



Ecological factors influencing invasive predator survival and movement: insights from a continental-scale study of feral cats in Australia

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Abstract Feral cats (*Felis catus*) pose a significant global threat to biodiversity, primarily through predation, disease and competition. A key gap in parameterizing models for improving management decisions for feral cat control relates to factors that drive feral cat survival and movement in the wild. Our study objective was to conduct the first continental-scale analysis of survival rates and displacement distances

for feral cats. We collated data on 528 feral cats from telemetry studies in naturally-vegetated landscapes across Australia. Using Cox-proportional hazards models, we investigated the effects of sex, presence of larger predators (dingoes, *Canis familiaris* and introduced foxes, *Vulpes vulpes*), presence of introduced prey (rabbits, *Oryctolagus cuniculus*), body mass, landscape productivity and feral cat density on feral cat survival. We also analysed the effects of sex, body mass and landscape productivity on feral cat displacement using linear mixed model analysis. Feral

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cat survival was positively associated with presence of dingoes and increasing body mass, whereas there was no clear association between feral cat survival and sex, presence of rabbits, or cat density. Presence of foxes had a strong negative effect on feral cat survival, but the hazard ratio was associated with considerable uncertainty. Net displacement of male feral cats was nearly two times further than that of females, and the proportion of feral cats making long-distance movements was greater in landscapes with low productivity. Increasing body mass of feral cats was positively related to net displacement, with heavier cats moving further. Analysis of metadata from telemetry studies can provide valuable insights into wildlife survival rates and movement behaviour. Our findings will help inform the development of effective management strategies and improve feral cat management for biodiversity conservation.

Keywords Feral cats · Survival rate · Net displacement · Fox · Dingo · Rabbit · Landscape productivity · Body mass · Australia

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Introduction

Free-ranging cats (*Felis catus*) pose a global threat to biodiversity conservation (Doherty et al. 2015; Medina et al. 2011). Cats are widely distributed around the world, and their generalist behaviour and diet makes them adaptable and capable of surviving diverse and extreme conditions (Fitzgerald 1988; Lepczyk et al. 2023; Loss et al. 2013; Medina et al. 2011; Nogales et al. 2013). Free-ranging cats are considered feral (hereafter ‘feral cats’) if they persist in self-sustaining populations living independently of humans (Lepczyk and Calver 2022).

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Feral cats have been listed as one of the 100 worst invasive species in the world, being responsible for at least 14% of global bird, mammal and reptile extinctions on islands (Lowe et al. 2000; Medina et al. 2011). They negatively impact native species through predation (Medina et al. 2014), disease transmission (Dubey 2002) and competition (Glen and Dickman 2005; Medina et al. 2014).

Developing effective invasive predator management strategies requires robust estimates of the species' survival, reproduction, and movement rates (e.g., Byers et al. 2019; Fleming et al. 2006; Moro et al. 2018). Reproductive behaviour and fecundity of feral cats are well documented; average litter sizes are similar worldwide, while the timing of breeding season varies with latitude (Woinarski et al. 2019). An increasing databank of GPS and radio-tracking studies also provide estimates of home range sizes in many regions (Bengsen et al. 2015; Kays et al. 2020; Nottingham et al. 2022).

In contrast, the survival rate and behaviour of feral cats are poorly understood. Studies of baseline survival rates for feral cats have focused either on cats in anthropogenic landscapes (excluding pet cat studies; Devillard et al. 2003; Gehrt et al. 2013; Nutter et al. 2004; Schmidt et al. 2007) or on remote islands (Danner et al. 2010; Van Aarde 1983). A summary of these studies and the estimated survival rate for each population is provided in Table S1. Alternatively, studies have examined the effects of management and diseases on feral cat survival (e.g., Andersen et al. 2004; Jones and Downs 2011). Very little is known about feral cat survival in naturally-vegetated landscapes, or how environmental conditions and demographic characteristics such as sex and body mass affect feral cat survival rates.

Telemetry studies (i.e., remote data collection from freely moving animals through radio or satellite collaring) have shown that feral cats can move long distances (10 to over 200 km, e.g., Jansen et al. 2021; McGregor et al. 2016; Roshier and Carter 2021). But the influence of sex, body mass and habitat types on the likelihood or frequency of these movements is not well known. Information on such long-range movements in telemetry studies may be limited by lack of consistent long-distance data, costs involved in collar deployment and data acquisition, operating life of collars, satellite signal reception, potential for technical failures and loss of collars (Matthews et al. 2013).

A better understanding of feral cat survival rates and long-distance movement behaviour is important for improving management decisions around feral cat control. Survival rates can be used to provide baseline rates for designing or improving management programs and to parameterize population models (Bull et al. 2009; Gaillard et al. 1998; Stenseth et al. 2001). Likewise, long-distance movements influence recolonization rates and recovery from lethal population management programs (e.g., Comer et al. 2018). Understanding movement between locations can provide information on how populations respond to habitat conditions and their distribution, as displacement and population dynamics are interlinked (Bowler and Benton 2005; Dunning et al. 1995).

Here we conduct the first continental-scale analysis of feral cat survival and long-distance displacements, with a focus on feral cats in naturally vegetated landscapes. Feral cats in Australia remain a major driver of species decline since their introduction by European settlers in the eighteenth century (Abbott 2008; Fisher et al. 2014; Woinarski et al. 2015). There are numerous examples of feral cats causing localised population declines and extirpations of prey species on both mainland Australia and offshore islands (Burbidge and Manly 2002; Risbey et al. 2000).

We hypothesised that feral cat survival rates are higher for female cats (e.g., Danner et al. 2010; Kaeuffer et al. 2004; Schmidt et al. 2007) and are negatively influenced by the presence of foxes and/or dingoes due to antagonistic interactions or dominance from these larger predators (Kennedy et al. 2012; Molsher et al. 2017). Conversely, we hypothesised that introduced rabbits benefit cat survival (McGregor et al. 2020), as does cat density (indirectly), as densities may reflect prey abundance (Legge et al. 2017; Read and Bowen 2001). Further, we hypothesised that cats survive better in more productive landscapes: cat home-ranges are larger in landscapes with lower productivity (Bengsen et al. 2015; Nottingham et al. 2022). We hypothesised that larger cats survive better, as they can hunt a greater range of prey sizes (Moseby et al. 2020; although see Fleming et al. 2020). Similarly, we hypothesised that larger male cats may move further due to their polygynous mating system and feral cats moving longer distances in landscapes with low to moderate level of resources (Bengsen et al. 2015; McGregor et al. 2015; Mirmovitch 1995; Say and Pontier 2004). We therefore predicted

that survival rates would be higher for female cats, be lower in the presence of foxes and dingoes, higher in areas with introduced rabbits, be positively associated with landscape productivity, cat density and increasing body mass. We further predicted that the average distance between a feral cat's first and final capture locations would be higher for males and vary among body mass classes and habitat types.

Materials and methods

Data collection

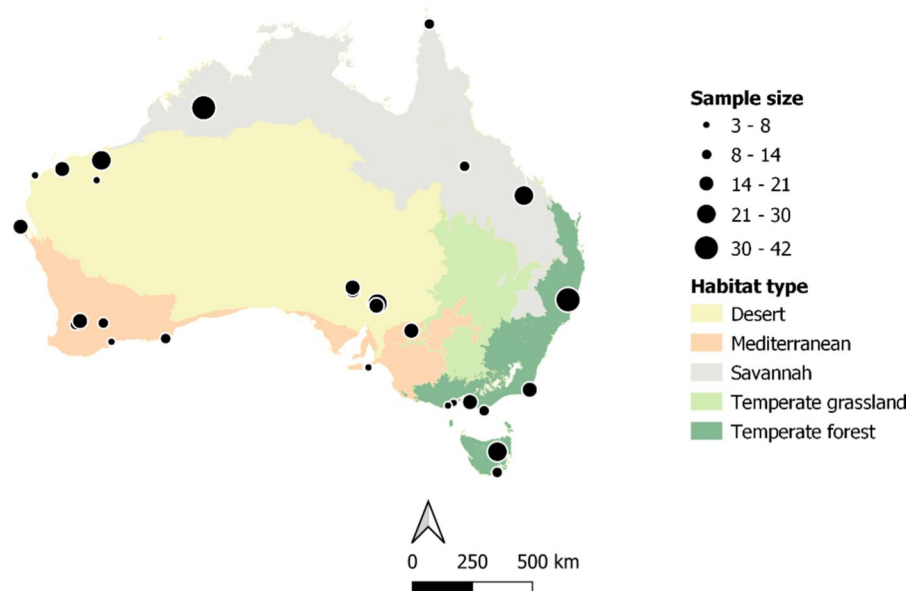
We compiled data on the fate and net displacement of individual feral cats from published and unpublished telemetry studies on 528 feral cats (354 male and 173 female) from thirty-one study areas across mainland Australia and offshore islands (Fig. 1). We define 'net displacement' as the linear distance between a feral cat's first and final capture locations. This value is a conservative estimate of the maximum distance moved, and does not distinguish between dispersive and home ranging movements as we were unable to collate all movement locations for individuals.

The telemetry studies were conducted from 2006 to 2023, with the study area defined as the area where a researcher collared cats for their study. All studies had a minimum of three cats from a single

study area. We collated information on the first and last recorded capture dates and location for cats, along with their sex, body mass, status at last capture (dead, alive or unknown), cause of death (if known), and the presence/absence of foxes, dingoes and rabbits in the study location (researcher reported, Figs. S1, S2, S3). For the studies that were trialling the effectiveness of feral cat baiting programs, we used the last known alive date before poison baits were deployed, if the cat died from bait intake. Similarly, cats that were trapped, shot or euthanized by the researchers were also considered to be alive until their date of death.

To test our predictions, we used two metrics for landscape productivity: (1) we classified Australia into distinct habitat types, with each study falling into one of these simplified categories: deserts, Mediterranean, Savannahs, temperate forests and temperate grasslands (Fig. 1; adapted from Australian Government Department of Sustainability, Environment, Water, Population and Communities 2012); (2) we obtained yearly rainfall data for the 12 months before a feral cat's last capture and a 50-year rainfall average for the study locations using the R package 'cropgrowdays' (Baker and Mortlock 2022). We then derived the rainfall deviation for each cat as $\text{rainfall deviation} = (\text{rainfall in the last year} - 50 \text{ year rainfall average}) / 100$. We included rainfall deviation as cat population densities can fluctuate with varying

Fig. 1 Data collected from telemetry studies on feral cats (*Felis catus*) across Australia. Different colours represent the different habitat types. Black dots denote study locations, with dot size indicating the number of cats collared



rainfall conditions in different environments in Australia (Legge et al. 2017).

We categorised feral cats into body mass classes, based on their weight during initial capture and the groupings in Jones and Coman (1982). We also distinguished between the sexes where sample sizes allowed (>125 individuals of each sex). This resulted in five classes: very small (male ≤ 2.2 kg, female ≤ 1.9 kg), small (male 2.2–3.2 kg, female 1.9–2.5 kg), medium male (3.2–4.5 kg), medium female (2.5–4.5 kg), and large cats (>4.5 kg). Only four (0.7%) of the large cats were female. Cat density estimates were obtained from field observations for each study where available ($n=7$). When unavailable, we derived predicted densities from published studies in the same area ($n=2$) or from Legge et al. (2017) ($n=20$). We removed very small cats from the compiled dataset as this class was not adequately represented across different habitat types ($n=23$). There were no cats from temperate grasslands (Fig. 1). Cats with an unknown status at last capture were not included in survival analyses.

Data analysis

Survival analysis

We used two different metrics to evaluate the effects of covariates on feral cat survival. Hazard ratios (HR) can be considered as an estimate of relative risk, which is the risk or probability of an event happening (i.e., death of a feral cat) relative to a reference group. A HR of 1 implies equal hazard in each group (i.e., an equal likelihood of dying), a HR >1 means increased risk relative to the reference group (i.e., an increased likelihood of dying) and HR <1 means reduced risk in comparison to the reference group (i.e., a reduced likelihood of dying). For example, $HR = \text{Hazard}_{\text{female cats}} / \text{Hazard}_{\text{male cats}} = 2$ means that the risk of dying for female cats is two times higher than male cats (i.e., female cats have lower chance of surviving).

We used a mixed-effects Cox proportional hazard model (Therneau 2022) to evaluate the effects of sex, body mass (as a continuous variable), habitat type, presence/absence of foxes, dingoes and rabbits, rainfall deviation and cat densities on the hazard ratio for feral cats. We fitted an additive model with all predictors to test our hypotheses. Study area was fitted as a

random effect (intercept) to account for potential clustering in the dataset. We accounted for differences in study duration by truncating the data to 52 weeks (Fig. 2). All feral cats that were collared for more than 52 weeks (9.2% of individuals) were classed as ‘alive’ at the end of the 52-week period.

We tested for the assumption of Cox models that the hazard ratio predicted by the model is proportional over time by plotting the scaled Schoenfeld residuals over time, where a non-zero slope over time can indicate a lack of proportionality (Cox and Oakes 1984; Kalbfleisch and Prentice 2002). The assumption of proportional hazards was supported for our predictor variables. We checked for correlations between predictor variables using pairwise Pearson’s correlation test. We removed habitat type from our hazard analysis as savannahs and presence of dingoes were moderately correlated ($R=0.54$, Fig. S4) and including both caused model instability; all study sites in savannahs had dingoes present. We also used a generalised linear mixed model with a binomial distribution to test for any association between the monitoring duration of feral cats (in weeks) and their status during the last capture (0 = dead, 1 = alive, NA = unknown), study area was included as a random effect.

Annual feral cat survival rate is the proportion of feral cats alive at the end of 52 weeks. We estimated the annual survival rate of feral cats using the duration of monitoring and status at last capture for each collared individual. We estimated annual survival probabilities separately for sex, body mass class, presence/absence of foxes, dingoes and rabbits, and habitat types using Kaplan Meier estimates (Kaplan and Meier 1958). The estimates also account for feral cats that were collared for <52 weeks by taking their survival into account until their last known event.

For both the hazard ratio and survival rate analyses, we considered that p values <0.05 indicated that estimates were significantly different for categories within each predictor variable. Note that confidence intervals around the estimated survival rates tend to increase over time, due to the decreasing sample size (i.e., censored individuals that were monitored for <52 -week). We provide both metrics for the effects of covariates on survival, as the effects of covariates may be misinterpreted from providing just hazard ratios. Hazard ratios estimate relative differences in the ratio of risk involved among animals as a function of covariates but cannot estimate changes in survival

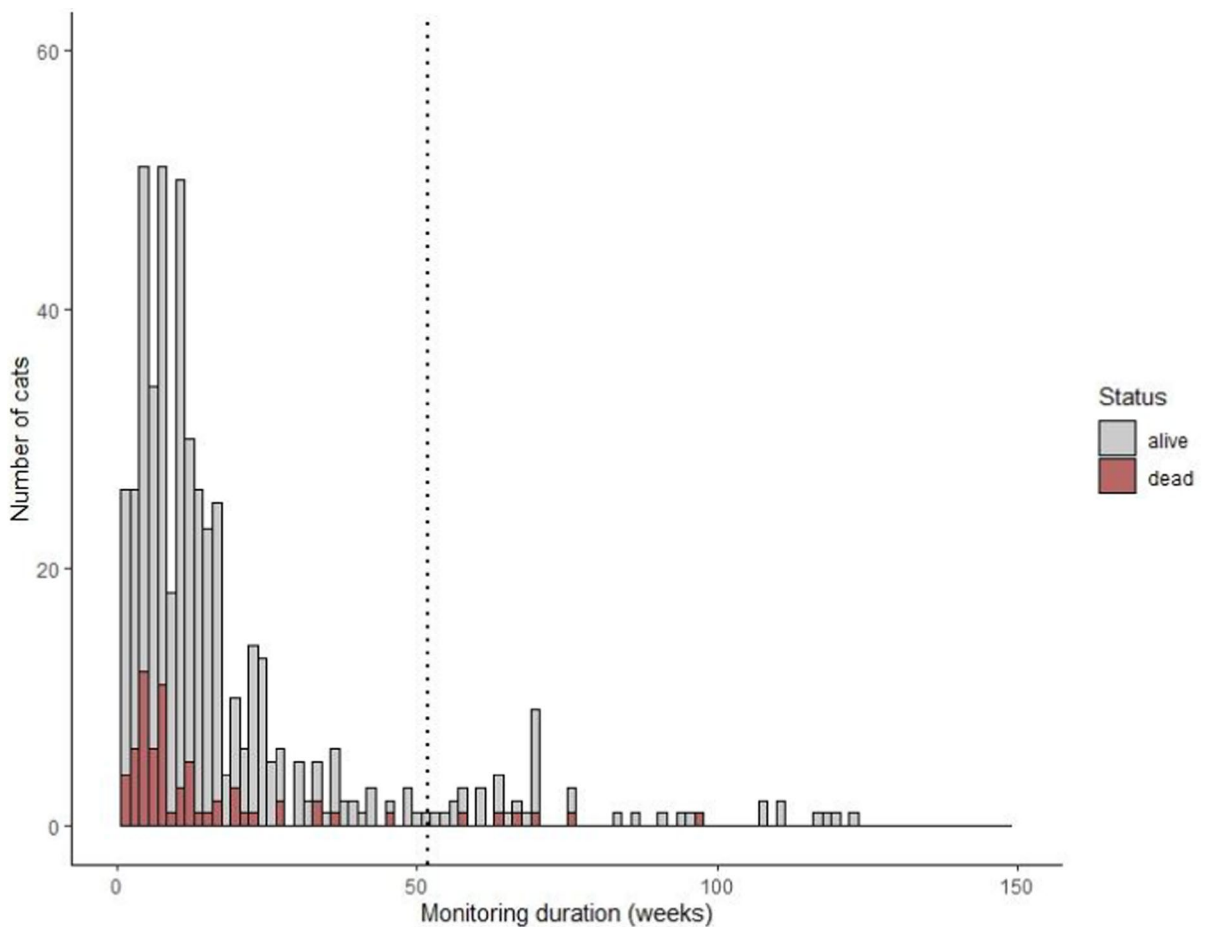


Fig. 2 Duration of cats collared (in weeks) in the collated dataset. Dotted line represents the 52-week period

over time. A high hazard ratio may not translate to a high difference in survival rates, as survival rates estimate the proportion of feral cats that survive after a particular time (Sashegyi and Ferry 2017; Spruance et al. 2004). Providing survival curves along with hazard ratios assists to interpret the effects of covariates on feral cat survival.

Net displacement

We used generalised linear mixed models to estimate mean net displacement for feral cats. Study area was included as a random effect (intercept). We compared net displacement for sex, body mass classes and habitat types. We log-transformed displacement to improve normality. We then back-transformed the modelled net displacement along with confidence

intervals (CIs) to interpret differences between feral cat demographics. We used 95% confidence intervals to provide a range of possible mean distances and not as a dichotomous inference of significance versus non-significance (Austin and Hux 2002; Payton et al. 2003).

We also looked at the proportion of feral cats that moved long distances between the first and last capture date. Using the largest observed home-range for a feral cat in a recent global meta-analysis (32.32 km² stable home-range, Nottingham et al. 2022), we calculated the diameter or maximum distance moved by a feral cat within its home-range to be 6.41 km (assuming a circular range). We therefore defined long-distance movement by feral cats as any displacement over 7 km. We acknowledge that we may have incorporated some intra-range movements in

our ‘long distance movements’ as some cats may have home-ranges larger than 32 km² or non-circular ranges (e.g., Comer et al. 2018; Roshier and Carter 2021). Furthermore, range size is influenced both by the analytic approach and definition of a stable home range. We tested for any association between monitoring duration and net displacement using Pearson’s coefficient and generalised linear models. All analyses were carried out in ‘survival’ (v 3.5-0; Therneau 2014), ‘coxme’ (v 2.2-18.1; Therneau 2022) and lme4 (v 1.1-31; Bates et al. 2015) in R (v4.2.1; R Core Team 2021).

Results

Survival analysis

We ran Cox-proportional hazards analysis on 498 cats. Sixty-three cats died within 52 weeks of collaring (12.6%), with most of these deaths attributed to unknown causes (n=56). Some showed signs of being depredated (n=4) or possibly killed by snakes (n=3). A further 389 cats (78%) were monitored for <52 weeks but were alive at the conclusion of the tracking period (Fig. 2). There was a significant but weak negative association between the monitoring duration of feral cats and their status during the last capture (R²m=0.36, p<0.01).

There was no association between sex and relative risk of mortality for feral cats (p=0.77; Fig. 3). Fox presence was estimated to double the risk to feral cats, but the associated uncertainty was high, and the effect was not significant (HR=2.04, p=0.12; Fig. 3). In contrast, there was strong evidence of an association between dingo presence and relative risk for feral cats (p=0.03; Fig. 3). The presence of dingoes was strongly associated with reduced risk to feral cats (i.e., positively associated with survival, HR=0.38; Fig. 3). There was no clear association between rabbit presence and relative risk of feral cats (p=0.30, Fig. 3).

Body mass of feral cats had a strong negative association with the risk of feral cats dying, with relative risk decreasing with every kilogram increase in weight (HR=0.66, p=0.01; Fig. 3). The model also supported a small but significant effect of increasing yearly rain deviation on relative risk to feral cats, with relative risk increasing when rainfall in the last 12 months was higher than the long-term average (HR=1.17, p=0.05). There was no relationship between cat density and relative risk to feral cats (p=0.32, Fig. 3).

The average annual survival rates for feral cats in these telemetry studies was 0.75. Consistent with our hazard ratio analysis, survival rates were similar for male and female cats (p=0.74, Fig. 4a). Survival rates for feral cats were significantly different

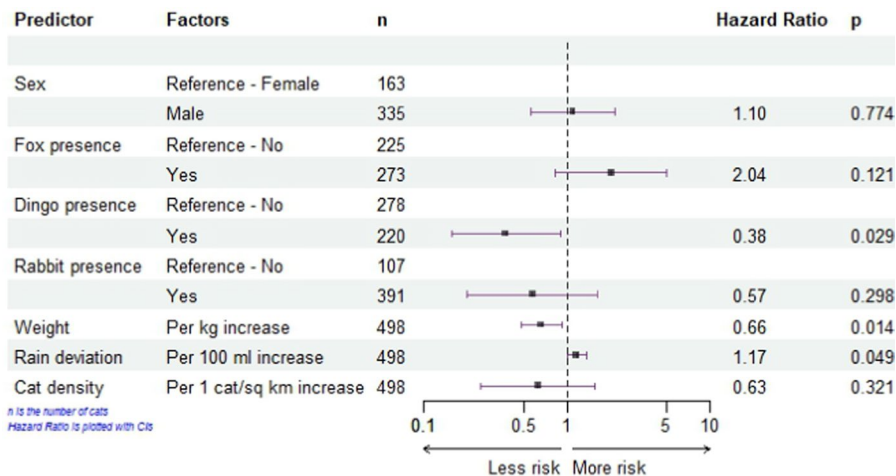


Fig. 3 Model summary for the additive Cox proportional-hazards model on feral cat survival. Data are from cats across Australia (n=498). Reference in factor refers to the reference levels for each predictor. Hazard ratios are plotted with confi-

dence intervals. Hazard ratios >1 indicates increased hazard (lower survival) for the predictor compared to the reference level whereas hazard ratios <1 indicates reduced risk (higher survival). P indicates the level of significance

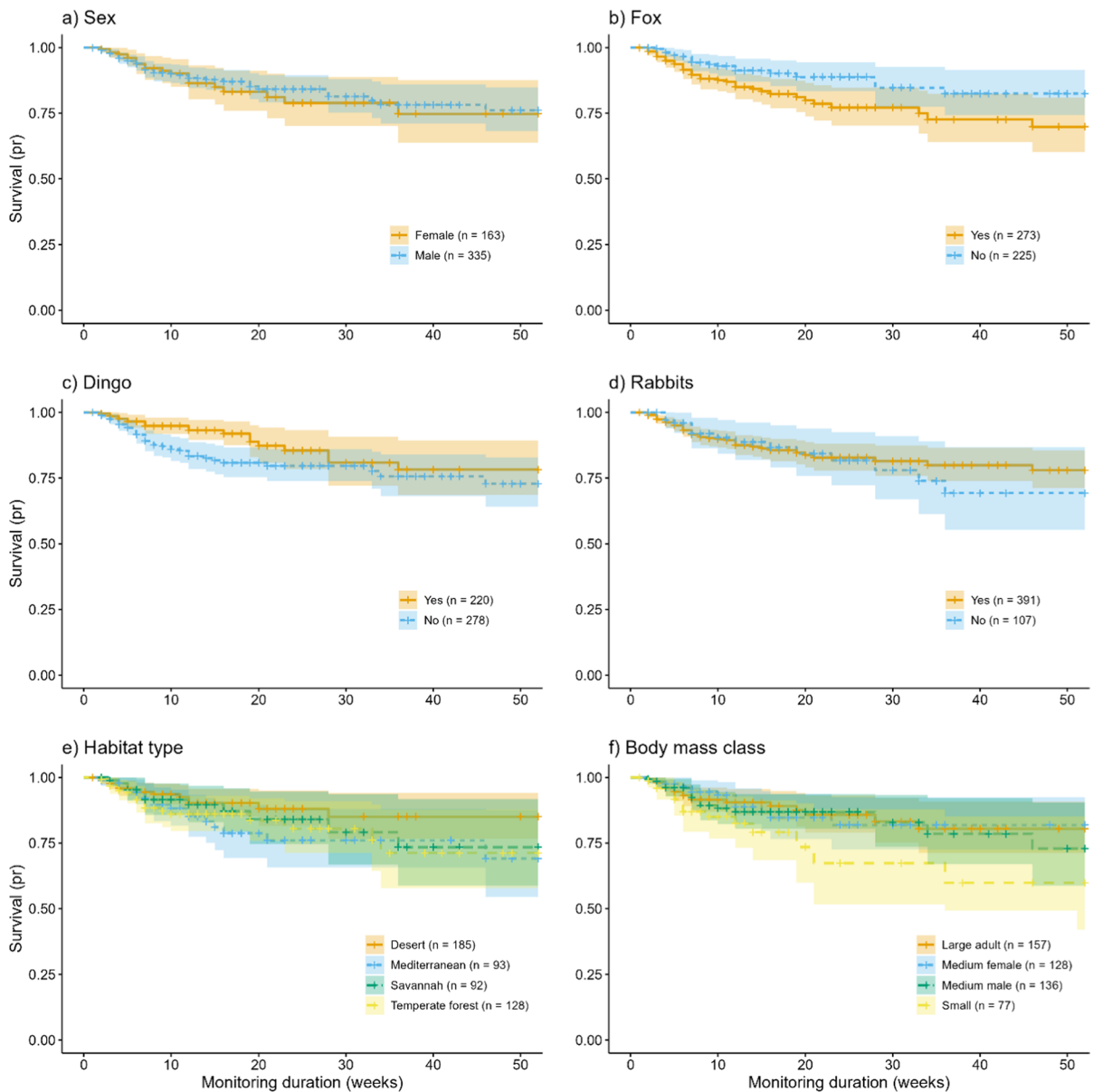


Fig. 4 Annual survival curves and confidence intervals for feral cats based on their sex, age categories, presence/absence of foxes, dingoes and rabbits, and habitat types. Lines indicate the survival function (i.e., the proportion of individuals surviv-

ing between landscapes with and without foxes ($p=0.01$, Fig. 4b). An opposite relationship was observed for dingoes: feral cat survival was significantly higher in landscapes with dingoes present ($p=0.02$; Fig. 4c). There were no significant differences in survival rates for landscapes with rabbits present compared to landscapes with no rabbits ($p=0.78$, Fig. 4d).

ing). Crosses indicate when an individual was ‘censored’ (due to conclusion of the tracking period or dropping out of the study)

Feral cat survival rates were similar across all habitat types ($p=0.29$, Fig. 4e). Cats in deserts tended to have the higher average survival rates, but confidence intervals around all estimates overlapped substantially throughout the monitoring period (Fig. 4e). There were no significant differences ($p=0.14$, Fig. 4f) in feral cat survival rates among body mass categories

(classed based on their sex and weight). Small cats tended to have the lowest survival rate, but confidence intervals were wide and substantially overlapped the estimates for the other body mass classes.

Net displacement

First and last capture locations were available for 447 cats. We found a significant but weak positive log–log relationship between the monitoring duration and net displacement of feral cats ($R=0.17$, $p<0.01$, Fig. S5). Displacement of cats over their tracking period varied from 0.01 to 164 km. The distribution of displacement was right skewed, with approximately 78% of the cats moving <5 km and only 7% moving more than 25 km. The mean displacement of male feral cats was nearly two times further (2.33 km, CI: 1.48–2.47) than that of female cats (1.21 km, 0.90–1.63). Displacement distances varied among habitat types. Feral cats in Savannah and Mediterranean habitat had overall higher mean displacement compared to desert and temperate forests. The effect of life-stage class also differed among different habitat types (Fig. 5).

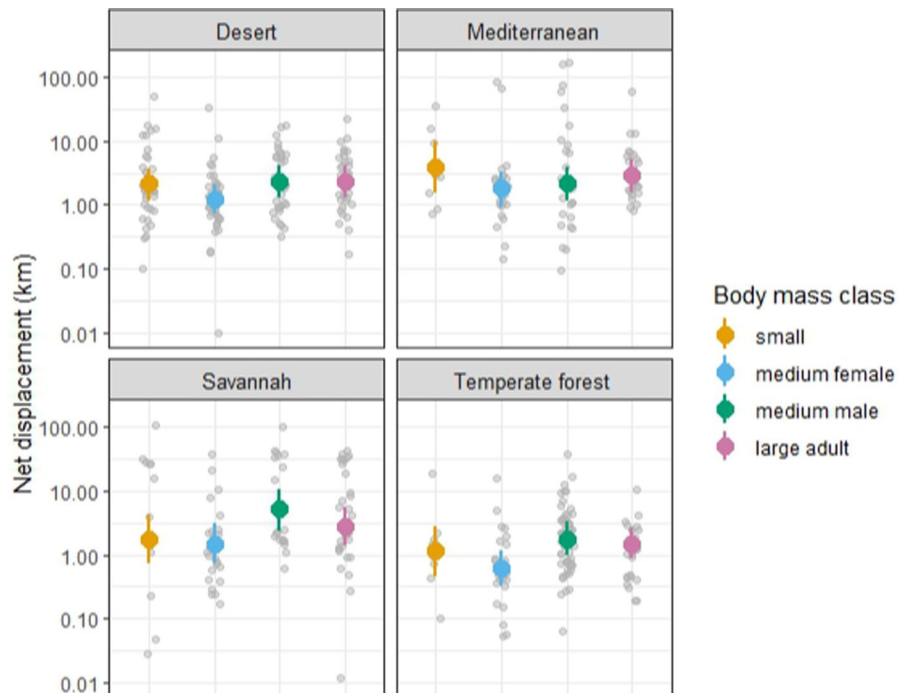
Mean displacement was lowest for medium female cats in all habitat types. For this class, mean

displacement was <2 km in all habitat types, with the lowest estimate in temperate forests (0.61 km, Fig. 5). Except in Mediterranean habitats, small cats also had lower mean displacement compared to adult male cats and large cats (Fig. 5). The highest mean displacement was for medium male cats in savannahs (5.06 km); this was nearly two times further than the distances for other body mass categories in the same habitat type. Mean displacement estimates and confidence intervals for feral cats in all habitat types and classes are provided in Table S2.

Among the feral cats that moved long distances between their first and final locations (more than 7 km, $n=74$), a much higher proportion were male (75.6%). When categorised by body mass and sex, the smallest proportion were medium female cats (12.1%) followed by small cats (22.9%), large cats (27%) and medium male cats (37.8%).

The proportion of cats that made long-distance movements was lowest in temperate forests, with only 8% of cats in temperate forests moving more than 7 km (10 of 124 individuals). This was followed by 12.7% of feral cats in desert (18 of 141 individuals) and 17.5% in Mediterranean forests (16 of 91 individuals) making long-distance movements. The

Fig. 5 Estimated mean displacement and confidence intervals for feral cats in each body mass class and habitat type. Y-axis is net displacement in the power of 10 and grey points show the raw distribution of net displacement



highest proportion of long-distance movements were observed in savannahs (32.9%, 30 of 91 individuals).

Discussion

Our study is among the first to conduct a continental-scale study of feral cat survival rates and long-distance movements. Contrary to our predictions, the presence of dingoes had a positive relationship with the survival of feral cats in telemetry studies. Greater body mass also had a positive relationship, whereas higher than average rainfall deviation had a negative relationship with feral cat survival. There was also some evidence that fox presence had a large negative effect, but this was associated with considerable uncertainty when the influence of other factors was also accounted for. Our predictions that net displacement would vary between body mass classes and habitat types were supported, with male cats in our study showing nearly two times higher net displacement than female cats and feral cats moving further in landscapes with lower productivity.

A better understanding of predator interactions and their functional roles in varying ecosystems can improve conservation outcomes (Ritchie and Johnson 2009). Not accounting for these complex interactions may reduce the efficacy of management actions or result in adverse outcomes such as the ‘release’ of other invasive mesopredators or prey species (Bergstrom et al. 2009; Courchamp et al. 1999). In many parts of Australia, lethal control of dingoes and foxes is widely conducted, with the aim of protecting livestock (both species) or wildlife (foxes). In contrast, feral cat control is primarily aimed at protecting native fauna with regulations for lethal control varying from state to state. Hence, it is important to understand whether these dominant predators affect feral cat survival.

There was some evidence for a negative association between fox presence and feral cat survival: the survival rate for feral cats was significantly higher in the absence of foxes (univariate survival analysis). Similarly, the multivariate hazard model estimated that the average relative risk to feral cats doubled when foxes were present, but this effect was associated with considerable uncertainty and so was non-significant. Foxes and feral cats are opportunistic predators and are sympatric over much of their

range in Australia (Woinarski et al. 2019). Foxes may potentially affect feral cat survival through direct predation or competition for food due to their large overlap in diet (Fleming et al. 2022; Molsher et al. 2017; Risbey et al. 2000). Our finding of higher feral cat survival rates in the absence of foxes is consistent with studies that have shown increased cat activity in areas with reduced fox activity (Marlow et al. 2015; Read and Bowen 2001; Risbey et al. 2000) and higher cat densities in areas where foxes are controlled (Rees et al. 2023). The discrepancy between the univariate survival model and multivariate hazard model may be because the hazard model simultaneously accounted for the effects of other predictors. As discussed below, fox presence was associated with several other predictors; further controlled experiments are needed to better quantify the effects of foxes and fox management on feral cat survival.

Contrary to our predictions, the presence of dingoes was associated with lowered risk to feral cats, with risk of mortality for feral cats in the telemetry studies nearly 2.5 times lower in landscapes with dingoes. Likewise, survival rates for feral cats were significantly higher in landscapes with dingoes. Note that we cannot infer any relationship between dingo presence and the survival of young or small feral cats, because these animals were not present in our telemetry dataset. Previous studies on interactions between dingoes and feral cats have mainly reported negative relationships (Brawata and Neeman 2011; Kennedy et al. 2012) or no clear relationships (Fancourt et al. 2019; Hunter et al. 2018; Letnic et al. 2009). A possible explanation might be that dingoes indirectly benefit feral cats through their negative effects on foxes. Moseby et al. (2019) tested potential mesopredator release using field data and prior information on predators and found a strong negative association between foxes and dingoes, but only a weak negative effect of dingoes on cats. Fancourt et al. (2019) also found no significant negative impact of dingoes on feral cat presence in the absence of foxes. The predator cascade hypothesis predicts that top-down suppression of mesopredators by large predators may indirectly benefit smaller predators (Levi and Wilmers 2012; Prugh and Sivy 2020). For example, foxes in North America can be released from top-down control by coyotes when wolves suppress coyote populations (Levi and Wilmers 2012; Newsome and Ripple 2014). Similar agonistic interactions may occur between competing

predators in Australia. Concurrent estimates of fox and dingo abundance in our study areas (Stobo-Wilson et al. 2022) would provide more insights into how these dominant predators influence feral cat survival.

We found no relationship between rabbit presence and relative risk to feral cats. While survival rates were slightly lower for feral cats in landscapes without rabbits, absence of rabbits may not pose a strong relative risk to feral cats as they are known to switch their diet to native prey when rabbit populations are low (McGregor et al. 2020). Our correlative approach meant that we were unable to fully distinguish the effects of foxes, dingoes and rabbit presence from each other, or from habitat type. For example, there are no foxes or rabbits in the tropics in northern Australia, whereas all study areas within savannahs in our study had dingoes present. Similarly, foxes and rabbits are common in south-east Australia where dingoes are heavily controlled through exclusion fencing and various lethal population management programs (Fleming et al. 2006). This highlights the need for more causal experiments to test the effects of manipulating the predator and introduced prey guild on native fauna.

We found that heavier feral cats were at significantly lower risk of dying. Likewise, small cats in our study had the lowest survival rate compared to large cats, and to medium male and female cats (albeit non-significantly). High levels of mortality among small cats or cats under 1 year old have also been observed in other studies (Jones and Coman 1982; Van Aarde 1983). Small cats may be vulnerable to territorial aggression from dominant feral cats (Corbett 1979; Langham and Porter 1991). However, our inference for this demographic is limited by our telemetry-based dataset (juvenile cats are usually too small to trap and fit with tracking collars). A growing body of evidence also shows that the individual traits of feral cats, particularly body mass, can influence their hunting behaviour and prey size (Dickman and Newsome 2015; Kutt 2012; Moseby et al. 2020). Note, however that Fleming et al. (2020) found that while older cats might pose a greater risk to large and ‘dangerous’ prey, small cats were also capable of hunting big prey. Nonetheless, their greater hunting experience and ability to handle larger prey may enhance the survival rate of heavier cats.

Although some studies have found higher survival rates for female cats (Danner et al. 2010; Schmidt

et al. 2007), this was not supported in our analysis. The difference in results might be because some of these studies were in urban areas. Causes of mortality in urban areas often differ from natural environments, with some of the primary causes of mortality in urban areas being attacks by stray or owned dogs and road-kills (Devillard et al. 2003; Nutter et al. 2004). Furthermore, urban areas often support very high cat densities (e.g., 972 cats per km² in Kaeuffer et al. 2004); cats in areas with such high densities have a different, matrilineal mating system (Crowell-Davis 2007) which might influence sex-related survival. Finally, the low sample sizes within some of these study populations may have increased the likelihood of sex-biased estimates (e.g., n=28 in Danner et al. 2010).

To our knowledge, ours is the first study to compare feral cat survival rates across different habitat types. Somewhat surprisingly, survival rates were similar across all habitats. Feral cats in deserts tended to have higher average survival rates, but confidence intervals overlapped substantially. Feral cat home-range size is often negatively associated with landscape productivity (Bengsen et al. 2015). That is, they have much larger home ranges in less productive arid and semi-arid environments (e.g., Roshier and Carter 2021) compared to temperate areas (e.g., Buckmaster and Dickman 2012). While higher landscape productivity might facilitate smaller home-ranges, it does not appear to confer higher survival rates, perhaps due to increased disease risk in wetter conditions (Brazier et al. 2014). Consistently, we also found that relatively wetter conditions increased the relative risk to feral cats. We found no significant effect of increasing cat density on feral cat survival. This may suggest that in areas with higher feral cat densities, cats are still able to find enough prey such that their survival is not negatively affected by increased intraspecific competition for prey.

Consistent with our predictions, we found differences in net displacement for feral cats among sex and body mass classes. Male feral cat displacement in our study was nearly two times higher than female cats. These findings were consistent with studies showing larger home-ranges for male cats than females, and feral cats having a polygynous mating system where males often displace further (Bengsen et al. 2015; Liberg et al. 2000). The highest proportion of long-distance movements in our study were

observed among medium male cats, followed by large cats and small cats. Heavier cats, particularly large male cats are known to have a significantly higher impact on native mammal populations (Moseby et al. 2020), and are more likely to travel beyond their territory to access areas of high resources (McGregor et al. 2016).

Displacement can also vary between environments, with cats often moving further in less productive landscapes (e.g., Edwards et al. 2001; Roshier and Carter 2021). A relatively high proportion of feral cats in Savannahs, Deserts and Mediterranean forests moved long distances. Although a lower proportion of cats in temperate forests moved long distances, some individuals still moved more than 10 km from their initial point of capture (maximum of 37.5 km). Such long-distance movements by feral cats may be driven by food resources (e.g., Edwards et al. 2001). McGregor et al. (2014) reported adult feral cats travelling up to 12 km to hunt on fire scars from intense burns and subsequently returning to their home-ranges. Long-distance movements can affect feral cat management programs as new individuals can return and re-occupy vacant territories even though populations may be suppressed in an area (e.g., Algar et al. 2013; Moseby et al. 2009). Management programs should perhaps occur at different scales in different environments given the high variability in long-distance movement among feral cats across different habitat types.

Conclusion

Spatial, temporal and individual variation in survival rates and movement behaviour have important implications for invasive predator management, but are often poorly understood and not included in management decisions. Our average estimated feral cat survival rates for Australia were similar or higher than some of those previously recorded in free-roaming cat populations in USA and Europe (refer to Table S1). Feral cats currently inhabit most of mainland Australia and many offshore islands, and are responsible for declines and extinctions of numerous native prey (Abbott 2008; Burbidge and Manly 2002; Woinarski et al. 2015). The rates we have calculated can be used to improve population models and simulate potential responses to management. Further we

have shown that the population dynamics of invasive predators like feral cats are likely driven not just by their density, but also by other ecological factors such as the predator community and landscape productivity. Integrating knowledge on how these factors affect survival and movement behaviour of feral cats could help improve the design of single-species and integrated invasive species management programs in Australia.

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Declarations

Conflict of interest The authors have not disclosed any competing interests.

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