RESEARCH ARTICLE

# The short-term effects of a routine poisoning campaign on the movements and detectability of a social top-predator

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Abstract Top-predators can be important components of resilient ecosystems, but they are still controlled in many places to mitigate a variety of economic, environmental and/or social impacts. Lethal control is often achieved through the broadscale application of poisoned baits. Understanding the direct and indirect effects of such lethal control on subsequent movements and behaviour of survivors is an important pre-requisite for interpreting the efficacy and ecological outcomes of toppredator control. In this study, we use GPS tracking collars to investigate the fine-scale and short-term movements of dingoes (Canis lupus dingo and other wild dogs) in response to a routine poison-baiting program as an example of how a common, social top-predator can respond (behaviourally) to moderate levels of population reduction. We found no consistent control-induced differences in home range size or location, daily distance travelled, speed of travel, temporal activity patterns or road/trail usage for the seven surviving dingoes we monitored immediately before and after a typical lethal control event. These data suggest that the spatial behaviour of surviving dingoes was not altered in

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ways likely to affect their detectability, and if control-induced changes in dingoes' ecological function did occur, these may not be related to altered spatial behaviour or movement patterns.

Keywords Behaviourally mediated trophic cascade  $\cdot$  Canis lupus dingo  $\cdot$  Detection probability  $\cdot$  Mesopredator release  $\cdot$ Passive tracking index  $\cdot$  Poison baiting  $\cdot$  Relative abundance indices . Sodium fluoroacetate or 1080

# Introduction

The ecological roles and functions of top-predators can be an important force contributing to ecosystem health and resilience (Estes et al. [2011](#page-11-0)). Such functions may include the suppression of sympatric mesopredators or prey. In many places, however, top-predator populations are subject to lethal control or harvesting in attempts to mitigate their economic, environmental and/or social impacts, such as predation of livestock or threatened fauna (e.g. Allen et al. [2013a](#page-10-0); Mech [2010;](#page-11-0) Ordiz et al. [2012\)](#page-11-0). Whether undertaken legally or illegally, lethal predator control is often achieved through broadscale distribution of poisoned baits (e.g. Glen et al. [2007b;](#page-11-0) Márquez et al. [2013](#page-11-0)). Although a reduction in the density of predators may be the most obvious direct result of lethal control programs, more subtle and indirect outcomes might accompany the removal of individuals from a population, such as spatial and/or temporal changes in space use by survivors (Haber [1996](#page-11-0)). While top-predator abundances may be resilient to lethal control programs due to compensatory population growth or reinvasion of areas subjected to control (e.g. Allen et al. [2013a;](#page-10-0) Stahler et al. [2013;](#page-12-0) Webb et al. [2011\)](#page-12-0), there remains the potential for top-predator populations to be functionally different in these areas due to altered social structures, prey preferences and/or movement behaviours (Haber [1996](#page-11-0); Salo et al. [2010\)](#page-12-0). Hence, the behaviour or

functions of top-predators may change subsequent to lethal control even if top-predator abundance recovers and remains relatively stable over time.

Understanding the direct and indirect effects of lethal control on top-predator populations is an important pre-requisite for interpreting the efficacy and ecological outcomes of toppredator control (Allen et al. [2012](#page-10-0); Haber [1996\)](#page-11-0). This understanding is also important for evaluating the reliability of common monitoring techniques used to assess the efficacy of lethal predator control programs, such as population indices or density estimates (Caughley and Sinclair [1994;](#page-11-0) Hone [2007\)](#page-11-0).

Mesopredator release theory (sensu Soulé et al. [1988](#page-12-0); but see Crooks and Soulé [1999](#page-11-0)) predicts that a reduction in higher-order predator densities can increase densities of lower-order predators—a numerical release. Alternatively or additionally, lethal top-predator control may also result in increased mesopredator activity through altering top-predator behaviour in ways that allow mesopredators to become more active at different times or places in the landscape—a behavioural release (Brown et al. [1999;](#page-10-0) Clinchy et al. [2013\)](#page-11-0). Mesopredator releases through either mechanism might have detrimental impacts for fauna at lower trophic levels under certain conditions (Crooks and Soulé [1999](#page-11-0); Johnson [2006](#page-11-0)). Much research has focused on indirect patterns in the numerical relationships between top-predators and mesopredators (Ripple et al. [2013](#page-11-0); Ritchie and Johnson [2009\)](#page-11-0). However, few studies focus on the actual behavioural responses of top-predators to lethal control, which remains an important research priority (Ordiz et al. [2012](#page-11-0); Salo et al. [2010](#page-12-0)). Some studies sample predators in areas with and without predator control, but when interpreting observed differences between these areas, many studies claim evidence for functional differences in controlled areas without attempting to measure the efficacy of, responses to, or outcomes of lethal control (Salo et al. [2010;](#page-12-0) but for examples, see Brook et al. [2012;](#page-10-0) Letnic and Koch [2010](#page-11-0)). The studies of Eldridge et al. ([2002](#page-11-0)), Kennedy et al. [\(2012\)](#page-11-0) or Allen et al. [\(2013a](#page-10-0)) provide noteworthy exceptions.

Efficacy assessments of lethal predator control programs ideally involve sampling predator populations in otherwise similar treated and untreated areas both before and after a lethal control event or over time (Glen et al. [2007b](#page-11-0); Hone [2007\)](#page-11-0). In many cases, sampling is undertaken using relative abundance indices, such as passive tracking indices, scat indices or camera-trap rates (e.g. Allen [2012c](#page-10-0); Brook et al. [2012](#page-10-0); Engeman [2005;](#page-11-0) Wang and Fisher [2012\)](#page-12-0). Although comparison of treated and untreated areas is a fundamental experimental design feature necessary for demonstrating the effects of lethal control on the target population (Allen et al. [2013b;](#page-10-0) Hone [2007](#page-11-0); Salo et al. [2010\)](#page-12-0), the utility of relative abundance indices for this purpose relies on the assumption that predator detectability is equal between treatments and remains unchanged by the control program itself (e.g. Archaux et al. [2012;](#page-10-0) MacKenzie et al. [2006;](#page-11-0) Pollock et al. [2002\)](#page-11-0). If the control program alters the detectability of surviving predators, then it may be difficult to separate the control-induced numerical responses from behavioural responses, potentially resulting in confounded conclusions about the 'control effect'. That observed index values post-control might reflect either behavioural and/or numerical responses is well known (e.g. Fleming et al. [2001](#page-11-0); Kennedy et al. [2012](#page-11-0)), but there remains a dearth of studies attempting to identify these effects separately. Rare or solitary top-predators experiencing typically low contact rates with conspecifics might be expected to respond indifferently to the loss of conspecifics, whereas common or social top-predators might be expected to respond more strongly.

In this study, we use a widely applicable approach to investigate the behavioural responses of Australian dingoes (Canis lupus dingo and other wild dogs) to a typical poisoning campaign which achieved moderate levels of population reduction. Dingoes are an appropriate model species for this study because they are a common social top-predator routinely exposed to lethal control for the protection of livestock and some threatened species (Fleming et al. [2001](#page-11-0)). Sodium fluoroacetate is the most common toxin used in poison baits for predator control in Australia; strychnine use has declined remarkably but is still used in some places, and para-aminopropiophenone is presently under consideration for registration as a broad-scale canid control toxin in the near future (APVMA [2008](#page-10-0); Fleming et al. [2006](#page-11-0); Southwell et al. [2013](#page-12-0)). Our primary aim was to determine what (if any) behavioural responses might occur following typical lethal control programs using manufactured poison baits. We use GPS tracking collars to investigate spatial and temporal changes in the movements of surviving dingoes. We discuss our results in light of what these changes may mean for fauna at lower trophic levels and for detecting predators using road-based relative abundance or activity indices, such as those commonly used to study dingoes (reviewed in Allen et al. [2013b\)](#page-10-0).

## Methods

Study site and environmental conditions

The study occurred within a 50-km radius of 30°31′33.45′′S, 140°39′37.72′′E on Quinyambie Station, a 12,000 km2 beef cattle-producing property located in the arid and sandy Strzelecki Desert of northern South Australia. The site has a mean annual rainfall of  $~160$  mm [\(www.bom.gov.au](http://www.bom.gov.au)), and landform is comprised of low parallel sand dunes and clay swales (Fitzsimmons [2007](#page-11-0)). Vegetation is dominated by lowgrowing hopbush (Dodonaea viscosa), buckbush (Salsola kali) and a variety of grasses and burrs including kerosene grass (Aristida spp.) and copperburr (Sclerolaena spp.) (Kutsche and Lay [2003\)](#page-11-0). A substantial amount of previous dingo research has occurred at the site, which includes additional description of the study area (e.g. Allen [2010](#page-10-0); Allen et al. [2013a;](#page-10-0) Allen and Leung

[2012;](#page-10-0) Brawata and Neeman [2011](#page-10-0); Caughley et al. [1980](#page-11-0); Letnic and Koch [2010;](#page-11-0) Newsome et al. [2001;](#page-11-0) Pople et al. [2000\)](#page-11-0).

Like much of Australia, northern South Australia received record-breaking rainfall between November 2009 and March 2011. Above-average monthly rainfall events occurred multiple times at the site during this period, with approximately 5 years' worth of rainfall occurring over the 12–15 months preceding the study (Fig. 1); the latest significant fall prior to our study occurring in February 2011, which inundated most swales with floodwaters for several weeks. The vegetation response to such high and ongoing rainfall was outstanding, with many extant vegetation types responding positively (B. Allen, personal observations). This facilitated rapid population growth of invertebrates, birds and small mammals, with subsequent increases in dingo activity and abundance during this period (Allen [2011\)](#page-10-0). In chronological order, successive plagues of locusts (Chortoicetes terminifera), beetles, crickets and native (Notomys fuscus) and introduced (Mus musculus) rodents were experienced. Rabbits (Oryctolagus cuniculus) had been in high abundance in 2008 but were reduced to near-undetectable levels by the end of 2009 because of drought (Allen [2010\)](#page-10-0). At the time of the present study, rabbits were distributed across the entire study site but were still in relatively low densities due to recurring bouts of Myxoma virus and Rabbit Haemorrhagic Disease during 2010 (P. Jonas and G. Conners, Quinyambie Station Mangers, pers. comms.). Cattle had not been grazed in the study area since September 2009, when the site was completely destocked following several severe dust storms at the climax of the drought. The present study began as crickets and rodents (predominately introduced house mice and native dusky hopping-mice) were super-abundant and surface water was readily available, in the latter stages of the annual autumn breeding season for dingoes. Cattle were stocked in areas adjacent to, but not within, the immediate study area during the study period.

#### Dingo capture, tracking and control

Dingoes were captured towards the centre of a  $1,400 \text{ km}^2$ cattle grazing area with soft-catch foot-hold traps (#3 'Jakes', JC Conner, USA) over three nights in early May 2011 using a

variety of lures and decoys. Traps were checked daily both in the late afternoon and in the morning between dawn and 1100 h. Captured dingoes were assessed for trap-related injuries (such as swelling, lacerations, broken or dislocated bones) that might affect their subsequent movement behaviour once released. Standard weights and measures were taken, including age, which was estimated visually from an assessment of tooth wear and breeding status, noting the date of capture and the annual breeding season of dingoes. Captured dingoes were fitted with Sirtrack® Argos-linked GPS datalogging collars (~450 g) before release. Collars were programmed to record a GPS point each 30 min continuously from the time of activation. To avoid undue human influence on dingo behaviour, no attempts were made to physically locate collared dingoes until after baiting had occurred. On the few occasions when collared dingoes were located, they were approached discretely and were not disturbed. In most cases, collars were recovered several months after the present study period when dingoes were humanely shot from a helicopter.

Not all expected GPS points were obtained (Table [1](#page-3-0)). Some minor technical errors with some collars on some occasions resulted in variable timing between sequential GPS points, which has the potential to produce unreliable values when calculating travel distances (or the sum of straight line-lengths between sequential GPS points). The GPS data were, therefore, filtered to include only GPS points where the time interval between sequential points was 25–35, 55–65, 85–95 or 115–125 min. This meant that only GPS points recorded at  $\sim$ 30-min intervals were included, allowing for maximum gaps of up to  $\sim$ 2 h between sequential points. Time gaps over 2 h were excluded because greater time gaps between GPS points increase the likelihood of underestimating the true distance travelled between sequential GPS points (Rowcliffe et al. [2012\)](#page-12-0). Less than 2.6 % of all the GPS data obtained was excluded by this process. Because the time gap between sequential points still varied between  $\sim$ 30–120 min, we further converted 'travel distance' values to 'speed of travel' (a function of distance and time) to account for the time gaps. These two measures are different but not independent; we report the results for both measures. Positional accuracy of filtered GPS



Fig. 1 Monthly rainfall trends at Quinyambie Station, 2001–2011. Arrow indicates approximate study period

<span id="page-3-0"></span>

points was assessed using the HDOP values (range  $1-15$ ) recorded with each GPS point, with a lower value indicating a more accurate GPS location. Results from the present study were consistent with previous studies at the site, with mean HDOP values ranging between 1.57 and 1.97 (Table [1\)](#page-3-0) or <50 m on-ground error (see also Allen [2012a](#page-10-0)). Hence, all filtered GPS points were considered suitable for inclusion in all analyses, and none were excluded based on HDOP values.

Approximately 65 % of Quinyambie Station (or  $\sim$ 8,000 km<sup>2</sup>) is not baited and is rarely subject to opportunistic shooting. The study was conducted in the southern portion of the property where opportunistic shooting occurs regularly and where baiting has occurred twice each year since March 2008 (Allen [2012b](#page-10-0); Allen et al. [2013a](#page-10-0)). Our study occurred in 2011, when manufactured Doggone®-substrate baits were distributed on the 14th and 18th June along all roads and around all livestock watering points within the  $1,400 \text{ km}^2$ baited area, at a rate of one bait every 300–500 m according to local practise and poison-use regulations (Bird [1993\)](#page-10-0). Such trail-based baiting practises are typical of those used in most arid and semi-arid areas of the Australian rangelands (Allen et al. [2013a](#page-10-0); Fleming et al. [2012](#page-11-0)). The study period was divided into pre-baiting (28th May–13th June), duringbaiting (14th–30th June) and post-baiting periods (1st–17th July). Bait uptake data indicated that most baits were taken within 17 days after baiting, by the end of the 'during-baiting' period (Allen [2011\)](#page-10-0). Pre- and post-baiting periods were, therefore, limited to equal periods of 17 days duration in order to provide consistent sampling timeframes for analyses, and to minimise any potential influence of seasonal behavioural changes (associated with annual breeding patterns) on the results. Comparisons of movement behaviours were made between pre- and post-baiting periods to avoid the dailychanging circumstances as extant dingoes consumed remaining baits during baiting.

## Dingo movements

Spatial analyses of changes in dingo behaviour and detectability (or the likelihood of a dingo being detected on or near a road) were undertaken in ArcView v9.3 (ESRI Inc.), using the extensions XTools Pro v7.0 (Data East LLC [2005](#page-11-0)), Hawths Tools v3 (Beyer [2004\)](#page-10-0) and Home Range Tools v3 (Rodgers and Carr [1998](#page-11-0)).

### Home range size changes

To investigate changes in dingo home range sizes, we compared their mean 100 % minimum convex polygons (MCPs; Harris et al. [1990;](#page-11-0) Mohr [1947\)](#page-11-0) between pre- and post-baiting periods (i.e. a '17-day home range') using a two-sample twotailed  $t$  test (assuming unequal variances). MCPs were chosen over other home range estimators for their ease of calculation,

their widespread use for describing dingo home range sizes (e.g. Claridge et al. [2009](#page-11-0); Robertshaw and Harden [1985;](#page-11-0) Thomson et al. [1992\)](#page-12-0), and because we were not concerned with utilisation distributions within the home range for this analysis. We chose 100 % MCPs over 95 % MCPs (or others) because we sought to include all GPS points, and because forays outside the 17-day home range seldom occurred for any dingo during this period (see below). Although calculation of MCPs technically requires the use of independent GPS points (Harris et al. [1990](#page-11-0)), and our GPS points were not independent, the use of non-independent points seldom changes the resulting MCP values in any substantial way (Gese et al. [1990;](#page-11-0) Swihart and Slade [1997\)](#page-12-0). Regardless, our primary interest was not in determining the size of the 17-day home range per se, but in detecting relative changes in the size of the home range between pre- and post-baiting periods. For this purpose, the consistent application of any home range estimation technique suffices.

#### Home range location changes

To investigate changes in the location of dingo home ranges, we calculated the distance between the centroids of the 100 % MCPs used in the home range size change analyses between pre- and post-baiting periods. We further assessed whether or not these centroids remained within the overall core area of each dingo. Core areas were calculated as the 50 % adaptive kernel (AK; h: 1; Harris et al. [1990](#page-11-0); Katajisto and Moilanen [2006;](#page-11-0) Seaman and Powell [1996](#page-12-0)) of all GPS points from the time of collaring until the time of death (typically several months; Table [1\)](#page-3-0). Overall home range sizes were similarly calculated as the 90 % AK. The use of AKs was chosen over other home range estimators because, for this analysis, we were concerned with utilisation distributions within the home range, which cannot be identified using MCPs. All the available GPS data for each dingo (from the time of collaring until the time of death) were used to generate AK home range data for this analysis because size and utilisation of home ranges are known to vary seasonally (Allen [2009;](#page-10-0) Purcell [2010](#page-11-0)), and using all the available data attempted to capture this variability.

## Daily activity period changes

To investigate changes in the daily activity periods of dingoes, we plotted the overall mean distance travelled (all dingoes pooled) for each hour of the day for both pre- and post-baiting periods. This analysis was repeated using overall mean speed of travel for each hour of the day for both pre- and post-baiting periods. Preliminary results indicated that collared dingoes at the site exhibited three activity peaks at 1700, 000 and 0700 h (see below), so two-sample two-tailed  $t$  tests (assuming unequal variances) were used to identify reductions or increases

in dingo activity at these dusk, midnight and dawn times between pre- and post-baiting periods.

# Daily distance travelled and speed of travel

To investigate changes in the daily distance travelled by dingoes, we compared the overall mean distance travelled per day (i.e. the mean of the daily sum of line-lengths between sequential GPS points) for the pre- and post-baiting periods using repeated measures ANOVA. The same analysis was performed to assess changes in mean daily speed of travel between periods.

## Road or trail usage

To investigate changes in the time dingoes spent on roads or trails, we compared the mean percentage of GPS points within 250 m of a road between pre- and post-baiting periods. We further compared the mean percentage of days that roads were contacted between pre- and post-baiting periods. Both analyses were undertaken using repeated measures ANOVA. Considering average dingo travel speeds (see below) and the 30-min duty cycle of the collars, we arbitrarily considered GPS points within 250 m of a road to reflect road use in order to capture true road use that might otherwise have been 'missed' had road use been restricted to within 100 or 50 m of a road.

# **Results**

# Dingo capture, tracking and control

A total of 13 dingoes (6 male and 7 female, weighing 13–19 kg) and a 5-kg male red fox (Vulpes vulpes) were captured in 70 trapnights during the 4th–6th May 2011 (Table [1](#page-3-0)). No other species were inadvertently captured. Each dingo was fitted with a collar and released before 11.15 a.m. the morning after capture. Most dingoes were caught on the front foot, and processing time was typically <10 min (median=7 min). Ten dingoes were released with minor swelling to the captured paw only, and the three others were released with minor lacerations to the captured paw also (where dingoes had bitten themselves on the toes below the padded jaw of the trap). Such trap-related injuries are common and considered minor (Fleming et al. [1998;](#page-11-0) Marks et al. [2004\)](#page-11-0). No broken or dislocated bones were observed. Thus, there were no trap-related injuries considered to significantly affect the subsequent movement behaviour of dingoes by the time the pre-baiting period commenced 3 weeks later on the 28th May 2011, nor were these injuries considered to influence the calculation of overall AK values. Three dingoes (and their collars) could not be located at the conclusion of the study, and two collared dingoes died from bait ingestion in the 'during-baiting' period (Allen [2011\)](#page-10-0). Data from two recovered collars were corrupted and unusable. Thus, seven collared dingoes survived

baiting and provided data suitable for analysis (Table [1\)](#page-3-0). Additional bait-efficacy data not reported here demonstrated a 30–40 % reduction in the relative abundance of dingoes depending on how efficacy was measured (Allen [2011\)](#page-10-0). Such reductions are consistent with previous control programs at the study site and elsewhere (Allen et al. [2013a;](#page-10-0) Fleming et al. [2001;](#page-11-0) but see Thomson [1986\)](#page-12-0).

## Dingo movements

The 17-day 100 % MCP home range sizes of dingoes ranged between 12.42 km<sup>2</sup> (Dingo17) and 194.42 km<sup>2</sup> (Dingo28) in the pre-baiting period and  $16.65 \text{ km}^2$  (Dingo24) and 78.23  $km^2$  (Dingo28) in the post-baiting period (Table [2\)](#page-6-0). Consistent changes in mean home range size were not apparent ( $t = 0.96$ , df=6;  $p = 0.37$ ); home range sizes increased. decreased or remained unchanged for different dingoes. The distance between home range centroids between the pre- and post-baiting periods ranged from 192 m for Dingo21 to just over 40 km for Dingo28. However, tracking data showed that Dingo28 went on an exploratory foray soon after capture and release, was absent from his home range during the prebaiting period and returned back to his home range during the post-baiting period, giving the misleading impression of a home range shift. Ignoring this extraterritorial foray (the only one observed during the study), the greatest distance between centroids was only 3.01 km. Pre- and post-baiting centroids remained within the 90 % AK home range of all seven dingoes, and within the 50 % AK core area of three of these dingoes (Fig. [2\)](#page-6-0), suggesting that despite apparent changes to the location of 17-day 100 % MCPs, no dingo shifted their home range grossly in response to baiting, but rather continued occupying the same home range area.

Dingo activity was highly variable between days over time (Fig. [3](#page-7-0)), but within days, collared dingoes at our study site exhibited crepuscular peaks of activity at 1700 (dusk) and 0700 h (dawn) in the pre-baiting period (Fig. [4](#page-8-0)). Three activity peaks were observed at 1700, 000 and 0700 h in the post-baiting period, suggestive of a behavioural shift from decreased morning to increased night-time activity between early June and early July. No demonstrable changes in dusk activity peaks were observed between pre- and post-baiting periods (for mean distance travelled  $t=0.83$ , df=7,  $p=0.43$ ; for speed of travel  $t=$ 0.33, df=9,  $p = 0.75$ ). Dawn activity peaks were demonstrably lower post-baiting (for mean distance travelled  $t = 3.06$ , df=11,  $p=0.01$ ; for speed of travel  $t=3.35$ , df=10,  $p=<0.00$ ) when activity during the hour after midnight was higher (for mean distance travelled  $t = -2.25$ , df=9,  $p = 0.05$ ; for speed of travel  $t =$  $-2.20$ , df=10,  $p = 0.05$ ; Fig. [4](#page-8-0)).

We found no difference in the overall mean daily distance travelled by dingoes between pre- and post-baiting periods  $(F_{1,6}=0.26, p=0.63)$ , the average speed dingoes travelled each day  $(F_{1,6}=0.003, p=0.96)$ , the percentage of GPS points on

Dingo ID	90 % $AK^{\wedge}$	50 % $AK^{\wedge}$	$100\%$ MCP pre*	100 $\%$ MCP post $\sim$	Change in 100 % MCP
Dingo17	31.51	6.56	12.42	23.48	189.05 %
Dingol <sub>8</sub>	29.33	3.13	70.03	17.65	174.80%
Dingo21	18.89	2.97	24.43	22.46	$18.06\%$
Dingo23	32.23	5.21	30.02	32.50	18.26 $%$
Dingo24	16.57	4.61	30.94	16.65	146.19%
Dingo28	286.33	28.53	194.42	78.23	159.76%
Dingo29	29.77	4.17	36.33	37.98	14.54 $%$
Mean	63.52	7.88	56.94	32.71	$12.42 \%$

<span id="page-6-0"></span>Table 2 Overall home range (90 % adaptive kernel) and core area (50 % adaptive kernel) sizes, and changes in the 17-day 100 % MCP home range size (km2 ) for seven dingoes in the Strzelecki Desert of northern South Australia between pre- and post-baiting periods in 2011

 $^{\wedge}$ Based on all available data from the time of capture until the time of death, \*based on data from the pre-baiting period only and  $\sim$  based on data from the post-baiting period only

roads ( $F_{1,6}$ =0.02,  $p$ =0.89) or the percentage of days that roads were contacted  $(F_{1,6}=1.28, p=0.30;$  $(F_{1,6}=1.28, p=0.30;$  $(F_{1,6}=1.28, p=0.30;$  Table 3). Despite fine-scale spatial variation in road usage for some dingoes between the preand post-baiting periods (i.e. some dingoes used different sections of the road more or less frequently between periods; Fig. [5\)](#page-9-0), such behaviours did not affect the overall time that dingoes spent on roads or the frequency that roads were contacted. Thus, with the exception of a moderate shift from dawn to night-time activity, all other measures showed no consistently significant change in dingo movement behaviours between pre- and postbaiting periods.

# Discussion

Our study demonstrates that lethal predator control does not influence the movement behaviour of surviving top-predators



Fig. 2 The overall home range (90 % AK; *light grey*) and core area (50 % AK; dark grey) sizes of six dingoes in the Strzelecki Desert of northern South Australia in 2011, showing the 17-day 100 % MCP home ranges during the pre-baiting (solid lines) and post-baiting (dashed lines) periods, inclusive of centroids for the 100 % MCPs in the pre-baiting (triangle) and post-baiting (squares) periods. Stars indicate the location of artificial water sources

in a consistent manner. Despite moderate population reductions in the order of 30–40 % (Allen [2011\)](#page-10-0), we found no demonstrable or consistent differences in home range size or location (Table 2; Fig. 2), daily distance travelled (Table [3](#page-9-0)) or road usage (Fig. [5](#page-9-0)) between the 17-day pre- and post-baiting periods for the seven surviving adult dingoes we monitored. The only change detected was a moderate reduction in dawn activity peaks, coincident with increased night-time activity (Fig. [4\)](#page-8-0). These data reveal no substantial effects of baiting on the movement behaviour of individuals that survived typical levels of lethal control under the conditions present during the study.

Because no data were available from collared dingoes in a paired site not exposed to lethal control, we cannot exclude some plausible alternative explanations for our observations. First, it is possible that normal seasonal changes in dingo activity (associated with whelping) may have masked baiting-induced changes, or have been responsible for the observed shift from dawn to night-time activity. However, dingoes' use of roads typically declines at this time of year (e.g. Allen et al. [2013a;](#page-10-0) Corbett [2001;](#page-11-0) Purcell [2010\)](#page-11-0) in preparation for birth and pup-rearing, which means that if baiting did affect dingoes' road use, then it increased it (to the same magnitude as the seasonal reduction), in order for the observed behaviours to appear stable (Fig. [5](#page-9-0)). Age, season and social status may also affect home range size. However, by the time of breeding in late autumn, all dingoes are approximately 12 months old or older, most juveniles unable to assimilate into a pack have already been killed or expelled, and home range sizes contract to a relatively stable size in preparation for whelping (Allen [2009;](#page-10-0) Corbett [2001](#page-11-0); Fleming et al. [2001;](#page-11-0) Purcell [2010](#page-11-0)). Given the relatively short duration of our study conducted at the latter end of the annual breeding season, seasonal factors are unlikely to have influenced our home range size and location analyses in any substantial way. Although individual changes in location of the 17-day 100 % MCP home range were apparent (Table 2), comparison with the overall 90 % AK home ranges (inclusive of several months of data subsequent to baiting) indicate that home range

<span id="page-7-0"></span>

Fig. 3 Daily variation in the mean daily distance travelled (bottom) and mean speed of travel (top) for eight dingoes in the Strzelecki Desert of northern South Australia between May 2011 and February 2012

locations did not shift in response to baiting (Fig. [2](#page-6-0)). Second, it is possible that rapid reinvasion of uncollared individuals may also have 'filled the hole' in any observed behavioural changes that might have occurred as a result of dingo control. In other words, dingo behaviour might have changed as a result of baiting, but immigrating dingoes moved into the area and prevented the opportunity for resident survivors to alter their behaviour. However, data from a variety of other measures collected simultaneously at the site (e.g. bait uptake data) typically recorded significant reductions in dingo activity on roads (Allen [2011\)](#page-10-0), suggesting that rapid immigration did not occur during the post-baiting period, or that immigrating dingoes avoided roads, which is unlikely (Allen [2009](#page-10-0); Corbett [2001](#page-11-0); Mahon et al. [1998\)](#page-11-0). Third, and perhaps most likely, given that poison-baiting occurs twice annually at the site, and the dingoes we monitored were typically resident adults (Allen [2012b](#page-10-0); Table [1\)](#page-3-0), these individuals have probably been exposed to but not taken baits in the past, and it is conceivable that they have experience with losing conspecifics. In other words, the individual dingoes we monitored may be indifferent to the loss of other related or adjacent dingoes, which may explain the non-responses we observed. Regardless, our results suggest that the overall population-level effect of baiting on dingo behaviour was negligible, and that the

movement behaviours of the surviving dingoes we monitored were not altered in consistent ways that may affect their ecological function or detectability.

Though many studies investigate the potential ecological outcomes of lethal dingo control (reviewed in Glen et al. [2007a](#page-11-0) or Letnic et al. [2012,](#page-11-0) but see Allen et al. [2013b](#page-10-0)), few use techniques able to measure the actual behavioural responses of dingoes, such as continuous follows (or direct and continuous first-hand observation), GPS tracking collars or camera traps. Nevertheless, it has been postulated that lethal dingo control results in social destabilisation, with surviving dingo populations being unable to repel expected increases in mesopredator (e.g. fox or feral cat Felis catus) abundance (e.g. Wallach et al. [2009](#page-12-0)). For example, after pooling singlesurvey camera-trap data across multiple years, seasons and sites, Brook et al. ([2012](#page-10-0)) reported that dingoes occupying baited areas had lower dusk activity levels than dingoes occupying nearby unbaited areas, coincident with higher activity levels of cats at dusk in baited areas. However, an absence of any data on control efficacy or pre- and post-baiting observations meant that the study could not reliably determine actual behavioural shifts of predators to lethal control. In contrast, Wang and Fisher [\(2012\)](#page-12-0) record a positive spatial relationship between dingoes and cats in baited areas, and negative temporal relationships only at certain times of the year. Only individual dingoes from small and newly formed packs are

<span id="page-8-0"></span>Fig. 4 The daily activity patterns (distance travelled, bottom; speed of travel, top) of seven dingoes in the Strzelecki Desert of northern South Australia during prebaiting (solid line) and postbaiting (dashed line) periods in June–July 2011. Vertical bars indicate standard errors of the mean for each hour



needed to kill invading foxes and feral cats (Moseby et al. [2012\)](#page-11-0), and because we found no change in the mean daily distance travelled by dingoes, the chances of a surviving dingo intercepting invading mesopredators might not be diminished by baiting. Indeed, our GPS tracking data do not show consistent changes in any dingo movement behaviours that would compromise the ability of dingoes to repel mesopredators, such as reduced daily distance travelled, speed of travel, home range size or a shift in home range location (Tables [2](#page-6-0) and [3,](#page-9-0) Figs. [2](#page-6-0) and [5\)](#page-9-0).

Dingo activity at dusk did not change as a result of baiting at our site either (Fig. 4), and the observed shift from dawn to night-time activity is most likely due to seasonal effects unrelated to baiting, as dingo populations reach peak whelping times during the post-baiting period in July (Corbett [2001\)](#page-11-0). At this time, dingoes tend to pups in their dens during the colder hours of the morning; dawn temperatures were typically below freezing, and as low as −6 °C during the study. This interpretation is supported by several first-hand observations. Dingo27 (a 13-kg adult female) died from bait ingestion in the duringbaiting period with four fully-grown foetuses just days away from birth (Table [1;](#page-3-0) Allen [2011\)](#page-10-0). Dingo18 (another 13-kg female that survived baiting) was located resting in her den

shortly after dawn on the 7th July when pups only a few days old were observed. Seven pups were later observed and photographed at the entrance of this den on the 2nd and 3rd August. Dingo29 and an uncollared male partner were also located resting in the sun at the entrance of a den shortly after dawn on the 8th July. These observations support the collar data, suggesting that, during whelping, adult dingoes primarily confine their activity within short proximity to (or even in) the den during dawn when temperatures are coldest and pups are most vulnerable to hypothermia.

That the surviving adult dingoes we monitored did not alter their movements consistently in response to lethal control supports a growing body of research indicating that poison baiting programs often target juvenile dingoes or the expendable surplus (e.g. Allen [2013\)](#page-10-0); juveniles contributing relatively little to overall social stability in canids (e.g. Boitani and Ciucci [1995](#page-10-0); Corbett [1988;](#page-11-0) Tallents et al. [2012](#page-12-0); Thomson [1992\)](#page-12-0). These observations do not suggest that lethal control changes dingo movement behaviour in ways that would facilitate an increase in mesopredator populations or impacts. A series of conclusive predator-manipulation experiments conducted at 12 sites across Australia (inclusive of the present study site) confirm that contemporary dingo control practises <span id="page-9-0"></span>Table 3 Changes in daily distance travelled, speed of travel and road usage for seven dingoes in the Strzelecki Desert of northern South Australia between preand post-baiting periods in 2011



do not necessarily result in mesopredator population increases (Allen et al. [2013a\)](#page-10-0).

That road-use by dingoes remained unchanged (e.g. Fig. 5) further suggests that baiting efficacy assessments involving comparisons of road-based relative abundance indices between treatments remain robust, given that detectability of survivors (or the likelihood of a surviving dingo using a road) did not change. Indeed, the percentage of GPS points on roads and the frequency that roads were contacted did not change between pre- and post-baiting periods in our study. However, fine-scale spatial changes in road-use did occur (Fig. 5). Although the lack of a nil-treatment area means that we cannot



Fig. 5 The overall 90 % AK home range size (*light grey*) and 50 % AK core area (dark grey) for three dingoes (Dingo23, top; Dingo17, centre; Dingo18, bottom) in the Strzelecki Desert of northern South Australia in 2011. Black dots represent GPS points within 250 m of roads in the prebaiting (left) and post-baiting (right) periods. Solid lines indicate the location of roads and stars denote water sources

be certain that such changes were baiting-induced, this result has important implications for the design of studies that sample dingoes on roads. Sand plots or camera-traps are often placed along roads at <1,000-m intervals (e.g. Allen et al. [2013a;](#page-10-0) Catling and Burt [1995;](#page-11-0) Letnic and Koch [2010](#page-11-0)); an approach which attempts to maximise the number of sampling points while accounting for logistical constraints and the likely home range size of the study animal. Our results, which suggest that within their home ranges, dingoes use different sections of the road at different times or seasons (as do dingoes elsewhere; Allen [2009](#page-10-0)), serve as reminder that sand plot or camera-trap sampling must be undertaken in a systematic way at large-enough spatial scales to detect population-level responses of predators to lethal control (Engeman [2005](#page-11-0); Glen et al. [2007a\)](#page-11-0). Should the temporal or spatial scale be too small (i.e. too few cameras or sand plots for a given area and period of interest), such studies risk obtaining severely biased results not representative of true population-level responses at larger scales relevant to predator management (Allen et al. [2013b;](#page-10-0) Allen et al. [2012\)](#page-10-0). Sampling predators off-road is unable to escape this problem of seasonal bias because avoidance of roads in a given season is akin to selection of off-roads in that same season. Considering that there is typically much greater area to place observation stations off-road for sampling activity than on-road, attempting to sample off-roads is also likely to be a highly inefficient means to observe activity, an effect already demonstrated for an ecological analogue of dingoes, the coyote (Canis latrans; Engeman et al. [2002\)](#page-11-0). Dingoes are also known to follow the tracks or paths of humans to off-road sampling stations (B. Allen, P. Fleming, personal observations), confounding attempts to 'hide' the sampling station

<span id="page-10-0"></span>off-road without influencing predator activity. Lethal control also primarily occurs along roads at our study site and in many other places (Allen et al. 2013a; Fleming et al. [2001\)](#page-11-0), which means that changes in animal activity or abundance might be most easily detected and expected to occur in close proximity to roads.

In conclusion, our results provide a useful step forward in our understanding of top-predator responses to lethal control and have important implications for dingo researchers and managers. Our results show that ongoing lethal control programs which remove moderate numbers of dingoes from areas routinely subject to control can fail to produce consistent spatiotemporal behavioural changes of a magnitude that may be necessary to elicit positive responses from mesopredators, and that lethal control does not always change the detectability of surviving dingoes on roads. This implies that should dingo control-induced mesopredator release be observed, it is likely to be a numerical response to lower dingo densities, not a behavioural response to altered dingo movements. Observed changes in predator abundance or activity indices post-baiting are also likely to reflect numerical changes more so than behavioural changes, provided population sampling is undertaken appropriately. Of course, surviving dingoes may respond differently to lethal control programs that achieve greater population reductions than ours, programs conducted in places where extant dingoes have not previously been exposed to lethal control, or programs conducted under different environmental conditions. Thus, there are likely to be cases where the movement behaviour of dingoes will change as a result of lethal control. Whether or not such changes ultimately result in sustained and undesirable outcomes for dingoes or fauna at lower trophic levels requires a much greater understanding of the direct and indirect behavioural effects of control (Allen et al. 2012). We encourage the use of GPS tracking data to explore the behavioural responses of predators to interventions. Such studies may include simultaneous GPS-tracking of sympatric predators exposed to lethal control, and we hope that our findings prompt future research assessing the actual responses of dingoes and other predators to lethal control in comparable ways.

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