

The demography and dynamics of an expanding, managed African wild dog metapopulation

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Long-term demographic data are central for the evaluation of endangered species recovery plans. We present the demography and dynamics of a managed African wild dog (*Lycaon pictus*) metapopulation in South Africa, based on life histories of 553 individually known animals collected between 1998 and 2006. This metapopulation, distributed across nine sites (and not including Kruger National Park), was established through reintroductions ($n = 9$ events) and maintained by periodic augmentation ($n = 10$ events) and translocation ($n = 20$ events). In total, 66 founders were used to establish subpopulations and mean founder group size was 9.6 animals. The metapopulation grew from 17 individuals in 1998 to a peak of 202 in 2005. Mean annual population density was 3.3 (S.E. 0.44) wild dogs/100 km², approaching the upper limit of densities reported from unmanaged populations. Mean size of breeding packs was 11.0 (S.E. 0.76), comparable to pack sizes in Kruger National Park (Kruger), the only viable unmanaged population in South Africa. Fecundity was lower than in Kruger – particularly in the older age classes. Pup survival to adulthood was 45% – nearly three times the survival rate for pups in Kruger. Mean annual population growth rate ($\lambda = 1.08$, S.E. 0.13) was higher than in unmanaged populations (range 1.000–1.038), with implications for population viability and management.

Key words: African wild dogs, demography, dynamics, *Lycaon pictus*, managed metapopulation, reintroductions, species recovery.

INTRODUCTION

The African wild dog (*Lycaon pictus*) has declined over the last century and by 1990 the species was listed as Endangered by IUCN (IUCN, 2009). At that time, its status in South Africa was tenuous, with the Kruger National Park (Kruger) harbouring the only remaining viable population in the country. A Population and Habitat Viability Assessment (PHVA) for wild dogs recommended the expansion of wild dog range in South Africa as an urgent conservation priority (Mills *et al.*, 1998). However, no sufficiently large contiguous patches of suitable habitat existed within which to establish such a population. One solution was to reintroduce wild dogs into the existing network of small, isolated, fenced reserves and parks (Mills *et al.*, 1998).

The establishment of small, isolated populations raised some general concerns. Small populations experience different pressures to large populations: the dynamics of the former are governed by the

fates of a few individuals and those of the latter by the law of averages (Caughley, 1994). Population persistence is strongly influenced by three types of variability, which all have greater impacts on small populations than large ones. Firstly, small populations are more susceptible to endogenous threats arising from the very nature of being small, including random demographic effects, such as skewed sex ratios, and high or low birth or death rates (Ballou, 1995), which may have disproportionately large effects when populations contain ≥ 25 individuals (Lande, 1993). Secondly, small populations are vulnerable to genetic stochasticity through increased likelihood of inbreeding and genetic drift due to reduced effective population size (Reed, O'Grady, Ballou & Frankham, 2003). Finally, exogenous threats include environmental variation, which in extreme cases can lead to catastrophic die-offs (Reed *et al.*, 2003). Environmental variations influence the dynamics of both large and small populations, but small populations may be particularly susceptible and less resilient

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(Reed & Hobbs, 2004). This is especially true for events such as disease epidemics that may have serious consequences, as has been shown for Ethiopian wolves *Canis simensis* (Laurenson, Shiferaw & Sillero-Zubiri, 1997; Marino, Sillero-Zubiri & Macdonald, 2006) and wild dogs (Ginsberg, Mace & Albon, 1995; Hofmeyr, Bingham, Lane, Ide & Nel, 2000).

In an attempt to circumvent these hazards of small populations, a decision was made to manage the wild dogs in ways that mirror the dynamics of a metapopulation (Mills *et al.*, 1998; Akçakaya, Mills & Doncaster, 2007). Metapopulations comprise constellations of discrete local breeding subpopulations that display asynchronous dynamics and that are connected through dispersal (Levins, 1969). Metapopulations display more demographic and genetic resilience than single populations – particularly when there is gene flow between patches that are susceptible to different environmental variation (Ballou, 1995). Given that most wild dog reintroduction sites would be separated by several hundreds of kilometres, with no guarantee of natural dispersal between them (*cf.* Somers, Gusset & Dalerum, 2012), human intervention would be required to ensure gene flow and recolonization of extinct populations (Mills *et al.*, 1998). A series of population viability models (VORTEX; Lacy, 1993) formed the basis of management recommendations to maximize the persistence of the network of reintroduced subpopulations – later known as the *managed metapopulation* (Davies-Mostert, Mills & Macdonald, 2009). These models, for which details are provided by Mills *et al.* (1998), were informed by life history data from long-term studies of unmanaged wild dog populations in Kruger and Moremi Game Reserve, Botswana (Moremi), and assessed founder effects *inter alia*, the required levels of gene flow (translocations; *cf.* Gusset *et al.*, 2009) necessary to reduce inbreeding to sustainable levels, and the benefits of disease vaccination programmes. A recovery target was set for at least nine packs in 10 years (Mills *et al.*, 1998).

Nearly a decade after the managed metapopulation strategy was initiated a network of nine small wild dog populations had been established throughout South Africa (Davies-Mostert *et al.*, 2009; Gusset, 2010). These populations were managed collectively with animals periodically transferred between sites to mimic gene flow, and monitoring efforts generated sufficient data to enable an investigation of life-history traits. Given

that the managed metapopulation comprised a series of small, discrete populations, each more susceptible than either Kruger or Moremi to threats arising from limited population size, it was anticipated that population dynamics would differ substantially from those used to inform the conservation strategy's development and implementation. Differences in life-history traits such as survival and fecundity might influence population persistence, and the management tools required to maximize it. In this study, our principal objectives were to: determine overall demographic rates for wild dogs within the managed metapopulation; determine those vital rates that had dictated its dynamics between 1998 and 2006; assess the success of the programme in achieving its stated targets; and compare our findings to the demography of larger, unmanaged populations, with special reference to the implications for management. The baseline data originating from this study will facilitate the assessment of long-term persistence of the managed metapopulation, and contribute to effective conservation planning for wild dogs.

STUDY AREA

The study area encompassed nine reserves participating in the managed wild dog metapopulation programme (Fig. 1). Wild dogs had been previously extirpated from all subpopulation sites prior to initial reintroductions. The reserves ranged in size from 50 to 900 km² (mean \pm S.E. = 437 \pm 93 km²) and were surrounded by electrified perimeter fences. Although all reserves fell within the Savanna Biome (Low & Robelo, 1996), various vegetation types were represented across the subpopulations (Table 1). All reserves contained populations of preferred wild dog prey, including impala (*Aepyceros melampus*) and greater kudu (*Tragelaphus strepsiceros*), but also nyala (*Tragelaphus angasi*) in KwaZulu-Natal. Populations of key wild dog competitors, namely lions (*Panthera leo*) and spotted hyaenas (*Crocuta crocuta*) (Creel & Creel, 1996; Mills & Gorman, 1997), were present in all study sites at varying densities. Reserves derived income from various combinations of ecotourism, live game sales and/or hunting, and biodiversity conservation was a primary objective of only two sites.

METHODS

Monitoring and management

Population dynamics were tracked in the nine subpopulations between January 1998 and Decem-

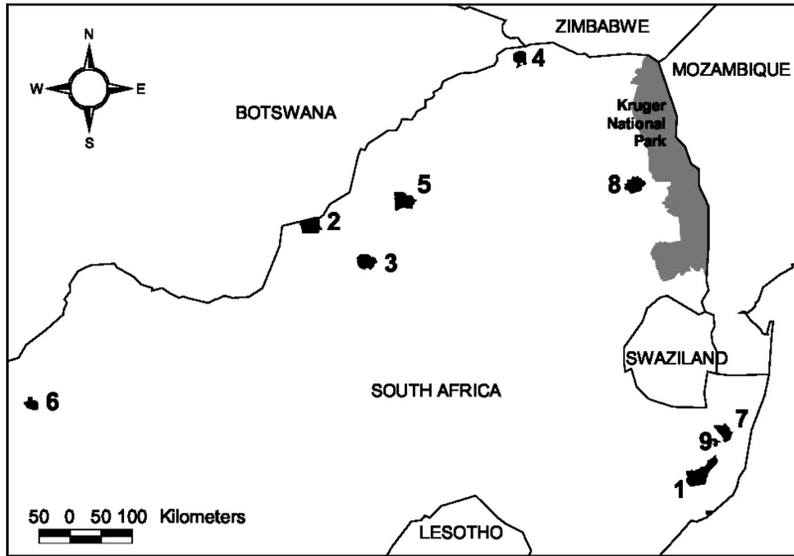


Fig. 1. The location of subpopulations within the managed African wild dog (*Lycaon pictus*) metapopulation in South Africa (1998–2007). Key: 1, Hluhluwe-iMfolozi Park; 2, Madikwe Game Reserve; 3, Pilanesberg National Park; 4, Venetia Limpopo Nature Reserve; 5, Marakele National Park; 6, Tswalu Kalahari Reserve; 7, Mkhuze Game Reserve; 8, Balule Game Reserve; 9, Thanda Game Reserve.

ber 2006. Reintroductions at two of the reserves had been made before the start of our observations. Wild dogs were first reintroduced into the 900 km² Hluhluwe-iMfolozi Park (Hluhluwe) in KwaZulu-Natal Province in 1980. By early 1998, this population comprised 12 individuals in three groups, survivors of the earlier reintroductions (Maddock, 1999; Somers, Graf, Szykman, Slotow & Gusset, 2008). Wild dogs were first reintroduced into the 600 km² Madikwe Game Reserve (Madikwe) in Northwest Province in 1995. Although the population grew to 24 animals by 1997, only three animals survived a rabies outbreak in early 1998 (Hofmeyr *et al.*, 2000). Between January 1998 and January 2007, seven additional subpopulations were established, encompassing a total area of c. 3930 km² (Table 1). In chronological order these included Pilanesberg National Park (Pilanesberg), De Beers Venetia Limpopo Nature Reserve (Venetia), Marakele National Park (Marakele), Tswalu Kalahari Reserve (Tswalu), Balule Game Reserve (Balule), Mkhuze Game Reserve (Mkhuze) and Thanda Private Reserve (Thanda).

Population size, structure and density

Population interventions included *reintroductions*, defined as the establishment of new subpopulations; *augmentations*, during which animals new to the metapopulation were added to existing

subpopulations; *translocations*, defined as the movement of animals between subpopulations; and *removals*, in which animals were permanently extracted from the metapopulation and sent to other conservation areas. Wild dogs were also recaptured as necessary following escapes onto neighbouring land, and these animals were either returned to their source population, translocated elsewhere in the metapopulation or removed entirely. Group size and composition (age and sex) were recorded for all interventions.

Post-release population dynamics were determined from photographic records, radio-telemetry studies and opportunistic sightings (Mills, 1992; Maddock & Mills, 1994; Somers *et al.*, 2008) (Table 1). Individual life histories were used to document changes in the structure and dynamics of each subpopulation, forming the basis of a database containing a timeline of all known individuals. Date of birth, age and sex were recorded for each individual, as were contemporaneous pack and subpopulation membership. We defined age classes as pups (<1 year), yearlings (1–2 years) and adults (>2 years) (Somers *et al.*, 2008) and recorded the status of each individual on the first day of each month as neonatal, present in group, dispersing, absent or dead. Records only for those wild dogs remaining within the metapopulation were made. Following Creel, Mills & McNutt (2004), we summa-

Table 1. Characteristics of the nine subpopulations within the managed African wild dog (*Lycyaon pictus*) metapopulation in South Africa.

Reserve name	Abbreviation	Province	Year wild dogs re-introduced	Vegetation type	Protected status	Reserve size (km ²)	Number of founders 1998–2006 ¹		Period of data used in this study	Data collection ²			Source of population data ³
							New	Meta		Photo	Radio	Opp	
Hluhluwe-Imfolozi Park	HIP	KwaZulu-Natal	1980	Natal Lowveld Bushveld	Provincial	900	8	7	1998–2004	✓	✓		Somers <i>et al.</i> (2008) (a)
Madikwe Game Reserve	MDGR	North West	1995	Mixed Bushveld / Kalahari Plains Thorn Bushveld	Provincial	620	11	9	1998–2006	✓	✓		(a)
Planesberg National Park	PNP	North West	1999	Mixed Bushveld	Provincial	500	11	2	1999–2000				van Dyk & Slotow (2003) (a)
De Beers Venetia Limpopo Nature Reserve	VLNR	Limpopo	2002	Mopane Bushveld	Private	320	16	4	2002–2006	✓	✓		(a)
Marakele National Park	MKNP	Limpopo	2003	Waterberg Moist Mountain Bushveld / Mixed Bushveld	National	740	16	1	2003–2006	✓	✓	✓	(a)
Tswalu Kalahari Reserve	TSKR	Northern Cape	2004	Shrubby Kalahari Dune Bushveld	Private	200	4	12	2004–2006	✓	✓	✓	(a)
Balule Game Reserve	BALU	Limpopo	2005	Mixed Lowveld Bushveld	Private	200	0	7	2005–2006	✓	✓	✓	(a)
Mkhuze Game Reserve	MZGR	KwaZulu-Natal	2005	Natal Lowveld Bushveld	Provincial	400	0	13	2005–2006	✓	✓	✓	(a)
Thanda Private Reserve	THDA	KwaZulu-Natal	2006	Natal Lowveld Bushveld	Private	50	0	4	2006		✓	✓	(a)
Total						3930	66	59					

¹New: animals new to the metapopulation; Meta: animals translocated from other subpopulations.

²Photographic records, radio-telemetry studies, opportunistic sightings.

³(a) Management reports and minutes from Wild Dog Advisory Group (WAG) meetings.

alized subpopulation and metapopulation characteristics on 1 January each year, the mid-point of the wild dog breeding cycle. This provided data on total population size, densities (wild dogs/100 km²), number of packs (defined as potential breeding units containing at least one unrelated adult pair), number of non-breeding groups, and mean pack and group sizes. We calculated mean annual population growth rate as $\ln(n_{t+1}/n_t)$ – the natural logarithm of the change in the population size (n) from one year (t) to the next ($t + 1$) – correcting population figures for newly reintroduced individuals to avoid inflated annual growth estimates.

We examined population size, growth and density in relation to one another, and to total land area available, which increased with time since metapopulation establishment. The effect of density on metapopulation composition was investigated by plotting densities of individuals and groups against total population size and annual population growth rate.

Breeding and reproduction

Pregnant females were identified by observation prior to giving birth and post-emergence litter size was determined by counting pups as soon as possible. Although this does not account for pre-emergence mortality, the constraints on counting pups were similar across subpopulations and enabled comparison of litter size and pup survival across sites and with other studies (Creel *et al.*, 2004; McNutt & Silk, 2008; Rasmussen, Gusset, Courchamp & Macdonald, 2008). It was not possible to determine variation in pre-emergence neonatal risks across sites. Pup survival (S_1) was calculated as the number of pups surviving to 12 months of age, as a proportion of L_E , the litter size at emergence.

We calculated age-specific fecundity (F_y) as $L_{EY} \times PF_E \times P_y$, where L_{EY} was the mean litter size at emergence produced by females of age y , PF_E was the proportion of females at emergence across all litters, and P_y was the proportion of females of age y that bred.

Sex ratios were determined for each age class (pups, yearlings and adults) on 1 June each year. Sex ratios at birth and 12 months were determined only from litters for which complete sex information was available. Calculations of adult sex ratios (≥ 24 months) included all individuals, as this sex information was complete.

We used Spearman's rank-order correlation coefficient (r_s) to determine the relationships between age of breeding female, parous status, litter size, pack size, number of pups surviving and proportion of pups surviving, unless otherwise indicated. We used two pack size variables to investigate the effect of pack size on pup production and survival: pack size at birth and mean pack size over the year (June–May, following Buettner, Davies-Mostert, du Toit & Mills, 2007). Parous status was positively correlated to age of breeding female ($r_s = 0.85$, $P < 0.001$, $n = 45$ litters) and, following McNutt & Silk (2008), female age was used in subsequent analyses, as it was the easier variable to determine.

Survivorship and mortality

We calculated age-specific survival for both sexes and overall by determining the proportion of individuals of a given age remaining in each subpopulation from one breeding season to the next, accounting for newly reintroduced animals. Individuals that disappeared from packs and were never detected again were presumed to be dead, or lost to the metapopulation through dispersal.

Apparent adult survival rate (restricted to females) was calculated by determining the proportion of females in each age group that survived from 1 June to the following breeding season. Pup and yearling survival rates were determined for all individuals (*i.e.* were not restricted to females) in order to reduce the biases described above.

We compared causes of mortality with those in larger, unmanaged wild dog populations, using only mortality data from radio-collared wild dogs to avoid detection biases resulting from differing levels of monitoring intensity (Woodroffe *et al.*, 2007). Pups were never radio-collared, so this approach precluded any comparison of pup mortality. We also examined known causes of mortality for un-collared individuals (adults, yearlings and pups), recognizing that these data cannot be directly compared with other areas without risk of confounding biases.

We recorded the incidence of catastrophic mortality events. These were defined as discrete events that caused mortality of multiple (>3) adults, pups or both, and were important for assessing population viability (Lande, 1993). Although simultaneous death of multiple pups can be common in large populations, we considered this more significant for populations comprising just one or two packs.

Dispersal

Wild dogs disperse in same-sex groups of either sex (McNutt, 1996). We identified dispersal events as primary (individuals leaving their natal pack) or secondary (individuals dispersing from non-natal groups), both of which were differentiated from pack fission, which occurred when mixed-sex groups fissured from existing packs to form potential breeding packs containing unrelated individuals of different sexes. Occasionally packs also split into mixed-sex but non-breeding (*i.e.* related) groups. The timing and group composition of all dispersal events were recorded, as well as whether dispersing groups were tolerated by management within their natal subpopulation, or were translocated elsewhere. Characteristics used elsewhere to describe dispersal – such as dispersal distance (McNutt, 1996), the duration of transience (Maehr, Land, Shindle, Bass & Hctor, 2002) or whether new pack formation was achieved successfully (Somers *et al.*, 2008) – were considered irrelevant, as dispersing groups were often used in management interventions, which prevented natural pack formation. Dispersers are often the best choice of dogs for interventions because this mimics the natural state as closely as possible.

Comparison with larger, unmanaged populations

We compared metapopulation vital rates to those reported for larger, unmanaged populations in order to determine potential effects of intensive management on wild dog demography in small, fenced reserves. Following Creel *et al.* (2004), we used age-specific survivorship and fecundity to build an age-based Leslie matrix population model for the metapopulation overall, setting age at first parturition at 2 years. The deterministic population growth rate (λ) was calculated from the dominant eigenvalue of the Leslie matrix. We validated our parameters by modelling the deterministic population trajectory correcting for newly reintroduced animals and comparing this to census data (Reid, Bignal, Bignal, McCracken & Monaghan, 2004). We used elasticities to determine which vital rates had the greatest impact on population growth (Caswell, 2001), and compared these to results from the larger, unmanaged wild dog populations examined by Creel *et al.* (2004), which included the Selous Game Reserve in Tanzania (1991–1997), Kruger National Park in South Africa (1989–2002) and Moremi Game Reserve and adjacent Wildlife Management Areas in Botswana (1989–2002).

Statistical methods and data sources

Where sample sizes allowed, spatial and temporal differences in demographic variables were examined by comparing results across subpopulations and years. For the most part, these analyses were only possible for the five reserves that joined the metapopulation prior to 2004. Spearman's rank-order correlation coefficients (r_s) and partial rank-order correlation coefficients (partial r_s) were used to assess the degree of correlation among variables; other parametric and non-parametric tests (identified in text) were used to compare differences among variables. Such tests do not require that data are distributed normally; however, they do tend to be less sensitive (*i.e.* have reduced power) at detecting an effect of the independent variable on the dependent variable (Robson, 1994). The relationship between pack size and litter size was determined by regressing pack size at emergence against square-root of litter size for packs where only one female bred and litter size was known ($n = 50$). Leslie matrix calculations and population projections were conducted using the shareware POPTOOLS (Hood, 2006). For all statistical analyses SPSS 14.0 was used.

Metapopulation data were collated from subpopulation managers and researchers (Table 1) and represented all data for the period 1998 to 2007. Comparative data from larger, unmanaged populations were obtained from published literature (Fuller *et al.*, 1992; Creel *et al.*, 2004) and in many cases were derived from study areas comprising just a small proportion of the population. Such data are susceptible to selection biases, as field studies tend to be conducted in areas where study populations are at higher densities (Smallwood & Schonewald, 1996). As a result, density estimates in these populations are likely to be overestimates for the unmanaged populations they represent. The Kruger data provided an exception as population structure was calculated from photographic submissions from throughout the park, thus providing a minimum count (Maddock & Mills, 1994).

RESULTS

Management history

Sixty-six animals were used as founders in nine separate reintroductions and augmentations between January 1998 and January 2007. These included animals sourced from Kruger (2%),

captive stock (15%), wild-caught packs outside protected areas (58%) and pups born in holding enclosures while awaiting translocation and/or release (24%). Mean founding groups for initial reintroductions were larger than for augmentations, and comprised mostly adults and yearlings, with smaller numbers of pups.

Sixty-nine individuals were used in translocations between reserves, which took place on 20 occasions. These interventions were undertaken to establish new subpopulations ($n = 12$), redress demographic and genetic imbalances at the subpopulation level ($n = 4$), promote new pack formation at release sites ($n = 3$) or remove dispersal groups from areas offering no further breeding opportunities ($n = 1$). Groups were removed from the metapopulation on two occasions: the first when an entire pack was translocated from Pilanesberg to Zimbabwe in an effort to reduce predator pressure in the reserve; the second when a single-sex male group was translocated from Marakele to Kruger in an attempt to prevent inbreeding at Marakele.

Population size, structure and density

The managed metapopulation was instigated with just 17 individuals in two populations in January 1998, with the addition of 66 individuals subsequent to 1998. It reached a peak of 202 individuals across all age classes in 16 packs in January 2005 (Fig. 2). At this juncture, densities reached 5.4 wild dogs/100 km² – nine times higher than concurrent densities in Kruger (0.6 wild dogs/100 km²; Kemp & Mills, 2005). This was mainly a result of the successful breeding season in 2004, when 91 pups were born, of which 82 (90.1%) survived to 1 January 2005. The metapopulation contained

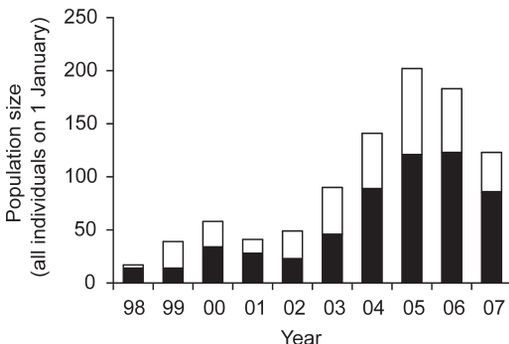


Fig. 2. Number of adult and yearling (black bars) and juvenile (white bars) African wild dogs (*Lycaon pictus*) in the managed metapopulation in South Africa on 1 January of each year (1998–2007).

143 adults and yearlings at the onset of the 2004 denning season.

Mean annual population density over the complete study interval was 3.3 (S.E. 0.42) wild dogs/100 km², with an increasing trend over time ($r_s = 0.77$, $P = 0.009$, 8 d.f.) and with the total area under metapopulation management ($r_s = 0.81$, $P = 0.005$, 8 d.f.). Mean annual densities for individual reserves were only weakly negatively related to reserve size ($r_s = -0.58$, $P = 0.08$, 8 d.f.), possibly because small population changes can lead to large density fluctuations in small reserves. Annual densities appeared to be highest at Venetia (6.5 wild dogs/100 km²) but this was not significantly higher than the other three reserves (Hluhluwe, Madikwe and Pilanesberg) where sample sizes were large enough to conduct this comparison (Kruskal-Wallis test; $H_{(4,33)} = 7.66$, $P = 0.054$, 3 d.f.).

The mean number of groups (breeding and non-breeding) per year was 10.8 (range 5–20, $n = 9$), and mean group size was 8.7 (S.E. 0.61). When only breeding packs were considered, this rose to 11.0 (S.E. 0.76), which was comparable to mean pack sizes reported for Kruger and northern Botswana (both 10.4), but larger than pack sizes in Selous (8.9; Creel *et al.*, 2004). Mean group size did not increase over time ($r_s = 0.39$, $P = 0.260$, $n = 9$), although, as expected, the total number of groups did ($r_s = 0.94$, $P < 0.001$, $n = 9$).

Corrected mean annual population growth rate calculated from life-table analysis was 0.08 (S.E. 0.13, range -0.63–0.53), indicating an average annual population increase of 8%, but with a large 95% confidence interval (CI 0.25). When newly reintroduced animals were included in the calculation of population growth, the figure was 0.22 (S.E. 0.14), indicating that periodic population augmentations may have an important positive effect on population trends (*cf.* Gusset *et al.*, 2009). After controlling for the size of area available, corrected annual population growth rate was positively correlated with both population size (partial $r_s = 0.73$, $P < 0.05$, $n = 9$) and density (partial $r_s = 0.79$, $P < 0.05$, $n = 9$), suggesting an absence of any inverse density-dependent effects on population dynamics, at least over the 9 years covered by this study.

Breeding and reproduction

The metapopulation contained on average 7.0 (S.E. 1.04, 48%) breeding females in each breeding season (range 3–11, $n = 9$ years) and, as

Table 2. Litter size, pup survival and yearling survival within the managed African wild dog (*Lycaon pictus*) metapopulation in South Africa (1998–2006).

Subpopulation	Litter size ¹		Pup survival ^{2,3}		Yearling survival ^{2,3}	
	<i>n</i>	Mean litter size ± S.E.	<i>n</i>	Mean pup survival	<i>n</i>	Mean yearling survival
Hluhluwe-iMfolozi Park	18	8.3 ± 0.79	18	0.66 (0.58–0.74)	17	0.67 (0.56–0.76)
Madikwe Game Reserve	10	9.8 ± 1.04	10	0.60 (0.50–0.70)	10	0.69 (0.56–0.80)
Pilanesberg National Park	6	8.3 ± 0.56	5	0.74 (0.61–0.88)	6	0.94 (0.75–0.98)
Venetia Limpopo Nature Reserve	6	8.3 ± 0.71	6	0.72 (0.58–0.84)	6	0.78 (0.61–0.90)
Marakele National Park	5	9.8 ± 1.59	5	0.76 (0.61–0.87)	4	0.60 (0.44–0.75)
Tswalu Kalahari Reserve	1	4.0	1	0.00	0	–
Balule Game Reserve	2	8.0	2	0.25	0	–
Mkhuze Game Reserve	2	7.0	2	0.36	0	–
Thanda Private Reserve ⁴	–	–	–	–	–	–
Managed metapopulation	50	8.6 ± 0.41	49	0.64 (0.60–0.69)	43	0.71 (0.65–0.76)

¹Litter sizes are provided only for packs where one female bred.

²Pup and yearling survival rates provided for litters with known fates.

³Values in brackets are exact binomial confidence intervals.

⁴No data are available for Thanda as two females bred during the only breeding season within the study period.

expected, the number of breeders increased over time as the number of subpopulations and packs within the metapopulation grew. Mean litter size was 8.6 (S.E. 0.41, $n = 50$) and within the range of mean litter sizes reported elsewhere (Fuller *et al.*, 1992; Creel *et al.*, 2004; McNutt & Silk, 2008; Rasmussen *et al.*, 2008). Litter sizes did not differ significantly among the five subpopulation reserves (Kruskal-Wallis test: $H_{(4,45)} = 1.59$, $P = 0.810$), or across the 6 years (2001–2006: $H_{(5,43)} = 6.4$, $P = 0.269$) with sample sizes adequate to make this comparison (Table 2).

As has been found in other study sites, pack size at the time of parturition was positively related to the number of pups emerging (linear regression, $F_{(1,56)} = 10.54$, $P = 0.002$, $R^2 = 0.143$), with larger packs producing larger litters (Creel & Creel, 2002; Buettner *et al.*, 2007; McNutt & Silk, 2008; Rasmussen *et al.*, 2008). In contrast to results from other populations, there was no relationship between pack size and the proportion of pups surviving to 3, 6 or 9 months (Buettner *et al.*, 2007) or to 1 year (Buettner *et al.*, 2007; McNutt & Silk, 2008) (Table 3). However, there was a significant

Table 3. Spearman's rank-order correlation coefficients (r_s) for the relationship between pack size at birth and pup survival of African wild dogs (*Lycaon pictus*) in the managed metapopulation in South Africa, represented as (i) the proportion and (ii) the number of pups surviving to 3, 6, 9 and 12 months.

(i) Proportion ¹ of pups surviving to	r_s	<i>P</i>	<i>n</i>
3 months	0.263	0.068	49
6 months	0.169	0.246	49
9 months	0.172	0.238	49
12 months	0.155	0.293	49
(ii) Number ² of pups surviving to	r_s	<i>P</i>	<i>n</i>
3 months	0.309	0.017	59
6 months	0.354	0.005	62
9 months	0.368	0.003	63
12 months	0.343	0.006	62

¹Includes only those litters that can be ascribed to one female, and have known fates.

²Includes all litters with known fates.

positive relationship between pack size and the actual number of pups surviving to these ages (Table 3), with the result that larger packs produced a larger number of yearling recruits than did smaller packs. This mirrors the direction of the non-significant relationship found by Gusset & Macdonald (2010).

Fecundity increased with age, with older females producing significantly more offspring than did younger females (Kruskal-Wallis test, $H_{(5,50)} = 20.76, P = 0.001$), and a larger proportion of older female age classes breeding (chi-square test, $\chi^2 = 68.4, P < 0.001, 5 \text{ d.f.}$). No data were available for females >9 years old.

Sex ratio of pups at emergence was biased towards males (80 of 136 pups = 0.59; exact binomial test; $P = 0.048$, for litters where sex information was complete). The proportion of males was not related to the age of the breeding female ($r_s = 0.08, P = 0.75, n = 19$) or litter size ($r_s = 0.10, P = 0.68, n = 19$) as suggested in earlier studies (Fuller *et al.*, 1992). The male-skewed sex ratios were equally apparent for yearlings (0.58; exact binomial test; $P = 0.01, n = 220$) and adults (0.56; exact binomial test; $P < 0.01, n = 569$), concordant with results of other studies (Fuller *et al.*, 1992).

Survivorship and mortality

Age-specific annual survival rates for females are provided in Fig. 3. In total, 45% of pups born into the metapopulation survived to adulthood (2 years). Adult survival was highest for 2-year-olds (0.91, exact binomial CI 0.82–0.96), with no significant trend towards decreasing survival rates with senescence ($r_s = -0.08, P = 0.844, n = 8$). Variance in survival rates was largest for the oldest age classes, as estimates were derived

from increasingly fewer individuals. Male survival tended to be higher than female survival for all age classes, although this was not significant (Mann-Whitney, U -test; $U = 82.5, P = 0.139, n = 8$).

The age-based Leslie matrix for females within the managed metapopulation was constructed as follows:

	0.00	0.00	0.09	0.58	1.43	2.08	3.00	2.73
	0.64	0.00	0.00	0.00	0.00	0.00	0.00	0.00
	0.00	0.71	0.00	0.00	0.00	0.00	0.00	0.00
A =	0.00	0.00	0.91	0.00	0.00	0.00	0.00	0.00
	0.00	0.00	0.00	0.64	0.00	0.00	0.00	0.00
	0.00	0.00	0.00	0.00	0.61	0.00	0.00	0.00
	0.00	0.00	0.00	0.00	0.00	0.67	0.00	0.00
	0.00	0.00	0.00	0.00	0.00	0.00	0.63	0.57

where the first row represents age-specific fecundity of females aged 0–8 years, and the diagonal represents annual survival rates for each age class. Annual population growth rate calculated from the dominant eigenvalue of the Leslie matrix ($\lambda = 1.10$) was higher than, but fell within the 95% confidence interval around the annual population growth rate calculated from corrected population figures using established methods ($\lambda = 1.08$). Deterministic population projections from the Leslie matrix corresponded to observed population sizes. Inter-annual variation in metapopulation growth rate was high, and 95% confidence intervals enclosed the value of $r = 0$, suggesting that metapopulation growth was unstable, with

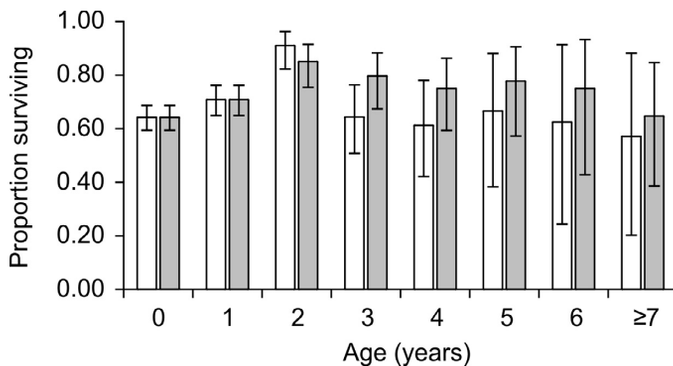


Fig. 3. Age-specific annual survival rates for female (white bars) and male (grey bars) African wild dogs (*Lycaon pictus*) in the managed metapopulation in South Africa. Data are pooled for animals ≥7 years to maintain sample sizes. Error bars show exact binomial confidence intervals.

Table 5. Summary dispersal statistics for African wild dogs (*Lycaon pictus*) in the managed metapopulation in South Africa (1998–2006).

Variable		Sex			All
		Males	Females	Mixed	
Number of dispersal events (groups)	Primary	20	16	1	37
	Secondary	4	3		7
Number of individuals dispersing	Primary	59	56		115
	Secondary	5	4		9
Mean size of dispersing groups (\pm S.E.)	Primary	2.9 \pm 0.37	3.4 \pm 0.48	4.0	3.1 \pm 0.29
	Secondary	1.3 \pm 0.25	1.3 \pm 0.33		1.3 \pm 0.18
Median age (months) at dispersal (range)	Primary	26.5 (8.5–62.5)	21.0 (13.5–62.2)		25.0 (8.5–62.5)
	Secondary	43.5 (33.0–53.5)	36.5 (36.5–56.5)	38.5 (33.0–56.5)	
Dispersals out of source subpopulations	Events	7	6	1	14
	Individuals	25	30		55
Number of individuals permanently leaving the metapopulation	Primary	19	20		39
	Secondary	1	1		2
Mean annual % of adults and yearlings emigrating		5.3%	7.8%		6.3%

was recorded, this following an outbreak of canine distemper at Tswalu in 2005, during which four adults and five juveniles died. Population augmentations staved off extinctions elsewhere.

Dispersal

Thirty-seven incidents of primary dispersal were recorded between 1998 and 2006, involving 115 animals (Table 5). Male and female dispersing groups were of similar size (2.9 males (S.E. 0.37) vs 3.4 females (S.E. 0.48), Mann-Whitney $U = 182.5$, $z = -0.7$, $P = 0.484$, $n_1 = 20$, $n_2 = 16$), and females were younger than males when they first left natal packs (median, 21.0 vs 26.5 months, $U = 1097.5$, $z = 3.1$, $P = 0.002$, $n_1 = 59$, $n_2 = 56$). One case of primary dispersal involved a mixed-sex group of four animals (two males, two females), although this group split by sex two weeks later. The annual incidence of primary dispersal was positively correlated with the number of breeding packs ($r_s = 0.78$, $P < 0.05$, $n = 8$) and wild dog density ($r_s = 0.95$, $P < 0.05$, $n = 8$) on 1 January each year (Table 5). However, per capita annual incidence of primary dispersal was not correlated to either wild dog density ($r_s = 0.43$, $P > 0.5$, $n = 8$) or the number of breeding packs ($r_s = 0.19$, $P > 0.05$, $n = 8$), suggesting an absence of any density effects on primary dispersal rates. Secondary dispersal was recorded in Hluhluwe ($n = 4$), Pilanesberg ($n = 2$) and Marakele ($n = 1$) (Table 5).

Dispersing groups were observed with unrelated

animals of the opposite sex in 29 (76%) of 38 primary and secondary dispersal events for which the outcome was known. Animals were also captured and introduced to artificially establish new packs either at the source site ($n = 1$) or at other metapopulation sites ($n = 5$). At least 41 individuals were confirmed to have naturally emigrated from the metapopulation, while another 27 individuals were removed to sites outside the metapopulation.

DISCUSSION

Understanding the influence of life-history traits on population regulation is especially important for managed populations, in which there is scope to target specific interventions to improve or stabilize population growth. The process of establishing a managed wild dog metapopulation in South Africa enabled the collection of demographic and life-history data over 9 years while the population was expanding. Several long-term studies of wild dog population dynamics in large unmanaged systems such as the Serengeti (Fuller *et al.*, 1992), Kruger (Maddock & Mills, 1994; Buettner *et al.*, 2007), Selous (Creel & Creel, 2002) and northern Botswana (Creel *et al.*, 2004; McNutt & Silk, 2008), provided a benchmark for the operation of natural wild dog population dynamics, and a means to assess whether various life-history traits in the managed metapopulation fell within the parameters of 'natural' populations. We observed broadly

similar traits to those in larger populations, suggesting that natural population dynamics can be achieved even in systems that are intensively managed. However, this study metapopulation displayed population densities, pup and yearling survival rates, as well as annual population growth rates higher than those recorded in established and unmanaged populations. Elevated rates could be attributed to the fact that subpopulations were being established in unoccupied areas with low levels of intra-specific competition, as has been found for other recolonizing carnivore populations (Pletscher, Ream, Boyd, Fairchild & Kunkel, 1997; Bales, Hellgren, Leslie & Hemphill, 2005).

Elevated population densities

Wild dog densities are low when compared to those of most other large predators (Creel & Creel, 1996) and are negatively correlated with lion densities in a range of ecosystems (Mills & Biggs, 1993; Creel & Creel, 1996, 1998, 2002). Wild dogs are able to coexist with these larger predators by avoiding them both spatially (Mills & Gorman, 1997; Creel & Creel, 2002) and temporally (Saleni *et al.*, 2007). Lion-induced mortality has hampered at least one previous wild dog reintroduction (Scheepers & Venzke, 1995) and has been flagged as a factor likely to limit reintroduction success (Mills *et al.*, 1998). Although we recorded lion-induced mortality in our study (Table 4), this was not, as would be expected, at a rate high enough to affect successful establishment of populations (Gusset *et al.*, 2008; Gusset, Stewart, Bowler & Pullin, 2010).

Wild dog densities were consistently higher than in Kruger and tended to be higher than in most of the other large unmanaged conservation areas, notwithstanding that high metapopulation densities were partially offset by the periodic addition of habitat in the form of new subpopulation sites. Mean pack sizes were comparable to those in unmanaged populations (Fuller *et al.*, 1992; Creel *et al.*, 2004) and it follows that the higher individual densities (*i.e.* wild dogs/100 km²) were a result of higher pack densities (*i.e.* packs/100 km²). Annual wild dog ranges have been reported to overlap only by 30–35% in Kruger (Reich, 1981) and between 10–50% in the Serengeti (Frame & Frame, 1976), thus higher pack densities were probably a consequence of an increase in range overlap within metapopulation reserves, a reduction in pack home range sizes, or both.

Carnivore ranging behaviour is influenced by

various factors including the ability to meet their ecological requirements (food acquisition, avoidance of predation) and also their social requirements (territoriality, mating). Elevated wild dog densities in the metapopulation suggest that packs were able to meet these requirements in smaller areas – at least during the period covered by this study – and that their home ranges were somehow constrained, either by perimeter fences or because release sites were resource-rich. There is evidence that wild dog hunting success in small fenced areas is improved by the propensity of wild dogs to capture prey against perimeter fences (Van Dyk & Slotow, 2003; Rhodes & Rhodes, 2004; Davies-Mostert, Mills & Macdonald, 2013). Prey availability might also have temporarily increased following herbivore reintroductions at some subpopulation sites. This would have increased densities and also provided naïve prey individuals that were more susceptible to predation (H.T. Davies-Mostert, unpubl. data 2010).

An alternative explanation for elevated wild dog densities is the statistical artefact of reintroducing wild dogs into areas smaller than their natural home range sizes. Only three of the nine subpopulation sites were larger than the average range of wild dog packs in Kruger (537 km²; Mills & Gorman, 1997) and four were actually smaller than the minimum range size reported (357 km²; Mills & Gorman, 1997). A breeding pair of wild dogs – the minimum required to meet the definition of a pack – in a 50 km² reserve (the size of Thanda, the smallest subpopulation site) equates to a density of 4.0 wild dogs/100 km². This is twice the density of wild dogs in Kruger when the population reached its peak in 1995, and does not account for any population growth following successful breeding. We suggest that elevated wild dog densities in the metapopulation can be at least partially explained by the fact that populations are constrained by electrified fences, which impede (but do not completely prevent; see Davies-Mostert *et al.*, 2012, 2013) dispersing individuals from moving out of their natal ranges. Elevated wild dog densities have created challenges in some subpopulation reserves due to the perception that they increase predation pressure on herbivore populations (Davies-Mostert *et al.*, 2009). Although a deeper understanding of the sustainability of predation will be useful for the future management of large carnivores in small areas (*e.g.* cheetahs (*Acinonyx jubatus*) and lions in South Africa), most reserves are unwilling to allow predation to con-

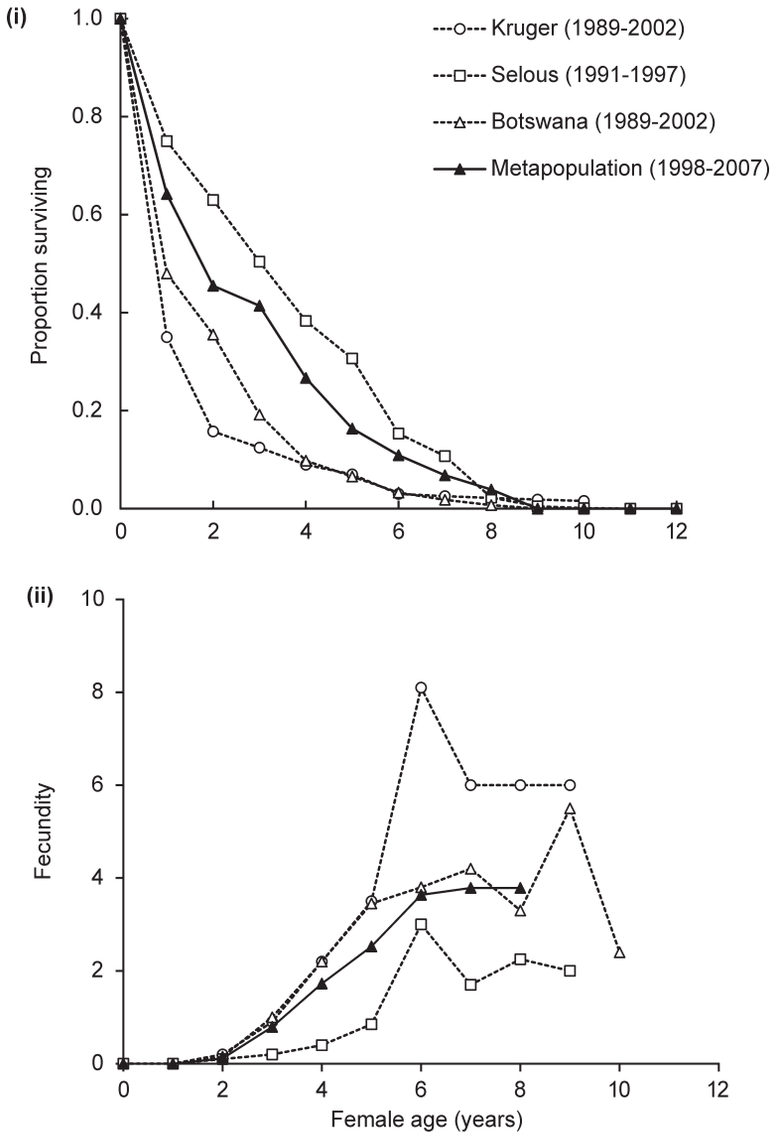


Fig. 5. Comparison of (i) female survivorship and (ii) age-specific fecundity in the managed metapopulation of African wild dogs (*Lycaon pictus*) in South Africa with those of unmanaged populations (Kruger National, Selous Game Reserve and northern Botswana; data from Creel *et al.*, 2004). Fecundity is measured as litter size divided by two, and age-specific fecundity incorporates the proportion of females breeding in each age class. Fecundity data for the metapopulation and Kruger were pooled across females ≥ 7 years to maintain sample sizes.

tinue unchecked and future field trials to investigate density dependence seem highly unlikely.

Elevated pup and yearling survival

It has been established that there are trade-offs between life-history traits within wild dog populations, with litter sizes being offset by pup survival, reproduction by adult survival, and juvenile survival rates offset by those of adults (Fuller *et al.*, 1992; Creel *et al.*, 2004; Gusset & Macdonald, 2010).

Survival rates for pups and yearlings in the metapopulation were higher than for both Kruger and northern Botswana (Fig. 5). This is important, firstly because survival of pups and yearlings has the largest effect on population growth (Creel *et al.*, 2004; this study), and secondly because litter sizes in the metapopulation are not smaller than in unmanaged populations. Consequently, population growth in the managed metapopulation was considerably higher than for the unmanaged pop-

ulations investigated by Creel *et al.* (2004), for which values of λ ranged from 1.000 (northern Botswana) to 1.038 (Selous). However, the annual metapopulation growth rate was highly variable, and 95% confidence intervals enclosed the value of $r = 0$, suggesting that metapopulation growth was unstable, with the potential for large fluctuations between positive and negative growth. This is a characteristic of wild dog populations generally.

Other factors influencing survival and population viability

Causes of adult wild dog mortality were similar to those in unmanaged populations, but pup mortality attributed to human causes was much higher. While recognizing that mortality data for pups are inherently biased (Woodroffe *et al.*, 2007), most pup mortality was attributed to management interventions and probably reflects true differences between managed and unmanaged populations. Anthropogenic mortality of wild dogs tends to be additive to natural causes (Woodroffe *et al.*, 2007) and hence efforts to reduce management-induced pup mortality would be likely to result in an improvement in overall pup survival. This, in turn, would create an even larger discrepancy in survival rates between the metapopulation and unmanaged populations.

Catastrophic events such as those caused by disease or human persecution can have significant impacts on tiny wild dog populations (Ginsberg *et al.*, 1995; Ginsberg & Woodroffe, 1997; Hofmeyr *et al.*, 2000; Hofmeyr, Hofmeyr, Nel & Bingham, 2004). Although catastrophic disease outbreaks (canine distemper) led to the extinction of one subpopulation during the period covered by this study (Tswalu 2005), this represented just 3.6% and 4.9% of adults and pups in the metapopulation, respectively. After 2004, no subpopulation comprised more than 38% of the total metapopulation; therefore, even though small populations are more susceptible to catastrophic events, catastrophic mortality at the subpopulation level was unlikely to have major impacts at the metapopulation level (*cf.* Gusset *et al.*, 2008, 2010). This is one of the benefits of spreading risk across several isolated populations (Shafer, 2001).

Outlook for conservation management of large carnivores

The dynamics exhibited by this managed wild dog metapopulation were indicative of an expand-

ing population as packs were established in areas free from conspecifics. Although metapopulation dynamics were in part an artefact of direct management intervention, the high population growth rates we observed demonstrated that wild dogs were able to successfully re-colonize new areas given suitable habitat and prey base, and adequate protection from human conflict (Gusset, 2009). Similar successes have been described for other large-carnivore recovery programmes (lions, Hunter *et al.*, 2007; wolves (*Canis lupus*), White, Lemke, Tyers & Fuller, 2008; cheetahs, Marnewick, Hayward, Cilliers & Somers, 2009) and although this bodes well for reintroductions, high population growth rates can also contribute to the complexities of population management (Davies-Mostert *et al.*, 2009). This is particularly relevant in small, fenced reserves where there is little opportunity for population expansion. Continued fragmentation of landscapes is likely to necessitate the managed metapopulation approach to conserving wide-ranging animals and our study emphasizes the importance of protected areas (even relatively small ones) for the conservation of carnivores as an alternative to the very difficult question of coexistence with people.

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