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DOI

<https://doi.org/10.1016/j.biocon.2009.04.026>

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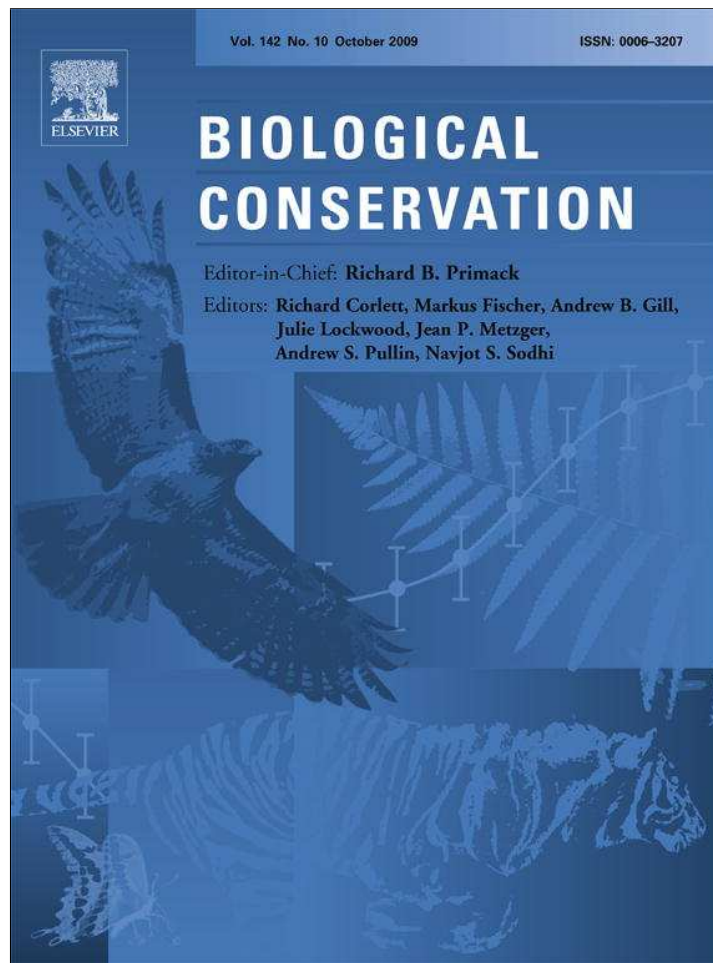
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Biological Conservation

journal homepage: www.elsevier.com/locate/biocon

Evaluating the demographic history of the Seychelles kestrel (*Falco araea*): Genetic evidence for recovery from a population bottleneck following minimal conservation management

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ARTICLE INFO

Article history:

Received 27 August 2008

Received in revised form 23 April 2009

Accepted 28 April 2009

Available online 23 May 2009

Keywords:

Bottleneck

Endemic island species

Genetic diversity

Museum DNA

ABSTRACT

An important requirement for biologists conserving vulnerable species of wildlife and managing genetic problems associated with small population size is to evaluate existing evidence regarding what is known of a species' recent population history. For endemic island species in particular, current genetic impoverishment could be due to either a recent population crash or a consequence of an evolutionary history of sustained isolation and small effective population size. Interpreting any given case can often be further complicated by incomplete or contradictory evidence from historical field surveys that might suggest a very different demographic history. Here, we use the case of the Seychelles kestrel (*Falco araea*), an island endemic previously listed as critically-endangered but now relatively common, to illustrate how genetic data from microsatellite genotypes of 100–150-year-old museum specimens reveals a recent and severe population crash since the 1940s to approximately eight individuals, before the population recovered. We re-interpret the historical population trajectory of the Seychelles kestrel in the light of the minimal intervention required for this species to recover. We examine different ecological explanations for the decline and apparently unassisted recovery of the Seychelles kestrel, review the evidence for similarly unaided recoveries elsewhere and discuss the implications of unaided population recoveries for future species conservation programmes. Demographic profiles from historical genetic signatures can provide highly informative evidence when evaluating past and future recovery efforts for endangered species.

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1. Introduction

The processes by which genetic and ecological factors contribute towards the extinction of wild populations, and how these factors affect endangered species endemic to islands, remain an important focus in conservation biology. Genetic factors such as loss of genetic diversity and inbreeding are believed to play a role in the increased susceptibility to extinction of island endemics (Frankham, 1998; Frankham et al., 2002; Spielman et al., 2004). However, an important additional concern for conservation biologists working to con-

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serve small island populations is a clear understanding of the recent population history of a species. In particular, has a species maintained a stable population size from historical times to the present day or has it endured a recent population crash and, if so, what impact has this had on contemporary levels of genetic diversity? Beyond providing verification of descriptions of a species' historical abundance, answers to these questions are important for managing the future recovery of an endangered species (Matocq and Villablanca, 2001). For example, prior knowledge that a species has experienced a recent dramatic reduction in effective population size can allow managers to anticipate the problems associated with inbreeding depression that are frequently seen in restored populations of endangered species, such as lowered reproductive fitness, reduced survival and increased susceptibility to disease (Swinnerton et al., 2004; Madsen et al., 1999; Keller et al., 2002).

Fortunately, the last decade has seen the development of genetic methods to measure ancestral levels of genetic diversity that

allow comparison with contemporary levels. Early pioneering work by Bouzat et al. (1998) and Nielsen et al. (1999) applied molecular genetic techniques to historical wild populations that were relatively abundant in the past. However, more commonly, conservation biologists managing endangered species endemic to islands have available to them information on the level of genetic diversity only in the contemporary population, allowing more than one interpretation. First, endemic island species are widely believed to have proportionately lower levels of genetic diversity than species with a continental distribution and this is considered to lead to increased levels of inbreeding and associated risks of extinction in endemic island populations (Frankham, 1997, 2005). Second, contemporary measures of genetic diversity have been shown to be lower in threatened species than in their non-threatened counterparts and this pattern has recently been shown to hold true within birds, suggesting that recent endangerment has played a role (Spelman et al., 2004; Evans and Sheldon, 2008). However, an alternative interpretation is that island species may generally be more robust to the genetic effects of recent population bottlenecks because they have evolved as isolated populations and, as a consequence, carry lighter genetic loads that are less likely to impact negatively on their ability to persist (Frankham et al., 2001; Bataillon and Kirkpatrick, 2000).

This debate surrounding endemic island species has drawn recent attention (Jamieson et al., 2006; Jamieson, 2007) and has generated renewed interest in genetic diversity in endemic island species (Groombridge, 2007). As a consequence of these different interpretations, often fuelled by a few well-documented cases where wild populations have recovered from extremely severe bottlenecks apparently unaffected by problems associated with inbreeding (Ardren and Lambert, 1997; Groombridge et al., 2000), island species can still present a dilemma for conservation management; when low levels of genetic diversity are revealed in a particular island endemic, is this a consequence of a recent event or an evolutionary history of sustained isolation? Furthermore, interpreting a given case of genetic impoverishment can often be complicated by incomplete or contradictory evidence from historical field surveys that fail to clarify the past population trajectory. Here, we use genetic markers to examine the population history of the Seychelles kestrel (*Falco araea*), an island species endemic to the Seychelles archipelago in the Indian Ocean. This island endemic was described in 1966 as Critically-Endangered (Vincent, 1966) but is now relatively common (IUCN, 2007), suggesting a recent and relatively unassisted population recovery.

Historical accounts described the Seychelles kestrel as 'tolerably common' in the 1860s (Newton, 1867) and 'frequently seen on nearly all the islands' in the 1930s (Vesey-Fitzgerald, 1940), whilst two contradictory reports describe the species as 'well-distributed' and 'rare' in the 1950s (Crook, 1960; Loustau-Lalanne, 1962). By 1966, the species was considered to be Critically Endangered and close to extinction (Vincent, 1966), with probably 'less than 30 birds' confined to the main island of Mahé (Gaymer et al., 1969). However, surveys in 1973 located 49 pairs and estimated a total population on Seychelles of 150–300 birds (Feare et al., 1974; Temple, 1977), and an estimated carrying capacity of approximately 370 pairs by 1981 (Watson, 1981). An evaluation of the status of the Seychelles kestrel in 1985 described the species as being far more numerous and widespread than previously thought, possibly due to under-recording in the past (Collar and Stuart, 1985). Since then, surveys in 2001–2002 suggested that the population had remained stable over the previous 25 years (Kay et al., 2002). Together, these field records raise the intriguing possibility that the Seychelles kestrel has recovered from a severe population bottleneck relatively unaided by conservation efforts, in comparison to the recovery of the Mauritius kestrel (*Falco punctatus*), a closely-related oceanic island endemic that was successfully rescued from

the brink of extinction by 25 years of intensive conservation management (Jones et al., 1995; Groombridge et al., 2001). More broadly, instances in which small populations appear to have recovered from a population bottleneck without the aid of intensive (and often long-term and expensive) intervention warrant particular attention, because theoretical expectations predict bottlenecked populations to have reduced genetic diversity and lowered evolutionary potential and to be at risk of problems associated with inbreeding (England et al., 2003; Frankham, 1998). Therefore instances where natural populations appear to buck this trend by showing a rapid and unassisted recovery require confirmation.

Importantly, single species studies of island endemics rarely have the benefit of a natural 'control' population or one with a precisely known bottleneck history (Bouzat, 2000), particularly those species without additional populations that may have been established through conservation management, for example by sequential translocations (Taylor and Jamieson, 2008). Here, however, a 'benchmark' against which to compare the genetic and demographic history of the Seychelles kestrel is provided by the Mauritius kestrel, whose contemporary population is well-documented to have declined to just a single breeding pair in 1974 (Groombridge et al., 2001). The Mauritius kestrel population declined during the 1800s due to habitat loss and degradation and was then reduced to a few individuals as a consequence of poisoning by DDT organochlorine pesticides (Safford and Jones, 1997). In the 1980s and 1990s an intensive captive-breeding and reintroduction programme successfully restored the population to approximately 800–1000 birds (Jones et al., 1995). Previous genetic work using microsatellite markers detected a dramatic loss of genetic diversity and a marked reduction in effective population size (N_e) in the Mauritius kestrel (Groombridge et al., 2000; Nichols et al., 2001). This work described a pattern of relative N_e across several other continental and island kestrel species; among these, the estimate for the contemporary Seychelles kestrel was very similar to that for the Mauritius kestrel, prompting the question of whether the Seychelles kestrel has experienced a similarly severe population crash.

In this paper, we illustrate how historical genetic data from microsatellite genotypes of 100–150-year-old museum specimens of the Seychelles kestrel reveal a temporal change in effective population size in comparison to contemporary populations. We interpret the population trajectory of the Seychelles kestrel in the light of this new genetic perspective and we review different ecological explanations of the decline and recovery of this species. Finally, we review other potentially similar examples of unassisted population recoveries and discuss the need to evaluate population genetic histories when conserving endemic island species.

2. Methods

2.1. Sample collection

Blood and feather samples from Seychelles kestrels were obtained from the main island of Mahé (Lat: -4.68 , Long: 55.47) and Silhouette (Lat: -4.48 , Long: 55.23) between January 1998 and May 2004 (see Fig. 1). Wild kestrels were located by field observation and, using play-back recordings of territorial male kestrel calls, caught using a traditional bal-chatri (Berger and Mueller, 1959). A blood sample (of 20–50 microlitre volume) was obtained from the brachial wing vein of captured individuals using a 0.3-ml U-100 micro-fine insulin syringe (Vet UK) and whole blood was mixed with 1 ml of absolute ethanol in a screw-topped rubber sealed microfuge tube and stored at -20 °C. All caught individuals were banded using a uniquely numbered metal band and two col-

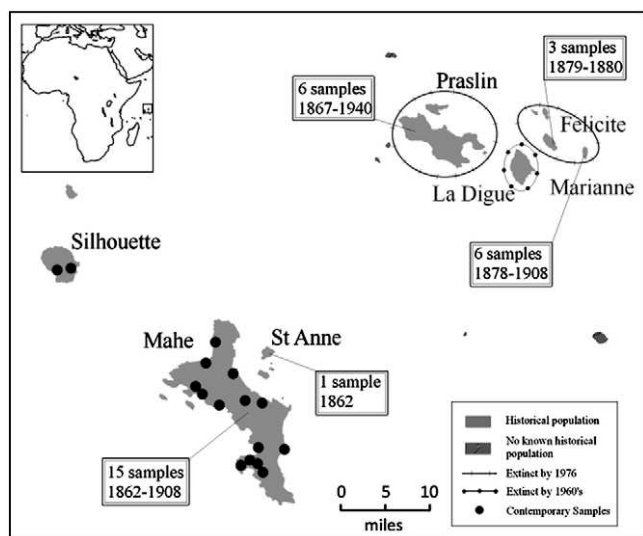


Fig. 1. Historical and contemporary range of the Seychelles kestrel, showing sampling locations for the contemporary population (circles). Boxes give the number of historical museum samples collected from each island and the range of dates for which location information was available from individual specimen labels.

oured bands to allow each bird to be individually identified in the field and to ensure no individual was sampled twice.

Samples of foot-pad tissue were obtained from all of the historical specimens of Seychelles kestrels ($n = 43$) that were available from museums worldwide. These specimens were collected in the Seychelles between ca 1840 and 1940. Previous genetic work using historical samples of the closely-related Mauritius kestrel identified foot-pad tissue samples to be the most reliable tissue type for DNA yield and for polymerase chain reaction (PCR) amplification (Groombridge et al., 2000; Nichols et al., 2001). Samples from historical specimens of Seychelles kestrels were obtained from the collections of the Natural History Museum (NHM, London, UK), the University Museum of Zoology (Cambridge, UK), the American Museum of Natural History (AMNH, New York, US), the Muséum National d'Histoire Naturelle (Paris, France) and the World Museum Liverpool. A foot-pad sample was excised from each skin using a fresh scalpel blade. Samples were stored dry at -20°C .

2.2. Microsatellite genotyping

Genomic DNA was extracted from whole blood samples of Seychelles kestrel individuals using the DNeasy Blood and Tissue Kit (Qiagen, UK). Individuals were genotyped at eleven microsatellite loci. Nine of the microsatellite loci used contained dinucleotide repeat regions and were originally isolated from the peregrine falcon (*Falco peregrinus*) by Nesje et al. (2000) (loci NVH-*fp*5, NVH-*fp*-46-1, NVH-*fp*-107, NVH-*fp*-89, NVH-*fp*-86-2, NVH-*fp*-31, NVH-*fp*-82-2, NVH-*fp*-13, NVH-*fp*-79-1; GenBank sequence accession numbers AF118420-AF118434). Two further microsatellite loci were used, respectively a hexanucleotide and tetranucleotide, both originally isolated in the European kestrel (*Falco tinnunculus*) (loci *Fpe*1, *Fpe*2; Jon Wetton unpublished data; see Dawney et al., 2009; GenBank sequence accession numbers EU518686 & EU518687).

PCR amplification of contemporary DNA was carried out in 10- μl volume reactions following 5'-end-labelling with a fluorescent dye (TET, 6-FAM or HEX) (Integrated DNA Technologies, Inc.). Each PCR reaction contained the following: approximately 50 ng of genomic DNA, 0.5 pmol of each primer, 1.5 mM MgCl_2 , 3.0 mM NH_4 buffer (Bioline, UK) and 0.75 units *Taq* DNA polymerase (Bioline, UK). PCR amplification was performed in a Tetrad thermal cycler (MJ Research). Cycling parameters were as follows: 94°C for

3 min, then 30 cycles of 94°C for 45 s, 58.5°C for 45 s, and 72°C for 45 s, with a final extension step of 72°C for 10 min. Each locus was amplified separately. The amplified products of some loci were pooled post-PCR but only those that had been fluorescently labelled with different dyes. PCR products were visualized using an ABI 3130 DNA Analyzer (Applied Biosystems, Warrington, UK). TAMRA size standards and filter prism set C was used and allele sizes were assigned using GENEMAPPER software (Applied Biosystems Inc.).

Genomic DNA was extracted from museum foot-pad samples of Seychelles kestrel individuals using the QiAmp DNA Micro Kit (Qiagen). The extraction protocol used was that supplied with the kit for DNA isolation from tissues, but with three modifications: (1) an additional digestion step was included in which after the finely chopped sample had been digested with Proteinase K during an overnight incubation at 55°C , (2) a further 20 μl of Proteinase K was added and the sample re-incubated at 55°C for a further 4 h to enhance full digestion of the sample, and (3) prior to extraction of the DNA: after each step of vortexing, samples were kept on ice instead of at room temperature as recommended in the protocol. All museum DNA extractions and PCR amplifications were carried out in a separate laboratory to that used for the modern DNA work. For PCR amplification of museum DNA for four loci (NVH-*fp*-5, NVH-*fp*-82-2, NVH-*fp*-107 and NVH-*fp*-13), the cycling parameters were adjusted such that the annealing temperature was reduced from 58.5°C to 55°C . All PCR amplifications for museum DNA samples were repeated three times. Negative DNA controls were carried out for all museum DNA samples. DNA extractions and amplification replicates were genotyped at all eleven loci in order to check for contamination from other individuals.

2.3. Statistical analysis

All loci were tested for deviation from Hardy-Weinberg equilibrium and linkage disequilibrium using GENEPOP 3.1 (Raymond and Rousset, 1995). When loci differed significantly from Hardy-Weinberg equilibrium, we applied the programme MICRO-CHECKER to identify the most likely reason for the departure (Van Oosterhout et al., 2004, 2006). Observations of heterozygote deficiency can result from the presence of null alleles, heterozygotes being mis-scored due to stutter bands or an artefact of non-amplification of the larger allele in heterozygotes (Wattier et al., 1998). Population genetic explanations can account for genuine heterozygote deficits, such as inbreeding or population structure. We calculated the mean number of alleles per locus and mean expected heterozygosity using CERVUS v3.0 (Kalinowski et al., 2007). Nei's gene diversity, allelic richness, pairwise F_{ST} values and pairwise tests for differentiation were carried out using FSTAT 2.9 (Goudet, 2001). The F -statistics F_{IS} , F_{ST} and R_{ST} were calculated using FSTAT over all loci, to partition heterozygote deficiency into its within and among population components, in order to assess genetic structure (Weir and Cockerham, 1984). F_{IS} measures the heterozygote deficit within populations and F_{ST} that among populations (i.e. the proportion of genetic variance that can be explained by differences among populations), whilst R_{ST} provides an additional measure of gene differentiation that is not dependent on the number of populations sampled (Slatkin, 1995). Given that many of the historical museum specimens have location information detailing the precise island within the Seychelles archipelago from which they were collected, we also examined the genetic data from the historical population for population substructure using the programme TESS (version 1.1; Chen et al., 2007), which integrates geographical information on sampled individuals. The spatial locations of all five islands (Fig. 1) were obtained using Google Earth (2007) and random spatial coordinates were generated within each island for each of the 31 historical specimens; the resulting neighbourhood was

modified to reflect the relative isolation of each island. TESS was run specifying a Markov Chain Monte Carlo model, an estimated number of clusters ranging from 1–7, with a burn-in period of 100,000 iterations and each model was run for 200,000 sweeps. A total of 200 runs was carried out for each model (the 20% highest likelihood runs were compared across the range of K values). Finally, we applied a Bayesian method to the full microsatellite genotype data set spanning ~164 years from ca 1840 to 2004, to detect temporal change in effective population size (N_e). This method, developed by Beaumont (2003), samples independent genealogical histories using importance sampling to estimate recent changes in N_e from temporally spaced gene frequency data and assumes that the sampling period is sufficiently short that the effects of mutations can be safely ignored. Whilst alternative Bayesian methods have been applied by Nichols et al. (2001) to estimate relative N_e for the Mauritius kestrel alongside other kestrel species, the method by Beaumont (2003) makes explicit use of historical collection dates for all of the individual museum skins, to combine historical and contemporary genetic data in a single Bayesian analysis to estimate ancestral and current N_e (N_A and N_0 , respectively, in Beaumont, 2003). Furthermore, Beaumont's (2003) method uses the likelihood based on allele frequencies, it takes into account unequal sample sizes across loci and sampling period. In implementing Beaumont's tmvp programme for the Seychelles kestrel data, we specified a mean generation time of four years and rectangular priors of 0–1000 for ancestral N_e and contemporary N_e . To explore the model's performance, we performed analyses stipulating different values of alpha, the smoothing parameter, ranging from $\alpha = 0.3$ –0.7 ($\alpha = 0.3$ in the final analysis). Beaumont (2003) previously applied this Bayesian method to existing microsatellite data for the historical and contemporary populations of the Mauritius kestrel (Groombridge et al., 2000; Nichols et al., 2001). Given the well-documented single-pair bottleneck that occurred in the Mauritius kestrel population in 1974 (Jones et al., 1995; Groombridge et al., 2001), we repeat here that temporal analysis for the Mauritius kestrel as a 'benchmark' for comparison with our temporal analysis for the Seychelles kestrel.

3. Results

Sixteen Seychelles kestrels were caught and blood-sampled between 1998 and 2004 (four of these individuals were those used for comparative purposes in an earlier population genetic study of the Mauritius kestrel (Groombridge et al., 2000; Nichols et al., 2001). Fig. 1 shows the location of each sampled bird and the current range of the Seychelles kestrel (Kay et al., 2002, 2004). A total of 43 historical museum specimens of Seychelles kestrel was sampled for DNA (see Appendix 1 for location and date of collection of each specimen). Each of the 43 museum DNA samples yielded microsatellite genotypes for between 2 and 11 loci (giving an overall success rate of 75% for successful microsatellite amplification across all samples and loci). Eighty percent of all museum DNA genotypes were repeated at least twice to check for allelic dropout.

Only two of the 11 microsatellite loci (*Fpe1* and *Fpe2*) were polymorphic in the contemporary Seychelles kestrel population, whereas five loci (*Fpe1*, *Fpe2*, *NVH-fp5*, *NVH-fp46-1* and *NVH-fp107*) were polymorphic in the historical museum samples and together contained a total of eight unique alleles not detected in the contemporary samples. None of the polymorphic loci within the contemporary population exhibited significant departure from Hardy Weinberg equilibrium. A heterozygote deficit was detected at two of the loci within the historical population. If the historical population were genetically structured, this might lead to a deficiency of heterozygotes when data from museum specimens ob-

Table 1

Comparison of measures of microsatellite genetic diversity for historical (1862–1936) and contemporary (1998–2004) populations of the Seychelles kestrel.

	Historical population	Contemporary population
Mean alleles per locus	2.180	1.450
Mean allelic richness	1.860	1.390
Mean expected heterozygosity	0.210	0.104
Nei's gene diversity	2.352	1.160

tained on different islands or from different time periods were pooled (Wahlund effect). In support of this idea, field studies of Seychelles and other Indian Ocean kestrels show strong site fidelity and poor dispersal ability, two characteristics that tend to generate structured populations (Watson, 1992; Burgess et al., 2008). The mean number of alleles per locus, allelic richness, mean expected heterozygosity and Nei's gene diversity were all higher in the historical population compared to the contemporary population (Table 1). Despite the observed reduction in allelic richness and heterozygosity, the historical and contemporary populations were not significantly differentiated (pairwise G test, $P = 0.383$). F_{IS} was higher (0.675) in the historical population than the contemporary population (0.300).

Locality information was available for 31 of the 43 historical museum specimens. They had been collected from five different Seychelles islands (Mahé [16], Praslin [6], Félicité [2], Ile Marianne [6] and St. Anne [1]; see Fig. 1), allowing measurement of genetic differentiation between islands (due to the close proximity of St. Anne and Mahé, samples from these locations were analysed together). Pairwise F_{ST} values ranged from 0 to 0.132 (tests for differences between islands were not significant). The weighted F_{ST} across all loci across the five islands was 0.104 (Rousset, 1996), and Goodman's R_{ST} was 0.151 (Goodman, 1997). When analysing the historical data using the programme TESS, the highest log-likelihood value was obtained for $K = 1$, suggesting the absence of population genetic structure. However, this finding should be treated with caution given the limitations imposed by the relatively small number of polymorphic markers used. Small sample size prevented a similar analysis for the contemporary population.

Fig. 2 gives the posterior distribution of the temporal change in historical and contemporary N_e for the Seychelles kestrel and Mauritius kestrel, as generated by the Bayesian analysis. The density of points is proportional to the probability density of population size at two different times, hence an off-diagonal distribution indicates a change in effective population size (N_e). The plot of the posterior distribution for the Mauritius kestrel (Beaumont, 2003, repeated here for comparison) provides very strong evidence of a recent decline in N_e ; the joint mode and 90% higher posterior density (HDP) limits for the marginals from the density estimation are $N_e = 957$ (390–1000) for the historical size and $N_e = 4.16$ (2.17–9.78) for the contemporary size, with a Bayes factor of >9900 in favour of recent population decline on Mauritius. In comparison, the posterior plot for the Seychelles kestrel shows a similar signature of recent reduction in effective population size, but with a joint mode of $N_e = 387$ for the historical size (90% HPD limits, 110–1000) and $N_e = 8$ for the contemporary population (90% HPD limits, 3.5–22.0), indicating a population crash that was only marginally less severe than that experienced by the Mauritius kestrel. Given that the Seychelles kestrel appears to have recovered mostly in just a portion of its ancestral range, we carried out a repeated analysis restricted to just the Mahé samples (ancestral $n = 14$) to estimate Mahé's ancestral N_e , to determine the main island's contribution to the overall ancestral N_e . As expected, this analysis produced a lower ancestral N_e of 264, approximately 68% of the total ancestral N_e .

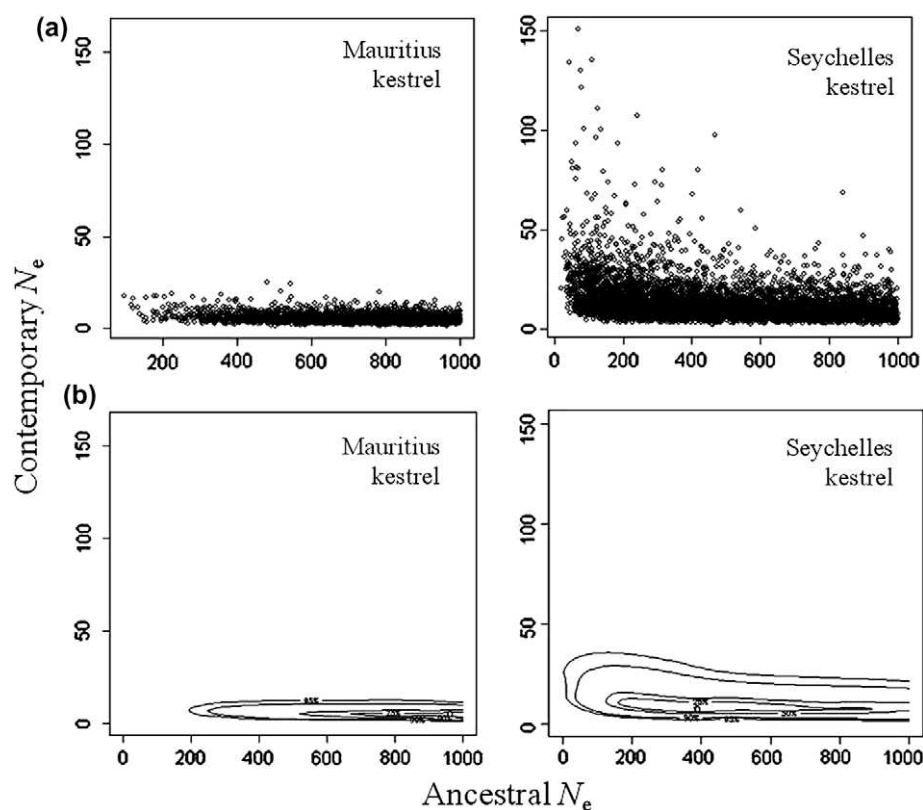


Fig. 2. Posterior distribution of the ancestral and contemporary effective population size (N_e) of the Seychelles kestrel and Mauritius kestrel, following the methods of Beaumont (2003). (a) The density of points is proportional to the probability density of population size at the two different times, hence an off-diagonal distribution indicates a change in N_e (note: the y-axis extends only to 150). (b) 25–95% higher posterior density limits of the posterior distribution. The joint mode is plotted as a single open circle.

4. Discussion

We consider the results for the museum samples to be very reliable; there was no sign of contamination, microsatellite genotype electropherogram peak heights were observed to be proportionally smaller than for equivalent contemporary genotypes from modern DNA sources (as would be expected from low copy number DNA template), replicates produced identical results (Pääbo et al., 2004), and we observed a higher number of polymorphic loci (including eight novel alleles) among the historical samples.

Our analysis of temporal genetic data for the Seychelles kestrel suggests that this species has endured a recent and severe population crash during the last 150 years to an estimated N_e of eight (3.5–22.0) individuals. A comparison of our genetic result for the Seychelles kestrel with a previous analysis of historical genotypes obtained for the Mauritius kestrel using the same suite of markers (Beaumont, 2003; Groombridge et al., 2000) suggests that the population crash experienced by the Seychelles kestrel may have approached the severity of that of the Mauritius species. Beaumont (2003) obtained a modal value of $N_e = 4$ for the contemporary Mauritius kestrel population, which is in close agreement with the lowest documented population size of four birds obtained from field surveys on Mauritius in 1974 (Jones et al., 1995; Groombridge et al., 2001). Consequently, our contemporary genetic estimate of an N_e of eight individuals adds credibility to claims made in the 1960s that the Seychelles kestrel was Critically Endangered and that the population on the main island of Mahé had declined at one point to 'less than 30 birds' (Vincent, 1966; Gaymer et al., 1969).

The genetic analysis described here is unable to determine precisely when the population crash experienced by the Seychelles kestrel occurred, but the distribution of collection dates for those individuals from which genetic data were obtained suggests that

the most severe part of the population bottleneck occurred after 1940 (the collection date of the most recent historical museum skin) and prior to the onset of intensive field surveys in 1973, which detected close to 50 pairs (Feare et al., 1974; Temple, 1977). The most recent field surveys of the Seychelles kestrel in 2001–2002 indicate that the population has remained stable since the early 1980s (Kay et al., 2002). We note that the low contemporary estimate of genetic N_e , described here as evidence of the severest extent of the population crash between the 1940s and 1970s, has been derived from the current population of Seychelles kestrels. Despite the recent recovery of the population to several hundred pairs, many future generations of mutation will be needed for the genetically estimated N_e to increase to ancestral levels.

If our genetic data for the Seychelles kestrel do indeed reflect the true historical trajectory of this island endemic, this case study provides some hope for the numerous other endemic island species whose populations are vulnerable to extinction. It would appear that the Seychelles kestrel has recovered from a severe population bottleneck with only minimal conservation efforts. The Seychelles kestrel population did not receive the level of intensive conservation management that occurred on Mauritius, where a 25-year long captive-breeding and reintroduction programme restored the Mauritius kestrel from just a handful of individuals in 1974 to over 800 birds by 2005 (Jones et al., 1995; Nicoll et al., 2004). By comparison, the Seychelles kestrel has received little active management; the species has been reintroduced by translocation to Praslin, but with limited success. In 1977, 13 kestrels were translocated from Mahé to Praslin where the species had previously been extirpated. By 1980 the Praslin population had grown to 10 pairs (Watson, 1989), but more recent surveys show the population has declined to just a handful of territories (Kay et al., 2004).

Several ecological factors may explain the unaided recovery of the Seychelles kestrel. First, in contrast to Mauritius, where widespread and intensive use of DDT organochlorines between 1949 and 1965 is known to have directly contributed to the rapid decline of the Mauritius kestrel population by reducing eggshell thickness and associated productivity (Safford and Jones, 1997), on Seychelles the use of DDT was less widespread, being confined largely to small-scale trials by the Seychelles Government to control a variety of agricultural pests and localised house-to-house spraying during the 1940s–1960s (Seychelles Department of Agriculture, 1947; Mathias, 1971). Consequently, the ecological bioaccumulation of DDT is likely to have been considerably lower across the Seychelles archipelago than on Mauritius, thereby allowing the Seychelles kestrel population to rebound quickly following the end of DDT use. Second, habitat loss on Seychelles has been less severe than on Mauritius. On Mauritius, logging over the last 200 years removed almost all of the native ebony forest (Cheke, 1987; Safford, 1993), and, whilst commercial sugarcane plantations now cover most of the island, the kestrel's intensively-managed recovery has benefited from its ability to occupy suboptimal habitats (Jones et al., 1995). By comparison, large tracts of native habitat still remain across the Seychelles. Where degradation has occurred, the natural habitat has been replaced with large invasive trees such as cinnamon (*Cinnamomum zeylanicum*) and coconut (*Cocos nucifera*), but these species have helped to maintain a forest canopy and support the gecko and lizard populations that comprise the kestrel's prey (Stoddart, 1984; Watson, 1992; Groombridge et al., 2004). Intriguingly, the contemporary Seychelles kestrel population appears to have been slow to expand back into its former historical range. However, both the Mauritius and Seychelles kestrels are known to show limited dispersal and this may go some way to explaining their current distribution (Burgess et al., 2008).

Our modal estimate of ancestral N_e for the Seychelles kestrel of 387, generated by genotyping of historical museum specimens collected between ca 1840 and 1940, is less than half the equivalent estimate for the Mauritius kestrel of 957 that was obtained by a similar analysis of historical genotypes of similarly-aged museum specimens from Mauritius using the same suite of microsatellite markers (Beaumont, 2003). One factor that may explain this result is the difference in total island area (and hence ancestral carrying capacity) between Seychelles (243 km²) and Mauritius (1650 km²), suggesting a markedly lower ancestral carrying capacity on Seychelles.

It should be noted that the estimation of the mode of the ancestral N_e is somewhat uncertain and depends both on the density estimation procedure and the number of points sampled in the Monte Carlo Markov Chain. In particular, for the Mauritius kestrel the estimated mode for ancestral N_e is close to the limit set by the prior, and is probably indistinguishable from this limit. It is likely in this case that if the prior limit were increased, then the modal N_e would be correspondingly higher. In contrast, in the case of the Seychelles kestrel it appears, for example, that the 25% HPD limit is well away from the upper limit of the prior, and thus the estimate of the mode is likely to be robust to any increase in the prior bounds (M. Beaumont, pers comm.). Our estimate of an ancestral N_e of 264 on Mahé alone provides further support for such relative differences in N_e in line with geographical area. The ancestral N_e estimate for Mahé accounts for 68% of the total ancestral N_e , which is broadly in line with island size; Mahé (144 km²) accounts for 64% of the total 224 km² area of those Seychelles islands believed to have formed the Seychelles kestrel's ancestral range.

As molecular genetic techniques become increasingly commonplace as an integral part of endangered species recovery programmes, there are more opportunities to compare evidence from demographic histories with temporal genetic data for other endan-

gered island species (Shepard and Lambert, 2008). The picture emerging from these studies suggests that island populations can largely be categorized into those that have experienced ancient bottlenecks that caused prehistoric loss of genetic variation, such as those due to founding events or prehistoric changes in the environment, and those that have experienced recent bottlenecks, such as population crashes that have taken place over the last few hundred years. For example, Paxinos et al. (2002) showed a prehistoric decline in genetic variation in the endangered Hawaiian goose (*Branta sandvicensis*) that was concurrent with growth in the prehistoric human population on Hawaii. In contrast, the picture is less obvious for the South Island saddleback (*Philesturnus carunculatus*) from New Zealand, for which genetic data from historical museum specimens suggest that low contemporary levels of genetic variation in the Big South Cape Island population appear not to be a consequence of a severe population crash caused by rat (*Rattus* spp.) infestations on the island in 1962. Instead, the low genetic variation is believed to be the consequence of a natural founding event (Taylor et al., 2007). Without this historical genetic information, managers working to conserve this saddleback population could have been misled by considering the ecological evidence alone.

Our genetic data provide evidence that the Seychelles kestrel has experienced a severe population crash within the last 70 years, in agreement with field surveys in the 1960s that indicated the species was 'critically endangered', and yet this island endemic appears to have recovered to a comparatively large population size in the absence of intensive conservation intervention. Similar examples of unaided species recoveries are extremely rare and many seemingly comparable 'unmanaged' species recoveries can often be explained by indirect intervention, such as broad-scale ecosystem management. For example, the endemic Rodrigues fody (*Foudia flavicans*) in the Mascarene Islands achieved a near 100-fold increase in population size from fewer than 20 individuals in 1968 to a population of over 900 by 1999 in the absence of taxon-specific management (Impey et al., 2002), because the population had responded well to over a decade of extensive reforestation. Elsewhere, populations of species previously thought to be extinct have "reappeared", for example the Bermudan petrel (*Pterodroma cahow*) and the New Zealand takahe (*Porphyrio hochstetteri*; Lee and Jamieson, 2001). The survival and subsequent growth of these populations has largely resulted from many years of ecosystem management, often in parallel with intensive captive-breeding and reintroduction programmes. In contrast, the Seychelles kestrel appears to represent an unusual example of a relatively unassisted recovery from a small population size by an island endemic despite extensive loss of genetic diversity.

Of the 794 so-called 'trigger species' identified by Ricketts et al. (2005) to be at imminent risk of extinction unless immediate and direct conservation action is taken, 39% are island species. Given the current pressure on global conservation resources, the undetected recovery of the Seychelles kestrel from a severe population crash may appear encouraging. However, outcomes such as this are likely to be the exception rather than the rule for other endangered island endemics.

Acknowledgements

This research was funded by a grant from the Royal Society to JG with further support by a Faculty Award from the University of Kent. The molecular genetic work was performed at the UK Natural Environmental Research Council's Molecular Genetics Facility. The research and associated fieldwork was approved by the Seychelles Bureau of Standards. Samples for the research were transported under CITES permits issued by the Conservation and National Parks Division, Department of Environment, Government of Seychelles. A. Huggins assisted with contemporary sample col-

lection. Fieldwork and logistical support was also provided by staff at Nature Seychelles, including R. Bristol, U. Bristol, and T. Vel. Additional contemporary samples were also provided by Nature Seychelles and the Nature Protection Trust of the Seychelles. Museum samples were kindly made available by the following collections: Natural History Museum, Tring; University Museum of Zoology, Cambridge; World Museum Liverpool; Museum National d'Histoire Naturelle, Paris; American Museum of Natural History, New York. We thank A. Frantz and M. Beaumont for discussions of the manuscript.

Appendix A

Museum	Catalogue #	Collector	Year	Location
1	Paris	MNHN-457	No data	–
2	NHM	1955.6.N20.1390	Bishop of Mauritius	–
3	NHM	1955.6.N20.1391	Bishop of Mauritius	–
4	NHM	1927.12.18.73	Lister	–
5	NHM	1946.75 ex dup	No data	–
6	NHM	[18]94.6.16.408	Ward	– Mahé
7	Cambridge	CAM-13/Fal/1	Newton	1862 Mahé
8	NHM	[18]74.5.13.2	Newton	1863 –
9	Cambridge	CAM-13/Fal/6	Brooks	1866 –
10	Cambridge	CAM-13/Fal/2	Wright	1867 –
11	Cambridge	CAM-13/Fal/4	Newton	1867 Mahé
12	Cambridge	CAM-13/Fal/7	Newton	1867 Mahé
13	NHM	1955.6.N20.1387	Newton	1867 Mahé
14	Cambridge	CAM-13/Fal/3	Newton	1867 Praslin
15	Cambridge	CAM-13/Fal/8	Newton	1867 Praslin
16	NHM	1955.6.N20.1386	W. A. Kennedy	1867 St. Anne
17	Liverpool	LIVER-T.1060	Wright	1861 –
18	Cambridge	CAM-13/Fal/5	Ward	1869 –
19	Paris	MNHN-496	Lantz	1878 Ile Marianne
20	Paris	MNHN-501	Lantz	1878 Mahé
21	Paris	MNHN-502	Lantz	1878 Mahé
22	Paris	MNHN-498	Lantz	1878 Praslin
23	Paris	MNHN-499	Lantz	1878 Praslin
24	Liverpool	LIVER-T.3979	Warry	1879 Felicite
25	NHM	1955.6.N.20.1388	Warry	1879 Felicite
26	NHM	1955.6.N20.1389	Warry	1880 Mahé
27	NHM	1843.11.17.28	No data	ca –
28	NHM	1927.12.18.72	Lister	1888 Mahé
29	AMNH	AMNH-538713	Mortimer	1903 Mahé
30	AMNH	AMNH-538714	Mortimer	1903 Mahé
31	AMNH	AMNH-538715	Thibault	1904 Ile Marianne
32	AMNH	AMNH-538716	Thibault	1904 Ile Marianne
33	AMNH	AMNH-538717	Thibault	1904 Ile Marianne
34	AMNH	AMNH-538718	Thibault	1904 Ile Marianne
35	AMNH	AMNH-538719	Thibault	1904 Ile Marianne
36	AMNH	AMNH-538710	Thibault	1904 Mahé
37	AMNH	AMNH-538711	Thibault	1904 Mahé
38	AMNH	AMNH-538712	Thibault	1904 Mahé
39	AMNH	AMNH-538720	Thibault	1905 Praslin
40	NHM	1906.12.21.185	Nicoll	1906 Mahé
41	NHM	1906.12.21.184	Nicoll	1906 Mahé
42	NHM	1965.M.1565	Meinertzhagen	1910 –
43	NHM	1946.75.11	Sapsworth & Goodfellow	1940 Praslin

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