It Takes Time to Be Cool: On the Relationship between Hyperthermia and Body Cooling in a Migrating Seaduck

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The large amount of energy expended during flapping flight is associated with heat generated through the increased work of the flight muscles. This increased muscle work rate can manifest itself in core body temperature (Tb) increase of 1–2°C in birds during flight. Therefore, episodic body cooling may be mandatory in migratory birds. To elucidate the thermoregulatory strategy of a short-distance migrant, common eiders (Somateria mollissima), we implanted data loggers in the body cavity of wild birds for 1 year, and report information on Tb during their entire migration for 19 individuals. We show that the mean body temperature during flight (TbMean) in the eiders was associated with rises in Tb ranging from 0.2 to 1.5°C, largely depending on flight duration. To understand how eiders are dealing with hyperthermia during migration, we first compare, at a daily scale, how Tb differs during migration using a before-after approach. Only a slight difference was found (0.05°C) between the after (40.30°C), the before (40.41°C) and the migration (40.36°C) periods, indicating that hyperthermia during flight had minimal impact at this time scale. Analyses at the scale of a flight cycle (flight plus stops on the water), however, clearly shows that eiders were closely regulating Tb during migration, as the relationship between the storage of heat during flight was highly correlated (slope = 1) with the level of heat dumping during stops, at both inter-individual and intra-individual levels. Because Tb at the start of a flight (TbStart) was significantly and positively related to Tb at the end of a flight (TbEnd), and the maximal attained Tb during a flight (TbMax), we conclude that in absence of sufficient body cooling during stopovers, eiders are likely to become increasingly hyperthermic during migration. Finally, we quantified the time spent cooling down during migration to be 36% of their daily (24 h) time budget, and conclude that behavioral body cooling in relation to hyperthermia represents an important time cost.

Keywords: body cooling, common eiders, endothermy, hyperthermia, migration, thermoregulation

INTRODUCTION

Homeothermic endotherms—animals that maintain a constant, regulated, body temperature—have the unique ability to maintain a high core body temperature (Tb) across a wide range of ambient temperatures (Ta; Crompton et al., 1978). Birds display the highest Tb of all endothermic animals, exceeding that of mammals, for example, on average by up to ≈2.5°C (Prinzinger et al.,
It has been proposed that such a high $T_b$ in birds is prevalent to enhance the performance of powered flight through $Q_{10}$—the factor by which the rate of a reaction increases for every 10 degree rise in temperature—driven effects on muscle contractile activity (Torre-Bueno, 1976). Although, birds are known to regulate body temperature to a high degree and within very narrow margins, there is building evidence that substantial deviations from normothermia naturally occur (Guillemette et al., 2016). Indeed, hypothermic events such as torpor is evident in many families of afrotropical birds while there are numerous studies reporting cases of controlled hypothermia as a strategy to conserve energy in temperate and arctic environments as well (McKechnie and Mzilikazi, 2011; Lewden et al., 2014). While $T_b$ may be lowered as an energy-saving mechanism in many bird species, the result of the high-energy demands imposed by flight, with flapping flight often being described as the costliest locomotion mode among vertebrates (Butler and WOakes, 1990; Butler, 2016), is associated with the highest levels of known heat production among endotherms (Clarke and Rothery, 2008). It is estimated that 80–96% of the metabolic energy used during flapping flight in birds and bats is dissipated as heat, while only 4–20% is converted into mechanical power (Carpenter, 1986; Speakman and Thomas, 2003). Such dramatic increases in $T_b$ during flight (Platania et al., 1986) require accurate regulation to avoid critical levels of hyperthermia being reached, and the associated deleterious physiological consequences such as excessive evaporative water loss (Boyles et al., 2011). Long-distance avian migrations can include non-stop flights of over 10,000 km, involving millions of wingbeats (Gill et al., 2005). Therefore, in flying birds, mechanisms for body cooling are likely to be an important feature of migration strategies, when flight time is considerably increased and the risk of hyperthermia may be highest.

To cope with the increases in $T_b$ during long-duration flights, a diverse range of behavioral and physiological strategies have evolved in birds as a means to maintain a constant $T_b$, and thus retain cellular homeostasis. Unlike endothermic mammals, however, birds lack sweat glands and thus rely on other mechanisms of effective heat loss. The respiratory tract, through panting or gular fluttering, plays an important role in evaporative heat loss in some species, while the plumage, skin, bill as well as the legs and feet are known important sites of heat exchange that may assist in alleviating the large water loss through the respiratory tract (Steen and Steen, 1965; Dawson, 1982; Giladi and Pinshow, 1999; Tattersall et al., 2016). Evaporative heat loss through the limbs has been well documented, notably in seabirds (Steen and Steen, 1965; Baudinette et al., 1976). The arteries and veins in the legs of many birds lie in close contact with each other, forming a rete mirabile (a net-like-complex of arteries and veins), and function as a countercurrent heat exchange system (Midtgård, 2008). In addition, many birds can adjust the blood flow through the legs by muscular contraction, and can thus regulate heat loss to some degree depending on environmental conditions (Kilgore and Schmidt-Nielsen, 1975).

The behavioral and physiological mechanisms employed by many birds to regulate body temperature are most pertinent during long-duration flights, chiefly those associated with migration when heat loss becomes a particularly challenging problem. Some species may increase their flight altitude to reach cooler temperatures or wind conditions (Torre-Bueno, 1976; Bäckman and Alerstam, 2001) whereas other species are known to fly predominantly at night-time in cooler temperatures which may be related to thermoregulation (Kerlinger and Moore, 1989). For many migratory birds, stopovers are a crucial component of migration and serve to replenish and restore energy reserves (Moore and Kerlinger, 1987), but such breaks from flight may also play a role in avoiding hyperthermic conditions and maintaining homeostasis.

Recently, Guillemette et al. (2016) demonstrated that common eiders (Somateria mollissima) experience large departures from normothermia during migratory flights. These increases in $T_b$ ranged on average between 0.2 and 2.4°C during migratory flights, and were likely due to the high wing-loading found in eiders (a large body to small wings ratio), the morphology of which is linked to their diving lifestyle. Indirect evidence that eiders proactively avoid deleterious levels of hyperthermia during flight is given by their migratory style, whereby they use a stop-and-go strategy where they perform many flights of short duration (15 min) during which $T_b$ increased by 1°C on average (Guillemette et al., 2016). In addition, about 60% of the migratory flights were stopped while $T_b$ was still increasing (Guillemette et al., 2016). Therefore, it appears that behaviorally mediated body cooling, might be an important component of the thermoregulatory strategy during migration for eiders.

In this paper, we evaluate the role of body cooling during migration stop-overs of common eiders while they migrate from breeding grounds to their late-summer molting sites. More specifically, we (1) compare daily $T_b$ during migration to $T_b$ prior to the migratory flights, and predict that daily $T_b$ during migration is higher overall than daily $T_b$ during the pre-migratory phase when flight time is significantly lower. We (2) predict that hyperthermia is a constraint to flight duration in migrating eiders, and therefore they should decrease their $T_b$ to within a normothermic range before engaging in any further flight activity. We thus tested, at both the inter- and intra-individual level, that the duration between flights is governed by the time it takes to become normothermic. Finally, we (3) quantified the rate of body cooling and the time to reach a minimum $T_b$, during stops between flights, in order to evaluate if it could influence migration speed.

**MATERIALS AND METHODS**

**Model Species and Population Studied**

Common eiders are large (2 kg) sea ducks that dive for food living in the benthos. They are characterized by short-pointed wings resulting in high wing-loadings and high flight speed (Day et al., 2004; Guillemette and Ouellet, 2005). The population studied is breeding in Baltic and molting in the Wadden Sea (Rigou and Guillemette, 2010). During migration, eider ducks usually follow the coast and rarely fly over areas of unsuitable feeding habitats (Guillemette, 2001; Guillemette et al., 2016). As with many other species of waterfowl, common eiders undergo a molt migration in late summer. They move from breeding habitats to...
their molting areas where they lose their wing feathers all at once leading to a period of flightlessness that last 36 days on average (Guillemette et al., 2007). The molt migration of this population in summer, involves movement from the breeding grounds in the central Baltic to the molting quarters located in the Wadden sea, covering a distance of 714 km (±286) on average.

The study was performed on Christiansø Island (55°19’N, 15°12’E), an old Danish fortress located in the southern Baltic Sea, 18 km from the Danish island of Bornholm. The general approach of our work involved the monitoring and deployment of data loggers on breeding females, using heart rate data to determine the start and the end of each flight, and computing variation of body temperature during and between flights. Using the mean eider groundspeed velocities estimated at 83.5 ± 0.3 km h⁻¹ by Day et al. (2004) and the total time spent flying (see below) during each migration day of each individual, we calculated the migration distance of each instrumented female. Using that information together with the molting movements of this Danish population, as described previously (Lyngs, 2000), we estimated the location and used the date of migration for all individuals to estimate air [15.5 ± (SD) 1.8] and (surface) water temperature [15.5 ± (SD) 1.9].

Deployment of Data Loggers

On Christiansø island, 45 common eiders were captured in 2003, 2004, and 2005 and implanted with custom made data loggers (DLs, Biometistics, A. J. Woakes) under license from Dyreforstingsbyrån (Danish Royal Veterinarian Corporation) and approved by the Canadian Council of Animal Care (# CPA 16-03-07-01). Only females were caught as males leave the colony when incubation starts. All surgical procedures were conducted indoors according to the procedure described by Guillemette et al. (2002). The 45 DLs were 36 mm long (±SD = 0.5) × 28 mm (0.2) wide × 11 mm thick (0.3) and weighed 21 g (0.3), that is 1.2% of body mass at implantation (Guillemette et al., 2007). Thirty nine (87%) experimental females returned to their molting areas where they lose their wing feathers all at once leading to a period of flightlessness that last 36 days on average (Guillemette et al., 2007). Thirty nine (87%) experimental females returned to

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Body Temperature during a Flight Cycle

A flight cycle is composed of flight followed by a period of time spent on the water before the next flight (stop). Body temperature (Tb) was recorded at the start (TbStart) and at the end (TbEnd) of each flight together with maximum Tb (TbMax), whenever it occurred during each flight (Figure 1). The Tb sensor time inertia was evaluated to be 3 min as we observed changes in Tb only after 3 min of flight. Thus, all flights <3 min were excluded from our analysis (n = 412). In addition, the minimum Tb (TbMin) occurring during each stop interval was recorded. We define the heat storage index (HSI) as Tb in relation to elapsed time (°C.h⁻¹). Here, we calculate the heat stored during flight for each time interval and hence report the interval specific HSI for each flight segment (0–5, 5–10 min). Similarly, we computed a body cooling index (BCI) by calculating the difference of Tb between the end of flight and Tb at 5 min increment after the end of each flight.

Data Analysis

First, we were interested to know how and if eiders ducks were able to regulate Tb during migration at a daily scale compared to periods before and after migration. Daily body temperature (TbDaily) during migration was calculated as the average of all Tb datum (5,400 per day) recorded during migration days of each individual. This average value was subtracted in a before-after fashion using a similar time interval (3 days), giving two individual deltas per individual. These deltas were averaged over the 19 females for which confidence intervals were computed using the bootstrap method (see main text for results).

Second, we were interested to know if body cooling between flights was sufficient to dump all the heat gained during flight. We started our analysis by relating the heat gain during flight with heat loss during stops at the intra-individual level. We

![Figure 1](https://via.placeholder.com/150)

**FIGURE 1** Schematic representation of a flight cycle during migration of common eiders. The red line depicts the heart rate of two flights, and the time interval between flights, used to quantify flight occurrence and flight duration of this study. The yellow line shows the (smoothed) variation of Tb in relation to time in the flight cycle. See text for definitions of acronyms.
calculated an intra-individual reduced-major axis (RMA) slope between the temperature gained during all flights performed in relation to the temperature lost between flights (stops) for each individual taken separately and then tested if the intra-individual slope averaged over all individuals differs from the theoretical value of one (i.e., complete $T_b$ regulation). The average slope and its associated confidence intervals (CI) was computed using a bootstrap and 10,000 re-samplings (Lunneborg, 2000). In a further step, we tested the complete regulation hypothesis at the inter-individual level by summing all gains and losses in $T_b$ (in relation to $T_b$Start) occurring during an average migration day for each individual in turn. From these data, a RMA equation was computed where the correlation coefficient was tested for significance using a permutation test.

Finally, we calculated the intra-individual correlation level between $T_{b\text{End}}$, $T_{b\text{Daily}}$, $T_{b\text{Max}}$ for all flights performed, which was averaged over all individuals with CIs computed from a bootstrap and ten thousand re-samplings.

**RESULTS**

Mean $T_b$ during flight increased positively with flight duration during the molt migration of the common eiders, from about 40.2°C for short flights to 41.5°C for longer flights (Figure 2). Mean $T_b$ of short flights (3–10 min) were within the mean daily variation of $T_b$ ($T_{b\text{Daily}}$) during migration days and only longer flights (>10 min) were outside the confidence intervals (Figure 2). Given that migration effort of these 19 females is slightly more than 200 min spent flying on average per day (Guillemette et al., 2016), we tested if average $T_b$ at a daily scale would be influenced by flight time.

This was achieved by comparing $T_{b\text{Daily}}$ during migration with periods before and after migration (Figure 3). We found that $T_{b\text{Daily}}$ before migration [40.41 ± (SD) 0.58°C] was slightly higher, on average, than during migration [40.36 ± (SD) 0.54°C], as the mean difference between the two temperatures (before and during migration) exclude zero [average difference = 0.05°C ± (bootstrap CI) 0.04]. In contrast, $T_{b\text{Daily}}$ after migration [40.30 ± (SD) 0.54°C] was lower on average than during migration [average difference = 0.06°C ± (bootstrap CI) 0.06]. Although, $T_{b\text{Daily}}$ varied little but significantly, on average, the shape of the frequency distribution was influenced by flight and migration. Unsurprisingly, during migration eiders experienced more $T_b$ between 41.4 and 42.2°C when compared to the before and after migration periods (Figures 3B,C).

We then tested the idea that body cooling between flights was a determining factor in the migration strategy of eiders, by relating the body heat gain during flight with body heat loss during stops, at the intra-individual level. We calculated an intra-individual RMA slope between the temperature increases during flights in relation to the temperature decreases between flights for each flight cycle taken separately, and then tested if the average intra-individual slope differs from the theoretical value of one (i.e., complete $T_b$ regulation). The RMA slope ranged between 0.75 and 1.35 among individuals, with a mean of 1.04 ± 0.06 (bootstrap CI) when averaged over all individuals (mean Pearson $r = 0.737$, Figure 4A). We further tested this hypothesis at the inter-individual level by summing all increases and decreases in $T_b$ occurring during an average migration day for each individual in turn; we also found a strong and linear relationship ($y = 0.985x - 0.11, r = 0.976, \text{permutation test, } p < 0.0001$; Figure 4B). Notably, cumulative daily individual variation of $T_b$ related to flight activity during migration was high (3–15°C).

The close regulation of $T_b$ observed (Figure 4) at the scale of a flight cycle led us to postulate that reduction of $T_b$ to a minimum level is likely to be mandatory, as it would reduce the likelihood of an occurrence of reaching an unacceptable level of hyperthermia for the subsequent flight. Despite the seemingly large variation between individuals, body temperature at the start of a flight ($T_{b\text{Start}}$) was positively related to body temperature at the end ($T_{b\text{End}}$) of a flight at the intra-individual level [average correlation = 0.263 ± (bootstrap CI) 0.095, Figure 5A]. A similar observation was made for $T_{b\text{Max}}$ [average correlation = 0.325 ± (bootstrap CI) 0.101] which supports our hypothesis (Figure 5B). In addition, at the inter-individual level (results not shown), a similar relationship with $T_{b\text{Start}}$ reveals a significant coefficient of correlation for both $T_{b\text{End}}$ ($r = 0.664, p < 0.05$) and $T_{b\text{Max}}$ ($r = 0.685, p < 0.05$).

During temporary migratory stop-overs, the body temperature of the eiders decreased at an exponential rate, which stabilized at about 45–50 min, on average, from the end of the preceding flight (Figure 6A). Indeed, the highest average cooling rate was during the first 10 min after landing, which itself was substantially higher following longer flights, reaching 6–7°C h⁻¹ (Figures 6B,C). Nevertheless, the average time required to cool down to $T_{b\text{Min}}$ [40.10 ± (SD) 0.56], the minimum level between flights [47.14 ± (SD) 21.73 min], is substantially shorter than the average duration of each stop between flights [72.1 ± (SD) 21.6 min], which raises the question of why birds are taking extra time during stops before resuming flight.
DISCUSSION

In a former paper, we have presented evidence that hyperthermia might dictate flight duration in a migrating bird (Guillemette et al., 2016). Here, we present evidence that body cooling is an integral part of the migration of common eiders by showing that daily fluctuations in $T_b$ caused by flight were precisely regulated at the scale of a flight cycle, despite large cumulative deviations in $T_b$ caused by the migration and associated flights.

We thus suggest below that such a level of regulation results from behavioral thermoregulation together with the intrinsic physiological flexibility of birds.

$T_b$ Daily and Heat Dumping during Stop-Overs

It is apparent that birds have higher body temperature tolerance in comparison to mammals (Prinzinger et al., 1991; McNab,
During molt migration, Baltic common eiders fly 205 min per day, on average (Guillemette et al., 2016), and individuals experience daily cumulative variations in $T_b$ ranging between 3 and 15°C (Figure 3B). Despite this large variation in $T_b$ associated with flight activity, $T_{b\text{Daily}}$ during migration days (40.36 ± 0.54°C) was only slightly different, by 0.05 degrees on average, when compared to before (40.41 ± 0.58°C) and after (40.30 ± 0.54°C) periods. The most distinctive features of migration were the time spent at $T_b$ values in the range of 41.4–42.4 and 40.0–40.2°C (Figures 2B–C), which correspond respectively to $T_b$ encountered during long flights (see Guillemette et al., 2016, Figure 1) and $T_b$ while idling during stops. The observation that eiders regulate $T_b$ around $T_{b\text{Min}}$ at 40.1°C during stops, idling between flights, suggests a selected body temperature target while waiting to perform the next flight. It is also interesting to note this temperature range to be at the low range of $T_{b\text{Daily}}$ confidence intervals (Figure 2) together with the fact that average $T_b$ during flights of 0–10 min are within the confidence intervals of $T_{b\text{Daily}}$. Altogether, it thus appears that migrating common eiders defend $T_{b\text{Daily}}$ with success and we suggest that a part of the absolute level of hyperthermia reached by an exercising animal, the time spent at a high level of $T_b$ is also an important point to consider when evaluating the potential and deleterious effects of hyperthermia.
Indeed, the return to normothermia was relatively rapid in migrating eiders (Figure 6A). Support for this interpretation is given by various cancer treatment therapies using hyperthermia in humans (Wang et al., 2013), which induces intracellular reactive oxygen species (ROS) production and mitochondrial ROS generation in a time-dependent manner. One advantage of maintaining $T_b$ at a low level before the next flight is to eventually reduce the level of hyperthermia reached during that subsequent flight. This hypothesis was supported by the data as we found a positive correlation between $T_{b_{\text{start}}}$ and $T_{b_{\text{end}}}$ and $T_{b_{\text{start}}}$ and $T_{b_{\text{max}}}$ (see Section Results).

**Body Cooling Rate and Idling Time**

It is expected that marine birds potentially dissipate heat at a high rate following landing, due to being on water. Indeed, the rate of cooling of migrating common eiders is high for the first 10 min after landing, particularly after longer flights (Figures 6B, C), with average rates of cooling ranging from 6 to 7°C h$^{-1}$. These estimates of body cooling are difficult to compare with any other species, given the paucity of data. From birds shot while flying, Platania et al. (1986) observed a decrease of $T_b$ varying between 1.2 and 3.6°C h$^{-1}$ for three species (222–449 g), 9.0°C h$^{-1}$ for the smallest species (40 g) and 1.7°C h$^{-1}$ for the largest species (765 g). Unfortunately, Platania et al. (1986) do not give any details about the medium (air vs. water) where these shot birds were cooling down when taking the measurements. One further complication is that we expect dead birds to cool down more...
slowly than live birds given they cannot divert blood flow to the periphery or legs to cool down more rapidly (see below).

Interestingly, the cooling rates observed during stop-overs of eiders are similar to the highest heat storage rates occurring at the beginning of flights (Figures 6B,C, indicating that body cooling and heat storage reach equivalent maximum values. In his review, McNab (2002) also reports similar values for ectotherms for both cooling and heat storage rates in air. However, it could be expected that body cooling in aquatic birds would occur at faster rates than heat storage, given the heat dissipation potential of sea water is much higher than air. In humans, cooling rates have been measured in circulated water baths (14 and 20°C water) after exercise in air (65% VO2Max at 38°C) and reaching a rectal temperature of 40.0°C (Proulx et al., 2003). They observed an exponential decrease of Tb with the largest decrease occurring in the first 10 min after immersion, similar to eiders during stop-overs (Figure 6A), with the difference that maximum (11.4−15.6°C h−1) and average (9−11.4°C h−1) cooling rates were much higher than average heat storage rates (3–4°C h−1) in the seven human subjects. Unfortunately, Proulx et al. (2003) did not report the maximum rate of heat storage though a maximum rate of 6°C h−1 was reported in a very similar experimental setting (González-Alonso et al., 1999). Thus, although humans are much larger animals, maximum body cooling in water seems to be much more rapid than maximum body cooling of migrating eiders, the other main difference being, of course, that humans are naked and deprived of fur or body feathers.

The naked legs of many bird species (Steen and Steen, 1965; Kilgore and Schmidt-Nielsen, 1975; Midtgård, 2008) have been shown to serve as controlled heat conduits, being of paramount importance for thermoregulation in both water (Steen and Steen, 1965; Kilgore and Schmidt-Nielsen, 1975; Midtgård, 2008) and air (Baudinette et al., 1976; Martineau and Larochelle, 1988). The evidence for this is stemming from calorific studies and blood flow measurements. For example, blood flow to the legs of mallards (Anas platyrhynchos) and fulmars (Macronectes giganteus) is reduced and heat conservation occurs (Johansen and Wesley Millard, 1973; Kilgore and Schmidt-Nielsen, 1975), when water temperature is reduced experimentally during rest. In contrast, heat dissipation can increase considerably where legs are serving as heat dumping organs during both flying and swimming, although the rate of heat loss can vary considerably given the varying experimental conditions used by the different studies. From a behavioral point of view, trailing or retracting the legs and feet in the plumage while flying (Udvardy, 1983; Neumann and Neumann, 2016) or swimming (Harris et al., 2010) is further modulating heat dissipation of birds.

Time as the Cost of Behavioral Thermoregulation

The fact that eiders stop flying when overheating can be seen as another example of behavioral thermoregulation. In his thorough review, McNab (2002) concluded that time is the major cost associated with behavioral thermoregulation of ectotherms although his book almost eludes the subject in relation to endotherms, most probably because examples were few (Smit et al., 2016). Here, we propose that cooling time is a major cost of the migration strategy of common eiders. As such, eiders take about 47 min on average to cool down by 1.1°C and reach TbMin, corresponding to an overall cooling rate of 1.4°C h−1, which is much lower than the max rate observed at the beginning of the cooling period (6–7°C h−1, Figure 6). This is readily explained by the (exponential) nature of the cooling curve and the fact that most flights of eiders during molt migration are short (16 min average, Guillemette et al., 2016) when the level of heat stored in the body is still relatively low (Figure 2). Interestingly, body cooling in eiders does not appear to be a passive cooling process as predicted by Newton’s law of cooling because the relationship between ln(Tb−Tb(eq)) and time (see Figure 6A insert), does not fit a linear relationship. In fact, it appears that cooling is initially enhanced and slows down progressively just before reaching the level of equilibrium. Although speculative, the initially increased rate of cooling may be indicative of an active processes of heat dissipation related to e.g., an increased blood flow through the legs of the birds in cooler water.

However, it remains that the 47 min to cool down is still lower than the average duration of stops between flights observed in our sample (72.1 ± 21, 6 min). This indicates that the time spent between flights is not entirely devoted to body cooling (Figure 1) and we hypothesized that such an extra time taken between flights is caused by unfavorable wind conditions. Nevertheless, assuming that cooling down to TbMin is a prerequisite to perform the next flight, the daily time spent cooling down on a typical migration day (with about 14 flights per day, Guillemette et al., 2016) would represent 523 min or 36% of the 24 h of the daily time budget. If we add the idling time (about 300 min or 20%) we obtain that eiders are spending altogether 56% of the time spent cooling and idling, between flights, during a typical migration day. This indicates that behavioral thermoregulation of common eiders during migration represents a major component of their time budget.

In conclusion, it is likely that the occurrence of hyperthermia in common eiders is due to their high wing loading and costs of flight, which in turn is a consequence of their morphology being adapted for diving. A fruitful future research direction would be to establish the prevalence of hyperthermia in other diving bird species during migratory flights, and whether some species show flight altitudinal changes in direct response to increases in Tb. It is likely hyperthermia is a hitherto underestimated contributing factor to migration strategies, and the possible daily distances that can be traveled. Moreover, hyperthermia may be one of major contributing factors of why other diving species with high loading (e.g., alcids) undertake a significant portion of their migrations swimming as opposed to flying (Olsson et al., 1999), with the constant contact of their feet with the water, coupled with the lower work-rate of the leg muscles compared to flight, ameliorating the effects of hyperthermia.
AUTHOR CONTRIBUTION
MG, DP conceived the ideas, conducted the experiments, prepared the manuscript and figures, and approved the manuscript. EP and SP prepared manuscript, assisted in analysis and approved the manuscript.

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REFERENCES
Bäckman, J., and Alerstam, T. (2001). Confronting the winds: orientation and flight behaviour of roosting swifts, Apus apus. Proc. R. Soc. Lond. B Biol. Sci. 268, 1081–1087. doi: 10.1098/rspb.2001.1622
Baudinette, R. V., Loveridge, J. P., Wilson, K. J., Mills, C. D., and Schmidt-Nielsen, K. (1976). Heat loss from feet of herring gulls at rest and during flight. Am. J. Physiol. Leg. Content 230, 920–924.
Boyles, J. G., Seebacher, F., Smits, B., and McKechnie, A. E. (2011). Adaptive thermoregulation in endotherms may alter responses to climate change. Integr. Comp. Biol. 51, 676–690. doi: 10.1093/icb/icr053
Butler, P. J. (2016). The physiological basis of bird flight. Philos. Trans. R. Soc. B Biol. Sci. 371:20150384. doi: 10.1098/rstb.2015.0384
Butler, P. J., and Woakes, A. J. (1990). “The physiology of bird flight”, in Bird Migration: Physiology and Ecophysiology, ed E. Gwinner (Berlin; Heidelberg: Springer), 300–318.
Carpenter, R. E. (1986). Flight physiology of intermediate-sized fruit bats (Pteropodidae). J. Exp. Biol. 120, 79–103.
Clarke, A., and Rothery, P. (2008). Scaling of body temperature in mammals and birds. Funct. Ecol. 22, 58–67. doi: 10.1111/j.1365-2435.2007.01341.x
Coulson, J. C. (2008). The population dynamics of the Eider Duck Somateria mollissima and evidence of extensive non-breeding by adult ducks. Ibis 126, 525–543. doi: 10.1111/j.1474-919X.1984.tb00278.x
Crompton, A. W., Taylor, C. R., and Jagger, J. A. (1978). Evolution of homeothermy in mammals. Nature 272, 333–336. doi: 10.1038/272333a0
Dawson, W. R. (1982). Evaporative losses of water by birds. Comp. Biochem. Physiol. Part A Physiol. 71, 495–509. doi: 10.1016/0300-9692(82)90198-0
Day, R. H., Rose, J. R., Prichard, A. K., Blaha, R. J., and Cooper, B. A. (2004). Environmental effects on the fall migration of eiders at Barrow, Alaska. Mar. Ornithol. 32, 13–24.
Giladi, I., and Pinshow, B. (1999). Evaporative and excretory water loss during free flight in pigeons. J. Comp. Physiol. B 169, 311–318. doi: 10.1007/s003600050226
Gill, R. E., Piersma, T., Hufford, G., Servranckx, R., and Riegen, A. (2005). Crossing the ultimate ecological barrier: evidence for an 11 000-km-long non stop flight from Alaska to New Zealand and Eastern Australia by bar-tailed godwits. J. Appl. Physiol. 94, 1317–1323. doi: 10.1152/japplphysiol.00541.2005
González-Alonso, J., Teller, C., Andersen, S. L., Jensen, F. B., Hyldig, T., and Nielsen, B. (1999). Influence of body temperature on the development of fatigue of facultative daytime hypothermia in a small passerine wintering at northern latitudes. IBIS 136, 321–329. doi: 10.1111/jb.12142
Lunneborg, C. E. (2000). Data Analysis by Resampling: Concepts and Applications. Pacific Grove, CA: Brooks/Cole.
Lyngs, P. (2000). Status of the Danish Breeding population of Eiders Somateria mollissima 1988–93. Dansk. Ornitol. Foren. Tidsskr. 94, 12–18.
Martínez, L., and Larochelle, J. (1988). The cooling power of pigeon legs. J. Exp. Biol. 136, 193–208.
McKechnie, A. E., and Melilakzi, N. (2011). Heterothermy in afrotropical mammals and birds: a review. Integr. Comp. Biol. 51, 349–363. doi: 10.1093/icb/icr035
McNam, B. K. (2002). The Physiological Ecology of Vertebrates. Ithaca, NY; London: Cornell University Press.
Midtgård, U. (2008). Heat loss from the feet of mallards Anas platyrhynchos and arterio-venous heat exchange in the rete tibiotarsale. IBIS 122, 354–359. doi: 10.1111/j.1474-919X.1980.tb00889.x
Moore, F., and Kerlinger, P. (1987). Stopover and food deposition by North American wood-warblers (Parulinae) following spring migration over the Gulf of Mexico. Oecologia 74, 47–54. doi: 10.1007/BF00377344
Neumann, C., and Neumann, C. (2016). Behavioural thermoregulation in the Common Swift during flight. Br. Birds 109, 10–13.
Olsson, O., Fransson, T., and Larsson, K. (1999). Post-fledging migration of common murreas Uria aalge in the Baltic Sea: management implications. Ecography 22, 233–239. doi: 10.1111/j.1600-0587.1999.tb00497.x
Pelletier, D., Guilleumette, M., Grandbois, J.-M., and Butler, P. J. (2007). It is time to move: linking flight and foraging behaviour in a diving bird. Biol. Lett. 3, 357–359. doi: 10.1098/rsbl.2007.0088
Platania, S. P., Grant, G. S., and Lee, D. S. (1986). Core temperature s of non-nesting common eiders: are females constrained by excessive wing-loading or by minimal flight muscle ratio? J. Appl. Physiol. 60, 1032–1039.
Poulin, R., Prefinier, A., and Schlescher, E. (1991). Body temperature in birds. Comp. Biochem. Physiol. Part A Physiol. 99, 499–506. doi: 10.1016/0300-9692(91)90122-S
Pruzk, C. L., Ducharme, M. B., and Kenny, G. P. (2003). Effect of water temperature on cooling efficiency during hyperthermia in humans. J. Appl. Physiol. 94, 1317–1323. doi: 10.1152/japplphysiol.00541.2002
Rigou, Y., and Guilleumette, M. (2010). Foraging effort and pre-laying strategy in breeding common eiders. Waterbirds Int. J. Waterbird Biol. 33, 314–322. doi: 10.1765/063.033.0307
Smits, B., Zietsman, G., Martin, R. O., Cunningham, S. J., McKechnie, A. E., and Hockey, P. A. R. (2016). Behavioural responses to heat in desert birds: implications for predicting vulnerability to climate warming. Clim. Chang. Respons. 3, 9. doi: 10.1007/s10589-016-0023-2

Harris, M. P., Daunt, F., Newell, M., Phillips, R. A., and Wanless, S. (2010). Wintering areas of adult Atlantic puffins Fratercula arctica from a North Sea colony as revealed by geolocation technology. Mar. Biol. 157, 827–836. doi: 10.1007/s00227-009-1365-0
Johansen, K., and Wesley Millard, R. (1973). Vascular responses to temperature in the foot of the giant fulmar, Macronectes giganteus. J. Comp. Physiol. 85, 47–64. doi: 10.1007/BF00694140
Kerlinger, P., and Moore, R. F. (1989). "Atmospheric structure and avian migration," in Current Ornithology, ed D. M. Power (New York, NY: Plenum Press), 109–141.
Kilgore, D. L., and Schmidt-Nielsen, K. (1975). Heat loss from ducks’ feet immersed in cold water. Condor 77:475. doi: 10.2307/1366094
Lewden, A., Petit, M., Milberge, M., Orso, S., and Vezina, F. (2014). Evidence of facultative daytime hypothermia in a small passerine wintering at northern latitudes. IBIS 136, 321–329. doi: 10.1111/jb.12142

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Speakman, J. R., and Thomas, D. W. (2003). “Physiological ecology and energetics of bats,” in Physiological Ecology and Energetics of Bats, eds T. H. Kunz and M. B. Fenton (Chicago, IL: The University of Chicago Press), 430–490.

Steen, I., and Steen, J. B. (1965). The importance of the legs in the thermoregulation of birds. Acta Physiol. Scand. 63, 285–291. doi: 10.1111/j.1748-1716.1965.tb04067.x

Steen, I., and Steen, J. B. (1965). The importance of the legs in the thermoregulation of birds. Acta Physiol. Scand. 63, 285–291. doi: 10.1111/j.1748-1716.1965.tb04067.x

Tattersall, G. J., Arnaout, B., and Symonds, M. R. E. (2016). The evolution of the avian bill as a thermoregulatory organ. Biol. Rev. 92, 1630–1656. doi: 10.1111/brv.12299

Torre-Bueno, J. R. (1976). Temperature regulation and heat dissipation during flight in birds. J. Exp. Biol. 65, 471–482.

Udvardy, M. D. F. (1983). The role of the feet in behavioral thermoregulation of hummingbirds. Condor 85, 281. doi: 10.2307/1367060

Wang, Z., Cai, F., Chen, X., Luo, M., Hu, L., and Lu, Y. (2013). The role of mitochondria-derived reactive oxygen species in hyperthermia-induced platelet apoptosis. PLoS ONE 8:e75044. doi: 10.1371/journal.pone.0075044

White, C. R., Green, J. A., Martin, G. R., Butler, P. J., and Grémillet, D. (2013). Energetic constraints may limit the capacity of visually guided predators to respond to Arctic warming. J. Zool. 289, 119–126. doi: 10.1111/j.1469-7998.2012.00968.x

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