Food availability and population structure: How do clumped and abundant sources of carrion affect the genetic diversity of the black-backed jackal?

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Abstract

Carnivores frequently come into conflict with humans in agricultural and livestock producing areas around the world. Understanding their fidelity and dispersal patterns in response to food availability is therefore important given the effort invested in conflict mitigation strategies. In this study, we investigated the influence of clumped and abundant sources of carrion on the genetic diversity of the black-backed jackal (*Canis mesomelas*) within six private game farms in the North West and Gauteng provinces of South Africa. It is predicted that clumped and abundant sources of carrion will increase immigration and thus genetic diversity in the local subpopulation. By quantifying the variability of microsatellite loci in black-backed jackals subjected to artificially increased carrion availability, and comparing them with individuals from control sites, we were able to describe patterns of historic gene flow within the total sampled population. The results of this investigation indicate that clumped and abundant sources of carrion promote genetic structuring ($F_{ST} = 0.0302$) which implies a lack of gene flow and a degree of isolation. Genetic artefacts of three populations could be identified through Bayesian clustering analysis of individuals based on their genetic identity. Individuals sampled from the two supplementary feeding sites could be assigned to one of two ancestral populations with an average population assignment of 69% and 82%, while individuals from the remaining four control sites, originate from a third population with percentage assignments of 63%, 46%, 53% and 42%. It is therefore likely that clumped and abundant sources of carrion in the agricultural landscape of South Africa can affect the population dynamics of the black-backed jackal and result in subpopulations with limited migration and dispersal when compared with the total population.
Introduction

It is generally recognised that carnivores play a fundamental role in the structure and function of an ecosystem (Ripple, et al., 2014; Ripple & Beschta, 2004). However, factors such as disease transmission and livestock depredation frequently promote conflict in areas where humans and carnivores exist in close proximity (Woodroffe, et al., 2005).

Understanding the ecological factors that drive the spatial organisation of free-ranging carnivores is therefore important when considering both conservation and management of species in the human-modified landscape. Thus this study follows a microsatellite-based approach to investigate the short term historic effects of four years of supplementary feeding on the genetic diversity of black-backed jackals (*Canis mesomelas*) at private game farms in South Africa.

Following the expectations of the resource dispersion hypothesis (Macdonald, 1983), an increase in localised food availability will often result in a breakdown in territorial stability and subsequently lead to an increase in local density (Johnson, et al., 2002; Johnson, et al., 2001). Indeed, anthropogenically derived sources of food, synonymous with agricultural and human modified landscapes, have been shown to strongly influence the spatial organisation of many omnivorous canids including the golden jackal (*Canis aureus* Rotem, et al., 2011), red fox (*Vulpes vulpes* Contesse, et al., 2004) coyote (*Canis latrans* Fedriani, et al., 2001) and dingo (*Canis lupus dingo* Newsome, et al., 2013). Furthermore, studies in both Namibia and South Africa have recorded the black-backed jackal at far greater abundances than expected in areas where scavenging opportunities are high and carrion availability is clumped, stable.
and abundant (Yarnell, et al., 2014; Jenner, et al., 2001; Hiscocks & Perrin, 1988). Studies using both radio-telemetry and behavioural observations in the Cape Cross Seal Reserve (CCSR) have also concluded that territorial boundaries of the black-backed jackal often overlap in close proximity to clumped, abundant resources such as seal colonies (Hiscocks & Perrin, 1988), and that home range sizes significantly increase with distance from the colony itself (Jenner, et al., 2001). As the social structure of the black-backed jackal is commonly reported to consist of a monogamous breeding pair, which holds and aggressively defends territory from transient individuals and neighbouring residents (Estes, 1991; Ferguson, et al., 1983), it is clear that an increase in local abundance of food can dramatically affect both the territorial behaviour and spatial organisation of this species. However, what remains unclear from contemporary observations is the effect that increased food availability has on the fidelity and dispersal of such subpopulations over time. Therefore by examining the genetic diversity of black-backed jackals in the game farms of South Africa, this study aims to elucidate the genetic consequences of clumped and abundant sources of food on the dispersal of a free-ranging canid within a human-modified landscape.

The black-backed jackal is a medium sized canid (5-15 kg) with two discrete distributions that span the majority of the Southern African sub-region, and parts of Eastern Africa (Skinner & Chimimba, 2005; Estes, 1991). This study focuses on the southern African subspecies (C. m. mesomelas), henceforth “black-backed jackal”, due to the high rate of human-carnivore conflict associated with this region (Thorn, et al., 2012). As a vector of rabies and canine distemper (Bellan, et al., 2012; Zulu, et al., 2009), and an opportunistic hunter of small game and livestock (Estes, 1991), the black-backed jackal is frequently
controlled as a pest species throughout its range (Thorn, et al., 2012; Ginsberg & Macdonald, 2004). With an omnivorous diet consisting of small mammals, livestock, forage and carrion (Klare, et al., 2010), this species is considered a generalist carnivore that is able to undertake diet switching in response to changes in local food availability (Kamler, et al., 2012; van der Merwe, et al., 2009; Rowe-Rowe, 1983; Fourie, et al., 2015; Humphries, et al., 2016). Therefore, to further investigate the effect of food availability on the population dynamics of the black-backed jackal, this study used carrion stations, known as vulture restaurants, to measure the historic effect of artificially increasing scavenging material on the gene flow and variation in genetic diversity within and between local subpopulations. Vulture restaurants were originally introduced in participating game farms and nature reserves across South Africa with an aim to supply declining vulture species with a safe and consistent source of carrion which originates from hunted or slaughtered livestock destined for the human food chain. Subsequent analysis has shown that the regular deposition of carcasses at these sites has resulted in an unintentional increase in the local abundance of many scavenging carnivores, including the black-backed jackal (Yarnell, et al., 2014). As the abundance of black-backed jackals residing in close proximity to vulture feeding sites are often far in excess of those in the surrounding area (pers. obs.), it is predicted that clumped and abundant sources of carrion will have resulted in an increase in genetic diversity within local subpopulations as it is hypothesised that increased food availability increases migration.

Methods

Sampling and study sites
This study was undertaken in the North-West and Gauteng provinces of South Africa. Individual black-backed jackals (n = 65) were sampled for genetic material from six game breeding farms (Fig. 1) between March 2011 and September 2012 for an analysis of population structure. Two game farms, Site VR1 and Site VR2, had active vulture restaurants initiated approximately four years prior to sampling (n = 27 and 19 jackal DNA samples, respectively). The remaining four game farms, Site C1, C2, C3 and C4, acted as control sites with no additional scavenging material provided (n = 6, 6, 3 and 4). Carrion, consisting of recently deceased ungulates, was placed at each vulture restaurant on a regular basis with an average of 797 kg a month being recorded between 2008 and 2011 at site VR1 (Yarnell, et al., 2013). A non-invasive genetic recovery protocol was used to acquire genetic material from 63 recently deposited faecal samples along with two tissue biopsies opportunistically collected from the ear lobe of deceased individuals. The non-invasive genetic recovery protocol used in this investigation was specifically designed for use with this species and had previously been tested for adequate recovery of host DNA prior to undertaking analysis (James, et al., 2015). Tissue samples were placed in 1.5 ml of absolute ethanol (EtOH) after collection and stored at -20°C prior to transport to the UK for further analysis.

To sample faecal deposits for genetic source material, driven transects of 5 km were undertaken along the road networks within each site. Transect routes were chosen to maximise an even coverage of area and habitat types. Transect width was standardised at 2m from the edge of the road to minimise the variation in detection probability. All
Analysis. These markers were used to estimate the population structure and inbreeding coefficients of the black-backed jackal. Individual multilocus genotype profiles that matched were considered to derive from the same source and were hence removed prior to the analysis. Results were pooled by site for an analysis of population structure.
A chloroform extraction protocol was used in conjunction with a Qiagen DNeasy spin column method to isolate and purify DNA templates from faecal samples collected in the field. Faecal samples stored in S.T.A.R. buffer were defrosted in batches at 4°C prior to DNA extraction. Individual samples were homogenised by shaking, then 10 ml of sample was transferred to a sterile collection tube. One millilitre of ≥ 99.8 % chloroform-EtOH (GC) was

### Table 1. Microsatellite loci, 5′ modification, forward (F) and reverse (R) primer sequences (5′-3′), Tm and NCBI accession numbers (AN).

<table>
<thead>
<tr>
<th>Locus</th>
<th>5′ mod</th>
<th>F primer</th>
<th>Tm oC</th>
<th>R Primer</th>
<th>Tm oC</th>
<th>AN</th>
</tr>
</thead>
<tbody>
<tr>
<td>cme144</td>
<td>FAM</td>
<td>aaccttaagccacactttgca</td>
<td>57.9</td>
<td>acttgcccttggtttaagc</td>
<td>58.4</td>
<td>KU050829</td>
</tr>
<tr>
<td>cme136</td>
<td>FAM</td>
<td>aactggccaaacataacagc</td>
<td>58.5</td>
<td>ttcattaaccccttgcccctg</td>
<td>58.5</td>
<td>KU050830</td>
</tr>
<tr>
<td>cme206</td>
<td>HEX</td>
<td>cgagagcaacataggcatga</td>
<td>58.4</td>
<td>caaagtgtggtgagggcgtc</td>
<td>58.8</td>
<td>KU050831</td>
</tr>
<tr>
<td>cme196</td>
<td>HEX</td>
<td>aggaggacagaagacagaagg</td>
<td>57.5</td>
<td>atggatgtattgtaggggtg</td>
<td>58.0</td>
<td>KU050832</td>
</tr>
<tr>
<td>cme193</td>
<td>FAM</td>
<td>gagctctgtggaagagccttga</td>
<td>58.6</td>
<td>catctggtcctgtacttcaa</td>
<td>58.0</td>
<td>KU050833</td>
</tr>
<tr>
<td>cme210</td>
<td>HEX</td>
<td>ctttgcaatcattcatctttga</td>
<td>57.2</td>
<td>cccgaggtaacctttggtc</td>
<td>57.5</td>
<td>KU050834</td>
</tr>
</tbody>
</table>
then mixed with the sample solution and vortexed to form an emulsion. Emulsified samples
then underwent centrifugation at 1,000 x g for 3 min and the subsequent supernatant was
removed for further processing. A Qiagen blood and tissue extraction protocol was followed
to recover DNA from the supernatant. Spin columns were centrifuged at 1400 x g for 3 min
prior to elution, to remove excess EtOH and chloroform from the silica membrane, and were
stood to dry at room temperature for 5 min. DNA elution was undertaken using 75 μl of
warmed elution buffer at 54°C (James, et al., 2015).

PCR reactions were undertaken in 25 μl volumes containing approximately 40 ng of DNA
template, estimated in triplicate using a nanodrop 2000 spectrophotometer, 1 × Invitrogen
PCR buffer, 1.5 mM MgCl₂, 1 unit of Invitrogen hot start PlatinumTaq DNA polymerase
(Invitrogen cat No: 10966-018), 1 unit of Qiagen Q-solution, 0.5 μl/ng BSA, 0.2 mM dNTP
mix and 0.2 μM primer mix. Amplification conditions used on a Techne TC-4000 thermal
cycler consisted of an initial denaturation at 94°C for 2 min, followed by 35 cycles of 94°C for
1 min, 55°C for 45 s and 72°C for 1 min finishing with a final extension stage of 72°C for 5
min.

**Statistical analysis**

The probability for exact Hardy-Weinberg proportions, F-statistics and estimates of allele
frequencies between the six sampled subpopulations and each STRUCTURE-identified
cluster were calculated using the program GENEPOP v. 4.2.1 (Rousset & Raymond, 1995;
limited sample sizes (Pritchard et al., 2000). This analysis employs a Bayesian clustering algorithm to correlate microsatellite allele frequency dissimilarities between individuals with prior knowledge of sample location. The inclusion of sample location is specifically recommended when determining low levels of population structuring under small spatial scales, where a significant $F_{ST}$ value has been determined (Hubisz, et al., 2009). This
approach assigns individuals to the most relevant deme based on genetic dissimilarity between individuals and groups. The admixture model used in this analysis accounts for the possibility of admixture within clusters, as opposed to pure distributions of genotypes, while remaining robust to the absence of admixture. This method was employed to detect any indication of subtle population structure using the genotype data in this study. The number of subpopulations, $K$, was estimated to be between 1 and 6 using a burn-in of 10,000 runs; Markov Chain Monte Carlo simulation (MCMC) run length of 100,000 with 10 iterations per simulation. Pairwise $F_{ST}$ values between STRUCTURE-identified clusters were calculated using the program GENEPOP v. 4.2.1 (Rousset & Raymond, 1995; Rousset, 2008) and examined for significance using the exact $G$ test.

Identification of the number of distinct and genetically consistent groups within the sampled population was estimated using the rate of change in the log probability of the data between successive estimates of the number of populations, termed delta $K$ ($\Delta K$) (Evanno, et al., 2005). The estimation of $K$ was undertaken using the program Structure Harvester (Earl & von Holdt, 2012). The programs CLUMPP V1.1 (Jakobsson & Rosenberg, 2007) and DISTRUCT v1.1 (Rosenberg, 2004) were then used to produce graphical representations of the structure analysis. However, recent research suggests that unbalanced sample sizes from known localities may result in the identification of spurious clusters by the program STRUCTURE (Puechmaille, 2016), which is likely to result in an underestimation of $K$ using the delta $K$ method outlined in Evanno (2005). As resampling a subset of genotypes to correct for unbalanced sample sizes is not appropriate in this case due to the small sample size, the approach of identifying a true value for $K$ using the estimators MedMeaK,
MaxMeaK, MedMedK and MaxMedK over 20 repeats per estimation of $K$ was used (Puechmaille, 2016). The maximum value of $K$ was interpreted by the number of clusters that contained at least one sampling locality at membership coefficient threshold of 0.5. The R package Kestimator (Puechmaille, 2016) was used to calculate the estimators listed above.

We used a cut off assignment to test for the number of potential migrants within each structure-identified cluster (Sacks, et al., 2004). An arbitrary cut off assignment of 70% was selected due to the limited sample size, local spatial arrangement and cluster assignment probability. A chi$^2$ test was used to assess the difference in the proportion of migrants between clusters.

The statistical power to reject the null hypothesis of genetic homogeneity in this investigation was assessed by undertaking a power test using the program POWSIM (Ryman & Palm, 2006) at $F_{ST}$ values of 0.001, 0.0025, 0.01, 0.03 and 0.05. Effective population size ($N_e$), when simulated populations drifted apart, was maintained at 4000 and the number of simulations per run was set to 1000. It is generally regarded that power scores should be greater than 0.8 to be confident of adequate power.

Results

Hardy-Weinberg exact tests and fixation statistics
Genotype frequencies across all loci were found to be in general alignment with Hardy-Weinberg proportions at the total population level ($X^2 = 73.4136$, d.f. = 72, $p = 0.432$). When examined by locus, three of the 36 tests were shown to deviate significantly from Hardy-Weinberg proportions across the six sampling localities ($p < 0.05$). However, the exact Hardy-Weinberg test by population indicated that the majority of this deviation was partitioned to Site VR1 ($X^2 = 33.4919$, d.f. = 12, $p< 0.05$), showing a heterozygote excess at locus cme136 (Weir and Cockeram $F_{IS} = -0.2203$, $p < 0.05$). Weir & Cockerham fixation statistics indicated that a degree of sub-structuring was apparent in the total population as highlighted by the multi-locus $F_{ST}$ estimate of 0.0302 (Table 2). Significant genetic differentiation was apparent between sample sites when examining the variation in allele frequencies between sites using the exact G test ($X^2 = 49.8182$, df = 12, $p < 0.05$).

Table 2. Weir & Cockerham fixation statistics for individual and combined loci across all localities.

<table>
<thead>
<tr>
<th>Locus</th>
<th>$F_{IS}$</th>
<th>$F_{ST}$</th>
<th>$F_{IT}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>cme144</td>
<td>0.0819</td>
<td>-0.0080</td>
<td>0.0746</td>
</tr>
<tr>
<td>cme136</td>
<td>-0.1783</td>
<td>0.0067</td>
<td>-0.1705</td>
</tr>
<tr>
<td>cme206</td>
<td>-0.0024</td>
<td>0.0834</td>
<td>0.0812</td>
</tr>
<tr>
<td>cme196</td>
<td>0.0875</td>
<td>-0.0019</td>
<td>0.0858</td>
</tr>
<tr>
<td>cme193</td>
<td>0.0223</td>
<td>0.0062</td>
<td>0.0284</td>
</tr>
<tr>
<td>cme210</td>
<td>-0.1823</td>
<td>0.1146</td>
<td>-0.0468</td>
</tr>
<tr>
<td>All:</td>
<td>-0.0272</td>
<td>0.0302</td>
<td>0.0039</td>
</tr>
</tbody>
</table>
Isolation by distance

Analysis of the entire microsatellite data set found no statistical correlation between Euclidian distance and pairwise F\textsubscript{ST} values at the population level (r = -0.1836, p = 0.75833). Furthermore, no evidence of isolation by distance could be ascertained at the individual level when the correlation between distance matrices was compared to simulated values under the absence of spatial autocorrelation (simulated p-value: 0.707, Fig. 2).

Figure 2 approximately here.

Analysis of population structure

The analysis of genetic variation within and between individuals and sites using the Evanno method (2005) indicated that the number of ancestral populations genetically represented in the contemporary data set can be inferred as K = 3 (Fig. 3).

Figure 3 approximately here.

STRUCTURE analysis indicated that the population structuring, highlighted by the inbreeding coefficient (F\textsubscript{ST} ≈ 0.03), was largely partitioned between feeding site VR1 and feeding site VR2, being consistently dissimilar to each other and the remaining four sites in individual population assignment. Individuals from the remaining four control sites (C1, C2, C3 and C4) showed variable population assignment probabilities, thus a high degree of genetic
admixture was inferred across these sites. The analysis of MedMeaK, MaxMeaK, MedMedK
and MaxMedK indicates that the true value of $K = 3$.

Allelic richness, observed and expected heterozygosity, $F_{IS}$ and the Hardy-Weinberg test for
heterozygote excess and the proportion of potential migrants for each STRUCTURE-identified
cluster are shown in Table 3. Contrary to our predictions a greater proportion of migrants
were found in the STRUCTURE-identified cluster that included the four control sites (Cluster
3) when compared with the two supplementary feeding sites ($\chi^2 = 13.091, df = 2, p < 0.05$).

Table 3. Genetic diversity estimators and proportion of migrants for each STRUCTURE-
identified cluster.

<table>
<thead>
<tr>
<th>Cluster</th>
<th>Site</th>
<th>N</th>
<th>Ar</th>
<th>HO</th>
<th>HE</th>
<th>Overall FIS</th>
<th>HWE (p-value)</th>
<th>Migrants (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>VR1</td>
<td>27</td>
<td>47</td>
<td>103</td>
<td>105.2368</td>
<td>0.0217</td>
<td>0.7322</td>
<td>25.9</td>
</tr>
<tr>
<td>2</td>
<td>VR2</td>
<td>19</td>
<td>36</td>
<td>65</td>
<td>64.3377</td>
<td>-0.0103</td>
<td>0.6633</td>
<td>36.8</td>
</tr>
<tr>
<td>3</td>
<td>C1, C2, C3, C4</td>
<td>19</td>
<td>43</td>
<td>91</td>
<td>82.8843</td>
<td>-0.1009</td>
<td>0.3053</td>
<td>57.8</td>
</tr>
</tbody>
</table>

All pairwise FST values for each STRUCTURE-identified cluster (Table 4) were shown to be
significantly different ($p < 0.05$).

Table 4. Pairwise FST values for each STRUCTURE-identified cluster.

<table>
<thead>
<tr>
<th>Clusters</th>
<th>Sites</th>
<th>Pairwise FST</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 + 2</td>
<td>VR1 + VR2</td>
<td>0.0329</td>
</tr>
<tr>
<td>1 + 3</td>
<td>VR1 + (C1, C2, C3, C4)</td>
<td>0.0274</td>
</tr>
<tr>
<td>2 + 3</td>
<td>VR2 + (C1, C2, C3, C4)</td>
<td>0.0647</td>
</tr>
</tbody>
</table>
Analysis of statistical power

Power analysis undertaken using the program POWSIM indicated that the suite of microsatellite loci used in this investigation were suitable for differentiating population structure at FST values of 0.03 and above, with a Fisher’s exact test statistic > 0.8. Power analysis with FST values of 0.001, 0.0025, 0.01, 0.03, and 0.05 were computed as 0.0760, 0.1660, 0.7580, 0.9980 and 1.000 respectively.

Discussion

Carnivore spatial organisation is rarely, if ever, homogeneous in space and time. Resource-based explanations of spatial organisation are able to describe such variation by exploring the relationship between the availability of resources (e.g. food) and the fitness cost associated with territorial defence (Johnson, et al., 2001; Johnson, et al., 2002). Theoretical models that link resource dispersion with spatial organisation describe plasticity in territory size and stability when the distribution of food is heterogeneous across the environment (Macdonald, 1983; Johnson, et al., 2002). Thus, traditional explanations of the resource dispersion hypothesis place emphasis on the selective advantage gained by reducing territorial defence when the availability of food exceeds the requirements of the individual and group. For example, studies have concluded that populations of free-ranging red foxes residing in close proximity to human settlements are more likely to exist at higher densities than their rural counterparts due to the overabundance of anthropogenically derived sources of food (Bino, et al., 2010). However, the underlying mechanisms by which such populations are formed and maintained have been heavily veiled by their complexity. In this
study we found evidence for a small degree of genetic structuring within the population as a whole (Table 2). Furthermore, a Bayesian analysis of population structure showed that black-backed jackals at supplementary feeding sites were genetically distinct relative to the total population (Fig. 3). However, contrary to our predictions, individuals from the remaining four control sites could not be accurately assigned to a single population of origin based upon their genetic identity alone, and showed a far greater number of potential migrants relative to the supplementary feeding sites (Table 3), which suggests a degree of historic gene flow between these sampling locations. In addition, no evidence of spatial auto-correlation could be detected across the total population (Fig. 2), providing further evidence of a discontinuous population across the sampled area. We believe, therefore, that the results of this study show that far from increasing migration as predicted; clumped, abundant and stable sources of carrion can cause population structuring in the black-backed jackal by reducing gene flow between these sites. However, it should be noted that, while the identification of population sub-structuring is highly indicative of barriers to gene flow within the sampled population, evidence of slight outbreeding (Table 2) suggests that the genetic composition of the total breeding population has not been captured in its entirety. Despite this shortfall, the result of this study provides an informative estimation of the parameters of a population in flux and describes the genetic consequences of a population responding to increased food availability in the resource rich agricultural landscape.

Competitive exclusion offers an attractive explanation for the degree of population structuring seen in this study. Once the carrying capacity of the environment has been reached, it is intuitive that a relative increase in competition for food would prompt
terrestrial behaviour and limit or reduce migration and gene flow. Furthermore, due to the
large diversity of alternate sources of prey available to the black-backed jackal within the
agricultural landscape of South Africa (Kamler, et al., 2012), long distance commuting
behaviour, as observed at the CCSR (Jenner, et al., 2001), may not be a cost effective
strategy in this system. Furthermore, investigations into movement patterns of the dingo,
which reside at resource-rich refuse sites in central Australia, have shown that individuals do
not always remain at refuse sites indefinitely. This indicates that further selective pressures
above those predicted by the resource dispersion hypothesis, such as group hunting, may
play an important role in the social structure of the Canidae (Newsome, et al., 2013).
However, given that approximately 24-33% of offspring of territory-holding black-backed
jackals have been recorded as delaying dispersal to provide alloparental care to subsequent
kin (Ferguson, et al., 1983; Moehlman, 1983; Moehlman, 1986; Moehlman, 1987; Estes,
1991), a more likely explanation for the results of this study is that following a substantial
increase in local food availability the offspring of individuals in close proximity to
supplementary feeding sites have reduced dispersal rates, due to the high carrying capacity
of the environment and reduced competition for resources between siblings, resulting in
the formation of genetically distinct clusters of individuals. Previous studies have shown that
pup survival rate is positively correlated to both food availability and alloparental care
(Estes, 1991; Moehlman, 1987). Furthermore, the mechanisms dictating whether an
individual chooses to disperse from its natal range or to remain and act as a helper has been
correlated to food availability, competition for available resources and persecution
(Moehlman, 1987; Ferguson, et al., 1983; Minnie, et al., 2016). Therefore offspring that have
failed to disperse from their natal range, in combination with an increase in dispersing
offspring following disturbance from persecution at the control sites (Minnie, et al., 2016),
would explain, at least in part, the degree of population structuring seen in this study. However, although previous studies have suggested that a breakdown in territorial stability can result from clumped and abundant sources of food (Hiscocks & Perrin, 1988; Johnson, et al., 2002; Bino, et al., 2010), by sampling faeces for genetic material, a prominent territorial marker in many mammalian species, it is possible that transient individuals may have eluded genetic identification and potentially induced a sampling artefact to the analysis. Furthermore, the limited number of microsatellite loci used in this investigation has the potential to induce a type-two statistical error in this analysis as statistical power is often reduced when both sample size and microsatellite loci are limited in number. To date, only six microsatellite markers have been published for the black-black Jackal, which is relatively few by current standards. However, despite the limited resolution these markers offer for population structure analysis, they appear to be sufficient for identifying weak differentiation ($F_{ST}=0.03$), which we regard as still biologically meaningful. It is therefore recommended that future studies focus on the characterisation of further microsatellite loci with the aim of undertaking pedigree analysis using high quality tissue samples to accurately infer relatedness between individuals at supplementary feeding sites.

**Conclusions**

Many previous studies have shown that excess food availability can dramatically affect the population dynamics of carnivores (Hiscocks & Perrin, 1988; Fedriani, et al., 2001; Jenner, et al., 2001; Johnson, et al., 2001; Bino, et al., 2010; Rotem, et al., 2011; Newsome, et al., 2013; Yarnell, et al., 2014). An increase in the abundance and density of local
subpopulations is therefore expected following a substantial increase in carrion availability.

The results of this study indicate that anthropogenically provisioned resources (e.g. carrion) results in genetically identifiable groups of black-backed jackals that show a degree of historic isolation from the surrounding population. Whether through kin selection or the principles of competitive exclusion, the formation of a structured population in response to excess carrion is not unexpected given the assumed territorial breakdown described by the resource dispersion hypothesis. However the degree of genetic admixture at site VR1 suggests that immigration may play a substantial role in the formation of this cluster. Yet the ability to identify genetically distinct groups, in response to a vastly increased local carrying capacity, provides additional insight into the group dynamics of a monogamous and territorial carnivore in the human-modified landscape.

Acknowledgements

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locate faecal samples in the field. DEFRA import permits numbers for genetic material: TARP/11/392, TARP/2012/252 and TARP/12/404.
References


Figures

Figure 1. A map depicting the six study sites and the three subpopulations of black-backed jackals sampled in this investigation. Subpopulations are denoted by black circles.

Figure 2. Genetic distance as a function of geographic distance between individual black-backed jackals showing the initial correlation (dot) and the distribution of simulated data under the absence of Isolation by distance.

Figure 3. A graphical representation of population structure. Individual black-backed jackals are represented by vertical lines, with the population assignment represented in grayscale, $k = 3$.

Supplementary material A. Maps depicting the spatial arrangement of faeces collected for genetic analysis within each game farm site. Faecal deposits of the black-backed jackal are denoted by black circles and carrion stations are represented by white circles.

Supplementary material B. Individual black-backed jackal population assignment values for each structure identified cluster.
A map depicting the six study sites and the three subpopulations of black-backed jackals sampled in this investigation. Subpopulations are denoted by black circles.

Fig. 1
167x149mm (72 x 72 DPI)
Genetic distance as a function of geographic distance between individual black-backed jackals showing the initial correlation (dot) and the distribution of simulated data under the absence of Isolation by distance.

Fig. 2

167x141mm (72 x 72 DPI)
A graphical representation of population structure. Individual black-backed jackals are represented by vertical lines, with the population assignment represented in grayscale, $k = 3$.

Fig. 3

197x54mm (95 x 95 DPI)
Cover image

531x366mm (72 x 72 DPI)