Torpor Patterns in Desert Hedgehogs (*Paraechinus aethiopicus*) Represent Another New Point along a Thermoregulatory Continuum

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ABSTRACT

Documenting variation in thermoregulatory patterns across phylogenetically and geographically diverse taxa is key to understanding the evolution of endothermy and heterothermy in birds and mammals. We recorded body temperature (T_b) in free-ranging desert hedgehogs (Paraechinus aethiopicus) across three seasons in the deserts of Saudi Arabia. Modal T_b 's (35°-36.5°C) were slightly below normal for mammals but still warmer than those of other hedgehogs. The single maximum T_b recorded was 39.2°C, which is cooler than maximum T_b 's recorded in most desert mammals. Desert hedgehogs commonly used torpor during winter and spring but never during summer. Torpor bouts occurred frequently but irregularly, and most lasted less than 24 h. Unlike daily heterotherms, desert hedgehogs did occasionally remain torpid for more than 24 h, including one bout of 101 h. Body temperatures during torpor were often within 2°-3°C of ambient temperature; however, we never recorded repeated bouts of long, predictable torpor punctuated by brief arousal periods similar to those common among seasonal hibernators. Thus, desert hedgehogs can be included on the ever-growing list of species that display torpor patterns intermediate to traditionally defined hibernators and daily heterotherms. Extant hedgehogs are a recent radiation within an ancient family, and the intermediate thermoregulatory pattern displayed by desert hedgehogs is unlike the deeper and more regular torpor seen in other hedgehogs, suggesting that this may be a derived—as opposed to ancestral—trait in this subfamily.

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We suggest that this family (Erinaceidae) and order (Eulipotyphla) may be important for understanding the evolution of thermoregulatory patterns among Laurasiatheria and mammals in general.

Keywords: body temperature, Ethiopian hedgehog, Eulipotyphla, heterothermy, hibernation.

Introduction

Describing variation in thermoregulatory and body temperature (T_b) patterns and determining the distribution—both phylogenetically and geographically—of those patterns in mammals and birds is vital to understanding the evolutionary history of endothermy. Of special interest is how various derivations of heterothermy relate to the evolutionary history of endothermy. Patterns of heterothermy have often been considered distinct (Geiser 1998), with daily heterothermy and hibernation (seasonal heterothermy) differentiated by parameters including length of torpor bouts and metabolic rate during torpor (Geiser and Ruf 1995). Increased interest in the topic and advances in technology have led to an increase in research on species from tropical, subtropical, and desert regions, and exceptions that fall in-between the normal classifications of heterothermy are common enough (Lovegrove et al. 2001; Lovegrove and Genin 2008; Geiser and Mzilikazi 2011; Geiser and Martin 2013) that many researchers now view the metabolic plasticity between normothermy and hibernation as a continuum (Canale et al. 2012; Boyles et al. 2013; van Breukelen and Martin 2015) arising from ancestral heterothermy (Lovegrove 2012a).

Differentiating between the two competing views of heterothermy (distinct categories vs. a continuum) will become easier as more data on T_b of phylogenetically and geographically diverse species become available. One such phylogenetically important order is the Eulipotyphla (solenodons, shrews, moles, hedgehogs, and moonrats), which occupy an interesting (albeit contentious) position in the mammalian phylogeny (Mouchaty et al. 2000; Douady et al. 2002). They are most commonly placed sister to the rest of Laurasiatheria (Douady et al. 2002; Bininda-Emonds et al. 2007), diverging 75–85 million years ago (Douady and Douzery 2003; Hallström and Janke 2010). The Eulipotyphla have been singled out as important in understanding the evolution of heterothermy in mammals because of their early divergence from the other laurasiatherians (Lovegrove 2012*b*).

Among the Eulipotyphla, the thermal physiology of hedgehogs (subfamily Erinaceinae) has long interested researchers

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(Dmi'el and Schwarz 1984; Soivio et al. 1968). Hedgehogs typically maintain normothermic T_b between 33° and 35°C (Shkolnik and Schmidt-Nielsen 1976; Fowler and Racey 1990; Król 1994; Hallam and Mzilikazi 2011), which is intermediate between monotremes, metatherians, and more basal placental mammals (T_b of 30°-33°C; Grant 1983; Grigg et al. 2003; Nicol and Anderson 2006; Lovegrove and Genin 2008) and the more typical mammalian T_b of 37°-38°C. Even among the Eulipotyphla, hedgehogs display unusually low $T_{\rm b}$ (Whittow et al. 1977; Campbell et al. 1999). While a large body of literature exists on thermoregulation of hedgehogs, relatively little work has been published on T_b patterns of this taxon in the wild, especially in arid environments. Laboratory and semicaptive studies suggest that the European hedgehog (Erinaceus europaeus) probably undergoes the longest and deepest torpor bouts among hedgehogs (Dmi'el and Schwarz 1984; Soivio et al. 1968; Fowler and Racey 1990; Webb and Ellison 1998), but other hedgehog species also display relatively long torpor bouts characterized by low T_b (Hallam and Mzilikazi 2011; Mouhoub-Sayah et al. 2012). The four species for which T_b data are available come from a variety of climates (temperature and subtropical) and habitats (woodlands and grasslands), and all use torpor.

We recorded $T_{\rm b}$ of free-ranging desert hedgehogs (*Paraechinus aethiopicus*) across three seasons in a harsh desert environment. Desert hedgehogs are unusual among the hedgehogs because of their adaptations to extremely arid environments; thus, their thermoregulation is likely to be unusual as well. Given that heterothermy seems ubiquitous among the hedgehogs, we expected heterothermy in desert hedgehogs as well. However, given the unusual ecology of this species relative to other hedgehogs, we had no a priori expectations about the form of heterothermy.

Methods

Study Species

The desert hedgehog (150–600 g) is an arid-adapted insectivore found throughout the nonmountainous regions of the Arabian Peninsula and in extreme northern Africa. Relatively little is known about the natural history, behavior, and ecology of this species compared with European hedgehogs and several species and hybrids commonly kept as pets. Courtship behavior most commonly occurs during late winter and into early spring (Yamaguchi et al. 2013). They are mostly nocturnal, and seasonal activity levels of captive animals decrease considerably during midwinter (Al-Musfir and Yamaguchi 2008), and T_b appears to drop when exposed to low ambient temperatures (as cited by Corbet 1988), so it has been assumed they hibernate. It is also known that desert hedgehogs bask during cool winter days, presumably to help control T_b (Abu Baker et al. 2016)

Experimental Design

We hand captured desert hedgehogs over two sessions in sandy habitats on or near date farms of the Unizah province, Saudi Arabia (26.136°N, 43.975°E). We started each capture period just after sunset in winter and around midnight in summer. The

first collection period was in summer 2013 (June 2013). The second collection period was in winter 2013–2014 (November 2013). We captured 20 animals in each season.

We temporarily housed the hedgehogs in an animal facility at the Department of Zoology of King Saud University in Riyadh. We used temperature-sensitive data loggers set to record at 1-h intervals (0.0625°C resolution, iButtons, DS1922L, 4.2 g with wax, Maxim Semiconductors, Dallas) to measure core T_b . We did not have proper equipment available to fully calibrate data loggers. Previous experience with this model of data logger indicates that most are accurate within values claimed by the manufacturer (0.5°C) and that precision within a batch is high (J. G. Boyles, personal observation). However, a weakness of this model is that a small proportion of them deviate outside these values. Therefore, before implanting the data loggers, we compared their recorded temperatures to measured air temperatures to remove any data loggers with a clear offset. Thus, we are confident that our measurements are within ± 0.5 °C of actual T_b values. We then coated the data loggers with biologically inert wax and had them implanted intraperitoneally by a licensed veterinarian following standard procedures. The veterinarian implanted data loggers after anesthetizing animals with isoflurane and sutured the incision with absorbable catgut. Animals were given a long-acting, broad-spectrum antibiotic (Oxytetracycline, Pfizer) to minimize chances of postsurgery infection and were held for at least 7 d to ensure recovery from surgeries. Finally, to facilitate recapture, we clipped a few spines and glued (Torbot Liquid Bonding Cement, Torbot) radio transmitters (RI-2B, 10 g, Holohil Systems) to the dorsal skin. Combined, the data logger and transmitter were well below 5% body mass of all individuals. We released the animals in the same area where we had captured them and attempted to recapture them approximately 2 mo after the first capture period and 5 mo after the second capture period. The first period was completed wholly during summer. The second period began in early winter and ended in late spring/early summer. Recapture was lower during summer because hedgehogs more commonly left the area or were killed on roads. We euthanized animals using ether to retrieve data loggers. These capture procedures were conducted under permits issued by the Saudi Wildlife Authority. Experimental protocols were approved by the Animal Use and Care Committee of the University of Pretoria (ethics clearance EC029-

We calculated basic descriptive statistics for each individual (modal, minimum, and maximum T_b) as well as the heterothermy index (HI), which is a biologically meaningful modification of a simple standard deviation that quantifies the amount of variation around the modal T_b instead of the mean T_b (Boyles et al. 2011b) and allows for comparison of thermoregulatory patterns of species with diverse physiologies (Boyles et al. 2013). During summer, we recorded hourly air temperature (T_a) in the shade using an iButton. We recorded T_a in the same way during the second study period but lost the data, so we downloaded air temperature from a weather station approximately 27 km northwest of the study site (Prince Nayef bin Abdulaziz Regional Airport). Air temperatures on the study site may therefore be slightly

different than reported, and we make only qualitative comparisons between T_a and T_b during winter. We compared modal, minimum, and maximum $T_{\rm b}$ and HI values between seasons and sexes using PROC GLM in SAS University Edition (SAS Institute, Cary, NC), after verifying that the data met all assumptions of that test. We used PROC MIXED to evaluate the effect of maximum T_a on maximum T_b . For all analyses with seasonal comparisons, we considered data for the first study period as summer (June 16-August 16, 2013), and we split the second study period into winter (January 7-March 15, 2014) and spring (March 16-May 30, 2014). We chose to split the winter and spring data on these dates somewhat arbitrarily on the basis of a noticeable shift in thermoregulatory patterns of every individual around mid-March. This also coincides with the period of increased activity reported for this species in captivity (Al-Musfir and Yamaguchi 2008). Because the data are from the same individuals, winter and spring data are not completely independent, but we could not account for individuals in the analyses because of the inclusion of summer data, which are from different individuals. We avoid quantification of torpor patterns based on the torpor cut-off method (Boyles et al. 2011a) and use only approximate descriptions of torpor patterns for the duration of torpor bouts. The data are available from the authors for analyses associated with future meta-analyses.

Results

We successfully recovered data for five individuals (three males, two females) during the first (summer) capture session and 13 individuals (nine males, four females) during the second (winter/ spring) capture session. There was more variation in descriptors of thermoregulatory patterns among seasons than between sexes (table 1; fig. 1). Body temperatures were normally distributed when pooled by season but were clearly skewed when pooled by hour within season (fig. 2). Generally speaking, T_b's were highest in late afternoon during summer (fig. 2). Mean modal T_b 's were significantly higher during summer (P < 0.0001) than spring or winter, during which time modal T_b 's were nearly identical (35.8°

vs. 35.9°C; P = 0.8890; table 1). Mean maximum T_b 's varied significantly among seasons ($F_{2,25} = 112.2, P < 0.0001$) but were again similar in winter and spring (37.0° vs. 36.9°C; P = 0.5713). During summer, maximum T_b was significantly affected by maximum $T_a(F_{1,212} = 15.64, P = 0.0001)$, but there was no significant effect of individual ($F_{4,212} = 2.22$, P = 0.0684). Ignoring individual then, the equation for a simple regression of the effects of maximum T_a on maximum T_b is $T_b = 33.06 + 0.0998 T_a$ ($R^2 =$ 0.047, P = 0.0012). The single warmest T_b recorded was 39.2°C. Minimum T_b 's were significantly different in all seasonal comparisons (P < 0.0001 in all comparisons) in the expected order (summer minimum T_b > spring minimum T_b > winter minimum T_b ; table 1). The single minimum T_b recorded during winter was 10.9°C, and the tenth percentile was 14.9°C at 0900 hours during winter (fig. 2). During winter and spring, the lowest T_b 's were recorded early morning and were slightly out of phase with T_a (fig. 3). During summer, the lowest T_b 's were recorded just after sunrise.

Every individual commonly entered torpor during winter and spring, but none did so during summer (fig. 1). HI values were significantly different in all seasonal comparisons (P < 0.0001 in all comparisons) in the expected order (summer HI < spring HI < winter HI; table 1). Among all the included descriptors of thermoregulatory patterns, only HI values were significantly different between sexes ($F_{1,25} = 11.33$, P = 0.0025) and in the season × sex interaction ($F_{2,25} = 5.28$, P = 0.0122). The HI values for males and females were similar in summer and spring but were significantly higher for females (9.45°C) than males (5.85°C) in winter (P = 0.0003). All but 10 recorded torpor bouts were less than a daylong, but the longest bout was more than 101 h (fig. 1; estimated conservatively excluding cooling and warming phases of the bout). Although the number of torpor bouts longer than 24 h was small, they appear more common among females (six bouts by four individuals) than males (four bouts by nine individuals). There were several individuals that used multiday torpor bouts well into spring, including one individual that was torpid for ~50 h during May 18-20, 2014. Similar to shorter bouts in spring and in contrast to longer bouts in winter, T_b

Table 1: Summary statistics of thermoregulatory patterns in desert hedgehogs (Paraechinus aethiopicus) in Saudi Arabia

Season	n	Modal T_b	Heterothermy index	Minimum $T_{\rm b}$	Maximum $T_{\rm b}$
Summer:					
Male	3	$36.4 \pm .25$	$.76 \pm .19$	$34.0 \pm .40$	$38.6 \pm .57$
Female	2	$36.5 \pm .00$	$.82 \pm .07$	$34.0 \pm .84$	$39.2 \pm .01$
Combined	5	$36.5 \pm .18$	$.78 \pm .14$	$34.0 \pm .50$	$38.9 \pm .52$
Winter:					
Male	9	$35.8 \pm .23$	5.85 ± 1.76	13.7 ± 2.47	$37.0 \pm .16$
Female	4	$35.9 \pm .00$	$9.45 \pm .69$	$11.8 \pm .69$	$37.0 \pm .14$
Combined	13	$35.8 \pm .20$	6.96 ± 2.28	13.1 ± 2.23	$37.0 \pm .15$
Spring:					
Male	9	$35.9 \pm .26$	$3.63 \pm .86$	$22.3 \pm .32$	$36.9 \pm .27$
Female	4	$35.9 \pm .10$	$4.75 \pm .25$	$22.3 \pm .29$	$36.9 \pm .28$
Combined	13	$35.9 \pm .22$	$3.98 \pm .90$	$22.3 \pm .30$	$36.9 \pm .26$

Note. All euthermic and torpid data are included, and values are presented as means ± SD. T₁₀, body temperature.

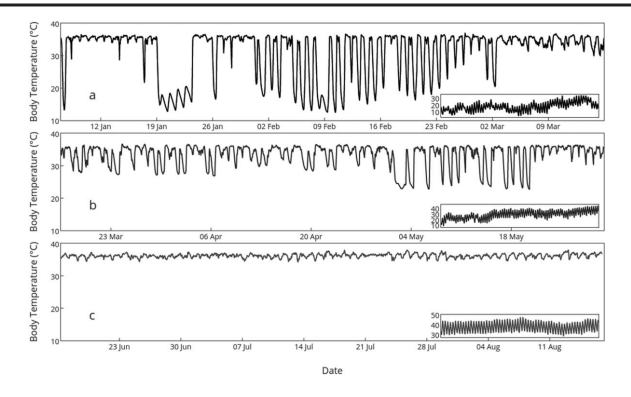


Figure 1. Representative body temperatures (T_b) of desert hedgehogs ($Paraechinus\ aethiopicus$) in Saudi Arabia during winter (January 7–March 15, 2014; a), spring (March 16–May 30, 2014; b), and summer (June 16–August 16, 2013). The temperature tracing depicted for winter was from a female and chosen to demonstrate both the normal pattern of T_b (as seen during most of February) and the longest torpor bout recorded in any individual (during mid-January). Data are T_b 's, and insets are ambient temperatures for the same periods (X-axes are not labeled for clarity). Note the different scales on the Y-axes of the insets.

remained above 22°C during these long spring torpor bouts. The lowest T_a recorded at the airport during any of these spring bouts was 22°C.

At some point during midwinter, nearly every individual went through a period of several days to several weeks with low nighttime $T_{\rm b}$'s but short torpor bouts (fig. 3). While we did not have $T_{\rm a}$ data from the site and specifically from within the burrows used by these animals, minimum $T_{\rm b}$'s often approached (and were occasionally below) $T_{\rm a}$ recorded at a nearby weather station. Unfortunately, without $T_{\rm a}$ data from the site, it is difficult to quantitatively analyze the $T_{\rm b}$ - $T_{\rm a}$ differential.

Discussion

The data on T_b patterns reported here are some of the first for a small desert mammal across multiple seasons. Desert hedgehogs maintain relatively low normothermic T_b 's during all seasons, with summer modal T_b 's of only 36.5°C (table 1). There was no indication of torpor during summer, but short torpor bouts with low T_b 's (<15°C) were common in winter, and short torpor bouts with moderate T_b 's (>22°C) were common in spring, when reproductive activity increases (Alagaili et al., forthcoming). Mammals often display larger variation in T_b in winter than summer (Chappell and Bartholomew 1981; Mustonen et al. 2007; Boyles et al. 2013), but the combination of strict homeothermy

during summer and short torpor bouts during winter is rarely reported for mammals (Ruf et al. 1989; Mzilikazi and Lovegrove 2004). More commonly discussed in the literature are species that (1) maintain homeothermy (or rarely use heterothermy) during summer and hibernate during winter (e.g., Zervanos and Salsbury 2003; Levesque and Tattersall 2010; Whiteman et al. 2015) or (2) use some form of heterothermy in all seasons (e.g., Geiser and Baudinette 1987; Bartels et al. 1998; Dzal and Brigham 2013). It is unclear whether these patterns are more common or are simply overrepresented in the literature because of geographic biases of researchers (McKechnie and Mzilikazi 2011). It seems logical that use of daily torpor during winter and homeothermy during summer should be common, especially among subtropical and desert species inhabiting environments that are hot during summer and either cold or offer variable food sources during winter. Among many likely candidates are those known to use short torpor bouts during winter but for which summer data are not available (e.g., Boyles et al. 2012; Cory Toussaint et al. 2010).

The daily cycle of $T_{\rm b}$'s was slightly out of phase with $T_{\rm a}$ during winter and spring, with $T_{\rm b}$'s still decreasing as $T_{\rm a}$'s were increasing in the morning (figs. 2, 3). The smallest $T_{\rm b}$ - $T_{\rm a}$ differentials were often several hours after the minimum $T_{\rm a}$ was recorded on a given day. This pattern was most pronounced on days when hedgehogs used torpor during winter, because it appeared that they allowed $T_{\rm b}$ to drop until the increasing heat of morn-

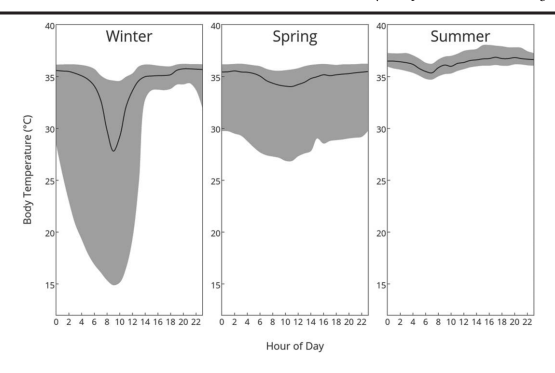


Figure 2. Body temperatures of desert hedgehogs (Paraechinus aethiopicus) across the day in winter, spring, and summer. Lines represent median T_b 's, and shaded areas are bounded by the tenth and ninetieth percentiles.

ing pushed it higher. The cyclical pattern was also seen during multiday torpor bouts but to a muted degree (e.g., figs. 1, 3); it is unclear whether T_b cycles during multiday torpor are driven by intrinsic or extrinsic factors. Occasionally, T_b dropped below recorded T_a during torpor bouts. We do not have behavioral observations of these animals during the periods when we were recording $T_{\rm b}$, so we cannot distinguish between two possible explanations for this pattern: either the weather station temperatures were slightly different than those at the study site, or hedgehogs were using burrows or other structures that buffered T_a (e.g., dead woody material, which we know they do use). Previous work on captive desert hedgehogs confirms that they are nocturnal and inactive during the day (Al-Musfir and Yamaguchi 2008), so the recorded T_b pattern is not surprising. From an energetic standpoint, it would be most efficient to cool into torpor as quickly as possible and remain in torpor as long as possible. Behavioral observations will be useful to determine whether the quick increases in T_b in the morning are related to adaptive behavior (e.g., basking) or reflect minimum available environmental temperatures, even in burrows.

Despite extensive laboratory work on thermoregulation of desert mammals, data on summer T_b patterns of small desert mammals are surprisingly rare. During summer, most variation in T_b occurs above normothermy. The diurnal species studied in the field occasionally reach T_b 's greater than 41°C (Chappell and Bartholomew 1981; Elvert et al. 1999; Alagaili et al. 2017). The few nocturnal species studied in the field do show regular increases in T_b above normothermy but reach maximum T_b 's several degrees lower than diurnal species (Elvert et al. 1999; this study). It remains unclear whether these increases in T_b represent an adaptive water conservation technique or are simply a biophysical reality for a small animal in a hot environment (Walsberg 2000).

The torpor patterns we recorded for desert hedgehogs during winter are intermediate to those classically described for hibernators and daily heterotherms (Geiser and Ruf 1995). The duration of most torpor bouts we recorded for hedgehogs was similar to daily torpor bouts used by daily heterotherms, but torpid T_b 's were regularly lower than those of daily heterotherms, especially for an animal that weighs 150-600 g (Geiser and Ruf 1995). Furthermore, while most torpor bouts were less than 24 h long, several individuals underwent bouts of 2, 3, or 4 d, which is unusual among daily heterotherms. Body temperatures often approached T_a , suggesting that metabolic rates must have been low (Geiser and Mzilikazi 2011). Conversely, while minimum torpid T_b 's were similar to those of some hibernators, torpor bouts were unpredictable, and no individual ever maintained a pattern of long torpor bouts interrupted by occasional euthermic periods, as is characteristic of hibernators. The pattern we recorded in desert hedgehogs is unusual among the hedgehogs studied to date, which show longer and more predictable torpor bouts during winter (Fowler and Racey 1990; Hallam and Mzilikazi 2011; Mouhoub-Sayah et al. 2012). This may relate to the more arid habitat of desert hedgehogs compared with the other species and the unpredictability of food resources in this habitat.

In the strictest sense, one could argue that the desert hedgehog is an unusual species that mixes physiological characteristics of hibernators and daily heterotherms, bringing to three the number of genera known to use such a pattern of torpor. Along with desert hedgehogs, Patagonian opossums (Lestodelphys halli; Geiser and

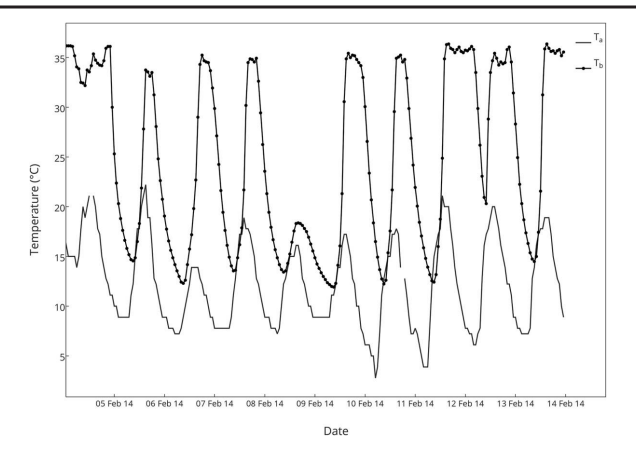


Figure 3. Representative winter body temperature (T_b) of a male desert hedgehog ($Paraechinus\ aethiopicus$) over a 10-d period during February 4–13, 2014. Ambient temperature (T_a) comes from a weather station approximately 27 km northwest of the study site (Prince Nayef bin Abdulaziz Regional Airport). Body temperatures were similar for females, but the T_b - T_a differentials were generally smaller during deep torpor than seen in males.

Martin 2013) and several sengis in the genus *Myurus* (also known as elephant shrews and locally as klaasneus) meet this strict criterion (Mzilikazi and Lovegrove 2004; Geiser and Mzilikazi 2011; Boyles et al. 2012). While ancestral hedgehogs appear in the fossil record shortly after the Cretaceous-Paleogene extinction event (O'Leary et al. 2013), the radiation of modern hedgehog genera was sudden and much more recent (Bannikova et al. 2014). The ecology of desert hedgehogs suggests they are derived from more temperate and semiarid species, so the unusual thermoregulatory pattern might also be derived. If so, this would be interesting because it has been suggested that this unusual pattern may be representative of an ancestral state in sengis and opossums (Geiser and Martin 2013).

A more liberal interpretation of the pattern of torpor we recorded in desert hedgehogs is that this pattern is not truly unique and represents another point along a gradient of thermoregulatory patterns (Boyles et al. 2013; van Breukelen and Martin 2015). If so, it strengthens the argument that greater variation in thermoregulatory patterns exists than traditionally realized and that desert hedgehogs are simply another exception to a strict classification scheme that separates daily heterotherms and hibernators (Canale et al. 2012). It is true that most arguments for a thermoregulatory continuum are based on measurements of

 $T_{\rm b}$, and other characteristics of thermoregulation (e.g., metabolic rates) need to be detailed more thoroughly in species with $T_{\rm b}$ patterns intermediate to daily heterotherms and hibernators.

The thermoregulatory pattern we measured herein adds interesting variation to thermoregulatory patterns seen among the Erinaceidae (hedgehogs and gymnures) and Eulipotypha and further solidifies suggestions that these taxa are interesting for thermoregulatory studies (e.g., Lovegrove 2012b). Even with the Erinaceidae, the opportunities to test predictions of evolutionary and ecological drivers of thermoregulation abound. For example, the divergence of gymnures (subfamily Galericinae) appears quite old, happening in the middle to late Eocene, while the extant hedgehogs radiated recently (Bannikova et al. 2014). Thus, within this single family, there is a diverse and contrasting evolutionary history (Bannikova et al. 2014). While conflated with phylogeny, there is also variation in habitat (forest, grassland, and desert) and latitude (tropical to high temperate) within this family. Finally, there is wide variation in thermoregulatory patterns, ranging from apparent homeothermy in lesser gymnures (Hylomys suillus; Genoud and Ruedi 1996) to deep hibernation in temperate hedgehogs. Despite the logistical difficulties associated with studying many species in this taxon, we encourage further study of thermoregulatory patterns of the Eulipotyphla to inform

our understanding of evolution of thermoregulation in laurasiatherians, eutherians, and mammals as a whole.

Acknowledgments

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