Phenotypic plasticity of post-fire activity and thermal biology of a free-ranging small mammal

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HIGHLIGHTS

• We present data on the short-term responses of a small mammal to a prescribed fire.
• Male and female brown antechinus decreased diurnal activity after the fire.
• Torpor use increased for both sexes in the post-fire environment.
• Females expressed more torpor and lower body temperatures than males.
• These responses may ensure survival by reducing foraging needs in a barren habitat.

ABSTRACT

Ecosystems can change rapidly and sometimes irreversibly due to a number of anthropogenic and natural factors, such as deforestation and fire. How individual animals exposed to such changes respond behaviourally and physiologically is poorly understood. We quantified the phenotypic plasticity of activity patterns and torpor use – a highly efficient energy conservation mechanism – in brown antechinus (Antechinus stuartii), a small Australian marsupial mammal. We compared groups in densely vegetated forest areas (pre-fire and control) with a group in a burned, open habitat (post-fire). Activity and torpor patterns differed among groups and sexes. Females in the post-fire group spent significantly less time active than the other groups, both during the day and night. However, in males only daytime activity declined in the post-fire group, although overall activity was also reduced on cold days in males for all groups. The reduction in total or diurnal activity in the post-fire group was made energetically possible by a ~3.4-fold and ~2.2-fold increase in the proportion of time females and males, respectively, used torpor in comparison to that in the pre-fire and control groups. Overall, likely due to reproductive needs, torpor was more pronounced in females than in males, but low ambient temperatures increased torpor bout duration in both sexes. Importantly, for both male and female antechinus and likely other small mammals, predator avoidance and energy conservation – achieved by reduced activity and increased torpor use – appear to be vital for post-fire survival where ground cover and refuges have been obliterated.

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1. Introduction

Anthropogenic environmental changes, such as habitat degradation and an increase in the variability of weather patterns, are having irreversible impacts on many ecosystems around the world. Often these changes are intertwined. For example, land clearing destroys post-fire refuges, replaces forests with more fire-prone vegetation and also increases ambient temperature (T_a) and therefore fire frequency [1,26]. The ensuing fires then further contribute to deforestation, placing even more pressure on ecosystems. The global increase in fires [44] and habitat fragmentation [16] are inflicting historically unprecedented levels of stress on individual animals, threatening the survival of populations and species.

Small terrestrial mammals are generally not able to flee from a fire, but they have been shown to survive in situ [2,31,42]. Fire-proof refugia, such as deep burrows and rock crevices, are vital to the persistence of small terrestrial mammals not only during, but also after a fire when the absence of ground cover increases predation pressure [2,10,33,42]. Therefore, small mammal populations often do not require recolonisation after a fire, provided the remaining individuals are able to survive in a habitat that is usually depleted of food, water, vegetative cover and refuges [2,11,42,47,53]. Indeed, many individuals that survive a fire are often subsequently killed by predators or starve to death [24,
The behavioural and physiological responses of individuals to fire-modified habitats and the subsequent reduction in vegetative cover will determine whether or not a population can persist and an enhanced phenotypic plasticity in any of these traits would be of great advantage [5,29].

Unfortunately, our understanding of how small mammals deal with post-fire challenges is limited. Some recent evidence suggests that heterothermic mammals, with fluctuating body temperatures (Tb) and adjustable energy expenditures, may have an adaptive advantage over homeothermic mammals that maintain a stable and high Tb and have continuously high energy requirements [2,42]. Heterothermic endotherms can save large amounts of energy and water by employing torpor, a controlled reduction of Tb and metabolic rate [36]. While mobility is generally reduced in torpor, torpid animals are able to respond to smoke, an early warning cue to an approaching fire front, by arouse from torpor to find or remain in a suitable refuge [43].

Behavioural and physiological responses of individuals to changes in their environment vary among species, which can greatly influence survival. Specifically, heterothermy has been posited to have enabled small mammals to survive the meteorite impact that decimated dinosaurs and also the current high levels of mammalian extinctions in Australia [14,21]. Therefore, the physiological flexibility afforded to heterotherms is likely advantageous in a rapidly changing environment [5]. These responses can also differ within a species and can be highly variable among different populations, individuals or even within an individual under different conditions [5,25,29]. For example, the duration and depth of torpor bouts as well as activity patterns have been observed to be different between sexes in a number of heterothermic mammals [12,20,35,38]. Some of these differences can be attributed to sexual dimorphism in body size and/or reproductive effort, which is energetically costly in both sexes. Males often have to invest more time in searching for a mate whereas females spend more energy on pregnancy and raising young [20]. Whether or not the sexes have varying adaptations in response to environmental catastrophes, such as fires, is largely unknown.

To reveal the key adaptations used by small mammals to survive a controlled fire we studied the brown antechinus (Antechinus stuartii), a small dasyurid marsupial mammal that occurs in south-eastern Australia, forages terrestrially and arboreally and also uses nests in a variety of locations such as tree hollows, rock crevices and logs [7,51]. Brown antechinus have an unusual life history: after a short two week mating period during the austral winter (August–early September) all of the males die whereas most females survive until after the young have been weaned in summer and some will live for a second breeding season [23,51,52]. Brown antechinus can use daily torpor throughout most of the year, often in response to unfavourable environmental conditions [35,42]. As brown antechinus typically consume insects [7] their food source is likely to become limited with cold ambient conditions and also after a fire [6,39,47], times when torpor expression would be beneficial. Because of the unique life history of this species, we aimed to determine whether and how male and female brown antechinus differ behaviourally and physiologically in dealing with the thermal and energetic challenges in a post-fire environment. We hypothesised that both sexes would show phenotypic plasticity and increase torpor use and decrease activity in response to a prescribed burn to save energy and ameliorate predation pressure to increase their chance of survival. However, we also expected that females would express more torpor than males as is typical in this species [12,35].

2. Material and methods

Permits to conduct this study were issued by the University of New England Animal Ethics Committee (AEC13-088) and the New South Wales National Parks and Wildlife Service (SL100791).

2.1. Study site

This study was undertaken during April–May 2014 (austral autumn), 3 months before the breeding season, at Guy Fawkes River National Park (30°04′58.6″S, 152°20′0.9″E) in Australia. A hazard-reduction burn was performed by New South Wales National Parks and Wildlife Service (NSW NPWS) on 8–9 May 2014. The affected area (our prescribed fire site) was 379 ha and was bordered by an escarpment and dirt roads. Our control site was situated in the unburned forest beyond these roads. Three treatment regimes were investigated in the study: (i) pre-fire group: prescribed area before the fire, (ii) post-fire group: prescribed area after the fire and (iii) control group: control area after the fire. As we were only given two weeks’ notice prior to the prescribed fire we only had time to capture and undertake surgeries on a limited number of individuals from the treatment site only. The amount of time for data collection in the pre-fire group was obviously also limited. Throughout the study period Tb was measured at 10 min intervals at both the prescribed fire site and the control site using temperature data loggers (±0.5 °C, iButton thermochron DS1921G, Maxim Integrated Products, Inc., Sunnyvale, California, USA). These loggers were suspended in trees ~1 m off the ground and placed in inverted Styrofoam cups to prevent direct sun exposure.

2.2. Study protocol

Brown antechinus were captured using aluminium box traps (Elliott Scientific Equipment, Upwey, Australia) baited with oats, peanut butter and honey. Bedding material was provided to prevent hypothermia in animals confined to traps overnight. Before the prescribed fire four male and four female antechinus were trapped in the prescribed fire site and immediately following the fire another four male and two female antechinus were captured in the control site. As one male and two females in the pre-fire group perished before/during the fire, we captured an additional two females just prior to the fire, which were included in the post-fire group only. Mean body mass of males was 27.7 ± 2.3 g (n = 8) and of females was 25.6 ± 4.2 g (n = 8). Individuals were implanted intraperitoneally with temperature-sensitive radio-transmitters that had individual frequencies (1.8 to 2.1 g, Sirtrack, Havelock North, New Zealand). Before implatation, transmitters were coated in inert wax (Paraffin/Evax, Mini Mitter, Respiromics Inc. OR, USA) and then calibrated in a water bath over a temperature range of 10 to 45 °C to the nearest 0.1 °C. All antechinus were weighed using an electronic balance to the nearest 0.1 g and a transmitter (1.8 to 2.1 g, Sirtrack, Havelock North, New Zealand). Brown antechinus were captured using aluminium box traps (Elliott Scientific Equipment, Upwey, Australia) baited with oats, peanut butter and honey. Bedding material was provided to prevent hypothermia in animals confined to traps overnight. Before the prescribed fire four male and four female antechinus were trapped in the prescribed fire site and immediately following the fire another four male and two female antechinus were captured in the control site. As one male and two females in the pre-fire group perished before/during the fire, we captured an additional two females just prior to the fire, which were included in the post-fire group only. Mean body mass of males was 27.7 ± 2.3 g (n = 8) and of females was 25.6 ± 4.2 g (n = 8). Individuals were implanted intraperitoneally with temperature-sensitive radio-transmitters that had individual frequencies (1.8 to 2.1 g, Sirtrack, Havelock North, New Zealand). Before implatation, transmitters were coated in inert wax (Paraffin/Evax, Mini Mitter, Respiromics Inc. OR, USA) and then calibrated in a water bath over a temperature range of 10 to 45 °C to the nearest 0.1 °C. All antechinus were weighed using an electronic balance to the nearest 0.1 g and a transmitter (<10% of body mass was chosen for each individual [34]. For details on the surgical procedure see Stawski et al. [43].

Field data were collected from four male brown antechinus in the pre-fire group, three males in the post-fire group and four males in the control group (Table 1). For the females, field data were collected from four females in the pre-fire group, four females in the post-fire group and two females in the control group (Table 1). Some of the

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<td>The number of days data were recorded for each individual in all three groups: pre-fire, post-fire and control. Also shown are the total (n = the number of individuals, shown in brackets) and mean number of days of data for each group.</td>
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collected data from the female brown antechinus have been published previously [42], but these are not included here and in the current study we present new physiological and behavioural data with an emphasis on phenotypic plasticity and comparing differences between sexes.

Each individual was radio-tracked daily to its nest except for four days during and after the fire as the study site was inaccessible. Coordinates of every nest site were recorded with a GPS. Once an individual was found a remote receiver/data logger with an antenna were placed in range of the transmitter signal [19]. Each logger was programmed to record the pulse rate of the transmitter once every 10 min, which was then converted into Tb using the calibration equations for each individual transmitter. Whenever animals moved nests the loggers were transferred to the new location to ensure they were in range of the transmitter signal.

Four remote cameras (HC600 Hyperfire, Reconyx, Inc., Wisconsin, USA) were placed along the road bordering the prescribed fire site closest to the nests of the study antechinus, recording the presence of predators (i.e. foxes, feral cats and wild dogs) along the road for three days before the fire and for eight days after the fire.

2.3. Torpor and activity definitions

The torpor onset Tb value of antechinus in the current study was calculated as 31.5 °C from equation 4 by Willis [49]. For antechinus this formula is particularly important as it provides a threshold estimate that detects shallow torpor bouts. Therefore, a Tb of 31.5 °C was used to calculate torpor bout entries and arousals for those that lasted longer than 30 min. The proportion of day spent torpid was calculated as the amount of time (min/per day, sunrise–sunset) each individual spent below the torpor onset Tb, but only for days when individuals did not change nests (i.e. the logger recorded the complete rest phase of an animal). Additionally, the depth of torpor bouts was measured as the absolute minimum Tb of each bout.

Activity periods were calculated from the time the individual's transmitter signal was absent on the logger to the time the signal returned for periods of longer than 30 min. These were calculated for three time frames: (i) whole day = 24 h from sunrise–sunset, (ii) daytime = sunrise–sunset, (iii) night time = sunset–sunset. Days when an individual changed nests and the logger did not record the end of the activity phase were excluded.

Figs. 1 (males) and 2 (females) provide example traces of recorded Tb and Tm over three days of the study period for an individual from each of the groups.

2.4. Data analysis

Statistical tests were undertaken in R (R v. 3.0.1, R Core Team, 2014) and StatistiXL (v 1.10, 2015). Means for each measured variable were first calculated for each individual and then an overall mean was derived from these individual means and are represented with ±1 standard deviation (SD); n = the number of individuals, N = the number of observations. A significance level (p) of < 0.05 was assumed. An analysis of variance (ANOVA; function ‘aov’) was undertaken to establish if Tb variables differed among the three sites. Linear mixed-effects models (package ‘nlme’) were fitted to test for differences among the treatment groups (pre-fire, post-fire and control) for the measured variables (activity, torpor bout duration, proportion of day spent torpid, minimum torpor Tb), with sex and daily minimum Tm as covariates, treatment:sex as an interaction term and individuals were included as a random factor. As a significant interaction was found between treatment and sex for all the measured variables (p < 0.0001) we performed separate analyses for males and females for each variable using the model stated above, but removing sex and the interaction term. For all models body mass was initially included, however, there was no significant effect of body mass on any of the variables so it was removed.

Percentages for the proportion of day spent torpid were arcsine transformed for analyses. Further, a residual plot to test for homoscedasticity and a normal Q–Q plot to test for normal distribution were used for all models. If there was a significant difference among the groups a post-hoc Tukey test (package ‘multcomp’) was performed to determine which groups were significantly different from each other. An analysis of covariance (ANCOVA; function ‘aov’) was performed separately for each of the groups to determine if any of the measured variables varied between the sexes, with daily minimum Tm as a covariate and individuals as a random factor. Least square linear regressions (activity and torpor bout duration against daily minimum Tm; function ‘Linear Regression’) for each of the treatment groups were compared using an ANOVA (function ‘Compare Linear Regressions’), separately for males and females. Data were pooled and regressed together if no significant differences were found in the slope and intercept among the treatment groups. As Tb was lower and more variable before the fire than after (Fig. 3), a significant relationship between torpor bout duration and daily minimum Tm could be established in the pre-fire group in both sexes. However, the narrower Tb range after the fire precluded a comparable analysis and therefore daily minimum Tm was included as a covariate when comparing torpor bout durations among all treatment groups as stated above.
night: $11.8 \pm 1.4\, ^\circ C$, $n = 15$; minimum day: $8.0 \pm 1.9\, ^\circ C$, $n = 15$; minimum night: $7.6 \pm 1.8\, ^\circ C$, $n = 15$).

Ground cover in the prescribed area before the fire and also the control area consisted of herbs, grass, Lomandra (a monocod with grass-like appearance), shrubs and also fallen timber. Especially along drainage lines, grasses and Lomandra formed dense mats totally obscuring the ground and any small animals underneath. After the fire about 70% of the ground cover was obliterated (NSW NPWS, personal communication) leaving only bare ground, but the mid- and upper-layers of the forest remained intact.

No mammalian predators were recorded by the trail cameras along the border of the prescribed area before the fire. During the first week after the fire three feral cats, two wild dogs and one fox were recorded, suggesting an influx of predators after the burn. Nevertheless, before the fire one male and one female antechinus from the pre-fire group were killed by a predator, most likely by a cat. However, after the fire no predation events were recorded in either the post-fire or control groups. Further, only one of the tagged individuals perished as a direct result of the fire.

3.2. Activity

Importantly, none of the individuals tracked in the burn site left the area after the fire. Antechinus were active on average between 6 and 13 h per day, with about a third of the activity occurring during the daytime and the remainder at night for individuals in the pre-fire and control groups (Fig. 4a,b). Radio-tracking revealed that when active during daytime hours animals regularly foraged in patches of matted grass and Lomandra where they could not be seen. Interestingly, the amount of time male antechinus spent active significantly increased at higher $T_a$ ($p = 0.005, R^2 = 0.2$, $f_{1,72} = 8.4, y = 0.5x + 3.2$), whereas there was no significant relationship for females ($p = 0.585, R^2 = 0.01, f_{1,74} = 0.3$). Consequently, when $T_a$ was lower in the pre-fire group females were active significantly longer over the whole day in comparison to males ($p = 0.003, f_{1,25} = 10.7$), whereas when $T_a$ was warmer in the control group males were active significantly longer than females ($p = 0.019, f_{1,33} = 6.0$; Fig. 4a,b).

3. Results

3.1. Habitat variables and predator numbers

Significant differences among the sites were found for mean ($p = 0.002, f_{2,31} = 8.0$), maximum ($p = 0.006, f_{2,31} = 6.1$) and minimum ($p = 0.008, f_{2,31} = 5.7$) daily $T_a$ (Fig. 3). However, for the post-fire and control animals that were measured at the same time in adjacent areas, all $T_a$ variables were statistically indistinguishable between the two sites (mean $= 11.1 \pm 1.2\, ^\circ C$, $n = 15$; maximum $= 16.5 \pm 1.7\, ^\circ C$, $n = 15$; minimum $= 7.4 \pm 1.4\, ^\circ C$, $n = 15$; Fig. 3). Conversely, the pre-fire group was measured before the burn when mean (8.3 ± 2.8 °C, $n = 8$), maximum (13.5 ± 2.6 °C, $n = 8$) and minimum (4.8 ± 2.8 °C, $n = 8$) daily $T_a$ were all on average ~3 °C lower (Fig. 3).
Both males and females in the post-fire group significantly reduced daytime activity, which did not differ between the sexes (p = 0.851, f1,73 = 0.04; Fig. 4a,b). However, while females in the post-fire group also reduced nocturnal activity, males in this group did not and therefore were active longer than females (p = 0.0002, f1,69 = 15.2; Fig. 4a,b).

For male antechinus the time spent active during the whole day differed significantly among the groups (p < 0.0001, t6,63 = 17.3; Fig. 4a) and activity duration was 1.7-fold longer for animals at the control (12.6 ± 3.5 h, n = 4, N = 28) site than individuals in both the pre- (7.9 ± 2.4 h, n = 4, N = 14) and post-fire (7.6 ± 0.6 h, n = 3, N = 31) groups. Importantly, at the fire site whole day activity did not change between before and after the fire. However, the partitioning between daytime and night time activity shifted after the fire. While pre-fire (2.7 ± 2.2 h, n = 4, N = 14) and control (4.7 ± 1.5 h, n = 4, N = 28) animals spent on average 28.2% and 23.4% of the total activity time during the day, diurnal activity was reduced to 4.4% in post-fire (0.3 ± 0.3 h, n = 4, N = 45) animals (Fig. 4a).

3.3. Torpor

After the fire, antechinus in the post-fire group expressed longer torpor bouts in comparison to the pre-fire and control groups (Fig. 5a,b). However, in all groups females were torpid longer than males (pre-fire: p < 0.0001, f1,16 = 173.1; post-fire: p < 0.0001, f1,69 = 495.4; control: p < 0.0001, f1,21 = 145.5) and torpor bouts were generally shallower in males than in females (pre-fire: p = 0.039, f1,18 = 4.9; post-fire: p < 0.0001, f1,64 = 67.4; control: p = 0.0154, f1,26 = 2.2; Fig. 6a,b). Furthermore, in both males (p = 0.037, R² = 0.4, f1,8 = 5.3, y = −15.5x + 159.7) and females (p = 0.011, R² = 0.5, f1,12 = 9.3, N = 45) group in comparison to both the pre-fire (11.6 ± 1.1 h, n = 4, N = 16) and control (10.3 ± 1.8 h, n = 2, N = 14) groups. This reduced level of activity encompassed both daytime and night time activity, but daytime activity more so. While pre-fire (3.3 ± 0.4 h, n = 4, N = 16) and control (2.4 ± 0.7 h, n = 2, N = 14) animals spent on average 28.2% and 23.4% of the total activity time during the day, respectively, diurnal activity was reduced to 4.4% in post-fire (0.3 ± 0.3 h, n = 4, N = 45) animals (Fig. 4b).
been previously proposed to play an important role in vertebrate resilience and the removal of ground cover. Antechinus reduced diurnal activity and increased torpor use to save energy and likely to avoid predation by foxes, wild dogs and feral cats. This flexibility in behavioural and physiological traits suggests that brown antechinus, and presumably other small heterothermic mammals, are able to adapt readily to sudden environmental changes and therefore enables them to survive catastrophic events. Indeed, phenotypic plasticity has been previously proposed to play an important role in vertebrate resilience in the light of climate change and habitat degradation [5,29].

Although the fire did not significantly change climate conditions in comparison to the control area and therefore thermoregulatory demands, the obliteration of ground cover increased visibility and thus exposure to predators. It is therefore not surprising that fire changed the activity patterns of brown antechinus in the post-fire group and both sexes reduced daytime activity to as little as 5% of pre-fire levels. Brown antechinus, like most small mammals, have been considered to be strictly nocturnal in the past [18,52]. However, from a thermoregulatory point of view activity during the daytime has energetic advantages especially in a cold climate, as the warmer temperatures and solar radiation from the sun reduce the amount of energy needed for thermoregulation while foraging [37]. Accordingly, our study clearly shows that male and female antechinus were active for 23 to 37%, respectively, during the daytime when ground cover was available in the post-fire and control groups. Daytime activity in other small dasyurids has been reported, but usually occurs in species that inhabit complexly structured habitats with shelters from visually hunting predators [30]. The kaluta (Dasykaluta rosamondae), for example, is currently the only known small dasyurid that is strictly diurnal in winter [20] and it occurs in areas densely covered with spinifex, a spikey grass that not only offers a visual but also a physical shield against most predators. Apparently, the presence of abundant ground cover is also important for brown antechinus [7], as they are often only found in habitats with low burn frequencies [27]. Our data suggest that this dense cover can be especially important during winter when shifting the activity period into the daytime can reduce thermoregulatory costs. However, since predator avoidance becomes crucial in a burnt landscape with limited protection, daytime activity would be risky and the observed reduction in daytime activity seen in the post-fire group would account for this.

Whereas females reduced overall activity, males in the post-fire group shifted their activity into the night and in comparison to pre-fire conditions males in both the post-fire and control groups maintained a high level of night-time activity. This was likely facilitated by the warmer Tb as activity in males increased with Tb which was possibly a response to higher food availability as they primarily feed on insects that often become more prevalent at warmer temperatures [39]. The overall higher levels of activity seen in males in the current study may be related to pre-mating season preparation to establish home ranges and indeed an increase in movements by male brown antechinus has been recorded as early as May [51], which corresponds to the timeframe of the current study. However, while a shift towards nocturnal activity somewhat reduces predation risk in comparison to diurnal activity, males the proportion of day spent torpid (p < 0.0001, fi,68 = 5.2; Fig. 5a) as well as mean torpor bout duration (p < 0.0001, fi,68 = 5.2) differed significantly among the groups. Males from the post-fire group were torpid -2.2-fold more each day in comparison to both of the other groups and torpor bouts were also -1.5-fold longer in the post-fire group (127.3 ± 38.4 min, n = 3, N = 26), in comparison to torpor bouts expressed by males from both the pre-fire (84.6 ± 37.1 min, n = 4, N = 11) and control (84.3 ± 26.8 min, n = 4, N = 20) groups. However, the depth of torpor bouts was the same for all groups (p = 0.228, fi,41 = 1.2; pre-fire = 31.0 ± 0.2 °C, n = 4, N = 8; post-fire = 30.8 ± 0.3 °C, n = 3, N = 22; control = 30.4 ± 0.6 °C, n = 4, N = 21; Fig. 6).

Similarly to the males, for females the proportion of day spent torpid differed significantly among the groups (p < 0.0001, fi,69 = 7.0; Fig. 5b) and females from the post-burn group were torpid -3.4-fold more each day in comparison to both of the other groups. Mean torpor bout duration of all bouts recorded for female antechinus also differed among the groups (p < 0.0001, fi,69 = 6.9) and torpor bouts were -2.5-fold longer in the post-fire (267.5 ± 61.9 min, n = 4, N = 57) group in comparison to torpor bouts expressed by females from both the pre-fire (118.0 ± 76.1 min, n = 4, N = 15) and control (94.2 ± 83.7 min, n = 2, N = 8) groups. Importantly, mean minimum Tb of female antechinus was 4 °C lower in the post-fire (26.0 ± 1.3 °C, n = 4, N = 44) group in comparison to both the pre-fire (29.7 ± 1.7 °C, n = 4, N = 14) and control (30.2 ± 1.3 °C, n = 2, N = 21) groups (p < 0.0001, fi,55 = 45.3; Fig. 6).

4. Discussion

The results of our study support our hypothesis and reveal that individual male and female brown antechinus responded behaviourally and physiologically to fire and the removal of ground cover. Antechinus reduced diurnal activity and increased torpor use to save energy and likely to avoid predation by foxes, wild dogs and feral cats. This flexibility in behavioural and physiological traits suggests that brown antechinus, and presumably other small heterothermic mammals, are able to adapt readily to sudden environmental changes and therefore enables them to survive catastrophic events. Indeed, phenotypic plasticity has been previously proposed to play an important role in vertebrate resilience in the light of climate change and habitat degradation [5,29].
physical protection at night is still limited after fire. Therefore, an overall reduction of both daytime and night time activity as observed in female antechinus appears to be a prudent strategy since animals resting in a secure location are less exposed to predation than when active [46]. Further, the major food resource of antechinus, insects, is often significantly reduced after a fire [6,47], suggesting that increasing energy savings is paramount. Importantly, this can only be achieved if daily energy demands can be lowered substantially, and in heterothermic endotherms an avenue for accomplishing this effectively is the use of torpor [36].

Torpor not only reduces energy demands, it also enhances predator avoidance [13,40,46,48]. This in turn appears to contribute to the lower extinction rates in heterotherm compared to homeotherm mammals [14,15]. While individual survival and longevity increase with the amount of time spent torpid [46], the state of torpor cannot continue forever, but must be interrupted for activity and especially reproduction [21,38]. Although torpor and reproduction are not mutually exclusive, certain phases during the reproductive cycle are often incompatible with entering torpor [22]. For example, in many species the hibernation season is often shorter in males, as they need to emerge earlier from hibernation to commence spermatogenesis and to secure territories for the mating period [3]. Similarly, male antechinus show increased levels of activity and metabolism, continue to grow and generally enter torpor rarely throughout winter (June–July), when the size of the testes is greatest [12,35,52]. The sexual differences for torpor use and activity patterns observed here are therefore not unexpected. The unusual reproductive strategy of male antechinus entails a high level of energy turnover that eventually culminates in complete male die-off [51,52] and therefore the time leading up to reproduction likely influences the behavioural and physiological responses to a catastrophic event such as fire. Consequently, while female antechinus minimised energy expenditure as well as predator exposure by considerably increasing torpor use after the fire, males increased torpor use only marginally, confirming our prediction that females would express more torpor than males. Nevertheless, the strategies employed by both sexes were successful at least in the short to medium term, as no deaths were recorded in the weeks after the fire event. Obviously, flexibility in behavioural and physiological traits in individuals of both sexes is advantageous, as it allows a population to respond to sudden changes in environmental conditions and food supply [5,28,29,41].

Even though the present study involved only a low-intensity burn, a large proportion of the habitat was severely degraded. Fires leave a fragmented patchwork of more or less suitable habitats that at least temporarily results in a significant reduction in habitat size and food availability, leading to increased levels of stress and decreased health in antechinus [9,16]. Fire regimes in Australia have been altered for millennia, with Aborigines regularly burning patches to maintain a variety of habitats for animals they could hunt [4]. However, these fire regimes have been radically changed since European settlement with drastic impacts on ecosystems contributing to the exceptionally high rate of mammal extinctions in Australia’s recent history [50]. While the negative impacts of modified fire regimes have also occurred in other parts of the world [17,44,47], in Australia such changes were founded by the introduction of new predators, namely feral cats and dasyus (A. swainsonii) antechinus were trapped along such drainage lines [8]. Interestingly, during our study radio-tagged antechinus were never observed to move into such refugia, even though the low intensity control burn left about 30% of the area untouched and further unburned habitat could be found in the control site just across a narrow forest trail. This could be a result of the unburnt areas already being occupied and defended, or a risk of venturing too far in the newly created open habitat and brown antechinus do generally show strong site fidelity [51]. Understanding the phenotypic plasticity of these mechanisms that link the fate of individuals with changes seen on a population level is vitally important for understanding fire ecology and to effectively manage fires in a conservation context.

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