

Conflicting management policies for the Arabian wolf *Canis lupus arabs* in the Negev Desert: is this justified?

ORLY COHEN, ADI BAROCAS and ELI GEFFEN

Abstract Conservation plans may conflict when both predator and prey in the same ecological system are threatened species. In this study we present a problematic case of conflicting conservation policies involving the Arabian wolf *Canis lupus arabs* and two species of gazelles (*Gazella gazella acaciae* and *Gazella dorcas*), all of which are threatened in Israel. By studying genetic subdivision using faecal DNA we evaluated the policy of treating the Arabian wolves in the Negev Desert as two separate populations. We analysed 95 wolf faecal samples from 12 feeding sites c. 20 km apart. Network analysis and Bayesian clustering were used for separating populations. Mark–recapture design, rarefaction and an urn model were applied to estimate wolf population size. We found that wolves in the central and southern Negev cannot be genetically separated, and their density is similar in both regions. Our results provide a better baseline for a unified management of wolves in the Negev. We call for the consideration of other factors influencing gazelle population size before adopting drastic measures such as wolf removal.

Keywords Carnivore removal, Israel, Middle East, non-invasive sampling, population size, population subdivision, scatology

This paper contains supplementary material that can be found online at <http://journals.cambridge.org/orx>

Introduction

A great deal of effort goes into protecting threatened and/or important species. In some cases several threatened species can be protected under the same programme. Nevertheless, when these threatened species include both predator and prey in the same ecological system there may be potential conflict of management plans. This is illustrated by examples from the Channel Islands off the Californian coast. These islands were previously densely populated by the island fox *Urocyon*

littoralis. The arrival and spread of the golden eagle *Aquila chrysaetos* to these islands in the 1980s, facilitated by the availability of the previously introduced feral pig *Sus scrofa* as prey, led to a drastic decline in fox numbers (Roemer et al., 2002), to the point that breeding programmes were established to save the fox from extinction. The dilemma was: should the golden eagle, which is a protected species, be removed to save another protected species, the island fox? To mitigate the decline in fox numbers, golden eagles were translocated, and feral pigs eradicated from the islands (Courchamp et al., 2003). A similar case in these Channel Islands is that of the threatened San Clemente loggerhead shrike *Lanius ludovicianus mearnsi*, which is predated by the island fox; a control programme was established to save the shrike (Roemer & Wayne, 2003).

Predator removal is commonly practised by wildlife managers. A recent review suggested that predator removal increased hatching and fledging success and breeding populations of birds (Smith et al., 2010). Predation by wolf *Canis lupus*, puma *Puma concolor* and coyote *Canis latrans* may be a significant mortality factor for medium or large ungulates (e.g. deer, caribou *Rangifer tarandus*, moose *Alces alces*) in some areas under certain conditions (reviewed in Ballard et al., 2001). A predator removal programme may not affect prey populations that are near carrying capacity but may significantly increase ungulate populations that are substantially below carrying capacity. Thus results of predator removal practice are confounded by numerous factors and only through intensive studies can predation be identified as a major limiting factor (Ballard et al., 2001). Wasser et al. (2011) studied the effect of predator removal on caribou populations and concluded that management should prioritize the control of human activities before implementation of more extreme actions such as removal of wolves. In other words, resource selection and its physiological consequences can significantly influence predator–prey interactions, and should be considered in management programmes that aim to manipulate complex ecosystems.

In this study we present a problematic case of conflicting conservation policies involving several threatened species. A small isolated population (20–30 individuals) of the mountain gazelle *Gazella gazella acaciae* inhabits a 6 km² acacia forest near Yotvata in the southern Arava Valley, Israel (Yom-Tov & Ilani, 1987; Dolev & Perevolotsky, 2004). A recent genetic survey of gazelles in the Middle East showed that this isolated population is genetically different

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from the mountain gazelle *Gazella gazella gazella* in northern Israel but similar to *G. gazella* from northern Saudi Arabia (Wronski et al., 2010). This Critically Endangered *G. g. acaciae* population has been in decline since the 1960s, presumably because of low calf survival, low genetic diversity, anthropogenic effects and climate change (Yom-Tov & Ilani, 1987; Dolev & Perevolotsky, 2004; IUCN, 2008a). The congeneric dorcas gazelle *Gazella dorcas* is a more common species in the Negev but at a regional scale it is categorized as Vulnerable (Dolev & Perevolotsky, 2004; IUCN, 2008b) because of intensive motorized hunting and habitat degradation caused by overgrazing and droughts (Mallon & Kingswood, 2001). The law in Israel protects both gazelle species and poaching in the south of Israel is rare. The main alleged cause for the decline of both the mountain and dorcas gazelles in the Negev is predation by the Arabian wolf *Canis lupus arabs* and the caracal *Caracal caracal*, a hypothesis based on anecdotal data (Shalmon, 2003, 2006). The mountain and dorcas gazelles differ in their laying-up period, which is longer in the mountain gazelle, suggesting greater vulnerability of mountain gazelle fawns to predation (Mendelssohn et al., 1995; Yom-Tov et al., 1995). In response to the decline in gazelle populations in the southern Negev the Israel Nature and Parks Authority (INPA; the government agency that manages all natural resources in Israel) instigated a wolf removal programme that eliminated 37 individuals from the area during 2005–2008. By mid 2006 the isolated and small population of the mountain gazelle was completely fenced and the wolf removal programme was terminated in early 2008.

Although *C. lupus* is categorized as Least Concern on the IUCN Red List (Mech & Boitani, 2010), the Arabian subspecies is threatened regionally and is vulnerable locally (Dolev & Perevolotsky, 2004; Sillero-Zubiri et al., 2004) because of systematic shooting and trapping (Mendelssohn, 1983; Cunningham & Wronski, 2010). In Israel the Arabian wolf is fully protected by law and is not persecuted except for specific management actions. Although Hefner & Geffen (1999) have studied some behavioural aspects of this subspecies, no reliable demographic data are available. Hefner & Geffen (1999) documented the dispersal of individuals over 50–150 km, suggesting a continuous admixture between wolves in the southern and central Negev and the likelihood that the wolves in the Negev Desert comprise a single population.

Israel is divided into four INPA districts that are relatively independent of each other but controlled by a single directing body. The 13,000 km² Negev Desert, which covers more than half of Israel, is divided into two INPA districts (Fig. 1). During 2005–2008 these two districts had different management policies for wolves. The Eilat district culled wolves by trapping or shooting (B. Shalmon, pers. comm.) and tightly controlled the access of carnivores to rubbish pits, although wolves may still have fed from

garbage containers inside settlements and army camps. The South district, on the other hand, did not cull wolves and did not control access of wolves to local rubbish pits. Consequently, while the Eilat district attempted to reduce the wolf population to a minimum, the management exercised by the South district counteracted this. The above actions taken by INPA raise a question: how can a single population be managed by two contradictory policies? To address this issue we studied the genetic population structure and estimated population size using faecal DNA. Specifically, we examined whether the wolf population in the two INPA districts has similar densities and genetic composition. The findings of our study offer a new and reliable baseline for modification of the current INPA management policy for the Arabian wolf in the Negev Desert. Furthermore, our results can shed light on the effects of different management policies on the population genetics of a threatened species.

Study area

The central and southern Negev is a rocky desert, a mixture of mountains, washes, and depressions. The 160 km Arava Valley, an elongated depression of the Rift Valley, borders the Negev in the east. Elevation in the mountainous regions is 400–1,000 m, falling to 50–150 m at the bottom of the Rift. The climate is hot and arid, with an annual rainfall of 50–100 mm, falling during the winter months of November–March (Evenari et al., 1982). There are two cities, Eilat (population 50,000) at the northern tip of the Red Sea, and Mitzpe Ramon (population 6,000) in the Negev highlands. All other settlements are agricultural communities (10 in the Eilat district and eight in the South district), mostly in the Rift Valley.

Methods

Sample collection

To sample wolf scats equally over the central and southern Negev Desert we established a grid of 12 feeding sites c. 20 km apart. Five feeding sites were established in the Eilat district (sampling an area of c. 2,129 km²) and seven in the South district (sampling an area of c. 2,910 km²; Fig. 1). In setting out this grid of feeding sites we also considered vehicle accessibility and other entry restrictions. Mean home range size of Arabian wolves in the Negev Desert is 34.6 ± SD 19.5 km² and mean daily travel distance is 13.7 ± SD 6.7 km (maximum recorded distance travelled in a day is 42 km; Hefner & Geffen, 1999). Considering this mean daily travel distance all wolves in this area could probably reach at least one of the feeding sites on a daily basis. Once a month INPA rangers placed a cow calf carcass

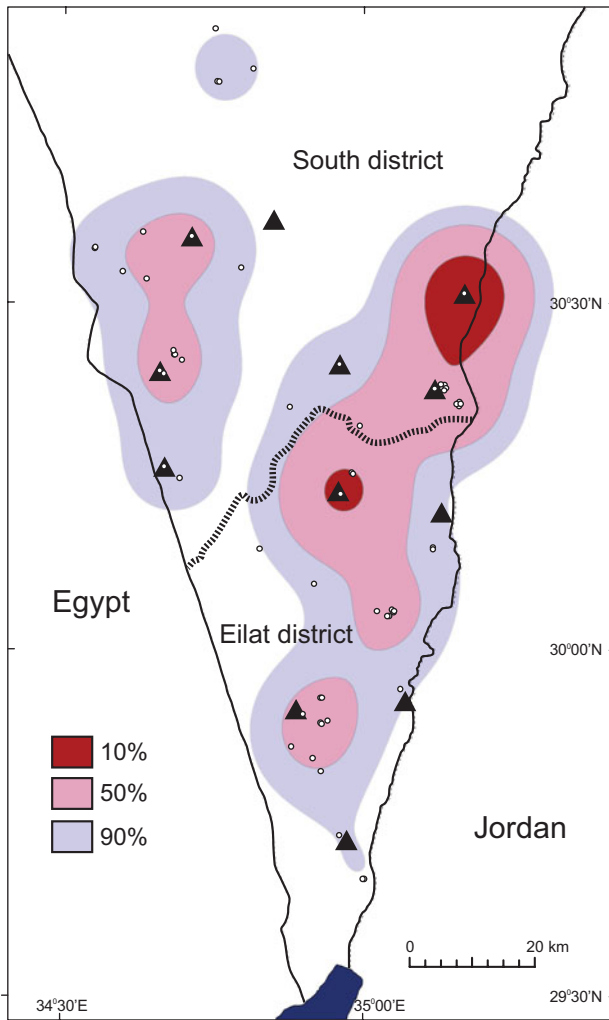


FIG. 1 Distribution of the 12 feeding sites (black triangles) used for collecting wolf *Canis lupus arabs* scats in the Negev Desert (see text for details). White circles denote locations where scats were collected. Kernel densities of 90, 50 and 10% for wolf scats are denoted. A single smoothing parameter of $h=10$ km was used in all kernel contours. The heavy dashed line indicates the boundary between the south and Eilat INPA districts.

at each feeding site and scanned the site for scats 1 week later. Each calf carcass was secured to prevent removal by predators. The meat provided at each feeding site was insufficient to support local wolves but provided an incentive for resident wolves to inspect the site on a regular basis and defecate nearby. Because it is difficult to find wolf scats by random searches, except around garbage dumps, this system facilitated the sampling of individuals that are not associated with human habitation on a regular basis. Scats found on the routes leading to the feeding sites were also collected. Each scat was placed in a paper bag and the location and date of collection recorded. In this hot, arid environment, scats become dry a few hours after defecation and storage in paper bags keeps them dry and protected from condensation.

Genetic analysis

We extracted DNA from faecal samples using the protocol in Reed et al. (1997). Diagnostic mtDNA segments were used to confirm wolf DNA. We used 10 polymorphic wolf microsatellites to identify individuals. Details of the molecular procedures are outlined in Supplementary Information 1.

We used two population genetic approaches to examine whether a spatial correspondence exists between genetic and spatial grouping (i.e. INPA districts) of wolves. Details of the relatedness network analysis are provided in Supplementary Information 1.

Bayesian clustering

To assign individuals to populations we used *STRUCTURE* 2.3.1 (Pritchard et al., 2000), which uses multi-locus genotypes (X) to infer population structure. The Bayesian model assumes K populations that have different allele frequencies within a set of independent loci. We applied the admixture ancestry model once with the independent allele frequency model and once with the correlated model. The number of populations (K) was modified from 1 to 10. For each value of K 10 independent simulations were run for each of the two allele frequency models. Each simulation had a burn-in period of 500,000 iterations followed by a sampling of 500,000 iterations. The likelihood of K (i.e. $\ln \Pr(X|K)$) and ΔK (Evanno et al., 2005) were used to infer the more likely number of populations. This algorithm also calculates a membership coefficient (Q) for each individual in each population. We used the highest Q value to assign each individual to a population.

Population size estimation

We used three different approaches to estimate the wolf population size in the Negev: rarefaction curves, capture-mark-recapture, and a simple urn model called *Capwire*.

Rarefaction curve In this technique the data are plotted as the number of unique multi-locus genotypes (i.e. individuals) as a function of sample size (number of scats analysed) and then fitted to a curve. The asymptote of the curve provides an estimate of the population size. Three fitting equations have been employed in similar previous studies: $y = ax/(b+x)$ (Kohn et al., 1999), $y = a(1-e^{-bx})$ (Eggert et al., 2003) and $y = a - a(1-1/a)^x$, which was developed by D. Chessel (Frantz & Roper, 2006). In all equations a is the asymptote, b determines the rate of decrease of the slope, x is the number of scats sampled, and y is the number of unique genotypes. We used *GIMLET* to generate rarefaction curves (Valiere, 2002). Because the order in which DNA samples are added has an influence on the shape of the curve

(Kohn et al., 1999) we used *R 2.10.1* (Ihaka & Gentleman, 1996) to randomize (1,000 iterations) the order of the DNA samples and projected the asymptote for each of these randomizations using each of the three equations. The median of all 1,000 iterations of the asymptote was taken as an estimate of the population size (Frantz & Roper, 2006). The 95% confidence interval (CI) corresponds to the 0.025 and 0.975 percentiles.

Capture–mark–recapture (CMR) model The study design (i.e. sampling at the feeding stations at monthly intervals) allowed us to easily adapt the genetic data to the CMR model. Identical multi-locus genotypes were considered to originate from the same individual and a history of captures and recaptures was assembled for each genotype. We divided the study period into 20 monthly sampling sessions. If the same genotype was sampled more than once in the same month we considered only one capture for that month. Analysis was performed with an open population model, using the *POPAN* module in *MARK* (White & Burnham, 1999). We used a set of four possible models: fully time dependent ($pt, \phi t, bt$) where p represents the probability of capture, ϕ represents the probability of an animal surviving between occasions and b is probability of entrance; constant catchability ($p, \phi t, bt$); constant survival rate (pt, ϕ, bt); and constant catchability and survival rate (p, ϕ, bt). We performed a goodness-of-fit using the *RELEASE* suite (Cooch & White 2007) to confirm that the most general sub-model adequately fitted the data. *MARK* uses AIC_c (Akaike's information criterion corrected for small sample size) for model selection.

Capture with replacement In non-invasive sampling individuals may be 'captured' several times within one sampling session. When using classical CMR models such multiple captures must be pooled, as an animal should not be captured more than once in one session. This potentially wastes data. *Capwire* (Miller et al., 2005) uses a method developed to accommodate this type of data and is based on a simple urn model. There are two capture models: the even capture probability model (ECM) in which every individual is equally likely to be captured in each sampling session with a probability of $1/\text{population size}$, and the two innate rates model (TIRM), which is more realistic, in which there are two types of individuals with different probabilities of capture. The software performs a likelihood-ratio test to determine which model is more suitable for the given data.

Results

Sample collection and microsatellite genotype

Over a period of 20 months from May 2007 to December 2008 a total of 334 faecal samples were collected and an additional three tissue samples were taken from wolves that

TABLE 1 The number of alleles, observed (H_{obs}) and expected (H_{exp}) heterozygosity, and the rate of allelic dropout and false alleles for each of the 10 loci used in this study.

Locus	No. of alleles	H_{obs}	H_{exp}	Allelic dropout	False alleles
CXX618	7	0.80	0.72	0.040	0.018
FH2137	14	0.87	0.84	0.062	0.174
FH2010	6	0.62	0.55	0.073	0.259
FH2175	9	0.76	0.73	0.068	0.066
CXX733	9	0.79	0.75	0.157	0.017
FH2079	7	0.67	0.68	0.128	0.164
CXX140	7	0.75	0.79	0.041	0.092
CXX250	5	0.93	0.76	0.025	0.123
CXX251	6	0.58	0.63	0.031	0.146
CXX758	7	0.55	0.74	0.049	0.073
Mean	7.7	0.73	0.72	0.068	0.113

had been shot by INPA rangers. Most scats (82%) were collected after the wolf control programme was terminated. Mitochondrial DNA for species identification was successfully amplified in 230 (69%) of the faecal samples, of which 95 were identified as wolf DNA, 93 were of red foxes *Vulpes vulpes*, 12 of dogs *Canis lupus familiaris*, nine of golden jackals *Canis aureus* and two of striped hyaenas *Hyaena hyaena*. In 5% of the samples that were identified, mtDNA of the prey (*Ovis aries*, *Capra hircus*, *Bos taurus*) rather than the predator was amplified.

Of the 98 wolf DNA samples (i.e. 95 faecal samples plus three tissue samples) we obtained a complete or nearly complete multi-locus genotype for 76 samples. The mean rate of allelic dropout was 0.068 and mean rate of false alleles was 0.113 for all replicates at all microsatellite loci. The probability of two individuals possessing identical genotypes (PI) was 1.40×10^{-10} for a randomly mated population and 1.635×10^{-4} for siblings. Error rates and PI values of each locus were calculated using *GIMLET* (Table 1, Supplementary Information 1). We identified 52 unique haplotypes of which 12 were sampled more than once, with a mean of 1.46 samples per individual. The maximum resample rate was six ($n=2$).

Population structure

We used the Bayesian model in the programme *STRUCTURE*. The highest $\ln \Pr(X|K)$ and ΔK were observed for $K=3$, indicating three genetic subpopulations (Figs 2 & 3a). This result was consistent for both correlated and independent allele frequency models. Most sampled individuals were easily assigned to a cluster as their estimated membership coefficient (Q) was > 0.7 for a specific population. However, three individuals (15, 7, and 11) could not be assigned because their Q values were ≤ 0.7 (Fig. 3a).

The distribution of two of the genetic populations (C_3 and C_1 ; Fig. 3b) did not show any clear spatial separation

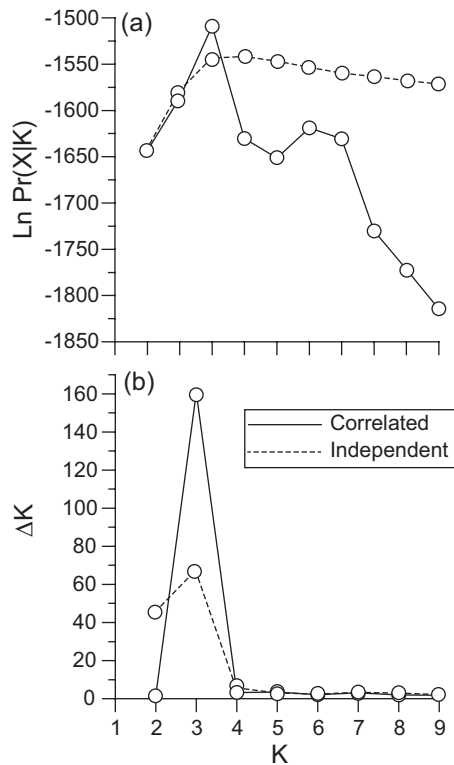


FIG. 2 Number of subpopulations inferred by *STRUCTURE*. (a) Mean $\text{Ln Pr}(X|K)$ over 10 runs for each K value. (b) ΔK for each K value, where $\Delta K = m[L(K+1) - 2L(K) + L(K-1)] / s[L(K)]$. Both $\text{Ln Pr}(X|K)$ and ΔK were calculated for the correlated and independent models.

between INPA districts. In Eilat district six individuals were of the C_1 cluster and 10 of the C_3 cluster, and in the South district 10 individuals were of the C_1 cluster and eight of the C_3 cluster (Fig. 3b). However, a third population (C_2 ; Fig. 3b) had most individuals ($n = 13$) in the South district and only one individual in the Eilat district. Thus, the division between districts was significantly dependent on genetic clustering ($\chi^2 = 8.114$, $P = 0.0172$) because of the uneven distribution of the C_2 cluster members between districts (Fig. 3b).

Population size estimation

The Kohn rarefaction equation gave the highest population estimate, of 176.8 wolves (95% CI 120.8–491.4), followed by that of Eggert with an estimate of 102.1 (95% CI 74.4–256.2), and Chessel with the lowest estimate of 86.2 wolves (95% CI 72.1–105.3; Table 2). The TIRM model of *Capwire* ($L(\text{TIRM}) - L(\text{ECM}) = 22.48$) gave an estimate of 142 wolves (95% CI 89–179). For the CMR model the goodness-of-fit test confirmed that the most general sub-model adequately fitted the data ($\chi^2_{11} = 3.447$, $P = 0.983$). The constant catchability and constant survival rate model ranked highest by AIC_c and gave an estimate of 173.4 wolves (95% CI 85.6–261.1; Table 3).

A separate population estimation for each district showed that although the number of wolves in the Eilat district was considerably lower than in the South district, the density of wolves was similar in both districts based on the rarefaction estimates but higher in the South district based on the CMR and *Capwire* estimates (Table 2). CMR models showed that the lowest AIC_c value was obtained by the constant catchability model for the South district and the constant catchability and survival rate model for the Eilat district (Table 3).

Discussion

Our results show that the wolf populations in the two INPA districts had a similar genetic composition. The Bayesian model in *STRUCTURE* identified three genetic subpopulations: two distributed across both districts, and one mostly in the South district (Fig. 3). However about half of the individuals of the latter population were found near the south-eastern corner of the South district, at the border between the districts. Additional sampling would probably reveal their presence in the north-eastern corner of the Eilat district. Furthermore, network analysis of relatedness shows that individuals of both communities are present in both INPA districts in approximately equal proportions (Supplementary Fig. S1). Taken together, the evidence for considering the wolves in the two districts as separate, distinct populations is weak. Overall, our data suggest that all the wolves in the Negev Desert comprise a single population, which should therefore be managed as a single conservation unit. This view is supported by the extensive dispersal distance of Arabian wolves in the Negev (Hefner & Geffen, 1999), implying that no isolation exists between wolf populations living throughout this region.

In these contradictory management approaches between the two INPA districts priority has been given to gazelles over wolves. This situation could have been avoided through a more balanced ecosystem approach that considers the ecological role of both species. The Arabian wolf is currently the top predator in this system, and the gazelle (mostly *G. dorcas*) is one of the largest native ungulates together with the Nubian ibex *Capra ibex nubiana*, onager *Equus hemionus* and Arabian oryx *Oryx leucoryx*. The latter two species were recently reintroduced into the Negev Desert. Both wolves and gazelles have key ecological roles in the Negev ecosystem (Mendelssohn & Yom-Tov, 1999).

Does the Arabian wolf exert a significant impact on the gazelle population? This issue has never been systematically studied in the Negev Desert. Our findings indicate that the number of wolves in the southern Negev (i.e. Eilat district) is c. 30–50, whereas 60–100 inhabit the central Negev (i.e. South district). Both gazelle species have been counted annually in the Negev Desert since the 1970s. The mountain

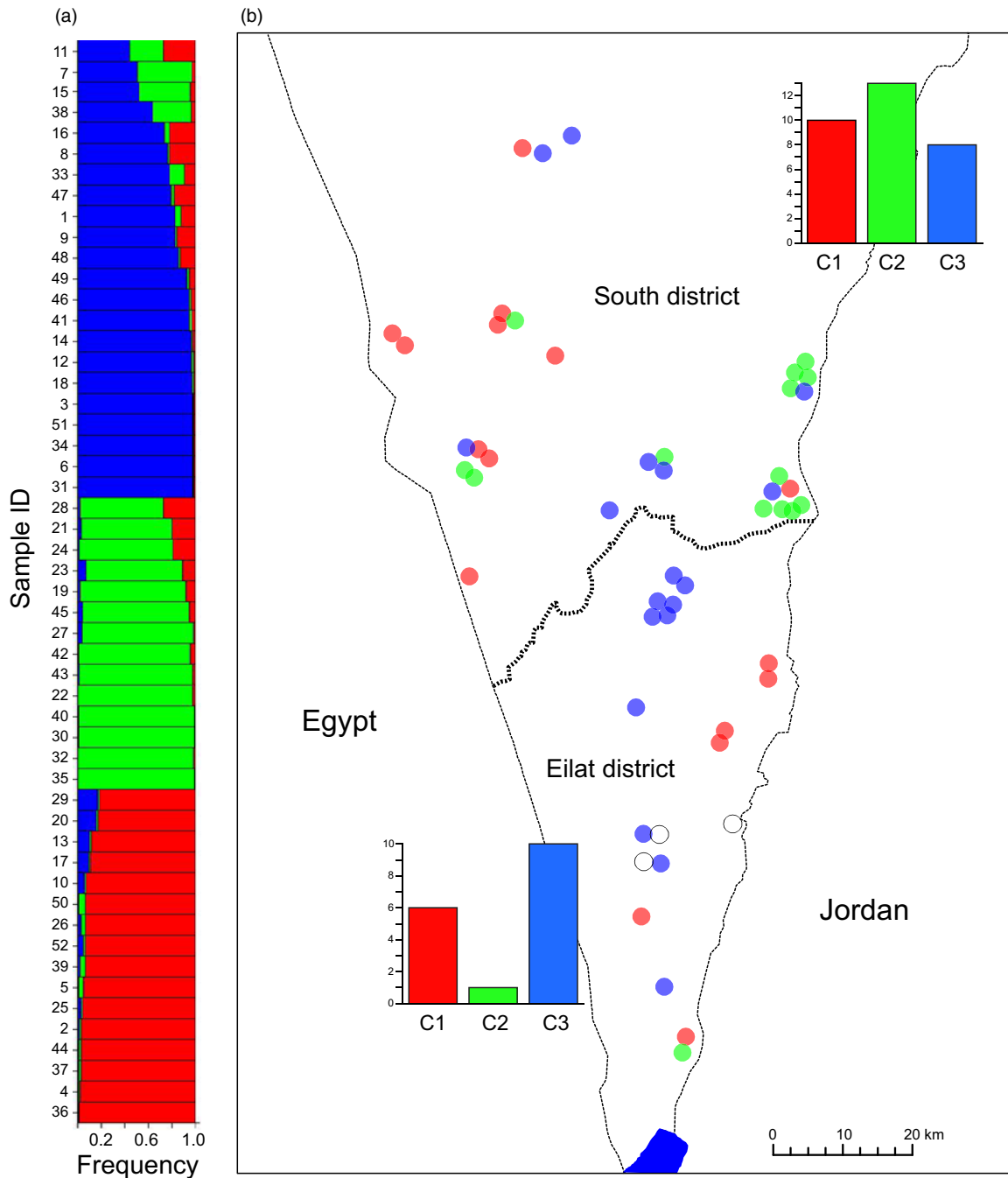


Fig. 3 Assignment of haplotypes by the *STRUCTURE* Bayesian algorithm. (a) Probability of belonging to each of three clusters ($K=3$) is denoted by shading (as in (b)). (b) Spatial distribution of haplotypes, assigned to the cluster with the highest probability. Empty circles are haplotypes that could not be assigned to a cluster. The frequency of each cluster (C1, C2, C3) in each of the districts is indicated by a bar chart. The heavy dashed line indicates the boundary between INPA districts.

gazelle population is restricted to a single locality in Eilat district and has numbered $c. \leq 40$ since 1978 (Yom-Tov & Ilani, 1987; Dolev & Perevolotsky, 2004). Thus, the significance of mountain gazelle to the overall Negev ecosystem is negligible. However, the dorcas gazelle is widespread in the Negev, with densities reaching 5 km^{-2} in areas where many acacias are present and 0.1 km^{-2} in areas without acacias (Baharav, 1980). Based on total counts the

population size of the dorcas gazelle in the Negev during 1980–1985 was estimated at 1,000–1,300 (Yom-Tov & Ilani, 1987; Mallon & Kingswood, 2001). During 1971–2009 the number of dorcas gazelles was 500–900 in the Eilat district (B. Shalmon, unpubl. data) and 150–550 in the South district (B. Shalmon & A. Tsoar, unpubl. data). Dorcas gazelle numbers in the Eilat district have fluctuated considerably in the past 40 years (Yom-Tov & Ilani, 1987); however, there

TABLE 2 Mean population size and mean density of the Arabian wolf *Canis lupus arabs* in the Negev Desert, and in the South and Eilat INPA districts separately (Fig. 1), based on five estimation methods (see text for details).

Area	MARK	Capwire	Rarefaction		
			Chessel	Eggert	Kohn
Negev Desert					
Mean population size (95% CI)	173.4 (85.7–261.1)	142.0 (89.0–179.0)	86.2 (72.1–105.3)	102.1 (74.4–256.2)	176.8 (120.8–491.4)
Density (100 km ⁻²)	3.4	2.8	1.7	2.0	3.5
South district					
Mean population size (95% CI)	98.9 (10.7–187.1)	86.0 (46.0–116.0)	44.5 (36.1–57.3)	60.6 (38.1–363.7)	106.0 (60.5–916.0)
Density (100 km ⁻²)	3.4	3.0	1.5	2.1	3.6
Eilat district					
Mean population size (95% CI)	47.6 (19.3–76.0)	33.0 (23.0–52.0)	33.3 (25.8–44.1)	33.1 (24.4–97.9)	55.4 (37.8–194.6)
Density (100 km ⁻²)	2.2	1.6	1.6	1.6	2.6

TABLE 3 Selection of MARK models (see text for details) for the Negev Desert population and INPA South and Eilat district populations separately.

Model	Parameters	AIC _c ¹	ΔAIC _c ²	WAIC ³	L ⁴ _{model}
Negev Desert					
Constant catchability & survival rate (p, φ, bt)	22	249.6	0.0	1.00	1.0
Constant catchability ($p, \varphi t, bt$)	40	377.3	127.7	0.00	0.0
Constant survival rate (pt, φ, bt)	41	389.3	139.7	0.00	0.0
Time dependent model ($pt, \varphi t, bt$)	57	1,094.7	845.1	0.00	0.0
South district					
Constant catchability & survival rate (p, φ, bt)	13	132.9	16.4	0.00	0.0
Constant catchability ($p, \varphi t, bt$)	14	116.5	0.0	0.91	1.0
Time dependent model ($pt, \varphi t, bt$)	16	121.1	4.6	0.09	0.1
Constant survival rate (pt, φ, bt)	26	314.6	198.1	0.00	0.0
Eilat district					
Constant catchability & survival rate (p, φ, bt)	10	151.9	0.0	1.00	1.0
Constant catchability ($p, \varphi t, bt$)	14	164.5	12.6	0.00	0.0
Constant survival rate (pt, φ, bt)	19	216.1	64.2	0.00	0.0
Time dependent model (pt, φ, bt)	21	261.9	111.0	0.00	0.0

¹AIC value corrected for small sample sizes ²Difference in AIC_c between the focal and the best model ³AIC weight ⁴Model likelihood

are no comparable annual data on wolf numbers to support the claim that these fluctuations are related to predation. Nor is reliable data on predation of fawn or adult mountain and dorcas gazelle species available for the Negev population. Analysis of 777 wolf scats from the southern Negev (Shalmon, 1986) revealed that the major dietary components of Arabian wolves are cow carrion (found in 62.5% of scats), fruit and vegetation (51.4% of scats) and human garbage (37.2% of scats). The frequency of indigenous prey, including gazelles, in the scats was only 6.3%. In North America Messier et al. (1995) showed that the number of wolves per area is linearly related to the density of deer-sized prey. In these populations, wolves consumed mainly large ungulates (northern Minnesota, Kunkel & Mech, 1994; Isle Royale National Park, Vucetich et al., 2002; Yellowstone National Park, Wright et al., 2006) In contrast, the Arabian wolves in the Negev seem to be dependent on human-related waste and domestic livestock and seldom rely on natural prey.

The implications of our results are that a major change in wolf management in both INPA districts is required. Firstly, Arabian wolves should not be culled without a robust corresponding study on the effects of such action. The Arabian wolf is a threatened species, living in an arid zone with limited resources (Dolev & Perevolotsky, 2004; Sillero-Zubiri et al., 2004). Its population fluctuates similarly to other species dwelling in this region, and may reach critical low levels. Furthermore, culling is indiscriminate and could thus inadvertently eliminate a distinct genotype and thereby reduce the evolutionary potential for survival of this population. Secondly, we suggest that reducing access to human-related waste and domestic livestock could adjust the wolf population size to this arid environment's natural carrying capacity by forcing the wolf population to feed more on natural prey. Thirdly, the effect of droughts and habitat modification by human activity on the dorcas gazelle population in the Negev should be systematically investigated before further culling of predators is carried out.

At present, both districts are modifying their management policies with respect to wolves. Wolf culling has been terminated, and new programmes are being developed for limiting the access of carnivores to human waste and agriculture surplus. We have also developed a study on wolf resource utilization, using GPS collars, to understand the dependency of wolves on human resources. Finally, we call for more detailed study on the population dynamics of gazelles in the Negev, especially in relation to effects of global warming, before instigating any further predator control programmes in this region.

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