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SHORT COMMUNICATION

No phylogeographic structure in the circumpolar snowy owl (Bubo scandiacus)

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Abstract The snowy owl (Bubo scandiacus) is a nomadic species with a circumpolar distribution. It has recently declined in the western Palearctic and may thus be worthy of special consideration for conservation. We investigated genetic structure in three well separated geographic regions within the snowy owls' breeding range. We sequenced two mitochondrial genes; the control region and cytochrome b, and two Z-chromosome introns; VLDLR-9 and BRM-15. We found no phylogeographic structure among the sampled regions, indicating high levels of gene flow in the recent past and possibly still today. Intra-population diversity did not vary between regions for the control region, but for Cyt b, North American birds had higher haplotype diversity than Scandinavian and eastern Siberian birds. Western Palearctic birds do not seem to be genetically deprived or inbred. Genetic diversity in the snowy owl was not lower than Scandinavian populations of three other owl species: tawny owls (Strix aluco), Tengmalm's owls (Aegolius funereus) and eagle owls (Bubo bubo).

Keywords Genetic structure · Gene flow · mtDNA control region · Phylogeography · Population genetics · Z-chromosome intron

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Introduction

The snowy owl has a circumpolar distribution and breeds mainly on the arctic tundra (Cramp and Simmons 1994). It is a food specialist, preying on rodents with fluctuating population cycles, such as lemmings (Lemmus and Dicrostonyx spp.) and voles (Microtus and Clethrionomys spp.). Consequently, snowy owls migrate to places where there are peak densities of these rodents, and thus show large fluctuations in local breeding populations (Alerstam 1990; Cramp and Simmons 1994). Snowy owls have declined in numbers in the western Palearctic in the 20th century due to illegal hunting and possibly reduction in rodent densities (Portenko 1972; Solheim 1994, 2004). In Norway, the snowy owl is listed as Vulnerable (Kålås et al. 2006), in Finland as Endangered (Rassi et al. 2001), and in Sweden as Critically Endangered (Gärdenfors 2005). World-wide, the species is not threatened (IUCN 2006), and in North America there are no indications of reduction in population sizes (del Hoyo et al. 1996). The global population of snowy owls today is estimated to about 290,000 individuals, and is reported to be stable (BirdLife International 2004).

The snowy owls capacity for long distance dispersal was demonstrated by Fuller et al. (2003), who tracked post-breeding movements of six adult snowy owls with satellite telemetry techniques. Maximum travel distance found was about 1,300 km during 11 days. The tracked birds did not remain in the nesting area after the breeding season, and they did not return to the same nesting area the next year. The potential for movement thus appears to be considerable in the snowy owl. One would therefore expect little or no barriers to gene flow in this species and a rather homogeneous population genetic structure across the breeding range.



Here we present the first analysis of the population genetic structure in snowy owls, using samples from three widely separated regions: Scandinavia, eastern Siberia and North America. We investigated whether there is genetic differentiation between these areas, and whether snowy owls in western Palearctic are less diverse than snowy owls in other areas. In addition, we compared the level of genetic diversity in snowy owls with three other owl species breeding in Scandinavia.

Methods

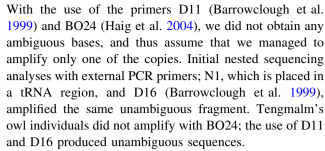
Samples and DNA extraction

Blood or tissue samples were taken from 14 snowy owl individuals from Scandinavia, 14 individuals from North America, and 12 individuals from eastern Siberia. Sample details are given in the Appendix. Most sampled birds were unrelated chicks or assumed breeding birds (i.e. at age 2Y+ and caught in the period May-August at breeding grounds). Exceptions were five Norwegian birds caught outside the breeding range or in a year of no recorded breeding, and the samples from North America, which were from wintering areas. We assume the latter ones belong to the breeding population in northern North America, an assumption supported by telemetry studies from Massachusetts where snowy owls caught during winter were tracked to Maine and Quebec during spring, apparently on their ways to breeding grounds (Smith 2005). Blood samples from 10 adult tawny owls (Strix aluco) and 10 adult Tengmalm's owls (Aegolius funereus), and down samples from 10 eagle owl chicks (Bubo bubo), were collected in Scandinavia (Appendix). Of these species, the eagle owl is the closest relative to the snowy owl, as the two species are now considered congeneric (Wink and Heidrich 2000; Sangster et al. 2004).

DNA was extracted from blood samples using QIA-amp® DNA Mini kit (QIAGEN), and from tissue and down samples using DNeasy® Tissue Kit (QIAGEN).

Sequencing

From the mitochondrial DNA (mtDNA) we sequenced a part of the control region corresponding to domain I and part of domain II (Marshall and Baker 1997), and the second half of the cytochrome *b* (Cyt *b*) gene. For the control region in the snowy owls, initial analyses with internal primers produced two sequences for each individual, differing in three sites. The occurrence of two copies of the control region in the mtDNA is known from other bird species, among them tawny owl (Brito 2005). We therefore assume the same is the case for snowy owls.



The 5' end of Cyt b was sequenced using modified versions of Sorenson et al.'s (1999) primer L15560 (5'-GCGACA AAATCCCATTCCACCC for snowy owl and eagle owl, 5'-GYGAYAARATCCCATTCCACCC for Tengmalm's owl and tawny owl), and H15646 (Sorenson et al. 1999).

We also sequenced the two introns BRM-15 and VLDLR-9, which are found on the Z-chromosome in a passerine bird (Hansson et al. 2005). In birds, Z-chromosomes are found in two copies in males, and one copy in females. In our study, individuals heterozygous for these loci were all males, and we thus assume the loci are located on the Z-chromosome in the analyzed species. We used the primers published by Hansson et al. (2005). Of the Z-introns, only VLDLR-9 displayed more than one polymorphic site; Tengmalm's owls were polymorphic at two sites (Table 1). No Tengmalm's owl males were polymorphic at both sites, and there were thus no ambiguities in any male genotype. For females of tawny owl and Tengmalm's owl, we obtained two different sequences for BRM-15, indicating that the primers also amplified a second sequence, most likely on the W-chromosome. We therefore used a specific internal forward primer (5'-AGTGTTTSAACTCTCCCTG GT) for these species and obtained single sequences for the second half of the original sequence. The Z-introns did not amplify on the eagle owl samples, probably due to low concentration of DNA extracted from the down samples.

PCR reactions were performed according to Wennerberg (2001). The cycle-sequencing reactions were carried out using the ABI PRISM BigDye Terminator v1.1 Cycle Sequencing Kit, and run on an ABI PRISM 3100 Genetic Analyzer following the manufacturer's instructions (Applied Biosystems). The sequences were aligned in SEQUENCHER 4.1.4 (Gene Codes Corporation) and edited in BIOEDIT 5.0.9 (Hall 1999). All sequences are deposited in GenBank (Accession numbers: EU410971–EU411039, EU436175–EU436319; Appendix).

Data analyses

Number of haplotypes, haplotype diversity and nucleotide diversity were calculated in ARLEQUIN 3.01. Allelic (haplotype) richness was estimated by rarefaction according to El Mousadik and Petit (1996) using the program Contrib 1.02 (Petit et al. 1998). We used a rarefaction size of 10 for the snowy owl regions and eight for the Scandinavian



Table 1 Characteristics of two mtDNA genes and two nuclear introns in four owl species

Species	Population Control region	Control	region							-	Cytochrome b	ne b							
		n chromo- somes	# Bases	n # Poly- chromo- Bases morphic somes sites		# Haplo- Allelic types richness ^a	Haplo- type diversity	SE I	Nucleotide diversity	SE	n chromo- somes	Bases 1	Poly- morphic sites	# Haplo- Allelic types richness	Allelic richness ^a	Haplo- type diversity	SE Nucleotide diversity	otide SE	
Snowy owl	All Eastern Siberia	40	510 510	37	33 11	8.32	66:0	±0.001 (±0.012 (0.012	±0.0010 40 ±0.0019 12		415	3	3	1.83	0.60	±0.009 0.0026 ±0.032 0.0027		±0.00032 ±0.00064
	es	14	510	23	12	8.01	86.0	±0.009 0.012	0.012	±0.0018	14	415	4	5	3.35	0.73	±0.027 0.0028		±0.00056
	Scandinavia 14	14	510	28	14	9.00	1.00	±0.007 (0.013	± 0.0019	14	415	2	2	1.00	0.53	$\pm 0.017 0.0025$		± 0.00053
Eagle owl	Scandinavia 10	10	512	11	2	1.00	0.56	±0.024 (0.012	± 0.0022	10	415 (0	1	0	0	0		
Tawny owl	Scandinavia 10	10	511	7	4	2.96	0.82	±0.022 (900.0	± 0.0012	10	415 (0	1	0	0	0		
Tengmalm's owl	Scandinavia	6	476	21	∞	6.22	0.97	±0.021 (0.014	±0.0028	10	415	8	9	4.98	0.91	±0.026 0.0041		±0.00097
Species	Population	BRM-15	i-15								VLDLR-9	6-							
		n n ind o	n n ind chromosomes ^b	# Bases	# Poly- # Bases morphic t	# Haplo- Haplo- types type diversi	Haplo- type diversity	SE	Nucleotide diversity	SE	n n ind chr	n # chromo- B somes ^b	ases	Poly-# morphic ty	# Haplo- Haplo- types type diversit	>	SE Nucleotide diversity	tide SE	
Snowy owl	All	40	63	343	1 2	2 0.	60.0	∓0.008	0.00027	±0.00009	40 62	3	354 (0 1	9	0	0		
	Eastern Siberia 12		20	343	1	1 0			0		12 20		354 (1	9	0	0		
	North America	4	23	343	1 2	2 0.	0.17	± 0.026	0.00048	± 0.00020 14	14 23	æ	354 (0 1	0	0	0		
	Scandinavia	41	20	343	1 2	2 0.	0.10	± 0.024	0.00029	± 0.00016	13 19	ж	354 0	1	2	0	0		
Eagle owl	Scandinavia		ı	I	1	ı			I		I	ı	'	1		1	I		
Tawny owl	Scandinavia	10	13	215	0 1	0 1			0		10 13	æ	355 1	2		0.385	$\pm 0.042 0.00213$		± 0.00060
Tengmalm's owl	Tengmalm's Scandinavia owl	10	14	208	0 1	1 0			0		10 14	<i>a</i>)	351 2	3		0.560	±0.040 0.00215		090000∓

^a Rarefaction size for allelic richness calculations was 10 for snowy owl regions and 8 for Scandinavian populations of four owl species. Scandinavian snowy owls had an allelic richness of 7.00 for the control region, and 1.00 for Cyt b, with a rarefaction size of 8

^b Number of chromosomes equals number of females (one chromosome) plus two times the number of males (two chromosomes)



populations of four owl species. Differences in nucleotide and haplotype diversity between regions and between Scandinavian populations of all four species were tested with *t*-tests, by hand, using mean and standard deviation values from ARLEQUIN (Table 1).

Median joining haplotype networks were constructed in NETWORK 4.1.1.2 (Fluxus Technology Ltd.).

We calculated both $F_{\rm ST}$ (Weir and Cockerham 1984) and $\Phi_{\rm ST}$ (Excoffier et al. 1992) values among the snowy owl regions in arlequin 3.01 (Excoffier et al. 2005). For $\Phi_{\rm ST}$ calculations we used Tamura and Nei's (1993) nucleotide substitution model for all loci, as this gave the lowest log likelihood scores in model available in arlequin. Tests for significance were performed with 3.000 permutations.

To test hypotheses about the phylogeographic pattern based on the control region in snowy owls statistically, we performed an automatic Nested Clade Phylogeographic Analysis (Templeton et al. 2005) using the program ANeCPA (Panchal 2007). This program incorporates TCS v1.18 (Clement et al. 2000) and GeoDis v2.2 (Posada et al. 2000).

We used the Bayesian version of LAMARC 2.1.2 (Kuhner 2006; Kuhner and Smith 2007), which assumes migration/ drift equilibrium, to calculate relative estimates of gene flow between snowy owl regions (M) and population growth rates (G) based on the control region sequences. In addition, we estimated long-term effective population size for snowy owls from estimates of theta (= $2N_e\mu$, where N_e is the effective female population size and μ is the mutation rate per sequence per generation) based on the control region and Cyt b sequences. The obtained estimates of long-term effective population sizes are uncertain for several reasons. Divergence rate for the control region is unknown for Strigidae, and furthermore, the fact that divergence rates appear to be higher on short than on longer time scales obscure such calculations (Ho and Larson 2006). We used the minimum and maximum divergence rates found in other bird species for the control region; 4% and 14% per Myr (Wenink et al. 1996; Drovetski 2003). For Cyt b we used a rate of 1% per Myr (Krajewski and King 1996). We calculated μ as [divergence rate/2/10⁶ * number of base pairs * generation time (in years)]. A generation time of 4 years (the birds probably do not breed until at least two-years old, Portenko 1972) and our sequence lengths (Table 1) correspond to mutation rates of $4.08 * 10^{-5}$ mutations per sequence per generation (4% divergence rate) and $1.43 * 10^{-4}$ (14%) divergence rate) for the control region, and $8.3 * 10^{-6}$ for Cyt b (1% divergence rate).

We ran three replicates for each run in LAMARC, with a burnin of 1,000 steps, followed by one initial chain of 10,000 steps; sampling 500 trees every 20 steps, and one final chain of 200,000; sampling 10,000 trees per 20 steps. We performed three runs and used the median values for the final estimates.

Results

Marker characteristics

We amplified a 510 base pair long fragment of the control region for snowy owl, in which 37 sites were polymorphic. In eagle owl, 11 of 512 sites were polymorphic, in tawny owl. 7 of 511 sites were polymorphic, and in Tengmalm's owl, 21 of 276 sites were polymorphic. All variable sites represented transitions except one A/C transversion in snowy owl. The different primer sets used for snowy owl, as described in Sect. 'Methods', amplified identical sequences. All individuals of each investigated species were similar in the conserved F block (as defined in Baker and Marshall 1997), which occurs at the end of the fragments used in the analyses. The investigated species differed by 3.3-20% in the F block, which is in the lower range of differences between more distantly related bird species (e.g. 10-25%, Baker and Marshall 1997; 7-59%, Ruokonen and Kvist 2002).

Base composition in the snowy owl control region was similar to other investigated bird species (Baker and Marshall 1997; Ruokonen and Kvist 2002), except for an unexpected high ratio of A compared to C (33.9% A, 28.6% C, 11.4% G and 26.1% T). All variable sites were found within domain I in snowy owl, eagle owl and tawny owl. Tengmalm's owl had one variable site within the start of domain II.

From the 5' end of Cyt b, a 415 base pair fragment was amplified for all species. The sequences did not contain stop codons or indels. Snowy owl and Tengmalm's owl had five variable sites, all transitions in third codon position. There was no amino acid variation within the two species. Tawny owl and eagle owl were monomorphic. Snowy owl differed from eagle owl by nine (6.6%) amino acid substitutions, from tawny owl by 13 (9.4%) and from Tengmalm's owl by 18 (13.1%) amino acid substitutions, which is what we expect for species within the same order (Haring et al. 2001).

Based on these characteristics and comparisons, we are confident that the mitochondrial markers were of truly mitochondrial and not nuclear origin, and that the amplified fragments in the different species were homologous.

There were 33 control region haplotypes, six Cyt *b* haplotypes, two BRM-15 haplotypes and one VLDLR-9 haplotype among the 40 snowy owl individuals (Table 1).

Genetic structure

Median joining haplotype networks given in Fig. 1 for snowy owls for the three variable loci (the control region, Cyt *b* and BRM-15) revealed no phylogeographic structure. The nested clade analysis confirmed the result of no



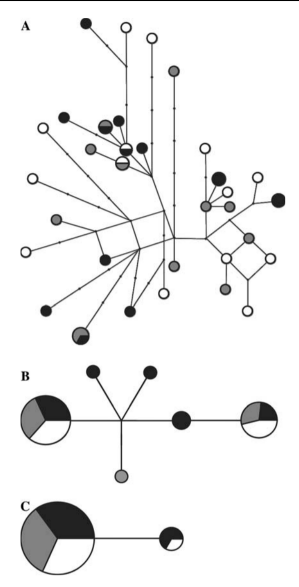


Fig. 1 Median joining haplotype networks for snowy owls revealing low levels of phylogeographic structure among three sampled regions (black = North America, grey = eastern Siberia and white = Scandinavia). (a) The mitochondrial DNA control region, (b) cytochrome b and (c) the Z-chromosome intron BRM-15. Sizes of pies indicate relative number of individuals with that haplotype. Black points on the branches in (a) represent hypothetical, non-sampled haplotypes. All lines in (b) and (c) represent single mutations

phylogeographic structure; there was no significant association between haplotype and geographic location in any clade, in any of altogether six nesting levels (all P-values > 0.20). The null hypothesis of no genetic structure could not be rejected, and there was no evidence of historical fragmentation, range expansion, long-range colonization or restricted gene flow.

Also F-statistics revealed low levels of phylogeographic structure. $F_{\rm ST}$ and $\Phi_{\rm ST}$ values for all loci comparing the snowy owl regions were similar; only $F_{\rm ST}$ values are given. Overall $F_{\rm ST}$ values were low and non-significant (-0.001

for the control region, -0.033 for Cyt b and 0.002 for BRM-15). Pairwise $F_{\rm ST}$ values were also non-significant, and all were negative except for the values comparing North America with eastern Siberia for two loci ($F_{\rm ST}=0.020$ for the control region, 0.077 for BRM-15).

Gene flow and long term effective population size

Gene flow estimated by LAMARC varied between runs, but was consistently lowest across the Atlantic Ocean. Median values of the gene flow parameter M was 853 (Credibility Interval: 279–1,014) from Scandinavia to eastern Siberia, 361 (24–1,048) from eastern Siberia to Scandinavia, 218 (27–483) from Scandinavia to North America, 166 (20–360) from North America to Scandinavia, 732 (246–1016) from North America to eastern Siberia, and 433 (119–944) from eastern Siberia to North America. One should however bear in mind that estimates of migration rates from LAMARC may not be accurate if population genetic structure is weak (LAMARC documentation), as is the case in this study.

These high gene flow estimates indicate that snowy owl constituted one panmictic population in the recent past and possibly still today (see Discussion). Therefore, we estimated the long term effective population size for all samples pooled. Theta was estimated to 0.332 (CI: 0.137–1.147), corresponding to a long-term effective population size of about 4,100 (1,700–14,100) females with 4% divergence rate, and 1,200 (500–4,000) females with 14% divergence rate. Based on Cyt *b*, theta was estimated to 0.0044 (0.0011–0.022), corresponding to a long-term effective population size of about 270 (70–1,330) females. The high diversity of estimates highlights the uncertainty of these calculations.

Genetic diversity and population growth

There was no difference between the snowy owl sampling regions in nucleotide diversity for either mitochondrial marker, nor in haplotype diversity for the control region (all tests non-significant) (Table 1). North America had significantly higher Cyt b haplotype diversity than eastern Siberia and Scandinavia (t = 3.40, df = 24, P < 0.01 and t = 6.3, df = 26, P < 0.01, respectively). Allelic richness was highest for Scandinavia for the control region, whereas North America had higher allelic richness for Cyt b (Table 1).

The Scandinavian snowy owl population had higher control region nucleotide and haplotype diversity than tawny owl (t = 2.78, df = 22, P < 0.01, and t = 8.71, df = 22, P < 0.01, respectively), and higher control region haplotype diversity than eagle owl (t = 20.33, df = 22, P < 0.01). In Cyt b, eagle owl and tawny owl were not polymorphic. Tengmalm's owl did not differ significantly



from snowy owl in nucleotide or haplotype diversities for mitochondrial loci (t = 0.30, df = 21, P > 0.05, and t = 1.57, df = 21, P > 0.05, respectively). Allelic richness showed the same pattern; Tengmalm's owl and snowy owl had the highest values (Table 1). For the Z-introns, all species showed too little variation for any comparisons to be relevant (Table 1). In summary, snowy owls from the different regions did not differ much in genetic diversity, and the snowy owl seemed to be more diverse than eagle owl and tawny owl, but at the same level as Tengmalm's owl.

The growth rate parameter (G) estimated based on control region sequences for all snowy owl individuals pooled was positive and relatively large (630, CI: 357–972), which is a strong indication of population growth (LAMARC documentation: http://evolution.gs.washington.edu/lamarc/). Of the Scandinavian populations of four species, snowy owl and Tengmalm's owl showed indications of population growth (G = 913, CI: 413–1,000 and G = 592, CI: 197–968, respectively), whereas tawny owl and eagle owl showed indications of a stable population, and population decline, respectively (G = 51, CI: -455-911, for tawny owl, G = -212, CI: -487-142, for eagle owl).

Discussion

Analyses of two mitochondrial and two nuclear loci provided no evidence of phylogeographic structure among snowy owls from Scandinavia, eastern Siberia and North America as shown by minimum spanning networks, low $F_{\rm ST}$ values and a nested clade analysis. Furthermore, no reduction in genetic diversity was revealed in Scandinavia compared to the other two investigated regions. Snowy owls in western Palearctic, where the species has declined the last 100 years, do thus not seem to suffer from inbreeding or reduced genetic diversity.

LAMARC analyses indicated high levels of gene flow, especially across the Palearctic and the Bering Strait. This is not surprising, given the snowy owls high potential for long distance flights (Fuller et al. 2003), and the Atlantic Ocean probably being the greatest barrier to gene flow. It is however difficult to assess whether gene flow occurs at present, or whether the estimated levels mirror historical events. The LAMARC analyses indicated that the snowy owl regions have been exchanging individuals at least recently.

Our inclusion of samples from non-breeding individuals (Appendix) could potentially lead to erroneous results since these may have been breeding in other areas than assumed. However, analyses excluding samples from non-breeding birds in Scandinavia did not change the results (not shown), and it seems unlikely that the North-American samples did not belong to the Nearctic breeding population

(see Smith 2005). The fact that North American samples were from wintering birds, and so may represent a more widespread sample of the breeding population than the Palearctic samples, may explain why the North American population had the highest Cyt *b* haplotype diversity. However, this hypothesis assumes some level of genetic structure, which we did not detect in our study.

The long-term effective population size estimates were low; the maximum estimate was 14,000 snowy owl females (upper credible interval limit for 4% divergence rate for the control region dataset). Although this number excludes males, juveniles and not-reproducing adults, this is far lower than today's census size of 290,000 individuals. However, the result corresponds well with the population expansion indicated in the LAMARC analyses.

Genetic analyses of other species with Holarctic distributions have also revealed recent population expansions. Reindeer (Rangifer rangifer) and Arctic fox (Alopex lagopus), particularly the 'lemming ecotype', have both been hypothesised to have been restricted to several refugia during the last interglacial period, when their tundra habitat was restricted (Flagstad and Røed 2003; Dalén et al. 2005). When the last glacial period started, their habitat increased, and the populations expanded and spread, creating a pattern of low genetic structure and relatively high levels of genetic diversity seen today. The haplotype networks of both of these species lack, similar to the snowy owl control region network, a central haplotype, and have several clades (Flagstad and Røed 2003; Dalén et al. 2005). The history of the three species may thus be parallel. Particularly the lemming-ecotype of Arctic fox resembles the snowy owl in habitat and prey choice. The snowy owls higher potential for dispersal through flights may have taken the mixing process farther than for Arctic fox and reindeer, and our results of no phylogeographic structure support this hypothesis.

Despite the restricted number and spread of samples for the other investigated owl species, the differences between species in levels of genetic diversity fit well with differences in phylogeographic history and behaviour. Tawny owl and eagle owl, which had the lowest diversity levels, and low or negative growth parameter estimates, are both relatively resident (Cramp and Simmons 1994). In addition, the sampled populations probably originate from a range expansion from a Pleistocene refugium in the Balkan region, as was suggested for Norwegian tawny owls (Brito 2005). Founder events during the expansion would have lead to lower genetic diversity. The diversity level in Tengmalm's owl, on the other hand, was about the same as in snowy owl. Tengmalm's owls have, like snowy owls, a nomadic behaviour due to the cyclic abundance of their prey (Mysterud 1970; Cramp and Simmons 1994). In contrast to snowy owl, however, Tengmalm's owls in



North America and Eurasia are genetically differentiated (Koopman et al. 2005). The North Pacific may function as a larger barrier to gene flow between the continents in this species, than does the Bering Strait for the snowy owl.

In conclusion, our results indicate no phylogeographic structure across the entire circumpolar breeding range of the snowy owl. The decline in breeding numbers seen in western Palearctic is not associated with reduced genetic diversity. Given the species' potential for long-distance breeding dispersal and lack of phylogeographic structure, it

seems as if snowy owls can be considered as one global panmictic population from a genetic perspective.

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Appendix: Information on samples from four owl species

Species	Region	Locality	Source ^a	Voucher	Sex ^c	Age	Date	GenBank a	accession no.		
				no.b				Control region	Cyt b	BRM-15	VLDLR-9
Snowy owl	Scandinavia	Värmland, Sweden	NRM	966087	F	-	1 February 1996	EU411003	EU436207	EU436255	
		Värmland, Sweden	NRM	976241	M	-	1 May 1997	EU411004	EU436208	EU436256, EU436257	
		Dalarne, Sweden	NRM	996648	M	-	1 November 1999	EU411005	EU436209	EU436258, EU436259	
		Västernorrland, Sweden	NRM	20006286	F	-	1 June 2000	EU411006	EU436210	EU436260	
		Norrbotten, Sweden	NRM	20006304	M	-	1 July 2000	EU411007	EU436211	EU436261, EU436262	
		Finnmark, Norway	TM	TM6	F	2Y	17 April 2000	EU411010	EU436214	EU436265	
		Finnmark, Norway	TM	TM117	F	3Y+	14 July 2003	EU411009	EU436213	EU436264	
		Finnmark, Norway	ANM	1100 (17811) ^d	M	2Y	July 2001	EU410983	EU436187	EU436246, EU436247	
		Captive, Norway	ANM	1619 (17812) ^d	F	_	2003	EU410984	EU436188	EU436248	
		Norway	ANM	1085 (17813) ^d	M	1Y	1997	EU410985	EU436189	EU436249, EU436250	
		Hedmark, Norway	ANM	1101 (<i>17814</i>) ^d	F	_	2000	EU410986	EU436190	EU436251	EU436291
		Finnmark, Norway	ANM	1572 (<i>17815</i>) ^d	F	_	1 July 2000	EU411008	EU436212	EU436263	
		Lapland, Finland	ZUO	31500	M	_	14 June 2000	EU411001	EU436205	EU436252, EU436253	
		Lapland, Finland	ZUO	970509	-	=	9 May 1997	EU411002	EU436206	EU436254	
Snowy owl	Eastern Siberia	Lopatka peninsula	SRE	-	M	Pullus	15 July 1994	EU410971	EU436175	EU436226, EU436227	
		Wrangel Island	SRE	_	M	Pullus	25 July 1994	EU410972	EU436176	EU436228, EU436229	
		Wrangel Island	SRE	=	F	Pullus	25 July 1994	EU410973	EU436177	EU436230	
		Wrangel Island	SRE	=	M	Pullus	25 July 1994	EU410974	EU436178	EU436231, EU436232	
		Wrangel Island	SRE	_	M	Pullus	25 July 1994	EU410975	EU436179	EU436233, EU436234	
		Wrangel Island	SRE	_	M	Pullus	25 July 1994	EU410976	EU436180	EU436235, EU436236	
		Wrangel Island	SRE	_	M	Pullus	25 July 1994	EU410977	EU436181	EU436237, EU436238	



Species	Region	Locality	Source ^a	a Voucher no.b	Sexc	Age	Date	GenBank a	ccession no.		
				no."				Control region	Cyt b	BRM-15	VLDLR-9
		Wrangel Island	SRE	=	M	Pullus	25 July 1994	EU410978	EU436182	EU436239, EU436240	
		Wrangel Island	SRE	=	F	Pullus	25 July 1994	EU410979	EU436183	EU436241	
		New Siberian Islands	SRE	-	F	Pullus	_	EU410980	EU436184	EU436242	
		New Siberian Islands	SRE	-	M	Pullus	1 August 1994	EU410981	EU436185	EU436243, EU436244	
		Wrangel Island	SRE	_	F	Pullus	25 July 1994	EU410982	EU436186	EU436245	
Snowy owl	North America	Illinois, USA	FMNH	334760 ^e	F	_	10 December 1987	EU410987	EU436191	EU436266	
		Minnesota, USA	FMNH	356917 ^e	M	_	27 April 1992	EU410988	EU436192	EU436267, EU436268	
		Wisconsin, USA	FMNH	384651 ^e	F	-	30 November 1993	EU410989	EU436193	EU436269	
		Illinois, USA		384661 ^e	M	-	29 November 1996	EU410990	EU436194	EU436270, EU436271	
		Minnesota, USA		385467 ^e	M	=	24 April 1997			EU436272, EU436273	
		Illinois, USA		386069 ^e	M	=	11 December 1996			EU436275	
		Minnesota, USA		396989 ^e	F	_	23 November 1999				
		Minnesota, USA		430331 ^e	M	_	29 November 2000			EU436278	
		Minnesota, USA		430332 ^e	F	_			EU436199		
		Illinois, USA		430528 ^e	M	_	28 October 2001			EU436280, EU436281	
		Minnesota, USA		435689 ^e	M	_	_			EU436282, EU436283	
		Minnesota, USA		436414 ^e	M	_	_			EU436284, EU436285	
		Captive, USA		437421 ^e	M	_	=			EU436286, EU436287	
	a	Minnesota, USA		438319 ^e	F	-	-		EU436204	EU436288	
Eagle owl	Scandinavia	Norway	NHMO		=		May/June 2005	EU411030			
		Nordland, Norway	NHMO		-		May/June 2005	EU411031			
		Nordland, Norway	NHMO		-		May/June 2005	EU411032			
		Nordland, Norway	NHMO		-		May/June 2005	EU411033			
		Nordland, Norway	NHMO		-		May/June 2005	EU411035			
		Nordland, Norway	NHMO		_		May/June 2005	EU411034			
		Nordland, Norway	NHMO		-		May/June 2005	EU411036			
		Nordland, Norway	NHMO		=		May/June 2005	EU411037			
		Nordland, Norway	NHMO		=		May/June 2005		EU436215 ^f		
		Nordland, Norway	NHMO	18699	=	Pullus	May/June 2005	EU411039			



Appendix continued

Species	Region	Locality	Source ^a	Voucher	Sexc	Age	Date	GenBank a	accession no.		
				no. ^b				Control region	Cyt b	BRM-15	VLDLR-9
Tengmalm's owl ^e	Scandinavia	Vest-Agder, Norway	NHMO	7531	F	2Y+	13 October 2003	EU411011	EU436217	EU436290 ^f	EU436306
		Vest-Agder, Norway	NHMO	7532	F	1Y	13 October 2003	EU411012			EU436307
		Vest-Agder, Norway	NHMO	7533	M	1Y	13 October 2003	EU411013	EU436219		EU436308, EU436309
		Vest-Agder, Norway	NHMO	7534	M	1Y	13 October 2003	EU411014	EU436220		EU436310, EU436311
		Vest-Agder, Norway	NHMO	7535	F	2Y+	13 October 2003	EU411015	EU436221		EU436312
		Vest-Agder, Norway	NHMO	7536	M	1Y	13 October 2003	EU411016	EU436222		EU436313, EU436314
		Vest-Agder, Norway	NHMO	7538	F	1Y	14 October 2003	EU411017	EU436223		EU436315
		Vest-Agder, Norway	NHMO	7539	F	1Y	14 October 2003		EU436224		EU436316
		Vest-Agder, Norway	NHMO	7540	F	1Y	14 October 2003	EU411018	EU436225		EU436317
		Vest-Agder, Norway	NHMO	7541	M	1Y	14 October 2003	EU411019	EU436218		EU436318, EU436319
Tawny owl	Scandinavia	Telemark, Norway	NHMO	6506	M	1Y+	5 November 1999	EU411023	EU436216 ^f	EU436289 ^f	EU436292, EU436293
		Akershus, Norway	NHMO	7017	F	Pullus	20 May 2004	EU411020			EU436294
		Akershus, Norway	NHMO	7018	F	Pullus	20 May 2004	EU411025			EU436295
		Akershus, Norway	NHMO	7019	M	Pullus	20 May 2004	EU411026			EU436296, EU436297
		Akershus, Norway	NHMO	9394	M	Pullus	22 May 2005	EU411027			EU436298, EU436299
		Akershus, Norway	NHMO	9395	F	Pullus	22 May 2005	EU411022			EU436300
		Akershus, Norway	NHMO	9396	F	Pullus	22 May 2005	EU411024			EU436301
		Aust-Agder, Norway	NHMO	19127	F	2Y	18 April 1994				EU436302
		Aust-Agder, Norway	NHMO	19128	F	3Y	18 April 1994	EU411028			EU436303
		Aust-Agder, Norway	NHMO	19129	F	3Y	18 April 1994	EU411021			EU436304
		Aust-Agder, Norway	NHMO	19130	F	3Y	18 April 1994	EU411029			EU436305

^a NRM: Swedish Museum of Natural History; TM: Museum of Tromsø, Norway; ANM: Agder Museum of Natural History, Norway, ZUO: Zoological Museum at the University of Oulu, Finland, SRE: Swedish-Russian tundra expedition 1994, FMNH: The Field Museum of Natural History, Chicago, NHMO: Natural History Museum of Oslo, Norway



^b Journal numbers from the DNA/tissue collection of NHM Oslo given in italic

^c Sex was determined genetically with the primers 2500F/2718R which has been shown to produce two bands for females (W + Z) and one for males (Z) on Tengmalm's owls (Fridolfson and Ellegren 1999). We tested snowy owl and tawny owl individuals with known sex (determined from morphological characters, Cramp and Simmons 1994) and found the expected pattern. We could not amplify any Z chromosome fragments for the eagle owl samples, probably due to low concentration of DNA extracted from the down samples, and could thus not determine their sex

^d Non-breeding bird (non-breeding year, or outside breeding season)

^e Birds caught on migration

^f All individuals of the species have the same haplotype

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