

## LJMU Research Online

Nowack, J, Stawski, C and Geiser, F

More functions of torpor and their roles in a changing world

http://researchonline.ljmu.ac.uk/9291/

Article

**Citation** (please note it is advisable to refer to the publisher's version if you intend to cite from this work)

Nowack, J, Stawski, C and Geiser, F (2017) More functions of torpor and their roles in a changing world. Journal of Comparative Physiology B, 187 (5-6). pp. 889-897. ISSN 0174-1578

LJMU has developed LJMU Research Online for users to access the research output of the University more effectively. Copyright © and Moral Rights for the papers on this site are retained by the individual authors and/or other copyright owners. Users may download and/or print one copy of any article(s) in LJMU Research Online to facilitate their private study or for non-commercial research. You may not engage in further distribution of the material or use it for any profit-making activities or any commercial gain.

The version presented here may differ from the published version or from the version of the record. Please see the repository URL above for details on accessing the published version and note that access may require a subscription.

For more information please contact researchonline@ljmu.ac.uk

http://researchonline.ljmu.ac.uk/

# *More functions of torpor and their roles in a changing world*

# Julia Nowack, Clare Stawski & Fritz Geiser

#### **Journal of Comparative Physiology B**

Biochemical, Systems, and Environmental Physiology

ISSN 0174-1578 Volume 187 Combined 5-6

J Comp Physiol B (2017) 187:889-897 DOI 10.1007/s00360-017-1100-y





Your article is published under the Creative Commons Attribution license which allows users to read, copy, distribute and make derivative works, as long as the author of the original work is cited. You may selfarchive this article on your own website, an institutional repository or funder's repository and make it publicly available immediately.



ORIGINAL PAPER



### More functions of torpor and their roles in a changing world

Julia Nowack<sup>1,2</sup> · Clare Stawski<sup>1</sup> · Fritz Geiser<sup>1</sup>

Received: 26 August 2016 / Revised: 26 November 2016 / Accepted: 26 February 2017 / Published online: 21 April 2017 © The Author(s) 2017. This article is an open access publication

**Abstract** Increased winter survival by reducing energy expenditure in adult animals is often viewed as the primary function of torpor. However, torpor has many other functions that ultimately increase the survival of heterothermic mammals and birds. In this review, we summarize new findings revealing that animals use torpor to cope with the conditions during and after natural disasters, including fires, storms, and heat waves. Furthermore, we suggest that torpor, which also prolongs longevity and was likely crucial for survival of mammals during the time of the dinosaur extinctions, will be advantageous in a changing world. Climate change is assumed to lead to an increase in the occurrence and intensity of climatic disasters, such as those listed above and also abnormal floods, droughts, and extreme temperatures. The opportunistic use of torpor, found in many heterothermic species, will likely enhance survival of these challenges, because these species can reduce energy and foraging requirements. However, many strictly seasonal hibernators will likely face the negative consequences of the predicted increase in temperature, such as range contraction. Overall, available data suggest that opportunistic heterotherms with their flexible energy

Communicated by F. van Breukelen.

This manuscript is part of the special issue Hibernation—Guest Editors: Frank van Breukelen and Jenifer C. Utz.

☐ Julia Nowack Julia.Nowack@vetmeduni.ac.at

- <sup>1</sup> Centre for Behavioural and Physiological Ecology, Zoology, University of New England, Armidale, NSW 2351, Australia
- <sup>2</sup> Department of Integrative Biology and Evolution, University of Veterinary Medicine, Vienna, Savoyenstraße 1, Vienna 1160, Austria

requirements have an adaptive advantage over homeotherms in response to unpredictable conditions.

Keywords Colonization  $\cdot$  Evolution  $\cdot$  Fires  $\cdot$  Heat  $\cdot$  Heterothermy  $\cdot$  Storm

#### Introduction

The reduction of energy expenditure, achieved by a controlled reduction of metabolic rate and body temperature  $(T_{\rm b})$ , during cold periods that often coincide with low food availability is frequently viewed as the ultimate benefit of torpor (Lyman et al. 1982). Indeed, many species use torpor mainly in response to such seasonal bottlenecks (Brigham et al. 2000). However, in the past decade, it has become evident that torpor has more functions than just energy conservation for adult mammals and birds during winter (reviewed in Geiser and Brigham 2012). For example, torpor can enhance fat storage during migration, or permit birds to remain resident during winter and maintain and defend their territories year round (Geiser and Brigham 2012), to name only a few additional functions of torpor (for more functions see Table 1). Since the original suggestion of "other functions" of torpor by Geiser and Brigham (2012), the list has continued to grow (Table 1, 2). Recently identified functions include coping with extreme heat (Bondarenco et al. 2014), facilitating (island) colonization (Nowack and Dausmann 2015), enabling mammalian survival at the Cretaceous-Palaeogene (K-Pg) boundary (Lovegrove et al. 2014a) and perhaps a prolonged lifespan (Lyman et al. 1981; Turbill et al. 2011). Importantly, one of the major recent findings adding to the varied benefits of torpor is that animals can use torpor to cope with detrimental conditions during and after natural disasters,

 Table 1
 List of the other functions of torpor that were already listed in Geiser and Brigham 2012 including additional supporting references that were published after 2012

such as fires and storms (e.g., Nowack et al. 2015; Stawski et al. 2015a). It has already been noted previously that the reduction in metabolic rate during torpor lowers the need of food intake, likely enabling animals to decrease the time spent actively foraging (Geiser and Brigham 2012). Heterothermic species (those that can employ torpor or those that increase/decrease their  $T_{\rm b}$  only slightly above/below normothermic levels) are less often subjects of extinction than homeothermic species (that maintain a stable high  $T_{\rm b}$ ), suggesting that the reduction of energy expenditure and its additional benefits substantially increase the chance of survival of individuals (Geiser and Turbill 2009; Hanna and Cardillo 2014). Hibernating mammals can survive for months without food and water intake in their hibernacula, where they are somewhat independent of the conditions on the earth's surface. Edible dormice (Glis glis) are an extreme example as they hibernate for 11 months in nonreproductive years, presumably to avoid predation during years when conditions are not favorable for reproduction (Bieber and Ruf 2009; Hoelzl et al. 2015). Similarly, eastern pygmy-possums (Cercartetus nanus) can hibernate for up to an entire year without feeding (Geiser 2007). We hypothesize that heterothermic mammals may have an advantage over homeothermic species as they can remain hidden and protected during short or long periods of torpor. Moreover, it is likely that opportunistic torpor will be advantageous in a world subjected to climate change. In this paper, we review the functions of torpor that have been reported since the original review by Geiser and Brigham (2012) and discuss whether and how some of these functions may help animals to deal with the predicted environmental changes.

# Heterothermy, extreme heat, and water conservation

Both severe cold and extreme heat result in thermal stress for organisms. From a short-term survival point of view, extreme heat is even more dangerous than cold, especially in small mammals (e.g., Welbergen et al. 2008). Prolonged exposure to ambient temperatures  $(T_a)$  above the thermo-neutral zone results in high rates of evaporative water loss during the daytime and consequently water stress, rather than or in addition to energy shortage, may be a major cue for torpor expression in desert species (Geiser 2004). While heterothermic species can decrease their  $T_{\rm b}$  and undergo torpor, they often can also thermoconform at high  $T_{a}$ s and increase  $T_{b}$  to hyperthermic values. Over 60 years ago, Schmidt-Nielsen et al. (1956) discovered that camels (Camelus dromedarius) exposed to extreme heat in a desert, increased daily amplitudes in  $T_{\rm b}$ from ~2 to >6 °C when water was restricted. Importantly, the increase in daily  $T_{\rm b}$  amplitude was achieved by both a predictive reduction in  $T_{\rm h}$  in the cooler morning as well as a further increase in  $T_{\rm b}$  during the warmer part of the day.

Function/challenge	Benefits of torpor	References
Fires	Allows individuals to remain inactive during a fire Allows survival on limited food resources Predator avoidance	Stawski et al. (2015a); Nowack et al. (2016a); Matthews et al. (2017)
Storms	Allows individuals to remain inactive during a storm	Willis et al. (2006); Nowack et al. (2015)
Colonization	Allows survival during raft/journey through harsh envi- ronments	Nowack and Dausmann (2015)
	Facilitates establishment of founder population	
	Facilitates survival in new climate	
Extreme heat	Reduces evaporative water loss	Schmidt-Nielsen et al. (1956); Song et al. (1997); Grimpo et al. (2013); Bondarenco et al. (2014)
	Delays/shortens the time period in which cooling is required	
Evolution of mammals	Allowed survival during global wildfires/nuclear winter following the meteoroid impact	Lovegrove et al. (2014a); Nowack et al. (2016a)
Longevity	Increases survival probability	Lyman et al. (1981); Carey et al. (2003); Turbill et al. (2011); Turbill et al. (2012); Blanco and Godfrey (2013); Blanco and Zehr (2015)
	Allows spreading of reproduction over several years	
	Reduces reactive oxygen (ROS)	
	Delays physiological aging	

Table 2 List of the newly identified other functions of torpor relevant to climate change

Small desert bats have an even greater heat tolerance than camels. While  $T_{\rm b}$  of camels increased to above 41 °C, skin temperature  $(T_{skin})$  in Mormopterus petersi increased to 45.8 °C during a heat wave. Therefore, these bats can tolerate the most extreme  $T_{\rm b}$  range known for mammals, ranging from  $T_{\rm skin}$  of 3.3 °C during torpor in winter to 45.8 °C during extreme heat (Bondarenco et al. 2014). Similarly, torpor use even within the thermo-neutral zone has been reported in eastern pygmy-possums (Cercartetus nanus) and golden spiny mice (Acomys russatus) at  $T_{a}$ s as high as 30°-35°C, whereby metabolic depression was associated with only small decreases in  $T_{\rm b}$  (Song et al. 1997; Grimpo et al. 2013). The advantage of thermoconforming at high  $T_{\rm a}$  is the concurrent reduction in metabolic rate and the associated decrease of evaporative water loss (see Geiser and Brigham 2012). Camels, for example, maintain a stable  $T_{\rm b}$  and use evaporative cooling when water is available, but use "adaptive heterothermy" (in birds termed 'facultative hyperthermia'; see Tieleman and Williams 1999), i.e., increased daily amplitudes in  $T_{\rm h}$ , when water stressed, thereby reducing water requirements enormously.

Torpor use to save water might be more essential for survival than the related energy savings (e.g., Cooper et al. 2005; Schmid and Speakman 2009; Dausmann 2014; Lovegrove et al. 2014b). Evaporative water loss can be decreased to below measurable levels during torpor (Withers et al. 1990) and the reduction in energy expenditure at high  $T_a$  does not only decrease evaporative water loss by reduced respiration, but also lowers the endogenous thermal load which further reduces the need for evaporative cooling (Cooper et al. 2005). Bats that undergo torpor during hot days in the desert displayed longer torpor bouts at a given  $T_a$  than bats from temperate and subtropical habitats (Bondarenco et al. 2013). A low  $T_b$  during torpor will delay the time until a critically high  $T_b$  is reached and shortens the time required to use evaporative cooling during the hot part of the day (Bondarenco et al. 2013).

#### **Torpor use and fires**

Heat is not only a challenge for a mammal's water budget, but also increases the frequency of wildfires. Regular wildfires during summer are a threat for many mammals and are especially dangerous for small-bodied terrestrial species that cannot easily flee. Nevertheless, post-fire surveys suggest that some small-to-medium sized mammals survive fires by hiding in rock cavities or underground borrows (Matthews et al. 2017). For example, short-beaked echidnas (Tachyglossus aculeatus) became inactive and entered multiday torpor during a fuel-reduction burn in their natural habitat (Nowack et al. 2016a). Although the echidnas mostly used tree logs that were not necessarily safe during a fire, the use of multiday torpor allowed them to remain inactive, reducing the risk of getting trapped by the fire while foraging. After the fire, echidnas in burnt areas decreased their activity and remained inactive for 2-5 days. In contrast to homeothermic animals, which have to resume foraging even under adverse conditions because of their constant high energy demands, heterothermic animals can compensate for lost foraging opportunities by remaining inactive and reducing energy demands through the use of torpor.

Fires not only pose an acute risk to animals, but also destroy food resources and leave the surviving animals in a fire-scorched landscape. How an individual will cope with conditions after a fire depends on the individuals' behaviour and physiology. Studies on the abundance of mammals after fires indicate that heterothermic species, such as yellow-footed antechinus (Antechinus flavipes), have higher survival rates than homeothermic species, such as bush rats (Rattus fuscipes) (Thompson et al. 1989; Recher et al. 2009). Antechinus are known to use daily torpor, whereas available data on bush rats suggest that they are homeothermic (Geiser and Körtner 2010). Post-fire studies on heterothermic species suggest that individuals reduce activity and increase torpor use for weeks after a fire to enhance survival (Stawski et al. 2015a; Nowack et al. 2016a; Matthews et al. 2017). In the short-beaked echidna, increased torpor use after the fire likely compensates for reduced food availability and destroyed nesting sites (Nowack et al. 2016a). Echidnas feed on ants and termites, which are limited after the fire has burned woody debris. However, in addition to reduced food availability, these animals also have to deal with diminished ground cover. Small animals will have less shelter during activity and, therefore, face an additional increased risk of predation (Radford 2012). Consequently, torpor use in the small marsupial brown antechinus (A stu*artii*,  $\sim 16-40$  g) is assumed to increase survival chances by limiting the time spent foraging while exposed to predators (Stawski et al. 2015a). The fact that neither echidnas, nor antechinus left their fire-destroyed habitats indicates that torpor use after fires allows animals to remain in scorched landscapes and reduces the need to re-populate habitats after a fire (Stawski et al. 2015a; Matthews et al. 2017). A long-term study on a population of brown antechinus (A. stuartii) found that the population had returned to normal activity and torpor use within a full year after a prescribed fire (Hume 2015).

Although increasing torpor use after a fire is beneficial for terrestrial mammals, such as the echidna and antechinus, microbats (insect eating bats) appear to respond differently. Torpor use and activity periods were monitored in lesser long-eared bats (Nyctophilus geoffroyi) 4 months after a severe wildfire and again 2 years later (Doty et al. 2016). These authors found that shortly after the wildfire, bats expressed shorter torpor bouts and longer normothermic periods in comparison with 2 years after the wildfire (Doty et al. 2016). As aerial insect abundance was significantly greater shortly after the fire and ambient conditions were warmer, it is likely that bats took advantage of the increase in insect numbers and also in the decrease in vegetation, which would have made foraging easier (Doty et al. 2016). As flying mammals, bats also have the advantage of being able to more easily flee a fire and then return to their home ranges once the fire has passed. These studies reveal that different animals respond in various ways to fire, highlighting the importance of obtaining data on diverse species in this context.

#### Torpor use and the evolution of mammals

The ability of hibernators to disappear from the surface and hibernate for months in underground burrows has also been suggested to have facilitated the survival of mammals during the meteorite impact at the K-Pg boundary, about 65.5 million years ago (Lovegrove et al. 2014a). This event killed the dinosaurs and many other vertebrate species by causing global wildfires (Morgan et al. 2013). The mammals that survived the meteoroid impact would have had to deal initially with global wildfires, followed by reduced light levels for a year or more, likely leading to a reduction of  $T_{a}$ , both of which would have significantly reduced food availability (Robertson et al. 2004). As we describe above, data on torpor use during and after fires support the hypothesis by Lovegrove et al. (2014a). Torpor use in protected shelters would not only have been beneficial for the survival of the animals during the cold, but also during the blazing fires (Stawski et al. 2015a; Nowack et al. 2016a). In this regard, it is particularly interesting that especially echidnas enter torpor in response to fires and were found as one of the few survivors after a severe wildfire (Nowack et al. 2016a; Matthews et al. 2017). Echidnas have been described as "protoendotherms" and have many ancestral traits, such as a relaxed thermoregulation and the ability to frequently use torpor (Grigg et al. 2004), and these traits likely resemble traits of ancestral monotremes at the time of the K-Pg boundary.

Hibernation in temperate zone animals is usually characterized by torpor bouts lasting up to 5 weeks, interspersed by regular arousal phases in which the animal rewarms to euthermic  $T_{\rm b}$ s for a few hours (Carey et al. 2003). Periodic euthermic phases are energetically costly and in temperate species, the greatest proportion (70-80%) of the energy expended during hibernation is used for these rewarming processes (Wang 1978). In contrast, some tropical species, such as tenrecs or lemurs, can undergo hibernation periods without this characteristic periodic arousal process, but in these species,  $T_{\rm b}$  closely tracks daily fluctuations in  $T_{\rm a}$ (Dausmann et al. 2004; Kobbe et al. 2011; Lovegrove et al. 2014a). The original suggestion that torpor use may have facilitated mammalian survival at the K-Pg boundary was based on the observation of long-term hibernation without regular interbout arousals in the common tenrec (Tenrec ecaudatus), another "protoendotherm" [termed "basoendotherm" by Lovegrove (2012)] (Lovegrove et al. 2014a). Torpor use with less frequent costly arousals would have led to even greater energy savings and could have enabled ancestral mammals to survive until conditions on the surface had eased (Lovegrove et al. 2014a).

#### Heterothermy and storms

Storms are experienced by animals worldwide and, unlike fires, can occur anywhere. However, evidence for torpor use during storms is rare, and to date has only been reported for two species, the hoary bat (Lasiurus cinereus) and the marsupial sugar glider (*Petaurus breviceps*) (Willis et al. 2006; Nowack et al. 2015). Pregnant hoary bats used torpor during a snow storm in the Canadian spring, not only reducing the need to forage during adverse conditions, but also delaying parturition until conditions were more favorable (Willis et al. 2006). Individuals stayed inactive for a maximum of 9.1 days and the longest torpor bout recorded lasted 5.6 days. However, torpor use might not have been caused by the storm itself, but rather by the low  $T_a$  of ~5 °C or by a combination of cold exposure and convective heat loss. In contrast, sugar gliders, which rarely used torpor before the storm, used highly synchronized torpor during spring in a warm, subtropical habitat during a cyclone. During this storm,  $T_a$  did not decrease below values observed on other nights, and therefore, torpor use likely occurred in response to high wind speed and rainfall (Nowack et al. 2015).

From the available data, it is not clear whether gliders or bats anticipated the storm, but since some sugar gliders remained completely inactive during the storm night, it appears that they responded proactively to environmental cues, such as changes in barometric pressure, that were preceding the storm. No data exist on perception of barometric pressure changes in sugar gliders. However, studies on hibernating brown bats (*Myotis lucifugus*) and eastern pipistrelles (*Pipistrellus subflavus*) have suggested that bats use changing barometric pressure as a cue for conditions outside their hibernacula. Bats emerge from torpor when falling barometric pressure indicates favorable foraging conditions (Paige 1995; Czenze and Willis 2015).

#### Torpor use and colonization

Torpor use during unpredictable situations, such as storms or fires, also support the hypothesis that torpor can be advantageous during over-water colonization events (Nowack and Dausmann 2015). This hypothesis proposes that land masses or tree logs that contain mammals could be washed into the ocean during a storm event (e.g., Simpson 1940). As a consequence, the animals would unexpectedly find themselves without access to food or drinkable fresh water for a long period of time. Opportunistic torpor use would allow animals to cope with these conditions until eventually arriving on a shore. Importantly, torpor use would not only be an advantage during the rafting event (Martin 1972; Kappeler 2000), but also after arrival in the new habitat (Nowack and Dausmann 2015). The arriving animals would face unfamiliar terrain and ambient conditions, including a lack of known food sources. Furthermore, torpor use could facilitate the establishment of a founder population, by enabling a reproductive diapause. Birth of viable offspring after arrival also could provide individuals of the opposite sex for reproduction (Nowack and Dausmann 2015).

#### **Risks of torpor use**

In contrast to threats such as predation or storms (Turbill et al. 2011; Nowack et al. 2015), a sit-and-wait strategy during fires is only advantageous when animals are already hiding in a fire-safe refuge. Natural disasters can happen fast and unpredictably, giving the animal limited warnings, such as a drop in barometric pressure or the smell of smoke. Depending on the severity of the fire, smoke might be perceived relatively late by the individual, giving it only a short time to react. In this regard, torpor use might not only be beneficial, but also risky. If torpid animals cannot respond quickly to environmental threats, this would make them vulnerable and impose costs to the otherwise beneficial use of torpor. However, recent studies have shown that torpid animals can perceive olfactory and acoustic signals with  $T_{\rm b}$ s below 20 °C (Luo et al. 2014; Stawski et al. 2015b; Nowack et al. 2016b). Dense smoke as well as the smell of smoke (without the toxic substances) leads to an immediate termination of torpor and, therefore, could stimulate flight behaviour of torpid animals into more suitable shelters. Mobile animals are expected to either flee from the burning area or to search for shelter in underground burrows, caves, or tree hollows (reviewed in Engstrom 2010). Furthermore, several studies have shown that torpid animals are able to have coordinated movements with  $T_{\rm b}$ s as low as 14.8–17.9 °C, although locomotion at these low  $T_{\rm b}$ s is slow (Warnecke et al. 2008; Warnecke and Geiser 2010; Rojas et al. 2012; Nowack et al. 2016b).

#### Longevity and torpor use

It has previously been suggested that predator avoidance could be one of the other functions of torpor (Geiser and Brigham 2012). This hypothesis has received further support from recent work on small mammals employing torpor after fires when predation pressure is likely to be increased, as described above. Along with allowing animals to avoid predation, it has also been proposed that torpor can increase longevity by enabling a "slow-paced" life history that is associated with increased survival rates during the hibernation season, slowed physiological aging, increased maximum longevity and long generation times (Lyman et al. 1981; Turbill et al. 2011, 2012, 2013; Ruf et al. 2012). Data on the relative length of telomeres (endcaps of DNA that shorten with every cell division) that are used as an indicator for physiological aging support this hypothesis (Turbill et al. 2012, 2013; Hoelzl et al. 2015, 2016). Meiotic cell divisions and erosion of telomeres via reactive oxygenspecies are arrested or reduced at low  $T_{\rm b}$ s experienced during torpor (Kruman et al. 1988; Marcand et al. 2000) and reduced telomere shortening during the hibernation season has been reported in various studies (Turbill et al. 2012, 2013; Hoelzl et al. 2015, 2016). A correlation between torpor use and life spans in strepsirrhine primate species suggests that the extent of torpor expression is strongly linked with the life expectancy (Blanco and Godfrey 2013; Blanco and Zehr 2015). Similarly, torpor use and longevity were found to be correlated in Turkish hamsters (Lyman et al. 1981).

#### Torpor and climate change

Climate change is expected to lead to an increase in occurrence and intensity of floods, storms, droughts, fires, and extreme temperatures (Christensen and Christensen 2003; CSIRO 2011). These events are expected to become the main causes of species extinction (McCain and King 2014). Wildfires are already increasingly being recognized as a major disturbance affecting many parts of the world, including Europe, Africa, America, Asia, and Australia. Progressing fragmentation, global warming, and human activities have led to frequent wildfires in ecosystems that were not classified as fire-prone (Moreira et al. 2001; Pechony and Shindell 2010). Furthermore, the occurrence of wildfires has increased in a number of regions over the past decades (e.g., Piñol et al. 1998). A gradually warming climate will also increase temperature extremes and, therefore, thermal stress on organisms, especially affecting desert species already coping with temperatures near their thermal limit. Heat waves already have had severe effects on many species and are especially challenging for species that do not shelter underground or in well-insulated nesting sites that offer protection from the heat, such as flying foxes (e.g., Welbergen et al. 2008).

Many seasonal hibernators, will likely face negative consequences as a result of the predicted increase in surface temperature. In the hazel dormouse (*Muscardinus avellanarius*), arousals from hibernation occurred more frequently during warmer years, causing substantial additional energy expenditure throughout the hibernation season (Pretzlaff and Dausmann 2012). Similarly, yellowbellied marmots (*Marmota flaviventris*) emerged earlier from torpor than two decades earlier and this difference was attributed to the warmer spring  $T_a$  (Inouye et al. 2000). On the one hand, this increases the risk for especially small seasonal hibernators to deplete fat reserves before the end of hibernation. Humphries et al. (2002) calculated that an increase in  $T_a$  of 10 °C will increase the energy requirements of hibernating bats (*Myotis lucifugus*) threefold. On the other hand, climate change will also result in shorter winters, potentially associated with other problems, such as a reduction or a shift in the geographical range of species that are reliant on longer winters (Humphries et al. 2002).

Many of the functions of torpor would allow heterothermic animals to cope with the assumed impacts of climate change and, therefore, should give them an adaptive edge over homeothermic species with their constantly high energetic requirements. However, to be advantageous during unpredictable events, such as fires and storms, torpor use needs to be a flexible, opportunistic response, not the highly seasonal strategy employed by many species (Boyles et al. 2011). Particularly, multiday torpor use, i.e., seasonal hibernation, often requires preparation in the form of energy storage, either fat accumulation or food hoarding (Carey et al. 2003). Therefore, strictly seasonal hibernators will be less likely able to use torpor effectively if the challenges do not coincide with the hibernation season. However, there is ample evidence that the opportunistic use of short bouts of torpor throughout the year is not only expressed by daily heterotherms, but also is used by some hibernating species (reviewed in Levesque et al. 2016).

Phenotypic plasticity is higher in species that have evolved in variable habitats and this plasticity has been proposed to play an important role in vertebrate resilience in the light of climate change and habitat degradation (Canale and Henry 2010). Many heterothermic species are highly flexible in adjusting their energy requirements seasonally and regionally (e.g., Bozinovic et al. 2007; Boyles et al. 2013; Ruf and Geiser 2015) and often show pronounced differences in torpor use among populations of one species, individuals of one population, or even within individuals between different years (reviewed in Levesque et al. 2016). This flexibility is usually an acclimation or adaptation to habitats (e.g., habitat resources) or body condition. Heterotherms tend to undergo short-term torpor when body condition is poor, whereas multiday torpor is often expressed when body condition is good (Hallam and Mzilikazi 2011; Kobbe et al. 2011). Importantly, the phenotypic plasticity of energy expenditure afforded by the opportunistic use of torpor may be crucial in dealing with climate change and other anthropogenic disturbances (Nowack et al. 2015; Stawski et al. 2015a). The responses of heterothermic mammals to fires and storms are highly flexible and specific to

individuals. While echidnas in an unburnt control habitat mostly used torpor synchronized in response to rainfall and  $T_a$  before the fire, echidnas adjusted  $T_b$  flexibly in response to varying resource distribution after the fire. Furthermore, brown antechinus (*A. stuartii*) displayed differences in torpor use between sexes (Stawski et al. 2016). This flexibility in behavioural and physiological traits suggests that heterothermic mammals are able to adjust readily to sudden environmental changes, and therefore, torpor use enables them to increase the likelihood to survive catastrophic events.

#### Conclusion

Despite the risks associated with torpor, there is increasing evidence that the use of torpor is advantageous in many different settings and scenarios. While the removal of parasites is likely an additional benefit of cold  $T_{\rm b}$ , most functions, such as predator avoidance, development under reduced food sources, or disappearance from the surface during storms or fires could well be primary functions of torpor and serve a similar benefit as torpor use during winter. Heterothermic species are often able to cope with a wide  $T_{\rm a}$  range, and perhaps, with the exception of strictly seasonal hibernators, many are likely to adjust better to a changing climate than homeotherms. Our review emphasises the many benefits of torpor use and suggests that especially the use of opportunistic torpor will give heterotherms an advantage over homeotherms during climate change.

Acknowledgements Open access funding provided by University of Veterinary Medicine Vienna.

**Open Access** This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

#### References

- Audet D, Fenton MB (1988) Heterothermy and the use of torpor by the bat *Eptesicus fuscus* (Chiroptera: Vespertilionidae): a field study. Physiol Zool 61:197–204
- Bieber C, Ruf T (2009) Summer dormancy in edible dormice (*Glis glis*) without energetic constraints. Naturwissenschaften 96:165–171
- Birkhead TR, Møller AP (1993) Sexual selection and the temporal separation of reproductive events: sperm storage data from reptiles, birds and mammals. Biol J Linn Soc 50:295–311
- Blanco MB, Godfrey LR (2013) Does hibernation slow the "pace of life" in dwarf lemurs (*Cheirogaleus* spp.)? Int J Primatol 34:130–147

- Blanco MB, Zehr SM (2015) Striking longevity in a hibernating lemur. J Zool 296:177–188
- Bondarenco A, Körtner G, Geiser F (2013) Some like it cold: summer torpor by freetail bats in the Australian arid zone. J Comp Physiol B 183:1113–1122
- Bondarenco A, Körtner G, Geiser F (2014) Hot bats: extreme thermal tolerance in a desert heat wave. Naturwissenschaften 101:679–685
- Boyles JG, Seebacher F, Smit B, McKechnie AE (2011) Adaptive thermoregulation in endotherms may alter responses to climate change. Integr Comp Biol 51:676–690
- Boyles JG, Thompson AB, McKechnie AE, Malan E, Humphries MM, Careau V (2013) A global heterothermic continuum in mammals. Global Ecol Biogeogr 22:1029–1039
- Bozinovic F, Muñoz JP, Naya D, Cruz-Neto A (2007) Adjusting energy expenditures to energy supply: food availability regulates torpor use and organ size in the Chilean mouse-opossum *Thylamys elegans*. J Comp Physiol B 177:393–400
- Brigham RM, Körtner G, Maddocks TA, Geiser F (2000) Seasonal use of torpor by free-ranging Australian owlet-nightjars (*Aegotheles cristatus*). Physiol Biochem Zool 73:613–620
- Buffenstein R, Jarvis JUM (1985) Thermoregulation and metabolism in the smallest African gerbil, *Gerbillus pusillus*. J Zool 205:107–121
- Callait M-P, Gauthier D (2000) Parasite adaptations to hibernation in alpine marmots (*Marmota marmota*). In: Heldmaier G, Klingenspor M (eds) Life in the cold: eleventh international hibernation symposium. Springer Berlin Heidelberg, Berlin, pp 139–146
- Canale CI, Henry PY (2010) Adaptive phenotypic plasticity and resilience of vertebrates to increasing climatic unpredictability. Clim Res 43:135–147
- Canale C, Perret M, Henry P-Y (2012) Torpor use during gestation and lactation in a primate. Naturwissenschaften 99:159–163
- Carey HV, Andrews MT, Martin SL (2003) Mammalian hibernation: cellular and molecular responses to depressed metabolism and low temperature. Physiol Rev 83:1153–1181
- Carpenter FL, Hixon MA (1988) A new function for torpor: fat conservation in a wild migrant hummingbird. Condor 90:373–378
- Christensen JH, Christensen OB (2003) Climate modelling: severe summertime flooding in Europe. Nature 421:805–806
- Chute R (1964) Hibernation and parasitism: recent developments and some theoretical consideration. Mammalian hibernation 2. Ann Acad Sci Fenn A 4 Biol 71:113–122
- Cooper CE, McAllan BM, Geiser F (2005) Effect of torpor on the water economy of an arid-zone marsupial, the stripe-faced dunnart (*Sminthopsis macroura*). J Comp Physiol B 175:323–328
- CSIRO (2011) Climate change impact on fire. http://www.csiro.au/ en/Outcomes/Climate/Adapting/Climate-Change-Fire-Weather. aspx. Accessed 07 Jan 2013
- Czenze ZJ, Willis CKR (2015) Warming up and shipping out: arousal and emergence timing in hibernating little brown bats (*Myotis lucifugus*). J Comp Physiol B 185:575–586
- Dausmann KH (2014) Flexible patterns in energy savings: heterothermy in primates. J Zool 292:101–111
- Dausmann KH, Glos J, Ganzhorn JU, Heldmaier G (2004) Physiology: hibernation in a tropical primate. Nature 429:825–826
- Doty AC, Stawski C, Law BS, Geiser F (2016) Post-wildfire physiological ecology of an Australian microbat. J Comp Physiol B 186:937–946
- Doucette LI, Brigham RM, Pavey CR, Geiser F (2012) Prey availability affects daily torpor by free-ranging Australian owlet-nightjars (*Aegotheles cristatus*). Oecologia 169:361–372
- Engstrom RT (2010) First-order fire effects on animals: review and recommendations. Fire Ecol 6:115–130

- Fietz J, Tomiuk J, Matuschka F-R, Richter D (2014) Seasonal prevalence of lyme disease spirochetes in a heterothermic mammal, the edible dormouse (*Glis glis*). Appl Environ Microb 80:3615–3621
- Fietz J, Langer F, Havenstein N, Matuschka F-R, Richter D (2016) The vector tick *Ixodes ricinus* feeding on an arboreal rodent the edible dormouse *Glis glis*. Parasitol Res 115:1435–1442
- Geiser F (2004) The role of torpor in the life of Australian arid zone mammals. Aust Mamm 26:125–134
- Geiser F (2007) Yearlong hibernation in a marsupial mammal. Naturwissenschaften 94:941–944
- Geiser F, Brigham RM (2012) The other functions of torpor. In: Ruf T, Bieber C, Arnold W, Millesi E (eds) Living in a seasonal world. Springer, Berlin, pp 109–121
- Geiser F, Körtner G (2010) Hibernation and daily torpor in Australian mammals. Aust Zool 35:204–215
- Geiser F, Masters P (1994) Torpor in relation to reproduction in the Mulgara, *Dasycercus cristicauda* (Dasyuridae: Marsupilia). J Therm Biol 19:33–40
- Geiser F, Turbill C (2009) Hibernation and daily torpor minimize mammalian extinctions. Naturwissenschaften 96:1235–1240
- Geiser F, Westman W, McAllan BM, Brigham RM (2006) Development of thermoregulation and torpor in a marsupial: energetic and evolutionary implications. J Comp Physiol B 176:107–116
- Giroud S, Turbill C, Ruf T (2012) Torpor use and body mass gain during pre-hibernation in late-born juvenile garden dormice exposed to food shortage. In: Ruf T, Bieber C, Arnold W, Millesi E (eds) Living in a seasonal world. Springer, Berlin, pp 481–491
- Giroud S, Zahn S, Criscuolo F, Chery I, Blanc S, Turbill C, Ruf T (2014) Late-born intermittently fasted juvenile garden dormice use torpor to grow and fatten prior to hibernation: consequences for ageing processes. P Roy Soc B 281:20141131
- Grigg GC, Beard LA, Augee ML (2004) The evolution of endothermy and its diversity in mammals and birds. Physiol Biochem Zool 77:982–997
- Grimpo K, Legler K, Heldmaier G, Exner C (2013) That's hot: golden spiny mice display torpor even at high ambient temperatures. J Comp Physiol B 183:567–581
- Grinevitch L, Holroyd SL, Barclay RMR (1995) Sex differences in the use of daily torpor and foraging time by big brown bats (*Eptesicus fuscus*) during the reproductive season. J Zool 235:301–309
- Hallam SL, Mzilikazi N (2011) Heterothermy in the southern African hedgehog, *Atelerix frontalis*. J Comp Physiol B 181:437–445
- Hanna E, Cardillo M (2014) Clarifying the relationship between torpor and anthropogenic extinction risk in mammals. J Zool 293:211–217
- Hiebert S (1993) Seasonal changes in body mass and use of torpor in a migratory hummingbird. The Auk 110:787–797
- Hoelzl F, Bieber C, Cornils JS, Gerritsmann H, Stalder GL, Walzer C, Ruf T (2015) How to spend the summer? Free-living dormice (*Glis glis*) can hibernate for 11 months in non-reproductive years. J Comp Physiol B 185:931–939
- Hoelzl F, Cornils JS, Smith S, Moodley Y, Ruf T (2016) Telomere dynamics in free-living edible dormice (*Glis glis*): the impact of hibernation and food supply. J Exp Biol 219:2469–2474
- Hosken JD, Withers PC (1997) Temperature regulation and metabolism of an Australian bat, *Chalinolobus gouldii* (Chiroptera: Vespertilionidae) when euthermic and torpid. J Comp Physiol B 167:71–80
- Hume T (2015) The physiology and behaviour of *Antechinus stuartii* in a post-fire landscape. Honours Thesis, Zoology, University of New England, Armidale, Australia
- Humphries MM, Thomas DW, Speakman JR (2002) Climate-mediated energetic constraints on the distribution of hibernating mammals. Nature 418:313–316

- Inouye DW, Barr B, Armitage KB, Inouye BD (2000) Climate change is affecting altitudinal migrants and hibernating species. PNAS 97:1630–1633
- Kappeler PM (2000) Lemur origins: rafting by groups of hibernators? Folia Primatol 71:422–425
- Kobbe S, Ganzhorn J, Dausmann KH (2011) Extreme individual flexibility of heterothermy in free-ranging Malagasy mouse lemurs (*Microcebus griseorufus*). J Comp Physiol B 181:165–173
- Körtner G, Brigham RM, Geiser F (2000) Metabolism: winter torpor in a large bird. Nature 407:318
- Körtner G, Pavey CR, Geiser F (2008) Thermal biology, torpor, and activity in free-living mulgaras in arid zone Australia during the winter reproductive season. Physiol Biochem Zool 81:442–451
- Kruman II, Ilyasova EN, Rudchenko SA, Khurkhulu ZS (1988) The intestinal epithelial cells of ground squirrel (*Citellus undulatus*) accumulate at G2 phase of the cell cycle throughout a bout of hibernation. Comp Biochem Physiol A 90:233–236
- Levesque DL, Nowack J, Stawski C (2016) Modelling mammalian energetics: the heterothermy problem. Climate Change Responses 3:7
- Levy O, Dayan T, Kronfeld-Schor N (2011) Interspecific competition and torpor in golden spiny mice: two sides of the energy-acquisition coin. Integr Comp Biol 51:441–448
- Liow LH, Fortelius M, Lintulaakso K, Mannila H, Stenseth NC (2009) Lower extinction risk in sleep-or-hide mammals. Am Nat 173:264–272
- Lourenço S, Palmeirim JM (2008) Which factors regulate the reproduction of ectoparasites of temperate-zone cave-dwelling bats?. Parasitol Res 104:127–134
- Lovegrove BG (2012) The evolution of endothermy in Cenozoic mammals: a plesiomorphic-apomorphic continuum. Biol Rev 87:128–162
- Lovegrove BG, Lobban KD, Levesque DL (2014a) Mammal survival at the Cretaceous–Palaeogene boundary: metabolic homeostasis in prolonged tropical hibernation in tenrecs. P Roy Soc B Biol Sci 281:20141304
- Lovegrove BG, Canale CI, Levesque DL, Fluch G, Reháková-Petrů M, Ruf T (2014b) Are tropical small mammals physiologically vulnerable to Arrhenius effects and climate change? Physiol Biochem Zool 87:30–45
- Luo J, Clarin B-M, Borissov IM, Siemers BM (2014) Are torpid bats immune to anthropogenic noise? J Exp Biol 217:1072–1078
- Lyman CP, O'Brien R, Greene G, Papafrangos E (1981) Hibernation and longevity in the Turkish hamster *Mesocricetus brandti*. Science 212:668–670
- Lyman CP, Willis JS, Malan A, Wang LCH (1982) Hibernation and torpor in mammals and birds. Academic Press, Inc., New York
- MacMillen RE (1965) Aestivation in the cactus mouse, *Peromyscus* eremicus. Comp Biochem Physiol A 16:227–248
- Marcand S, Brevet V, Mann C, Gilson E (2000) Cell cycle restriction of telomere elongation. Curr Biol 10:487–490
- Martin RD (1972) Review lecture: adaptive radiation and behaviour of the Malagasy lemurs. Philos T Roy Soc B 264:295–352
- Matthews JK, Stawski C, Körtner G, Parker CA, Geiser F (2017) Torpor and basking after a severe wildfire: mammalian survival strategies in a scorched landscape. J Comp Physiol B 187:385–393
- McAllan BM, Geiser F (2014) Torpor during reproduction in mammals and birds: dealing with an energetic conundrum. Integr Comp Biol 54:516–532
- McCain CM, King SRB (2014) Body size and activity times mediate mammalian responses to climate change. Glob Change Biol 20:1760–1769
- Moreira F, Rego FC, Ferreira PG (2001) Temporal (1958–1995) pattern of change in a cultural landscape of northwestern Portugal: implications for fire occurrence. Landsc Ecol 16:557–567

- Morgan J, Artemieva N, Goldin T (2013) Revisiting wildfires at the K-Pg boundary. J Geophys Res 118:1508–1520
- Morrow GE, Nicol SC (2009) Cool sex? Hibernation and reproduction overlap in the echidna. PLoS One 4:e6070
- Morton SR (1978) Torpor and nest-sharing in free-living *Sminthopsis crassicaudata* (Marsupialia) and *Mus musculus* (Rodentia). J Mamm 59:569–575
- Nagel A (1977) Torpor in the European white-toothed shrews. Experientia 33:1455–1456
- Nowack J, Dausmann KH (2015) Can heterothermy facilitate the colonization of new habitats?. Mamm Rev 45:117–127
- Nowack J, Rojas AD, Körtner G, Geiser F (2015) Snoozing through the storm: torpor use during a natural disaster. Sci Rep 5:11243
- Nowack J, Cooper CE, Geiser F (2016a) Cool echidnas survive the fire. P Roy Soc B 283:20160382
- Nowack J, Delesalle M, Stawski C, Geiser F (2016b) Can hibernators sense and evade fires? Olfactory acuity and locomotor performance during deep torpor. Sci Nat 103:73
- Paige KN (1995) Bats and barometric pressure: conserving limited energy and tracking insects from the roost. Funct Ecol 9:463–467
- Pechony O, Shindell DT (2010) Driving forces of global wildfires over the past millennium and the forthcoming century. PNAS 107:19167–19170
- Piñol J, Terradas J, Lloret F (1998) Climate warming, wildfire hazard, and wildfire occurrence in coastal eastern Spain. Clim Change 38:345–357
- Pretzlaff I, Dausmann K (2012) Impact of climatic variation on the hibernation physiology of *Muscardinus avellanarius*. In: Ruf T, Bieber C, Arnold W, Millesi E (eds) Living in a seasonal world. Springer, Berlin, pp 85–97
- Racey P, Swift S (1981) Variations in gestation length in a colony of pipistrelle bats (*Pipistrellus pipistrellus*) from year to year. J Reprod Fertil 61:123–129
- Radford IJ (2012) Threatened mammals become more predatory after small-scale prescribed fires in a high-rainfall rocky savanna. Austral Ecol 37:926–935
- Recher HF, Lunney D, Matthews A (2009) Small mammal populations in a eucalypt forest affected by fire and drought. I. Longterm patterns in an era of climate change. Wildl Res 36:143–158
- Robertson DS, McKenna MC, Toon OB, Hope S, Lillegraven JA (2004) Survival in the first hours of the Cenozoic. Geol Soc Am Bull 116:760–768
- Rojas AD, Körtner G, Geiser F (2012) Cool running: locomotor performance at low body temperature in mammals. Biol Lett 8:868–870
- Ruf T, Geiser F (2015) Daily torpor and hibernation in birds and mammals. Biol Rev 90:891–926
- Ruf T, Bieber C, Turbill C (2012) Survival, aging, and life-history tactics in mammalian hibernators. In: Ruf T, Bieber C, Arnold W, Millesi E (eds) Living in a seasonal world. Springer, Berlin, pp 123–132
- Schmid J, Speakman JR (2009) Torpor and energetic consequences in free-ranging grey mouse lemurs (*Microcebus murinus*): a comparison of dry and wet forests. Naturwissenschaften 96:609–620

- Schmidt-Nielsen K, Schmidt-Nielsen B, Jarnum SA, Houpt TR (1956) Body temperature of the camel and its relation to water economy. Am J Physiol 188:103–112
- Simpson GG (1940) Mammals and land bridges. J Wash Acad Sci 30:137–163
- Song X, Körtner G, Geiser F (1997) Thermal relations of metabolic rate reduction in a hibernating marsupial. Am J Physiol Regul Integr Comp Physiol 273:R2097–R2104
- 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 a reproductive female marsupial, Antechinus flavipes. Mamm Res 61:417–421
- Stawski C, Körtner G, Nowack J, Geiser F (2015a) The importance of mammalian torpor for survival in a post-fire landscape. Biol Lett 11:20150134
- Stawski C, Matthews JK, Körtner G, Geiser F (2015b) Physiological and behavioural responses of a small heterothermic mammal to fire stimuli. Physiol Behav 151:617–622
- Stawski C, Körtner G, Nowack J, Geiser F (2016) Phenotypic plasticity of post-fire activity and thermal biology of a free-ranging small mammal. Physiol Behav 159:104–111
- Thompson MB, Medlin G, Hutchinson R, West N (1989) Short-term effects of fuel reduction burning on populations of small terrestrial mammals. Wildl Res 16:117–129
- Tieleman BI, Williams JB (1999) The role of hyperthermia in the water economy of desert birds. Physiol Biochem Zool 72:87–100
- Turbill C, Bieber C, Ruf T (2011) Hibernation is associated with increased survival and the evolution of slow life histories among mammals. P Roy Soc B-Biol Sci 278:3355–3363
- Turbill C, Smith S, Deimel C, Ruf T (2012) Daily torpor is associated with telomere length change over winter in Djungarian hamsters. Biol Lett 8:304–307
- Turbill C, Ruf T, Smith S, Bieber C (2013) Seasonal variation in telomere length of a hibernating rodent. Biol Lett 9:20121095
- Wang LCH (1978) Energetic and field aspects of mammalian torpor: the Richardson's ground squirrel. In: Wang LCH, Hudson JW (eds) Strategies in cold. Natural Torpidity and Thermogenesis Academic Press, New York, pp 109–145
- Wang Z, Liang B, Racey P, Wang Y-L, Zhang S-Y (2008) Sperm storage, delayed ovulation, and menstruation of the female Rickett's big-footed bat (*Myotis ricketti*). Zool Stud 47:215–221
- Warnecke L, Geiser F (2010) The energetics of basking behaviour and torpor in a small marsupial exposed to simulated natural conditions. J Comp Physiol B 180:437–445
- Warnecke L, Turner J, Geiser F (2008) Torpor and basking in a small arid zone marsupial. Naturwissenschaften 95:73–78
- Welbergen JA, Klose SM, Markus N, Eby P (2008) Climate change and the effects of temperature extremes on Australian flyingfoxes. P Roy Soc B 275:419–425
- Willis CKR, Brigham RM, Geiser F (2006) Deep, prolonged torpor by pregnant, free-ranging bats. Naturwissenschaften 93:80–83
- Withers PC, Richardson K, Wooller R (1990) Metabolic physiology of euthermic and torpid honey possums, *Tarsipes rostratus*. Aust J Zool 37:685–693