Responding to the weather: energy budgeting by a small mammal in the wild

Taylor Hume, Fritz Geiser, Shannon E. Currie, Gerhard Körtner, and Clare Stawski

Abstract

Energy conservation is paramount for small mammals because of their small size, large surface area to volume ratio, and the resultant high heat loss to the environment. To survive on limited food resources and to fuel their expensive metabolism during activity, many small mammals employ daily torpor to reduce energy expenditure during the rest phase. We hypothesized that a small terrestrial semelparous marsupial, the brown antechinus Antechinus stuartii, would maximize activity when foraging conditions were favorable to gain fat reserves before their intense breeding period, but would increase torpor use when conditions were poor to conserve these fat reserves. Female antechinus were trapped and implanted with small temperature-sensitive radio transmitters to record body temperature and to quantify torpor expression and activity patterns in the wild. Most antechinus used torpor at least once per day over the entire study period. Total daily torpor use increased and mean daily body temperature decreased significantly with a reduction in minimum ambient temperature. Interestingly, antechinus employed less torpor on days with more rain and decreasing barometric pressure. In contrast to torpor expression, activity was directly related to ambient temperature and inversely related to barometric pressure. Our results reveal that antechinus use a flexible combination of physiology and behavior that can be adjusted to manage their energy budget according to weather variables.

Key words: antechinus, heterothermy, marsupial, semelparous, temperature, torpor

The ability of endothermic mammals to maintain and regulate a constant high body temperature ($T_b$) that is largely independent from ambient temperature ($T_a$) provides many advantages, including the capability to be active at night and throughout all seasons (Scholander et al. 1950; Tattersall et al. 2012; Withers et al. 2016). However, in small mammals, this can be problematic because their surface area to volume ratio is greater than large mammals. As heat loss is a function of both the relative surface area and the $T_b - T_a$ differential, small mammals use proportionally more energy for thermoregulation at low $T_a$, and therefore have higher metabolic demands relative to their body mass than large mammals when maintaining a high and constant $T_b$ (Tattersall et al. 2012; Withers et al. 2016).

To be able to cope with these energetic costs, many small mammals employ torpor, during which their metabolic rate, $T_m$ heart rate, and other physiological functions are substantially reduced (Ruf and Geiser 2015). Torpor is often expressed to conserve energy in response to adverse environmental conditions, such as low $T_a$, food reduction, and heavy rainfall (Boyer and Barnes 1999; Ruf and Geiser 2015). Torpor is also crucial for survival in response to sudden changes in the environment, such as fires and storms (Nowack et al. 2015; Stawski et al. 2017).
2015; Geiser et al. 2018). Furthermore, by lowering energy demands, torpor use can allow individuals to reduce foraging time, which may reduce predation and starvation risk (Stawski and Geiser 2010; Hanna and Cardillo 2014; Turbill and Stojanowski 2018).

Torpor expression fluctuates with changes in resource availability, competition, risks of predation, and the costs of dedicating energy to foraging. Animals may therefore trade-off time spent active by employing torpor to avoid these problems if they outweigh the benefits gained by foraging (Stawski and Geiser 2010; Levy et al. 2012). This ability to identify risks is important for small mammals to respond appropriately to current and approaching weather conditions; deciding whether it would be more beneficial for them to forage or to employ torpor on any given day. Endotherms are able to sense changes in $T_\text{a}$ using primary sensory neurons through the periphery in their skin (Glotzbach and Heller 1976). Some placental mammals can sense changes in barometric pressure, which often precedes rainfall or inclement weather that may interrupt food availability (Vickery and Bider 1981; Czence and Willis 2015); however, data for marsupials are not available.

To determine how small mammals respond to, or predict, variations in weather patterns, we quantified torpor expression and activity patterns of free ranging female brown antechinus Antechinus stuartii, during the Austral winter in relation to $T_\text{a}$, precipitation and changes in barometric pressure. Brown antechinus are small heterothermic insectivorous marsupials from the family Dasyuridae and are distributed along the eastern coast and ranges of Australia (Geiser 1988). They are short-lived marsupials that undergo an almost complete yearly population turnover (Woodley 1966; McAllan and Dickman 1986). Males die shortly after expending excessive energy during a brief 2-week mating period in late winter that leads to starvation and increased parasite loads, whereas most females die after their young have been weaned (Woodley 1966); however, one-third of females may survive to reproduce for a second breeding season (Rojas et al. 2014). We hypothesized that torpor would be employed extensively by females leading up to the energetically demanding reproductive period as they are typically food-limited at this time (Dickman 1989; Stawski and Rojas 2016). Furthermore, we predicted that torpor expression would be greater in response to challenging environmental conditions, such as low $T_\text{a}$ and rainfall.

### Material and Methods

Our study site was located at Guy Fawkes River National Park, NSW, Australia (30°04’58.6”S, 152°20’0.9”E); the work was conducted between 18 May and 27 June 2015 during the austral autumn/winter. This time period is ~2–3 months before the 2-week breeding period that occurs during August (known from trapping survey data conducted throughout the year at the study site). Permits from the NSW National Parks and Wildlife Services and the University of New England animal ethics committee (AEC13-088) were obtained before commencing research. $T_\text{a}$ was recorded every 10 min using a temperature data logger (Thermochron, DS 1921 G, Maxim, resolution 0.5°C, Rio Robles, CA, USA) tied to a small branch in the shade ~1 m above the ground in the study site. Rainfall and barometric pressure variables were obtained from a NSW National Parks and Wildlife Service weather station positioned ~2 km from the field site.

We used aluminium box traps (Elliott Scientific Equipment, Upwey, Australia) to capture a total of 17 female $A. \text{stuartii}$. The traps were baited with a mixture of peanut butter, oats, and honey and insulated with polyester fiber to prevent hypothermia during overnight trapping. During the afternoon, we set up 9 trap lines of 25 traps (a total of 225 traps each night). We set each trap ~4–5 m apart and checked them early the next morning. Captured study animals were later transported to the University of New England (130 km and ~2 h drive) where they were weighed to the nearest 0.1 g using electronic scales (mean capture body mass: 23.4 ± 2.2 g, $n = 17$) and housed individually in cages overnight from the day of capture. Individuals were implanted with a temperature-sensitive transmitter with a unique frequency for animal identification (2.0–2.3 g, Sirtrack, Havelock North, New Zealand) under isoflurane/oxygen anesthesia as described by Rojas et al. (2010). Transmitter pulse rate as a function of temperature was calibrated to the nearest 0.1°C between 15°C and 40°C, at 5°C increments in a temperature-controlled water bath. Prior to implantation, we coated each transmitter in wax (Paraffin/Elvax, Minimitter, Bend, Oregon, USA) and spray bandage (BSN Medical (Aust) Pty Ltd, Clayton, VIC, Australia). The reproductive age of the animals was recorded at time of surgery, however, behavior and physiology between first and second reproductive year females did not differ. For post-surgery recovery, the antechinuses were kept in a warm room and fed cat food, meal worms and water before being released at the site of capture the next day.

To locate each individual, we radio-tracked the antechinuses each morning over the initial 40 days, and then once a week for a further 30 days using receivers (ICOM IC-R10, Icom Inc. Osaka, Japan) and Yagi antennae (Sirtrack, Havelock North, New Zealand). Overall, antechinuses were tracked for a mean of 47.7 ± 12.5 days depending on the battery life of the transmitter (range of 28–69 days).

During radio-tracking, manual $T_\text{b}$ readings using a stopwatch were recorded immediately when a signal was detected to ensure immediate data collection of $T_\text{b}$ before data loggers were deployed. Receiver/ loggers with antennas were setup near nest sites to record transmitter pulse intervals every 10 min. Each device was equipped with a scanner receiver, logger and a 12 V battery (Kortner and Geiser 2000) that was sealed in a plastic container and wrapped in a plastic bag. H-frame antennae were tied to a tree in a position to maximize signal detection. The batteries were exchanged once a week and the pulse rate data were downloaded. Pulse rates were converted to $T_\text{b}$ using a second-order polynomial equation ($r^2 = 0.999$) based on the individual transmitter calibration data.

We defined torpor as $T_\text{b} ≤ 31.5°C$, as calculated on the basis of Willis (2007); Equation 4: $T_\text{b, min} = 1$ SE = (0.041)BM + (0.040)$T_\text{a}$ + 31.092; SE = standard error, BM = body mass, $T_\text{a, min}$ (for study period) to enable the detection of shallow torpor bouts. Torpor bout duration was calculated as the time $T_\text{b}$ remained below the torpor threshold for durations >30 min. As antechinus often displayed more than 1 torpor bout/day (maximum of 5 torpor bouts), the total daily torpor (TDT) use in minutes for each study day was calculated for each animal from sunrise to sunrise (data from the Australian Bureau of Meteorology) and was used for analyses. The duration of activity bouts was calculated from the time the individual was out of the range of the receiver/locator (reception range ~5–
10 m), indicating that they had left the nest, until the time they returned. For analyses, we used the total daily activity (TDA) duration summed for each day of the study period between sunrise and sunrise. Only the days that an individual left and returned to the same nest were included. However, it is possible that during longer activity bouts the animal may have used another nest location for a short period of time.

Statistical analyses were conducted using R (Version 3.4.3, 2018). A \( P < 0.05 \) was considered significant and means are presented with \( \pm \) standard deviation (SD). We ran a Pearson’s correlation test on the weather variables and as all had a correlation statistic \( < 0.7 \) (i.e., Henning et al. 2005) we included all of the weather variables and first-order interaction terms (barometric pressure \( \times T_a\text{ min} \) \( = -0.32 \); barometric pressure \( \times \text{rain} \) \( = -0.04 \); \( T_s \text{ min} \times \text{rain} \) \( = 0.26 \). Linear mixed effects models were fitted (package “nlme”; Pinheiro et al. 2018) to determine the effect of \( T_a\text{ min} \) (°C), rainfall (mm) and absolute barometric pressure (hPa) on the measured variables (TDT [min], daily mean \( T_b\text{ mean} \), TDA [min]). Body mass (g) at capture was included as a covariate and multiple measures for individuals by including individual (ID) as a random effect. We tested for normality (Q–Q plot) and homoscedasticity (residual plot). For the initial model, we included interactions between the weather variables. The best-fit model was determined using Akaiki information criterion (AIC; Akaiki 1974) and performing a log likelihood ratio test on the top 2 models (package “MuMIn”; Bartoli 2013). Linear mixed effects models were also fitted to determine if the measured variables differed on days of changing barometric pressure. The change in barometric pressure was categorized as increasing (1–11.9hPa), decreasing (1 to \(-10.4\text{hPa})\), or null (\(-0.9 \text{ to } 0.9 \text{ hPa}\)), by subtracting each day’s barometric pressure from the previous day. Following this, a post hoc Tukey test (package “multcomp”; Hothorn et al. 2008) was employed to determine the difference.

Results

The \( T_s \) throughout the study period ranged from a minimum of \(-1.5\) °C to a maximum of \(18.0\) °C and the mean was \(7.5\) °C \( \pm \) 3.6 °C (\( n = 62 \); Table 1). Rainfall (mm) was recorded on 25 of the total 70 study days with a mean daily rainfall of 1.6 \( \pm \) 4.2 mm (\( n = 62 \); Table 1). A total of 115.8 mm of rain fell throughout the entire study period and the maximum amount of rain that fell on a single day was 24.0 mm.

The mean daily \( T_b\text{ mean} \) varied with \( T_s \) throughout the study period (Figure 1) and for all individuals was 32.6 °C \( \pm \) 0.8°C (\( n = 17 \), \( N = 793 \)), with an absolute lowest \( T_b\text{ min} \) of 18.1°C and an absolute highest \( T_b\text{ max} \) of 40.9°C. Our best-fit model (Table 2) included \( T_s\text{ min} \) rainfall, barometric pressure, and the interaction between \( T_s\text{ min} \) and rainfall. Therefore, the effect of \( T_s\text{ min} \) was mediated by rain, such that \( T_b\text{ mean} \) was lowest on cold and dry days (Figure 2). In addition, \( T_b\text{ mean} \) increased with lower barometric pressure and was therefore significantly higher on days of falling barometric pressure (\( z = -4.3, P < 0.0001; n = 17, N = 793 \); Table 3).

Torpor was expressed by all antechinus studied and torpor use was recorded on \(82.2 \pm 11.6\% \) (\( n = 17 \)) of study days. The mean TDT use throughout the whole study period (including days of torpor and no torpor) was \(330.8 \pm 100.9\text{min} \) (\( n = 17 \), \( N = 793 \)); the maximum TDT during a single day was 1,200 min when average \( T_s \) was 7.4°C. The best-fit model for TDT (Table 2) revealed that TDT was affected also by \( T_s\text{ min} \) rainfall, barometric pressure, and the interaction between \( T_s\text{ min} \) and rainfall. This interaction term revealed that TDT was longer on cold and dry days (Figure 3). A change in barometric pressure also significantly affected the TDT; on days of falling barometric pressure, less torpor was employed (\( z = 4.3, P < 0.0001; n = 17, N = 793 \); Table 3).

The mean TDA duration (sunrise to sunrise) was \(549.7 \pm 96.3\text{min} \) (\( n = 17 \), \( N = 794 \), ranging from 40 to 1,220 min. Opposite to torpor use, activity increased at higher \( T_s \) (Figure 4). The best-fit model (Table 2) for TDA also included rainfall and barometric pressure. However, the best-fit model for TDA differed from TDT and \( T_s\text{ mean} \) and did not include the interaction between \( T_s\text{ min} \) and rainfall. Conversely to TDT, the amount of time ante- chinus were active throughout the day was greater on days of falling barometric pressure (\( z = -3.4, P < 0.0001; n = 17, N = 794 \); Table 3).

Discussion

Our study revealed that thermal and foraging biology of free ranging \( A.\text{stuartii} \), 2–3 months before the breeding period, were affected by prevailing weather conditions. Torpor and activity were linked in fe-

male antechinus as torpor use was negatively correlated whereas ac-

tivity was positively correlated with \( T_s \). These results support previous observations that torpor and activity are governed by changes in the environment (Rojas et al. 2014; Stawski et al. 2015). Interestingly, antechinus in our study employed more torpor during periods of less rain and increasing barometric pressure. Therefore, our results suggest that physiological and behavioral flexibility is an important aspect of an individual’s response to sudden changes in the environment, such as inclement weather.

Torpor was used almost daily (\(82.2 \pm 11.6\% \) of all study days) by all female \( A.\text{stuartii} \) during the study period. Torpor expression increased at lower \( T_s \), resulting in lower daily \( T_b\text{ mean} \). Furthermore, and somewhat unexpected, antechinus spent less time torpid when it was raining and at low and decreasing barometric pressure. We recorded rainfall on 25 of the total 70 study days with 9 of these days recording >5.0 mm of rainfall. During such periods of prolonged rainfall, foraging returns would likely be limited and heat loss would increase, especially if the fur is wet (Withers et al. 2016). Therefore, one might assume that the animals would employ longer and deeper torpor during the rain event, but this was not the case. Instead, the longest and the deepest bouts of torpor occurred on a day that had been preceded by 4 days of significant rainfall (>5.0 mm; Figure 1).

TDA duration was longer at higher \( T_s \)s and decreasing barometric pressure. This may, in part, explain the torpor patterns we recorded. It is likely to be profitable for antechinus to forage during the warmer conditions preceding a weather front as insect activity is typically higher (Pellegrino et al. 2013; McFarlane et al. 2015; Dagaeff et al. 2016). Kaspari and Weiser (2000) found that ants were more active during wetter seasons that promote humid microclimates and the release of nutrients as result of moist leaf litter

Table 1. Summary of weather variables including means with SD and absolute minimum and maximum values recorded

| Variable | Mean | Absolute minimum | Absolute maximum |
|----------|------|------------------|------------------|
| \( T_s \) (°C) | 7.5 \( \pm \) 3.6 | -1.5 | 18.0 |
| \( T_{s\text{ max}} \) (°C) | 12.8 \( \pm \) 2.3 | 6 | 18.0 |
| \( T_{s\text{ min}} \) (°C) | 3.6 \( \pm \) 2.9 | -1.5 | 12.0 |
| Rainfall (mm) | 1.6 \( \pm \) 4.2 | 0.0 | 24.0 |
(McColl 1975). Furthermore, some rodents increase activity during rainfall if the temperature is higher (Vickery and Bider 1981). Rainy days were warmer and antechinus often forage under thick ground cover to likely avoid getting wet. In addition, cloudless nights with high barometric pressure will be colder and antechinus will likely have reduced foraging opportunities suggesting that it is more beneficial for antechinus to decrease activity and conserve energy by using torpor. This observation suggests that antechinus begin to employ more torpor and reduce activity as barometric pressure increases and reduce torpor with decreasing barometric pressure that often signals inclement weather. Therefore, the results of our study indicate that antechinus are more energy restricted in response to cold conditions rather than rain alone.

Other species such as some bats and birds appear to perceive and in some cases predict changing weather patterns through sensing barometric changes, but the method by which species can predict these changes is still poorly understood (Heupel et al. 2003; Wikramanayake et al. 2006; Breuner et al. 2013; Bender and Hartman 2015; Streby et al. 2015). Changes in barometric pressure usually foreshadow changes in $T_a$ and other weather variables (Breuner et al. 2013). Therefore, an individual’s ability to detect changes in barometric pressure and predict the associated weather patterns would allow it to precisely manage its energy budget and/or take refuge during a storm. A laboratory study showed that a decline in barometric pressure stimulated an increase in food intake in birds, suggesting they were preparing for a storm (Breuner et al. 2015). This is particularly important for insectivorous animals as many insects respond to changes in barometric pressure by altering their activity and mating behaviors (Pellegrino et al. 2013; McFarlane et al. 2015; Dagaeff et al. 2016). Indeed, insectivorous bats appear to be able to track changes in barometric pressure and adjust their activity accordingly (Dechmann et al. 2017), and it is likely that antechinus can do the same.

Limited data suggest that the paratympanic organ of the middle ear in vertebrates may function as a sensor for changes in barometric pressure (von Bartheld and Giannessi 2011). To date, this organ has

### Table 2. The 4 best-fit models with variables (minimum ambient temperature $= T_{a_{min}}$ [°C], rainfall = rain [mm], absolute barometric pressure $= bp$ [hPa], body mass $= bm$ [g]) that explain differences in TDT use (min), mean daily body temperature ($T_b$ mean, °C), and TDA duration (min)

| Variable | Model | AICc | Delta | $P$ |
|----------|-------|------|-------|-----|
| $T_b$ mean | $T_{a_{min}} + \text{rain} + bp + bm + T_{a_{min}} \times \text{rain}$ | 2899.9 | 0.00 | 0.005 |
| | $T_{a_{min}} + \text{rain} + bp + bm$ | 2901.3 | 1.35 | |
| | $T_{a_{min}} + \text{rain} + bp + bm + T_{a_{min}} \times \text{rain} + T_{a_{min}} \times bp$ | 2909.6 | 9.65 | |
| | $T_{a_{min}} + \text{rain} + bp + bm + T_{a_{min}} \times \text{rain} + T_{a_{min}} \times bp + \text{rain} \times \text{bp}$ | 2920.9 | 21.02 | |
| TDT | $T_{a_{min}} + \text{rain} + bp + bm + T_{a_{min}} \times \text{rain}$ | 10488.5 | 0.00 | 0.264 |
| | $T_{a_{min}} + \text{rain} + bp + bm + T_{a_{min}} \times \text{rain} + T_{a_{min}} \times bp$ | 10489.3 | 0.81 | |
| | $T_{a_{min}} + \text{rain} + bp + bm + T_{a_{min}} \times \text{rain} + T_{a_{min}} \times bp + \text{rain} \times \text{bp}$ | 10489.9 | 1.34 | |
| | $T_{a_{min}} + \text{rain} + bp + bm$ | 10492.9 | 4.40 | |
| TDA | $T_{a_{min}} + \text{rain} + bp + bm$ | 10917.7 | 0.00 | 0.156 |
| | $T_{a_{min}} + \text{rain} + bp + bm + T_{a_{min}} \times \text{rain}$ | 10917.7 | 0.03 | |
| | $T_{a_{min}} + \text{rain} + bp + bm + T_{a_{min}} \times \text{rain} + T_{a_{min}} \times bp$ | 10918.5 | 0.81 | |
| | $T_{a_{min}} + \text{rain} + bp + bm + T_{a_{min}} \times \text{rain} + T_{a_{min}} \times bp + \text{rain} \times \text{bp}$ | 10918.9 | 1.20 | |

The $P$-value is from a log likelihood ratio test performed for the 2 top models. The best-fit model for each variable is indicated in bold.
primarily been found and described not only in birds, but also in 1 bat species (von Bartheld and Giannessi 2011). We were unable to find any literature on marsupials with regard to this organ, but it has been suggested that some mammals may be able to use the hair cells of the inner ear to detect pressure changes (Funakubo et al. 2010). Therefore, more research is needed to determine the mechanism by which marsupials may be able to detect changes in barometric pressure, the sensitivity of this mechanism, and if they are able to predict weather patterns using it.

Overall, our study suggests that *A. stuartii* alter torpor expression and activity according to prevailing and oncoming environmental conditions which may enhance their survival during weather changes. Being able to predict and respond to changes in the weather provides many survival benefits for small mammals like antechinus, primarily by balancing foraging and fat accumulation when conditions are favorable and employing torpor during adverse weather for energy conservation. This is particularly important for antechinus that have a single breeding period, as it is likely they attempt to build up fat reserves before the energy demanding reproductive period in late winter and early spring.

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**Authors’ Contributions**

T.H., F.G. and C.S. conceived the original research proposal. All authors conducted the fieldwork to collect the data. T.H. and C.S. analyzed the data and wrote the original article. All authors contributed to further drafts of the manuscript and accepted the final version.

**References**

Akaike H, 1974. A new look at the statistical model identification. *IEEE Trans Aut Control* 19:716–723.
Barton K, 2013. MuMIn: multi-model inference. R package version 1.9.13.
Bender MJ, Hartman GD, 2015. Bat activity increases with barometric pressure and temperature during autumn in central Georgia. *Southeast Nat* 14:231–224.

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**Table 3.** Means with SD of TDT use, mean daily body temperature (*T*<sub>b</sub> mean), and TDA duration on days of decreasing and increasing barometric pressure (BP)

| Variable       | Decreasing BP | Increasing BP | *P*       |
|----------------|---------------|---------------|-----------|
| *T*<sub>b</sub> mean (°C) | 32.9 ± 0.7    | 32.2 ± 0.9    | <0.0001   |
| TDT (min)      | 291.9 ± 80.2  | 373.5 ± 121.2 | <0.0001   |
| TDA (min)      | 595.1 ± 99.7  | 502.7 ± 101.5 | <0.0001   |

Bold *P*-value indicates a significant difference.

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**Figure 2.** 3D scatterplot of daily mean resting body temperature (*T*<sub>b</sub> mean) as a function of daily minimum ambient temperature (*T*<sub>a</sub> min) and rainfall.

**Figure 3.** 3D scatterplot of TDT use as a function of daily minimum ambient temperature (*T*<sub>a</sub> min) and rainfall.

**Figure 4.** 3D scatterplot of TDA duration as a function of daily minimum ambient temperature (*T*<sub>a</sub> min) and rainfall.
Boyler BB, Barnes BM, 1999. Molecular and metabolic aspects of mammalian hibernation. *Bioscience* 49:713–724.

Breuner CW, Sprague RS, Patterson SH, Woods HA, 2013. Environment, behavior and physiology: do birds use barometric pressure to predict storms? *J Exp Biol* 216:1982–1990.

Czene ZJ, Willis CRK, 2015. Warming up and shipping out: arousal and emergence timing in hibernating little brown bats *Myotis lucifugus*. *J Comp Physiol B* 185:575–586.

Dagaeff AC, Pocheville A, Nöbel S, Loyau A, Isabel G et al., 2016. *Drosophila* mate copying correlates with atmospheric pressure in a speed learning situation. *Anim Behav* 121:163–174.

Dechmann DKN, Wikelski M, Ellis-Soto D, Safi K, O’Mara MT, 2017. Determinants of spring migration departure decision in a bat. *Biol Lett* 13: 20170395.

Dickman CR, 1989. Demographic responses of *Antechinus stuartii* (Marsupialia) to supplementary food. *Aust J Ecol* 14:387–398.

Funakubo M, Sato J, Honda T, Mizumura K, 2010. The inner ear is involved in the aggravation of nociceptive behavior induced by lowering barometric pressure of nerve injured rats. *Eur J Pain* 14:32–39.

Geiser F, 1988. Daily torpor and thermoregulation in *Antechinus* (Marsupialia): influence of body mass, season, development, reproduction, and sex. *Oecologia* 77:395–399.

Geiser F, Stawski C, Doty AC, Cooper CE, Nowack J, 2018. A burning question: what are the risks and benefits of mammalian torpor during and after fire? *Cons Physiol* 6:e000787.

Glotzbach SF, Heller HC, 1976. Central nervous regulation of body temperature during sleep. *Science* 194:537–539.

Hanna E, Cardillo M, 2014. Clarifying the relationship between torpor and anthropogenic extinction risk in mammals. *J Zool* 293:211–217.

Henning J, Schmitzler FR, Pfeiffer DU, Davies P, 2005. Influence of weather conditions on fly abundance and its implications for transmission of rabbit haemorrhagic disease virus in the North Island of New Zealand. *Med Vet Entomol* 19:251–262.

Heupel MR, Simpfendorfer CA, Hueter RE, 2003. Running before the storm: blacktip sharks respond to falling barometric pressure associated with tropical storm Gabrielle. *J Fish Biol* 63:1357–1363.

Hothorn T, Bretz F, Westfall P, 2008. Simultaneous inference in general parametric models. *Biometrical J* 50:346–363.

Kaspari M, Weisser MD, 2000. Ant activity along moisture gradients in a neotropical forest. *Biotropica* 32:703–711.

Körtner G, Geiser F, 2000. Torpor and activity patterns in free-ranging sugar gliders *Petaurus breviceps* (Marsupialia). *Oecologia* 123:350–357.

Levy O, Dayan T, Rottes S, Kronfeld-Schor N, 2012. Foraging sequence, energy intake and torpor: an individual-based field study of energy balancing in desert golden spiny mice. *Ecol Lett* 15:1240–1248.

McAllan BM, Dickman CR, 1986. The role of photoperiod in the timing of reproduction in the dasyurid marsupial *Antechinus stuartii*. *Oecologia* 68:259–264.

McColl HP, 1975. The invertebrate fauna of the litter surface of a Nothofagus truncata forest floor, and the effect of microclimate on activity. *New Z J Zool* 2:115–134.

McFarlane DJ, Rafter MA, Booth DT, Walter GH, 2015. Behavioral responses of a tiny insect, the flower thrips *Frankliniella schultzei* Trybom (Thysanoptera, Thripidae), to atmospheric pressure change. *J Insect Behav* 28:473–481.

Nowack J, Rojas AD, Körtner G, Geiser F, 2015. Snoozing through the storm: torpor use during a natural disaster. *Sci Rep* 5:11243.

Pellegrino AC, Milgr P, Nardi C, Bezner-Kerr W, Guglielmo CG et al., 2013. Weather forecasting by insects: modified sexual behaviour in response to atmospheric pressure changes. *PLoS ONE* 8:e75004.

Pinheiro J, Bates D, DebRoy S, Sarkar D, R Core Team, 2018. *nlme*: linear and nonlinear mixed effects models. R package version 3.1–137.

Rojas AD, Körtner G, Geiser F, 2010. Do implanted transmitters affect maximum running speed of two small marsupials? *J Mammal* 91:1360–1364.

Rojas AD, Körtner G, Geiser F, 2014. Torpor in free-ranging antechinus: does it increase fitness? *Naturwissenschaften* 101:105–114.

Ruf T, Geiser F, 2015. Daily torpor and hibernation in birds and mammals. *Biol Rev* 90:891–926.

Scholander PF, Hock R, Walters V, Johnson F, Irving L, 1950. Heat regulation in some arctic and tropical mammals and birds. *Biol Bull* 99:237–258.

Stawski C, Geiser F, 2010. Fat and fed: frequent use of summer torpor in a subtropical bat. *Naturwissenschaften* 97:29–35.

Stawski C, Rojas AD, 2016. Thermal physiology of reproductive female marsupial, *Antechinus flavipes*. *Mammal Res* 61:417–421.

Stawski C, Körtner G, Nowack J, Geiser F, 2015. The importance of mammalian torpor for survival in a post-fire landscape. *Biol Lett* 11:20150134.

Streby HM, Kramer GR, Peterson SM, Lehman JA, Buchler DA et al., 2015. Tornadic storm avoidance behavior in breeding songbirds, *Carr Brid* 25:98–102.

Tattersall GJ, Sinclair BJ, Withers PC, Fields PA, Seebacher F et al., 2012. Coping with thermal challenges: physiological adaptations to environmental temperatures. *Compr Physiol* 2:2151–2202.

Turrell C, Stojanovski L, 2018. Torpor reduces predation risk by compensating for the energetic cost of antipredator foraging behaviours. *Proc R Soc B* 285:20182370.

Vickery WL, Bider JR, 1981. The influence of weather on rodent activity. *J Mammal* 62:140–145.

von Bartheld CS, Giannessi F, 2011. The paratympanic organ: a barometer and altimeter in the middle ear of birds? *J Zool* 285:402–408.

Wikramanayake E, Fernando P, Leimgruber P, 2006. Behavioral response of satellite-collared elephants to the tsunami in southern Sri Lanka. *Biotropica* 38:775–777.

Willis CRK, 2007. An energy-based body temperature threshold between torpor and normothermia for small mammals. *Physiol Biochem Zool* 80:643–651.

Withers PC, Cooper CE, Maloney SK, Bozinovic F, Cruz-Neto JP, 2016. *Ecological and Environmental Physiology of Mammals*. Oxford: Oxford University Press.

Woolley P, 1966. Reproduction in *Antechinus* spp. and other dasyurid marsupials. *Symp Zool* 15:281–294.