_a = 45$ °C were significantly lower than in cool-acclimated individuals (McKechnie and Wolf, 2004). Similarly, RMR at $T_a > T_b$ was lower in heat-acclimated rock doves (Columba livia) with a greater fraction of evaporation occurring cutaneously (Marder and Arieli, 1988). We also note that increased ventilatory rates increase heat gain from the environment when $T_a > T_b$. On the other hand, a possible disadvantage of reliance on CEWL was highlighted by a recent study evaluating the effect of humidity on evaporative heat loss in hot conditions. Evaporative cooling at $T_a > T_b$ in the sociable weaver (Philetairus socius), a passerine that relies heavily on panting, was less sensitive to elevated humidity compared to the Namaqua dove (Oena capensis), a columbid that relies primarily on cutaneous evaporation (Gerson et al., 2014).

To explore further the implications of reliance on either REWL or CEWL as the primary mode of heat dissipation for evaporative cooling and heat tolerance, we determined the upper thermoregulatory limits and maximum evaporative cooling capacities of four species of columbids varying approximately 5-fold in body mass ($M_b$). We predicted that, during acute exposure to $T_a > T_b$, columbids would: a) exhibit relatively gradual increases in EWL, RMR and $T_h$ with increasing $T_a$, and b) are able to tolerate higher maximum $T_a$ before becoming hyperthermic compared to passerines and other taxa that rely primarily on respiratory pathways for evaporative cooling. We did not directly compare thermoregulatory variables between columbids and passerines, because the lack of overlap in the $M_b$ ranges of the members of these two taxa for which data are currently available potentially confounds such comparisons.

**Materials and methods**

**Study sites and species**
The southern African component of the study took place at the same study sites and during the same periods as described by Whitfield et al. (2015). We measured EWL,
RMR and $T_b$ over a range of $T_a$ in three species from the family Columbidae, namely Namaqua dove (*Oena capensis* Linnaeus, ~40 g), laughing dove (*Spilopelia senegalensis* Linnaeus, ~100 g; formerly *Streptopelia senegalensis*) and Cape turtle dove (*Streptopelia capicola* Sundevall, ~153 g) (Hockey et al. 2005). All three species are granivorous, occur widely throughout sub-Saharan Africa in almost all habitats except forests, and are common year-round in the Kalahari desert, although their numbers decrease during dry periods (Hockey et al. 2005). The Australian component of the study involved data collection for crested pigeons (*Ocyphaps lophotes* Temminck 1822) in Gluepot Reserve, South Australia (S 33°46’ E 140°07’).

Birds were captured using Japanese mist nets at various times of the day, and initially held in cloth bags. All birds used in the study were adults and appeared to be in good condition. The mean body masses of *O. capensis* *S. senegalensis*, and *S. capicola* were 37.1 ± 3.2 g (mean ± SD; n = 29), 89.4 ± 13.0 g (n = 33) and 147.5 ± 17.6 g (n = 26), respectively. The mean body mass of *Ocyphaps lophotes* was 186.5 ± 16.5 g (n = 39).

All experimental procedures were approved by the Animal Ethics Committees of the University of Pretoria (protocol EC071-11) and the University of Adelaide (S-2013-151A), and the Institutional Animal Care and Use Committee of the University of New Mexico (12-1005370-MCC). Birds were captured under permits issued by the Northern Cape Department of Environmental Affairs (ODB 008/2013) and the Department of Environment, Water and Natural Resources South Australia (E26141-2).

**Gas exchange and temperature measurements**

Measurements of EWL, carbon dioxide production ({$\dot{V}_{CO_2}$}), $T_a$ and $T_b$ were conducted using the same general methods and experimental setup as described by (Whitfield et al., 2015). Birds were placed individually in sealable plastic chambers with volumes of 4 L (*O. capensis*) or 9 L (*S. senegalensis*, *S. capicola* and *Ocyphaps lophotes*). Depending on $T_a$ and the $M_b$ of the bird, flow rates ranging from 6 - 85 L min$^{-1}$ were used. Birds tended to remain calmer when flow rates were higher and chamber humidities were lower. As was the case in the study by Whitfield et al. (2015), the high flow rates we used meant that fractional depletion of oxygen within the chamber was below the resolution of the oxygen analyser we used (FC-10A, Sable Systems,
Las Vegas NV, USA), and oxygen consumption ($\dot{V}_{O_2}$) could therefore not be measured accurately. Core $T_b$ was measured during experiments using a temperature sensitive passive integrated transponder (PIT) tag injected into the abdominal cavity of each bird, and a PIT tag reader and portable transceiver system, following Whitfield et al. (2015).

Before each dove was placed in a respirometry chamber, we palpated its crop to determine the presence or absence of recently ingested food. We could not be certain whether or not birds were postabsorptive, and the lack of $\dot{V}_{O_2}$ measurements precluded the calculation of respiratory exchange ratio (i.e., $\dot{V}_{CO_2}/\dot{V}_{O_2}$) and hence inference of the metabolic substrate (Walsberg and Wolf 1995). For this reason, we converted all measurements of $\dot{V}_{CO_2}$ to metabolic rate (Watts) assuming RER = 0.85 (i.e., a mix of carbohydrate and lipid metabolism; Walsberg and Wolf 1995), using a thermal equivalence value of 24.4 J mL$^{-1}$ CO$_2$ (Withers, 1992).

**Experimental protocol and data analyses**

We used the same experimental protocol during which birds were exposed to progressively higher $T_a$ in a stepped profile as described by Whitfield et al. (2015) and Smith et al. (2015) and analysed our data in the same way as in the latter studies. Birds were exposed to $T_a$s of 25–40 °C in 5 °C increments, and $T_a$s of 40–62 °C in 2 °C increments. Birds were continuously monitored while in the chamber using a video camera and infrared light source. Birds were removed from chambers when they reached their heat tolerance limit, defined by one of two events: a) escape behavior sustained for more than 5-10 s, or b) thermal endpoint (i.e., extreme heat stress manifested as a loss of coordination or balance, and/or a rapid increase in $T_b$ to > 45°C; Whitfield et al. 2015). Data for active birds hence correspond with event a) and data for calm birds with event b). Thermal endpoints were taken as $T_a$ values associated with a loss of coordination or balance, sudden decreases in EWL and RMR, and/or uncontrolled increases in $T_b$ to values exceeding 45 °C (Whitfield et al., 2015). In the present study, $T_b$ in birds at their heat tolerance limits (i.e, either escape behaviour or thermal endpoint) was generally consistent with values observed in two columbids and a quail by Smith et al. (2015); in all cases $T_b$ exceeded 45 °C and/or increased at a rate of > 0.1°C min$^{-1}$ during the last 5 min of measurements. The $T_a$
associated with the onset of gular flutter for each bird in the chamber was also recorded.

All results are reported as whole-animal values and expressed as mean ± SD for calm birds only (i.e., event b above), unless otherwise stated. Mean \( T_B \) is the average across the last ~ten minutes at a given \( T_a \), whereas \( T_{B,\text{max}} \) is the single highest recorded \( T_B \) within the same ten minute period. Rates of evaporative water loss were converted to rates of heat loss using a latent heat of vaporisation of 2.41 J mg H\(_2\)O\(^{-1}\), corresponding to \( T_a = 40 \) °C (Tracy et al., 2010). In order to model relationships between EWL and high \( T_a \), we followed the approach of Whitfield et al. (2015) and fitted both segmented linear and 2\(^{nd}\)-order polynomial regression models to EWL vs \( T_a \) data, and compared Akaike Information Criterion (AIC) values in order to verify the validity of using segmented linear models for interspecific comparisons (McKechnie and Wolf, 2010). Segmented linear models provided a better fit for *S. senegalensis* (polynomial AIC = 592.5; linear segmented AIC = 590.7), *O. capensis* (polynomial AIC = 664.3; linear segmented AIC = 659.9) and *Ocyphaps lophotes* (polynomial AIC = 957.0; linear segmented AIC = 955.8), whereas a polynomial model provided a better fit for *S. capicola* (polynomial AIC = 468.5; linear segmented AIC = 472.1). Given the small differences in AIC values, we are confident that our approach of using segmented linear models to describe patterns of EWL and related variables at high \( T_a \) is justified, and use such models in all further analyses. We used generalized mixed-effect models with R package ‘nlme’ (Pinheiro et al., 2009) to determine the coefficients of EWL, RMR, EHL/MHP and \( T_B \) as a function of \( T_a \) above the respective \( T_a \) inflection points identified in the segmented analyses.

**Results**

**Namaqua doves**

In Namaqua doves, RMR decreased from 0.39 ± 0.07 W at 25 °C to 0.27 ± 0.03 W at 35 °C (Figure 1). At 56 °C and 60 °C, RMR was 0.35 ± 0.08 W and 0.34 ± 0.11 W, respectively (Figure 1). An inflection point in RMR occurred at \( T_a = 35.3 \) °C, above which RMR increased significantly with \( T_a \) (\( t_{1,52} = 2.557, P = 0.014 \)). Namaqua doves commenced gular fluttering at \( T_a = 55.1 ± 3.6 \) °C.
Figure 1. Resting metabolic rate (RMR) in calm Namaqua doves (*Oena capensis*; n=29), laughing doves (*Spilopelia senegalensis*; n=33), Cape turtle doves (*Streptopelia capicola*; n=26) and crested pigeons (*Ocyphaps lophotes*; n=39) as a function of air temperature (*T*<sub>a</sub>). Segmented regressions were used to estimate the inflection point in the relationship between RMR and *T*<sub>a</sub>, and the slopes and intercepts were calculated using linear mixed-effects models. Significant relationships are represented by asterisks: * = *P* < 0.05, ** = *P* < 0.01, *** = *P* < 0.001.

At *T*<sub>a</sub> < 40 °C, EWL was consistently low, averaging 0.15 ± 0.09 g h<sup>-1</sup> (Figure 2). Above the inflection at *T*<sub>a</sub> = 40.9 °C, EWL increased linearly and significantly (*t*<sub>1,55</sub> = 12.140, P < 0.001) by ~8-fold to 1.75 ± 0.38 g h<sup>-1</sup> at *T*<sub>a</sub> = 56 °C and 2.36 ± 0.69 g h<sup>-1</sup> at 60 °C (Figure 2). At these two *T*<sub>a</sub> values, rates of EWL were equivalent to 5.1-6.3 % of *M*<sub>b</sub> per hour. Over the same *T*<sub>a</sub> range, the slope of the relationship between mass-specific EWL and *T*<sub>a</sub> was 2.50 mg g<sup>-1</sup> h<sup>-1</sup> °C<sup>-1</sup>. The ratio of evaporative heat loss (EHL) of metabolic heat production (MHP) increased linearly and significantly from 0.30 ± 0.18 at *T*<sub>a</sub> < 40 °C to 3.41 ± 0.76 and 4.66 ± 0.42 at *T*<sub>a</sub> = 56 and 60 °C respectively (*t*<sub>1,40</sub> = 15.103, P < 0.001, Figure 3, Table 1).
Figure 2. Evaporative water loss (EWL) as a function of air temperature ($T_a$) in calm Namaqua doves (*Oena capensis*; n=29), laughing doves (*Spilopelia senegalensis*; n=33), Cape turtle doves (*Streptopelia capicola*; n=26) and crested pigeons (*Ocyphaps lophotes*; n=39) as a function of air temperatures ($T_a$). Segmented regressions were used to estimate the inflection point in the relationship between EWL and $T_a$, and the slopes and intercepts were calculated using linear mixed-effects models. Significant relationships are represented by asterisks: * = P < 0.05, ** = P < 0.01, *** = P < 0.001.

Table 1. Mean (± SD) maximum body temperature ($T_{b,max}$), rate of $T_b$ increase (~10 minute period), evaporative water loss (EWL), resting metabolic rate (RMR) and ratio of evaporative heat loss to metabolic heat production (EHL/MHP) in Namaqua doves, laughing doves, Cape turtle doves and crested pigeons at firstly, the highest air temperature ($T_a$) at which all species were tested (~56 °C). Only data from inactive birds were included.
Figure 3. Ratio of evaporative heat loss (EHL) to metabolic heat production (MHP) in calm Namaqua doves (*Oena capensis*; n=29), laughing doves (*Spilopelia senegalensis*; n=33), Cape turtle doves (*Streptopelia capicola*; n=26) and crested pigeons (*Ocyphaps lophotes*; n=39) as a function of air temperature (*T_a*).

Segmented regressions were used to estimate the inflection point in the relationship between EHL/MHP and *T_a* and the slopes and intercepts were calculated using linear mixed-effects models. For crested pigeons, the regression model for EHL/MHP vs *T_a* between 40 °C and 55.1 °C is provided in the figure; the solid line at *T_a* > 55 °C shows the non-significant model fitted above the inflection point at higher *T_a* values. Significant relationships are represented by asterisks: * = P < 0.05, ** = P < 0.01, *** = P < 0.001.

Mean *T_b* was 40.3 ± 0.9 °C at 25 °C < *T_a* < 35 °C, but increased significantly (*t_{1,64} = 9.211, p < 0.001*) at higher *T_a* (Figure 4). At *T_a* = 56 and 60 °C, mean *T_b* was 42.5 ± 0.6 °C and 43.1 ± 1.0 °C respectively (Figure 4). The relationship between *T_a* and the rate of *T_b* increase was not significant (*t_{1,61} = -0.40, p = 0.691*), with *T_b* either remaining approximately constant (rate of 0 °C min⁻¹) or increasing/decreasing slightly (rate typically between -0.1 and 0.1 °C min⁻¹).
Figure 4. Body temperature ($T_b$) in Namaqua doves (*Oena capensis*; $n=29$), laughing doves (*Spilopelia senegalensis*; $n=33$), Cape turtle doves (*Streptopelia capicola*; $n=26$) and crested pigeons (*Ocyphaps lophotes*; $n=39$) as a function of air temperature ($T_a$). Segmented regressions were used to estimate the inflection point in the relationship between mean $T_b$ and $T_a$ and the slopes and intercepts were calculated using linear mixed-effects models. Significant relationships are represented by asterisks: * = $P < 0.05$, ** = $P < 0.01$, *** = $P < 0.001$.

The maximum $T_a$ tolerated by Namaqua doves was 60 °C. All Namaqua doves at their heat tolerance limits were active (i.e., measurements were terminated because of activity), and therefore not included in the results above. The individuals that reached heat tolerance limits increased from zero at $T_a = 54$ °C to 39 % at $T_a = 60$ °C (Figure 5). In two of the birds that had reached their heat tolerance limits at 60 °C, mean $T_b$ was 44.2 and 44.5 °C and $T_{b\text{, max}}$ was 45.1 °C in both individuals.
Figure 5. Frequency of individuals that reached (black bars) and did not reach (white bars) heat tolerance limits (i.e., showed escape behaviour or reached thermal endpoints) during measurements at air temperatures ≥ 40°C for Namaqua doves (*Oena capensis*), laughing doves (*Spilopelia senegalensis*), Cape turtle doves (*Streptopelia capicola*) and crested pigeons (*Ocyphaps lophotes*).

** Laughing doves **

Calm laughing doves that reached their thermal endpoints at high *T*ₘs were included in the results below. There was a decrease in RMR from 0.89 ± 0.21 W at *T*ₘ = 25 °C to 0.56 ± 0.11 W at *T*ₘ = 40 °C (Figure 1). An inflection point in RMR occurred at *T*ₘ = 40.0 °C, above which RMR increased linearly and significantly (*t*₁,₃₈ = 6.637, *p* < 0.001; Figure 1), with RMR averaging 1.05 ± 0.27 W (Table 1), and 1.11 ± 0.36 W (Figure 1) at 56 °C and 58 °C respectively. Laughing doves began to gular flutter at *T*ₘ = 43.5 ± 2.3 °C.

At *T*ₘ < 40 °C, EWL was stable and low, averaging 0.46 ± 0.27 g h⁻¹ (Figure 2). Above 40 °C, EWL increased linearly and significantly (*t*₁,₃₈ = 13.642, *p* < 0.001) with increasing *T*ₘ by ~8-fold to 3.60 ± 1.17 g h⁻¹ (Table 1) and 3.65 ± 1.39 g h⁻¹ (Figure 2) at *T*ₘ = 56 and 58 °C respectively. At these higher *T*ₘs, however, a large amount of scatter in the EWL data was observed. The slope of the relationship between mass-specific EWL and *T*ₘ was 1.87 mg g⁻¹ h⁻¹ °C⁻¹. Above 35 °C, EHL/MHP
increased approximately linearly \((t_{1.38} = 8.247, p < 0.001)\) from \(0.62 \pm 0.11\) at \(35 ^\circ C\) to \(2.28 \pm 0.39\) at \(56 ^\circ C\) and \(2.27 \pm 0.76\) at \(58 ^\circ C\) (Figure 3). Though a linear regression was fitted for comparative purposes, EHL/MHP appeared to reach a plateau at higher \(T_a\)'s (Figure 3), suggesting that the relationship may be a nonlinear one.

Mean \(T_b\) averaged \(40.8 \pm 0.8 ^\circ C\) at \(T_a < 40 ^\circ C\), and increased linearly \((t_{1.28} = 8.14, p < 0.001)\) above this \(T_a\) (Figure 4). At \(T_a = 56\) and \(58 ^\circ C\), mean \(T_b\) was \(43.6 \pm 1.0 ^\circ C\) and \(43.7 \pm 0.8 ^\circ C\) (Figure 4) respectively. The was no significant relationship between \(T_a\) and the rate of \(T_b\) increase \((t_{1.15} = 1.67, p = 0.12)\), and considerable variation was observed above \(T_a = 56 ^\circ C\).

The maximum \(T_a\) tolerated by laughing doves was \(58 ^\circ C\), although one individual reached \(T_a = 60 ^\circ C\). The \(T_a\) at which individuals reached their heat tolerance limits was more variable than in the other three species, varying from \(44 ^\circ C\) to \(58 ^\circ C\) (Figure 5).

**Cape turtle doves**

In calm Cape turtle doves, RMR decreased from \(1.23 \pm 0.39\) W at \(T_a = 25 ^\circ C\) to \(0.81 \pm 0.16\) W at \(T_a = 35 ^\circ C\) (Figure 1). An inflection point occurred at \(T_a = 44.56 ^\circ C\), above which RMR increased linearly and significantly \((t_{1.26} = 7.219, p < 0.001)\) to a maximum of \(1.62 \pm 0.29\) W at \(T_a = 56 ^\circ C\) (Table 1, Figure 1). Cape turtle doves began to gular flutter at \(40.8 \pm 3.4 ^\circ C\).

At \(T_a < 40 ^\circ C\), EWL was stable and low, averaging \(0.48 \pm 0.21\) g h\(^{-1}\) (Figure 2). Above \(40 ^\circ C\), EWL increased approximately linearly with increasing \(T_a\) \((t_{1.26} = 9.379, p < 0.001)\). Water loss rates increased ~10-fold, averaging \(5.40 \pm 1.48\) g h\(^{-1}\) at \(T_a = 56 ^\circ C\) (Table 1). The slope of the relationship between mass-specific EWL and \(T_a\) was \(1.62\) mg g\(^{-1}\) h\(^{-1}\) °C\(^{-1}\). Above \(T_a = 35 ^\circ C\), EHL/MHP increased linearly \((t_{1.25} = 8.552, p < 0.001)\) from \(0.36 \pm 0.16\) at \(T_a = 35 ^\circ C\) to \(2.30 \pm 0.88\) at \(T_a = 56 ^\circ C\) (Figure 3). Average EHL/MHP increased only slightly at the higher \(T_a\)'s, but more linearly than in the other two southern African species (Figure 3).

Mean \(T_b\) averaged \(41.1 \pm 0.9 ^\circ C\) at \(T_a < 40 ^\circ C\), and increased linearly \((t_{1.25} = 13.63, p < 0.001)\) above this \(T_a\) (Figure 4). At \(T_a = 56\), mean \(T_b\) was \(44.7 \pm 0.3 ^\circ C\) (Table 1, Figure 4). The rate of \(T_b\) increase increased linearly \((t_{1.25} = 2.81, p < 0.01)\) with increasing \(T_a\), though considerable variation was observed.

The maximum \(T_a\) tolerated by Cape turtle doves was \(56 ^\circ C\). In most cases, measurements were terminated on account of activity, with just one individual
reaching its thermal endpoint at $T_a = 56 \, ^\circ C$ and included in the results above. The single individual that reached its thermal endpoint at 56 °C exhibited a mean $T_b$, mean $T_{b,\text{max}}$, rate of $T_b$ increase and EHL/MHP of 44.7 °C, 45.5 °C, 1.27 °C min$^{-1}$ and 1.62 respectively (Table 1).

**Crested pigeons**

Crested pigeons generally remained calm with little sign of agitation, and data from almost all individuals are included in the results below. Mass-specific RMR decreased from 1.52 ± 0.35 W at 30 °C to a minimum of 1.00 ± 0.11 W at 46 °C, with an inflection point at $T_a = 46.6 \, ^\circ C$ (Figure 1). At $T_a$ above the inflection point, RMR increased to 1.28 ± 0.45 W at $T_a = 56 \, ^\circ C$ and a maximum of 2.13 ± 0.95 W in four individuals that reached $T_a = 62 \, ^\circ C$ (Figure 1). The mixed model revealed a significant linear increase in RMR above the inflection point ($t_{1,46} = 4.36, p < 0.001$).

Crested pigeons commenced gular fluttering at $T_a = 48.0 ± 5.7 \, ^\circ C$. EWL remained low and stable at $T_a < 40 \, ^\circ C$, averaging 1.12 ± 0.46 g h$^{-1}$ (Figure 2). Above $T_a = 40.3 \, ^\circ C$, EWL increased linearly and significantly ($t_{1,89} = 19.211, p < 0.001$) by ~6-fold to 4.63 ± 1.59 g h$^{-1}$ at $T_a = 56 \, ^\circ C$ and reached a maximum of 7.26 ± 1.82 g h$^{-1}$ at $T_a = 62 \, ^\circ C$ (Figure 2). The slope of the relationship between mass-specific EWL and $T_a$ was 1.39 mg g$^{-1}$ h$^{-1}$ °C$^{-1}$. Minimum EHL/MHP was 0.51 ± 0.12 at $T_a = 38 \, ^\circ C$. EHL/MHP then increased linearly and significantly ($t_{1,70} = 23.032, p < 0.001$, Figure 3, Table 1) above a lower inflection point at $T_a = 38.8 \, ^\circ C$ to 2.57 ± 0.28 at $T_a = 54 \, ^\circ C$. A second inflection point then occurred at $T_a = 55.1 \, ^\circ C$, above which EHL/MHP was not significantly related to $T_a$ ($p = 0.95$; Figure 3). Values for EHL/MHP were 2.75 ± 0.54 and 2.40 ± 0.38 at $T_a = 60 \, ^\circ C$ and 62 °C respectively.

Mean $T_b$ was 40.6 ± 0.8 °C at $T_a < 40 \, ^\circ C$, but increased significantly ($t_{1,17} = 6.93, p < 0.001$) at higher $T_a$ (Figure 4). At $T_a = 60$ and 62 °C, mean $T_b$ was 44.3 ± 1.2 °C and 44.3 ± 0.8 °C respectively (Figure 4). The relationship between $T_a$ and the rate of $T_b$ increase was not significant ($t_{1,22} = 1.51, p = 0.15$), with $T_b$ either remaining approximately constant (rate of 0 °C min$^{-1}$) or increasing/decreasing slightly (rates typically between -0.1 and 0.1 °C min$^{-1}$).
Discussion
The picture that emerges from our data for four columbids is a lack of a clear relationship between $M_b$ and the maximum $T_a$ value tolerated during acute heat exposure. Although the three southern African species showed negative scaling, with the maximum $T_a$ tolerated during acute heat exposure being highest in the smallest species and *vice versa*, the larger Australian crested pigeon tolerated the highest $T_a$ values (60-62 °C) of any species in this study. A pattern of negative scaling would be expected if evaporative heat dissipation is limited by surface area / volume ratios, as would be predicted for birds that rely primarily on cutaneous evaporation at very high $T_a$.

The lack of a relationship between $M_b$ and maximum $T_a$ tolerated among these four columbids contrasts with the pattern among three ploceid passerines, where maximum $T_a$ reached scaled positively with $M_b$, ranging from ~48 °C in the 10-g scaly-feathered weaver to ~54 °C in the 40-g white-browed sparrow-weaver (Whitfield et al., 2015). Notwithstanding the small sample sizes involved (3-4 species per study) and the limited overlap in $M_b$ ranges, these data raise the possibility that the scaling of avian heat tolerance and evaporative cooling capacity during acute heat exposure may differ fundamentally depending on the primary mode of evaporative heat dissipation.

Body temperatures and thermal endpoints
The columbids investigated here were able to maintain $T_b$ at sub-lethal levels even at $T_a \approx 56$-62 °C during acute heat exposure, and showed more gradual increases in $T_b$ at high $T_a$ than were observed in the ploceids (Whitfield et al., 2015). The maximum $T_b$ values we recorded (both at $T_a = 60$ °C) were 45.8 °C in a Namaqua dove and 45.9 °C in a crested pigeon, values within the known avian lethal $T_b$ range (45.7 – 47.8 °C; (Arad and Marder, 1982; Brush, 1965; Dmi'el and Tel-Tzur, 1985; Randall, 1943). We did not, however, observe a loss of coordinated movement associated with high $T_b$ in the doves, which contrasts with the passerines in our previous study (Whitfield et al., 2015). The latter observation suggests lethal $T_b$ values in these four columbids are nearer the upper end of the range reported in the literature.

Thermal endpoints (i.e., $T_b$ and $T_a$ values associated with thermoregulatory failure in birds not displaying escape behaviour) were not as clear in the columbids investigated here as in the ploceid passerines Whitfield et al. (2015) examined.
previously. Whereas in the latter study 100% of white-browed sparrow-weaver and sociable weaver individuals, and 60% of scaly-feathered weaver individuals reached thermal endpoints, in the present study at no \( T_a \) did > 50% of individuals of any species reach thermal endpoints. The fact that we could not determine clear thermal endpoints in the current study reflected the tendency of the doves to become agitated and show prolonged escape behavior at high \( T_a \) to a greater degree than was the case for the passerines in Whitfield et al.’s (2015) study, with the result that we often had to remove birds from the chambers before their thermal endpoints in the absence of activity could be elicited.

At \( T_a \approx 56 \, ^\circ C \) (the highest \( T_a \) reached by all four species), hyperthermia was more pronounced in Cape turtle doves (mean \( T_b = 44.7 \, ^\circ C \)) than laughing doves (mean \( T_b = 43.6 \, ^\circ C \)) or Namaqua doves (mean \( T_b = 42.5 \, ^\circ C \)), and least pronounced in crested pigeons (mean \( T_b = 41.7 \, ^\circ C \)) (Table 1). This positive scaling relationship between the extent of hyperthermia and \( M_b \) among the three southern African species, but a much lower value for the larger Australian crested pigeon, approximately mirrors the pattern we found for overall heat tolerance, as quantified by the highest \( T_a \) tolerated. These observations also raise the possibility that the evolution of crested pigeons’ thermal physiology has taken place under a qualitatively and/or quantitatively different set of selection pressures compared to those experienced by the three African species. One climatic factor that may be relevant is the lower maximum \( T_a \) values typical of the Kalahari Desert compared to those that occur in many parts of the Australian arid zone.

Among the five of seven Cape turtle doves which exhibited hyperthermic \( T_b \) at \( T_a = 56 \, ^\circ C \), rates of \( T_b \) increase remained fairly low, which we interpret as indicative of facultative, regulated hyperthermia rather than thermoregulatory breakdown and lethal heat stroke (Leon, 2006). Laughing and Namaqua doves showed similar patterns of hyperthermic \( T_b \) combined with modest rates of \( T_b \) increase at \( T_a = 58 \, ^\circ C \) (mean \( T_b = 43.6 \, ^\circ C \)) and 60 \( ^\circ C \) (mean \( T_b = 43.8 \, ^\circ C \)), respectively. The hyperthermic \( T_b \) values in these columbids are within the range for avian facultative hyperthermia (reviewed by Tieleman and Williams, 1999), with water conservation thought to be the primary function of this physiological response. The latter authors noted that during acute exposure to heat (1 hr), birds ranging in \( M_b \) from 10-1000 g may reduce their total evaporative water loss by as much as 50% by becoming hyperthermic, whereas during chronic exposure (5 hr), only small birds (\( M_b < 100 \, g \)) benefit
(Tieleman and Williams, 1999). Whereas ploceid passerines exposed to high $T_a$s exhibited rapid increases in $T_b$ combined very high mean $T_b$ values (~45 °C) (Whitfield et al., 2015), hyperthermia in the columbids was characterized by a generally stable $T_b$ somewhat elevated above normothermic levels, with little or no change in rate of $T_b$ increase with increasing $T_a$.

In general, columbids appear to show shallower in increases in $T_b$ at high $T_a$ compared to passerines: an examination of changes in $T_b$ between $T_a = 35$ °C and $T_a = 48$ °C among six passerine species reveals a mean $T_b$ increase of 3.02 ± 1.37 °C (Tieleman et al., 2002a; Whitfield et al., 2015; Wolf and Walsberg, 1996; B. Smit et al., unpublished data), whereas the corresponding value for seven columbids is just 1.46 ± 0.19 °C (Hoffman and Walsberg, 1999; McKechnie and Wolf, 2004; Withers and Williams, 1990; present study). The smaller increase among columbids supports the notion that the CEWL-predominated evaporative cooling of the former taxon provides the physiological basis for more effective maintenance of $T_b$ below $T_a$ compared to the REWL-predominated evaporative cooling of the latter group, although the larger $M_b$ of the columbids compared to passerines potentially confounds this comparison.

**Evaporative water loss**

As expected on the basis of allometric scaling predictions (Bartholomew and Cade, 1963; Dawson, 1982; Williams, 1996), EWL at $T_a = 56$ °C (highest $T_a$ reached by all three species) generally increased with increasing $M_b$, although the value for crested pigeons was lower than the smaller Cape turtle dove (Table 1). The slope of mass-specific EWL vs $T_a$ was steepest in Namaqua doves and shallowest in crested pigeons, as expected on the basis of the scaling of this variable (McKechnie and Wolf, 2010). The slope for Namaqua doves was very similar to the value predicted by the allometric relationship reported by the latter authors, whereas the slopes for laughing doves, Cape turtle doves and crested pigeons were 20 %, 33 % and 30 % higher, respectively, than the predicted values.

At the highest $T_a$s to which birds were exposed in this study (56-62 °C), Namaqua doves dissipated heat more than twice as rapidly than the three larger species. Namaqua doves were dissipating ~470 % of their metabolic heat load at $T_a = 60$ °C, a value higher than those recorded for other columbids at the same $T_a$ (~286, 369 and 308 % in heat-acclimated rock doves, white-winged doves and mourning
doves, respectively (Marder and Arieli, 1988; Smith et al., 2015). The skin surface area to mass ratio for Namaqua doves is 25-42% greater than the other dove species, indicating that mass differences alone do not account for increased evaporative rates in the Namaqua dove. We hence conclude that Namaqua doves have an unusually pronounced capacity for evaporative cooling. The EHL/MHP value we observed in this species also exceeds by a substantial margin those reported for other species with pronounced heat tolerance (e.g., spotted nightjar *Eurostopodus argus*; Dawson and Fisher, 1969). It is likely that the relatively high surface area / volume ratio of Namaqua doves allows for very efficient cutaneous evaporative heat dissipation. Despite the comparatively large volumes of water necessary for this mechanism of heat dissipation, water storage within the crop (Williams and Koenig, 1980) may make it sustainable over short periods.

Once acute dehydration limits are better understood, assessing the period over which a bird can defend $T_b$ before the onset of lethal dehydration will be important, particularly in the context of extrapolating laboratory data to free-ranging birds. Studies of dehydration tolerance in the past have typically involved withholding water at $T_a$ well below body temperature ($\sim 25 ^\circ C$) and measuring $M_b$ loss over time scales of days to weeks (e.g., Williams and Koenig, 1980), an approach that does not permit the separation of water *versus* tissue loss. Rock doves, for example, have been shown to tolerate $M_b$ loss equivalent to 16–18 % after being deprived of water for 48 hr, as well as food for 24 hr of that period (Arad et al., 1989), conditions very different to rapid evaporative water loss during acute heat stress.

*Resting metabolic rate*

All four columbids examined showed only small increases in RMR at $T_a$ above thermoneutrality, and even when defending $T_b$ more than 15°C below $T_a$, the mean RMR of inactive birds never increased to more than $\sim 2 \times$ RMR (Figure 1). This observation is qualitatively similar to those made for other columbids during acute heat exposure (Marder and Arieli, 1988; Withers and Williams, 1990), as well as houbara bustards *Chlamydotis macqueenii* (Tieleman et al., 2002b). Marder and Arieli’s (1988) data for heat-acclimated rock doves are particularly striking in this regard, with RMR at $T_a = \sim 60 ^\circ C$ virtually identical to that at $30 ^\circ C \leq T_a \leq 40 ^\circ C$. The general absence of large increases in RMR at $T_a > T_b$ in columbids contrasts with the pattern typical of passerines (Dawson, 1954; Hinds and Calder, 1973; Tieleman et al.,
2002a; Weathers and Greene, 1998; Whitfield et al., 2015; Wolf and Walsberg, 1996) and other avian orders (Hinsley et al., 1993; Lasiewski et al., 1970; Marder and Bernstein, 1983; Weathers and Caccamise, 1975), and indeed the classic Scholander-Irving model of endothermic homeothermy (Scholander et al., 1950).

Contrary to expectations based on literature on the scaling of RMR in heat-stressed birds (Bartholomew and Cade, 1963; Weathers, 1981), Namaqua doves in our study exhibited a shallower slope of mass-specific RMR vs $T_a$ (when accounting for individual responses in a mixed model) than the three larger species. We suspect this result reflects the fact that panting/gular fluttering was delayed until much higher $T_a$ values (~55 °C) in Namaqua doves compared to the other two species (~44, 41 and 48 °C in laughing doves, Cape turtle doves and crested pigeons, respectively). Earlier work suggested that, in columbids, panting and/or gular fluttering commences at $T_b = 42$-43°C (Bartholomew and Dawson, 1954; Randall, 1943), observations that are supported by our data, with $T_b = 42$ °C corresponding to $T_a = ~ 55, 44$ and 42°C in Namaqua, laughing and Cape turtle doves, respectively. The observation that larger doves apparently needed to supplement CEWL with panting and/or gular fluttering at lower $T_a$ than smaller species likely reflects decreasing surface area / volume ratios with increasing $M_b$, although Marder and Arieli (1988) noted the absence of panting or gular flutter in some rock doves exposed to $T_a = 60$-65°C and low humidity. This observation might hold for other species that were habituated to human disturbance as were the rock doves.

As is the case for increases in $T_b$, columbids also appear to generally show smaller fractional increases in RMR at high $T_a$ compared to passerines. Among seven passerines, the mean ratio of RMR at $T_a = 48$ °C compared to $T_a = 35$ °C is 1.38 ± 0.22 (Tieleman et al., 2002a; Whitfield et al., 2015; Wolf and Walsberg, 1996); B. Smit et al., unpublished data), whereas the corresponding value for seven columbids is 1.07 ± 0.09 (Hoffman and Walsberg, 1999; McKechnie and Wolf, 2004; Withers and Williams, 1990; present study). The significantly smaller fractional increases in RMR among columbids compared to passerines between $T_a = 35$ and 48°C in the conventional analysis support the notion that evaporative cooling predominated by CEWL is more energetically efficient than REWL-predominated cooling. Although $M_b$ is again a confounding factor in this comparison, we argue that these differences likely reflect the metabolic cost of muscle contractions involved in panting (Dawson, 1982; Richards, 1970), and the concomitant rapid increases in RMR that typically
occur with increasing $T_a$ above the thermoneutral zone in passerines (e.g., Ambrose et al., 1996; Trost, 1972; Williams, 1999). A recent demonstration that the metabolic cost of lung ventilation in running birds is very low ($< 2\%$ of total metabolic rate; Markley and Carrier, 2010) reiterates the need to better understand the costs involved in respiratory evaporative heat dissipation, particularly in the context of how variation in these costs might contribute to inter- and intraspecific variation in the efficiency of evaporative cooling (Noakes et al., 2016).

Cutaneous vs respiratory evaporation: ecological and evolutionary implications

Our data suggest that variation in the primary avenue of avian evaporative heat loss may have important consequences for their capacity to tolerate acute heat exposure under both laboratory and natural conditions. For instance, data for columbids (present study) and ploceid passerines (Whitfield et al., 2015) suggest that the scaling of thermal limits during acute heat exposure may depend on the primary avenue of evaporative heat dissipation.

To understand the ecological significance of phylogenetic variation in avian evaporative cooling pathways, we need to extrapolate laboratory data on acute heat stress to natural conditions. The goal of the present study, like those of Whitfield et al. (2015) and Smith et al. (2015), was to quantify upper limits to heat tolerance and evaporative cooling capacity in a manner facilitating direct comparisons of standardized variables among species. The conditions birds experienced during our experimental protocol (rapid increases in $T_a$ combined with very low humidity values maintained via high flow rates) are unlikely to directly mirror conditions they routinely experience in natural habitats, although humidity values in desert habitats are often similar to those experienced by birds in our study (see e.g., http://www.bom.gov.au/ for data for Australia). Nevertheless, we would argue that the broad patterns identified here concerning the very efficient evaporative cooling and tolerance of high environmental temperatures by columbids reflect ecologically important differences. In hot desert environments in southern Africa and North America, for instance, columbids are often more active at higher $T_a$ compared to passerines (B. Smit, B.O. Wolf and A.E. McKechnie pers. obs). Moreover, the onset of panting/gular flutter in columbids typically occurs at considerably higher $T_a$ than in similarly-sized passerines (B. Smit, N. Pattinson, M. Thompson et al. unpublished data).
One factor critical in extrapolating the responses of arid-zone birds to acute heat stress under laboratory conditions to natural environments concerns the availability of drinking water. Most arid-zone columbids are strongly water-dependent and regular drinkers (Fisher et al., 1972; Maclean, 1996; Wolf et al., 2002), and the greater overall heat tolerance of columbids we have documented in the present study is probably tightly linked to the availability of water for rapid evaporative heat dissipation. In the absence of drinking water, heat tolerance in columbids may be compromised relatively quickly; for instance, our data suggest that at an environmental temperature of 48 °C and with no access to water, Namaqua, laughing and Cape turtle doves would experience EWL equivalent to 11 % of body mass after 4.6, 4.6 and 6.4 hours, respectively. Dehydration equivalent to 11-15 % of body mass is likely near the upper limit of avian dehydration tolerance (Wolf, 2000).

The data currently available on the relative roles of respiratory and cutaneous evaporation in avian thermoregulation at very high $T_a$ are largely restricted to two orders, Columbiformes and Passeriformes (Tieleman and Williams, 2002; Wolf and Walsberg, 1996), and patterns of EWL partitioning in other orders remain less clear. One reason behind the paucity of data for many orders is that measuring REWL and CEWL is more technically challenging than measuring total evaporation, and requires either a partitioned chamber in which REWL and CEWL are measured in separate compartments (e.g., Hoffman and Walsberg, 1999; Lasiewski et al., 1971; Wolf and Walsberg, 1996) or a mask system (e.g., Tieleman and Williams, 2002). Both these approaches require that birds be habituated to the experimental setup, and that chambers and/or masks be custom-built for particular study species. However, the pronounced differences in heat tolerance and evaporative cooling capacity between the only two orders that are relatively well-studied in this regard highlight the need for quantitative data on the contributions of REWL and CEWL to total evaporation in many more avian taxa. Such data are a prerequisite for fully understanding phylogenetic variation among birds in heat tolerance and evaporative cooling capacity.

Finally, data on avian thermoregulatory responses to acute heat stress and dehydration tolerance are relevant to modeling the impacts of more frequent and intense heat waves on arid-zone avifaunas. Absolute maximum air temperatures and the frequency of intense heat events are predicted to increase substantially in coming decades (IPCC, 2011), and catastrophic avian mortality events similar to those
documented historically, particularly in the Australian arid zones (Finlayson, 1932; Serventy, 1971; Towie, 2009; Towie, 2010) are likely to occur much more frequently than they have in the past (McKechnie and Wolf, 2010). Developing models that use data collected under laboratory conditions to predict the heat tolerance and hydration status of free-ranging birds in natural habitats is vital for predicting where and when these die-offs are likely to occur.

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Abbreviations
CEWL cutaneous evaporative water loss
EHL evaporative heat loss
EWL evaporative water loss
\( M_b \) body mass

MHP metabolic heat production

RER respiratory exchange ratio

REWLR respiratory evaporative water loss

RMR Resting metabolic rate

\( T_a \) air temperature

\( T_b \) body temperature

\( \dot{V}_{\text{CO}_2} \) carbon dioxide production

\( \dot{V}_{\text{O}_2} \) oxygen consumption

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