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Contrasting patterns of genetic diversity at three different genetic markers in a marine mammal metapopulation

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Abstract

Many studies use genetic markers to explore population structure and variability within species. However, only a minority use more than one type of marker and, despite increasing evidence of a link between heterozygosity and individual fitness, few ask whether diversity correlates with population trajectory. To address these issues, we analysed data from the Steller's sea lion, *Eumetopias jubatus*, where three stocks are distributed over a vast geographical range and where both genetic samples and detailed demographic data have been collected from many diverse breeding colonies. To previously published mitochondrial DNA (mtDNA) and microsatellite data sets, we have added new data for amplified fragment length polymorphism (AFLP) markers, comprising 238 loci scored in 285 sea lions sampled from 23 natal rookeries. Genotypic diversity was low relative to most vertebrates, with only 37 loci (15.5%) being polymorphic. Moreover, contrasting geographical patterns of genetic diversity were found at the three markers, with Nei's gene diversity tending to be higher for AFLPs and microsatellites in rookeries of the western and Asian stocks, while the highest mtDNA values were found in the eastern stock. Overall, and despite strongly contrasting demographic histories, after applying phylogenetic correction we found little correlation between genetic diversity and either colony size or demography. In contrast, we were able to show a highly significant positive relationship between AFLP diversity and current population size across a range of pinniped species, even though equivalent analyses did not reveal significant trends for either microsatellites or mtDNA.

Keywords: amplified fragment length polymorphism (AFLP), conservation genetics, demography, *Eumetopias jubatus*, genetic diversity, microsatellite, mtDNA, phylogeography, pinniped, Steller's sea lion, stock structure

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Introduction

Molecular genetic analysis of population structure is now a commonplace tool in the armoury of those wishing to understand the dynamics of natural populations. Currently, two classes of marker dominate those used in this context: maternally inherited mitochondrial DNA (mtDNA) and presumed neutral microsatellites (Zhang & Hweitt 2003; Schlotterer 2004). These markers provide

contrasting views of a given scenario, the mitochondrial sequences allowing reconstruction of maternal lineages while the microsatellites give a joint window of both maternal and paternal contributions. When combined, these contrasting views can become synergistic, with the mitochondrial markers uncovering patterns of maternally directed natal site fidelity, while the microsatellites help to quantify levels of paternal gene flow among subpopulations (e.g. Waits *et al.* 2000; Miller-Butterworth *et al.* 2003). Given this, it is perhaps surprising that studies combining both markers are the exception rather than the rule.

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In addition to differences in mode of inheritance, markers can also differ in their rate of evolution. Mitochondrial sequences tend to evolve faster than nuclear sequences while microsatellites evolve much faster than, for example, protein isozymes (e.g. Ellegren 2000; Ballard & Whitlock 2004; Schlotterer 2004). Rapidly evolving markers tend to be most useful for capturing recent demographic patterns and generally offer greater resolution due to their higher allelic/haplotypic diversities but may saturate over longer periods of time (Selkoe & Toonen 2006). However, deeper patterns such as residual signals of glacial refugia could potentially benefit from the use of more slowly evolving markers such as amplified fragment length polymorphisms (AFLPs). Arguably, even fewer studies compare markers that evolve over different timescales relative to those that compare nuclear and mitochondrial markers.

The primary aim of many studies of this nature is to understand current patterns of gene flow and genetic diversity in the context of historical patterns of demographic expansion and contraction, for example by identifying putative population bottlenecks or vicariant events that created isolated subpopulations. However, recent work has highlighted the possible importance of genetic diversity in determining the health of both individuals (e.g. Colman *et al.* 1999; Acevedo-Whitehouse *et al.* 2003) and perhaps by implication, populations. Indeed, a long-standing question in conservation genetics concerns the extent to which populations carrying high genetic diversity perform in some sense better than populations with low diversity, either as a consequence of identified bottlenecks or perhaps as an inherent property of a species. Here, use of contrasting types of markers is desirable, since it is quite possible for high, recently acquired microsatellite diversity to mask longer-term patterns in the underpinning additive genetic variability upon which selection is most likely to act.

Little is currently known about the link between genetic diversity and fitness at the population level, although relatively heterozygous vertebrate populations have been shown to experience lower parasite loads (Whiteman *et al.* 2006), improved body condition (Knaepkens *et al.* 2002), faster growth rates (Rowe *et al.* 1999; Cena *et al.* 2006) and greater survivorship (Saccheri *et al.* 1998; Shikano & Taniguchi 2002; Andersen *et al.* 2004). A recent meta-analysis of both plant and animals (Reed & Frankham 2003) suggests that genetic diversity could explain as much as 15–20% of variation in population fitness. However, the majority of studies to date have used allozymes, which may not be selectively neutral. In addition, although heterozygosity has been linked to a number of population fitness traits, surprisingly few studies have looked for a link between genetic diversity and the rate at which natural populations grow or decline.

Although a positive relationship between population heterozygosity and viability seems intuitive, it may not

always be that simple. Two recent studies have examined how levels of heterozygosity vary during the course of well-documented demographic challenges. Vazquez-Dominguez *et al.* (1999) found that heterozygosity increased during population declines in the spiny pocket mouse, and Valsecchi *et al.* (2004) found that Mediterranean striped dolphins dying early in an epizootic were significantly less heterozygous than those dying later. These studies imply that natural selection may sometimes remove relatively homozygous individuals from populations during demographic declines, raising the counter-intuitive possibility that, at least in the short term, declining populations may in fact be more heterozygous than stable ones. Furthermore, the purging of genetic load during population bottlenecks could generate a scenario in which relatively homozygous populations do well in the face of challenges.

The Steller's sea lion, *Eumetopias jubatus*, provides an opportunity to explore range-wide patterns of genetic diversity and to study the relationship between genetic variation and population viability in a natural metapopulation of a vertebrate species. The largest of the extant otariids, this species is distributed across the North Pacific Rim and throughout the Bering and Okhotsk Seas (Fig. 1). The worldwide population was once estimated to number about a quarter of a million animals (Kenyon & Rice 1961), but by 1989 the count had fallen to a little over 100 000 (Loughlin *et al.* 1992). This steep decline attracted worldwide attention and led to the Steller's sea lion being listed as Threatened under the US Endangered Species Act in 1990. Genetic studies using mtDNA (e.g. Bickham *et al.* 1996) led to the recognition of two well-differentiated stocks, eastern and western, which were listed as Threatened (eastern) and Endangered (western) in 1997. Subsequently, a larger study using mtDNA argued for the partitioning of the western stock to yield an additional Asian stock. Biparentally inherited microsatellite markers yield qualitatively similar findings (Hoffman *et al.* 2006), but also suggest that two genetically distinct subpopulations may exist within the western stock (O'Corry-Crowe *et al.* 2007). More recently, on the basis of morphological differences between skulls from the western and eastern stocks, Philips *et al.* (in press) elevated these to subspecies, designated the western Steller's sea lion (*Eumetopias jubatus jubatus*) and Loughlin's northern sea lion (*Eumetopias jubatus monteriensis*), respectively.

The reasons why certain Steller's sea lion populations have experienced a precipitous decline while others have remained stable or even increased are not readily apparent. Suggested causes of the decline include changes in food availability caused by overfishing and/or a regime shift in the North Pacific Ocean, legal and illegal shooting and predation (Loughlin & York 2000; Atkinson *et al.* 2008). Whatever the reason or reasons might be, it is clear from the highly subdivided nature

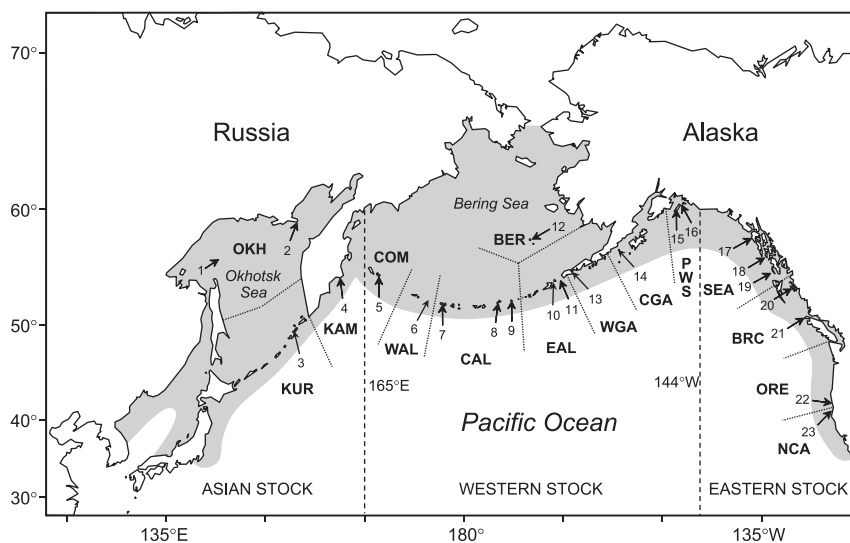


Fig. 1 Map showing the locations of 23 Steller's sea lion rookeries sampled in this study. The grey area indicates the current distribution of the species. Stocks and regions are as defined by Baker *et al.* (2005). For details of regions and rookeries, including the numbers of individuals genotyped, see Table 1.

of this metapopulation that the extirpation of rookeries could lead to an erosion of overall genetic variability within the species.

Here we analyse a data set comprising 285 Steller's sea lions genotyped at AFLP loci together with previously published data sets for microsatellites (598 individuals genotyped at 13 loci) and mtDNA (1559 individuals sequenced at 238 bp of D-loop HVR-1). Our aims were to assess overall levels of genetic diversity, to test whether AFLPs show the same signal of genetic structure as the other two marker types, to examine the relationship between genetic diversity and both colony size and rate of decline, and finally to place observed levels of genetic diversity in the context of other pinniped species.

Materials and methods

Tissue sample collection and DNA extraction

We utilized 285 tissue samples that were collected as part of a previous study (Baker *et al.* 2005) from pups at their natal rookeries ranging from Iony Island in the Okhotsk Sea to St. George Reef in northern California (Table 1, Fig. 1). Samples were obtained from rear flipper punches and stored individually in the preservative buffer 20% dimethyl sulphoxide (DMSO) saturated with salt. Total genomic DNA was extracted using a standard phenol-chloroform protocol (Sambrook *et al.* 1989).

AFLP genotyping

The AFLP protocol was similar to that used by Vos *et al.* (1995) and is described in detail by Dasmahapatra *et al.* (online early). Briefly, 100–400 ng of genomic DNA was first digested using *TaqI* (5 U in a 10- μ L volume at 65 °C for

2 h) and then with *EcoRI* (5 U in a 20- μ L volume at 37 °C for 2 h). *TaqI* and *EcoRI* adapters were then ligated onto the digested DNA using T4 DNA ligase (1 U in a 50- μ L volume at 37 °C for 3 h), and the resulting products diluted 10-fold in 10 mM Tris-HCl and EDTA (0.1 mM, pH 8.0). For the pre-amplification, 5 μ L of ligation mix was added to 50 μ L polymerase chain reaction (PCR) containing Tris-HCl (10 mM, pH 8.3), MgCl₂ (1.5 mM), KCl (50 mM), dNTPs (0.2 mM), *Taq* polymerase (1 U) and 50 ng each of the *TaqI*-C and *EcoRI*-A pre-amplification primers (the primer sequences were 5'-GATGAGTCCTGACCGAC-3' and 5'-GACTGCGTACCAATTCA-3', respectively). Following 30 pre-amplification cycles (30 s at 94 °C, 60 s at 50 °C and 60 s at 72 °C), the products were diluted 10-fold with 10 mM Tris-HCl and EDTA (0.1 mM, pH 8.0). For the selective amplification, 2.5 μ L of the diluted pre-amplification product was added to a 12.5- μ L reaction containing Tris-HCl (10 mM, pH 8.3), MgCl₂ (1.5 mM), KCl (50 mM), dATPs, dTTP and dGTP (0.2 mM each), dCTP (0.04 mM), α -³³P-dCTP, *Taq* polymerase (0.2 U), *TaqI* selective primer (30 ng) and *EcoRI* selective primer (5 ng). Samples were subjected to 13 selective amplification cycles (30 s at 94 °C, 60 s at 65 °C, reducing by 0.7 °C each cycle, and 60 s at 72 °C), followed by a further 23 cycles (30 s at 94 °C, 60 s at 56 °C and 60 s at 72 °C). PCR products were resolved by electrophoresis on standard 6% polyacrylamide sequencing gels and detected by autoradiography. AFLP profiles were assessed and scored manually by an experienced operator (J.H.). Only clear, polymorphic bands that could be scored in all individuals were included, these being recorded as 1, present and 0, absent. Eight different selective primer combinations were used (Table 2) to generate 238 AFLP loci that could be scored unambiguously across all of the samples.

Although AFLPs tend to be highly reproducible (Vos *et al.* 1995; Jones *et al.* 1997), as with other genetic markers

Table 1 Numbers of Steller's sea lion samples genotyped at AFLPs, microsatellites and mtDNA control region (see methods for details). Stocks and regions are as defined by Baker *et al.* (2005)

Stock	Region	Rookery	Number of samples genotyped			
			AFLPs	Microsatellites	mtDNA	
Asian	Sea of Okhotsk-OKH	1. Iony Island	15	25	100	
		2. Yamsky Island	15	25	80	
Western	Kuril Islands-KUR	3. Lovushki Island	10	15	39	
		Kamchatka Peninsula-KAM	4. Kozlova Cape	10	25	59
			5. Medny Island	15	25	126
	Commander Islands-COM	6. Buldir Island	9	12	45	
		7. Kiska Island	25	24	72	
	Western Aleutian Islands-WAL	8. Seguam Island	10	24	31	
		9. Yunaska Island	10	22	40	
		10. Akutan Island	10	56	85	
	Eastern Aleutian Islands-EAL	11. Ugamak Island	10	100	99	
		12. Walrus Island	10	13	42	
	Bering Sea-BER	13. Clubbing Rocks	10	19	35	
	Western Gulf of Alaska-WGA	14. Chowiet Island	10	25	32	
	Central Gulf of Alaska-CGA	15. Fish Island	10	25	47	
		16. Seal Rocks	10	50	102	
	Eastern	Southeastern Alaska-SEA	17. White Sisters Island	15	9	49
18. Hazy Island			10	26	103	
19. Forrester Island			15	10	215	
British Columbia-BRC		20. N. Danger Rocks	10	10	10	
		21. Triangle Island	10	8	13	
Oregon-ORE		22. Rogue Reef	16	25	84	
Northern California-NCA		23. St. George Reef	20	25	51	
Entire range			285	598	1559	

Table 2 Numbers of AFLP loci generated by eight AFLP selective primer combinations

TaqI primer (5'-3')	EcoRI primer (5'-3')	Total no. of loci	No. of polymorphic loci
GATGAGTCCTGACCGA-CAC	GACTGCGTACCAATTC-AGC	33	7
GATGAGTCCTGACCGA-CAG	GACTGCGTACCAATTC-ATG	31	6
GATGAGTCCTGACCGA-CGA	GACTGCGTACCAATTC-ACA	16	3
GATGAGTCCTGACCGA-CCA	GACTGCGTACCAATTC-AAC	29	3
GATGAGTCCTGACCGA-CCA	GACTGCGTACCAATTC-AGC	24	4
GATGAGTCCTGACCGA-CCA	GACTGCGTACCAATTC-ATG	33	4
GATGAGTCCTGACCGA-CTG	GACTGCGTACCAATTC-ATG	40	4
GATGAGTCCTGACCGA-CAG	GACTGCGTACCAATTC-ACA	32	6
Total		238	37

genotyping errors can easily accrue (Bonin *et al.* 2004; Hoffman & Amos 2005; Pompanon *et al.* 2005; Meudt & Clarke 2007). Consequently, we estimated the genotyping error rate for our data set by independently re-genotyping and blind scoring 24 individuals (almost 10% of the samples). The error rate per reaction was quantified following Bonin *et al.* (2004) as the number of mismatching genotypes divided by the number of bands compared.

Data analysis

The final AFLP character matrix consisted of 67 830 binary characters representing the presence and absence genotypes of 285 individuals at 238 loci. To examine patterns of genetic structure, we used the program AFLP-SURV version 1.0 (Vekemans 2002) to calculate pairwise F_{ST} values among rookeries and regions, to generate F_{ST} matrices for each of

1000 bootstrapped data sets, and to conduct a permutation test for overall genetic differentiation using 10 000 permutations of the data set. A consensus neighbour-joining (NJ) tree was then generated using the Neighbour, Consense and Fitch modules in PHYLIP (Felsenstein 1993). The significance of the correlation between pairwise geographical and genetic distance matrices was assessed using Mantel tests with 10 000 iterations implemented in the Mantel Nonparametric Test Calculator version 2.0 (Liedloff 1999). To explore range-wide patterns of genetic diversity, Nei's gene diversity was calculated for each rookery using AFLP-SURV (Vekemans 2002).

We next conducted a Bayesian cluster analysis using Structure 2.2.3 (Pritchard *et al.* 2000; Falush *et al.* 2007). This program uses an iterative approach to cluster the genotypes into K populations without knowledge of the population membership of individuals. The approach essentially subdivides the data set in a way that maximizes Hardy-Weinberg equilibrium and linkage equilibrium within the resulting clusters. The membership of each individual in a population is then estimated as q , which varies between 0 and 1 with the latter indicating full population membership. We ran five independent runs for $K = 1-10$ using 1×10^6 Markov chain Monte Carlo (MCMC) iterations after a burn-in of 1×10^5 , specifying the correlated allele frequencies model and assuming admixture. The most likely number of populations was evaluated using both the maximal value of $\ln P(D)$, a model-choice criterion that estimates the posterior probability of the data, and ΔK , an ad hoc statistic based on the second order rate of change of the likelihood function with respect to K (Evanno *et al.* 2005).

To enable comparisons across markers, we also analysed data from 13 highly polymorphic microsatellites ($n = 598$, Hoffman *et al.* 2006) and from a 238-bp section of the mitochondrial D-loop HVR-1 region ($n = 1559$, Baker *et al.* 2005; J. W. Bickham, unpublished data). The three data sets overlapped considerably, with the individuals typed at AFLPs being a subset of those genotyped for microsatellites, which were in turn a subset of the much larger sample of individuals typed for mtDNA. Genetic differentiation among rookeries was estimated at both of these markers using Wright's F -statistics (Wright 1951) calculated in Arlequin 2.0 (Schneider *et al.* 2000). In addition, in response to recent concerns raised about the reliability of F_{ST} in highly polymorphic systems, we also calculated D for our microsatellite data set following the method of Jost (2008), equation 14. Overall, results using this approach were very similar to, but less significant than those obtained using classical F_{ST} , suggesting that F_{ST} has adequate resolution in our system. For comparability with other studies, we therefore used F_{ST} . Nei's gene diversity was also calculated for microsatellites using FSTAT 2.9.3 (Goudet 1995) and for mtDNA using DNAsp (Rojas & Rojas 1995).

To explore the relationships between genetic diversity at each of the three markers and population size, we constructed a series of general linear models (GLMs). Population sizes were log transformed both to minimize heteroscedasticity and because population growth trajectories tend to be exponential. Population size estimates were available for the period 1957–2006 inclusive, but the number and timing of records for each of the rookeries varies greatly, from as few as five to as many as 35 observations (Table 3). To standardize our procedure, we therefore defined two critical years: 1960, reflecting the best balance between a date before the declines begin yet where data coverage is still adequate, and 2006, reflecting the current status. For each of these two time points, we estimated the most likely population size by means of linear extrapolation/interpolation in plots of \log (population size) on year using all available data for that colony. The r^2 values of the regressions are shown in Table 3 and averaged 0.614. Population size estimates are, of necessity, somewhat crude but visual inspection of the graphs suggests they are adequate for our purposes. Initially, we constructed GLMs of gene diversity at each of the three different markers fitting \log (population size) as a continuous predictor variable. However, because of the presence of three genetically distinct stocks with on average very different population trajectories (the eastern population was originally small but is now increasing, while the Asian and western stocks were initially large but have since declined), we also constructed additional GLMs of gene diversity fitting both \log (population size) and stock, the latter as a factor with three levels corresponding to the Asian, Western and Eastern stocks.

Next, to explore relationships between genetic diversity and the rates at which different Steller's sea lion rookeries have declined, we calculated the growth trajectory of each rookery as the \log of the gradient of year on population size. We then constructed GLMs of growth trajectory fitting gene diversity as a single, continuous explanatory variable. As with the GLMs of genetic diversity, we then additionally controlled for stock membership by fitting stock as an additional predictor variable (as a factor with three levels) in GLMs of growth trajectory. All GLMs were fitted using R (R development team 2005) as full models and then simplified following Crawley (2002) by stepwise deletion of non-significant terms (strictly, terms whose deletion did not cause a significant reduction in the proportion of the null deviance explained by the model).

Next, we used two different approaches implemented in the program Bottleneck 1.2.02 (Piry *et al.* 1999) to test whether any of the three Steller's sea lion stocks have experienced a recent reduction in effective population size or a genetic bottleneck. The first of these approaches exploits the fact that during a bottleneck, alleles are lost more rapidly than heterozygosity at neutral markers, generating a transient 'heterozygosity excess'. This was assessed using the micro-

Table 3 Summary of population size estimate data for 23 Steller's sea lion rookeries spanning the period 1957–2006 inclusive. The gradient and r^2 values refer to the regressions of time on log population size, with positive gradients indicating population growth and negative gradients indicating decline

Stock	Rookery	No. of observations	First observation	Last observation	Gradient	r^2
Asian	Iony Island	7	1974	2002	0.010	0.661
	Yamsky Island	12	1974	2003	0.006	0.397
	Lovushki Island	23	1967	2001	-0.012	0.361
Western	Kozlova Cape	17	1982	2003	-0.021	0.420
	Medny Island	35	1967	2002	-0.024	0.703
	Buldir Island	13	1968	2004	-0.058	0.942
	Kiska Island	12	1979	2004	-0.048	0.858
	Seguam Island	12	1979	2004	-0.028	0.506
	Yunaska Island	13	1979	2006	-0.033	0.779
	Akutan Island	21	1965	2006	-0.024	0.726
	Ugamak Island	18	1969	2006	-0.028	0.579
	Walrus Island	5	1982	1994	-0.054	0.881
	Clubbing Rocks	18	1957	2006	-0.007	0.460
	Chowiet Island	15	1957	2004	-0.029	0.829
	Fish Island	15	1957	2006	-0.017	0.638
	Seal Rocks	16	1973	2006	-0.016	0.537
	Eastern	White Sisters Island	13	1979	2004	0.006
Hazy Island		13	1979	2005	0.015	0.889
Forrester Island		12	1979	2005	0.004	0.235
North Danger Rocks		9	1971	2002	0.013	0.633
Triangle Island		7	1971	2998	0.012	0.874
Rogue Reef		23	1977	2001	0.016	0.782
	St. George Reef	11	1990	2001	0.010	0.140

satellite data set for each of the stocks separately under a range of mutation models ranging from the infinite allele model (IAM) through the two-phase mutation model (TPM) with 70%, 90%, 95% and 99% single-step mutations (with a variance of 30%), to the stepwise-mutation model (SMM). Statistical significance was assessed using the Wilcoxon test. The second test implemented using Bottleneck was one for a shift away from an L-shaped allele frequency distribution to one with fewer alleles in low frequency categories (Luikart & Cornuet 1997).

Finally, we analysed genetic diversity in a representative panel of other pinnipeds, both to provide a context for interpreting the diversity seen in our focal species, and also to learn the extent to which the levels of diversity exhibited by our three classes of marker correlate with likely demographic history. Our panel of species includes some that have been heavily exploited to near extinction and either recovered or stayed endangered, and others that have expanded greatly to become some of the most abundant large mammals on the planet. For this analysis, we collated published and unpublished data on microsatellites, mitochondrial D-loop sequences and AFLP markers from as many pinniped species as were available (Table 4). Recognized subspecies were treated separately, as were the Western and Eastern Atlantic populations of the grey seal, *Halichoerus grypus*. To avoid ascertainment bias, micro-

satellite data were only accepted if based on markers that were derived from the species being analysed. We also excluded markers that were out of Hardy–Weinberg equilibrium and/or exhibited high frequencies of null alleles. For each of 12 species/subspecies, average observed heterozygosity was calculated over all of the microsatellites that met our criteria. Mitochondrial D-loop sequences (with the 5' end of tRNA-Pro) together with haplotype frequencies were available for 19 species/subspecies. For these species, a section corresponding to positions 39–327 in the *Arctocephalus pusillus* mitochondrial genome (AM181018) was used to calculate haplotype diversity. The length of sequence varied among the species due to the presence of indels and in some cases, only sequence data for a slightly shorter section were available. Within each species, sequences were aligned using ClustalW and by eye, and sequence diversity π (Nei 1987) was calculated using MEGA 4 (Tamura *et al.* 2007). AFLP genotypes were generated for 14 different pinniped species for which adequate samples were available. We chose a target sample number of five as the best compromise between generating representative profiles and including as many species as possible. Five is probably too small for many classes of marker, but for AFLPs the low sample size is partly compensated for by the large number of traits (= bands) that can be scored. AFLP diversity was calculated as the

Table 4 Genetic diversity at AFLP markers (proportion of polymorphic loci), microsatellites (observed heterozygosity) and mtDNA control region sequence diversity in a variety of pinniped species

Family	Species	Proportion of polymorphic AFLP loci	Observed microsatellite heterozygosity (no. of loci, no. of samples)	Sequence diversity, π , at mtDNA D-loop (sequence length used, no. of samples)
Phocidae	Crabeater seal, <i>Lobodon carcinophaga</i>	—	0.813 (6, 25) ¹	—
	Grey seal, <i>Halichoerus grypus</i> (Eastern Atlantic population)	0.133	0.784 (5, 805) ²	0.014 (327, 1025) ³
	Eastern Atlantic Harbour seal, <i>Phoca vitulina vitulina</i>	0.056	0.238 (6, 50) ⁴	0.0054 (320, 159) ⁵
	Western Atlantic Harbour seal, <i>Phoca vitulina concolour</i>	—	0.390 (5, > 40) ⁶	0.012 (320, 18) ⁵
	Eastern Pacific Harbour seal, <i>Phoca vitulina richardsi</i>	—	—	0.014 (320, 38) ⁵
	Western Pacific Harbour seal, <i>Phoca vitulina stejnegeri</i>	—	—	0.015 (320, 12) ⁵
	Harp seal, <i>Pagophilus groenlandicus</i>	0.215	—	—
	Hawaiian monk seal, <i>Monachus schauinslandi</i>	—	—	0.0001 (337, 50) ⁷
	Hooded seal, <i>Cystophora cristata</i>	0.168	—	0.030 (334, 123) ⁸
	Leopard seal, <i>Hydrurga leptonyx</i>	0.144	0.626 (7, 21) ¹	—
	Northern elephant seal, <i>Mirounga angustirostris</i>	—	—	0.004 (299, 150) ^{9, 10}
	Southern elephant seal, <i>Mirounga leonina</i>	0.066	0.597 (2, 263) ¹⁰	0.021 (301, 48) ¹⁰
	Spotted seal, <i>Phoca largha</i>	—	—	0.024 (335, 66) ¹¹
	Weddell seal, <i>Leptonychotes weddellii</i>	—	0.737 (17, 96) ¹	—
	Otariidae	Antarctic fur seal, <i>Arctocephalus gazella</i>	0.113	0.744 (15, 20) ^{12, 13}
Australian Sea lion, <i>Neophoca cinerea</i>		—	—	0.016 (288, 194) ¹⁵
California sea lion, <i>Zalophus californianus</i>		0.109	0.602 (9, 58) ¹⁶	0.020 (283, 52) ¹⁷
Cape fur seal, <i>Arctocephalus pusillus</i>		—	—	0.031 (285, 105) ¹⁸
Galapagos fur seal, <i>Arctocephalus galapagoensis</i>		0.055	—	—
Galapagos sea lion, <i>Zalophus californianus wollebacki</i>		0.037	0.677 (15, > 20) ^{19, 20}	0.005 (285, 336) ²¹
Guadalupe fur seal, <i>Arctocephalus townsendi</i>		—	—	0.021 (212, 32) ²²
Juan fernandez fur seal, <i>Arctocephalus philippii</i>		—	—	0.031 (298, 28) ²³
Northern fur seal, <i>Callorhinus ursinus</i>		0.130	—	—
South American fur seal, <i>Arctocephalus australis</i>		0.069	—	—
Steller sea lion, <i>Eumetopias jubatus</i>		0.063	0.507 (6, 20) ²⁴	0.011 (196, 2599) ²⁵
Subantarctic fur seal, <i>Arctocephalus tropicalis</i>		—	—	0.044 (299, 103) ²⁶
Obeniidae	Atlantic walrus, <i>Odobenus rosmarus rosmarus</i>	—	0.800 (7, 57) ^{27*}	—
	Pacific walrus, <i>Odobenus rosmarus divergens</i>	0.152	—	—

1, Davis *et al.* (2002); 2, Allen *et al.* (1995); 3, Amos, unpublished data; 4, Coltman *et al.* (1996); 5, Stanley *et al.* (1996); 6, Goodman (1997); 7, Kretzmann *et al.* (1997); 8, Coltman *et al.* (2007); 9, Weber *et al.* (2000); 10, Hoelzel *et al.* (1999); 11, Mizuno *et al.* (2003); 12, Hoffman *et al.* (2008); 13, Hoffman (online early); 14, Hoffman, unpublished data; 15, Campbell (2003); 16, Hernandez-Velazquez *et al.* (2005); 17, Maldonado *et al.* (1995); 18, Mathee *et al.* (2006); 19, Wolf *et al.* (2005); 20, Hoffman *et al.* (2007); 21, Wolf, unpublished data; 22, Weber *et al.* (2004); 23, Goldsworthy *et al.* (2000); 24, Huebinger *et al.* (2007); 25, Bickham, unpublished data; 26, Wynen *et al.* (2000); 27, Buchanan *et al.* (1998); *data from Foxe Basin population but similar observed microsatellite heterozygosity values were also found for a range of other populations by Andersen *et al.* (1998).

proportion of polymorphic bands. Population size estimates were obtained from the Seal Conservation Society website (www.pinnipeds.org) and where a range was given, we took the average of the upper and lower estimates.

Since the taxa (populations and species in the case of intraspecific and interspecific analyses respectively) are related, we also explored the use of phylogenetic correction. For this we chose the program Continuous as implemented in BayesTraits 1.0 (Pagel 1997; Pagel 1999). This program accepts as input a phylogeny plus data from two variables and then uses either a likelihood-based approach or Monte Carlo Markov Chain to estimate the degree to which the variables are correlated given the phylogeny. We chose to

implement the likelihood option in which the likelihood of the data given the phylogeny is calculated twice, once under the assumption of independence and a second time under the assumption that a correlation is present. Twice the difference between these likelihoods can then be interpreted as a chi-squared value with one degree of freedom.

Results

We genotyped 285 Steller's sea lions sampled from 23 natal rookeries representing 15 regions and three stocks (Fig. 1, Table 1) at eight selective AFLP primer combinations, yielding 238 putatively homologous loci (= bands) that could be scored unambiguously (Table 2). The calculated

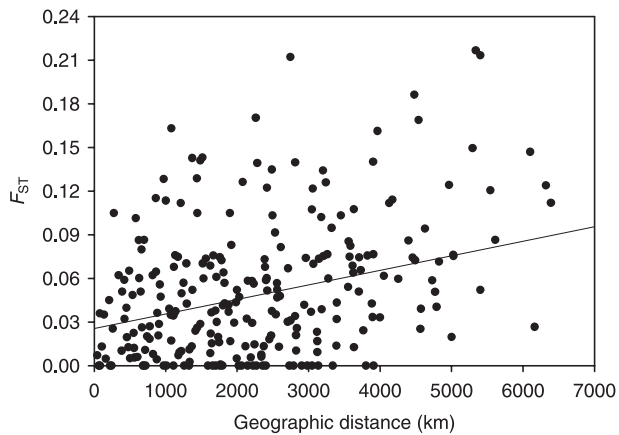


Fig. 2 The relationship between geographical and genetic distance calculated using AFLPs among 23 Steller's sea lion rookeries. The linear regression line is shown to indicate the underlying trend ($r^2 = 0.135$).

genotyping error rate was low at 0.012 per band (11 differences observed out of 888 band–band comparisons). Of the discrepancies observed between the two sets of genotypes, four (36.4%) were attributed to scoring or data entry errors and the remaining seven (63.6%) were due to the stochastic appearance or disappearance of bands as similarly documented by Bonin *et al.* (2004). Overall, levels of AFLP variability were low, with only 37 out of the 238 loci scored (15.5%, Table 2) being polymorphic in our large and geographically diverse sample. To facilitate interspecific comparisons, Milot *et al.* (2007) proposed quantifying AFLP variability using $P_{5\%}$, the proportion of loci where at least 5% of individuals carry the minor genotype. $P_{5\%}$ for our data set is 5.9%, far lower than the normal range of values reported for vertebrates (summarized by Milot *et al.* 2007) and is comparable with values obtained for wandering (5.1%) and Amsterdam (2.1%) albatrosses which were interpreted by Milot *et al.* (2007) as being extremely low.

Genetic structure and isolation by distance

A statistically significant pattern of genetic differentiation was observed across the species range (overall F_{ST} among rookeries = 0.050, $P < 0.001$ using 10 000 permutations of the AFLP data set). Pairwise F_{ST} values correlated positively with the geographical distance among rookeries (Fig. 2, Mantel test, $r = 0.367$, $n = 23$ colonies, $P < 0.001$), yielding a similar pattern to that obtained previously using 13 microsatellite loci (Hoffman *et al.* 2006). Also concordant with previous analysis of the same samples using microsatellites, no relationship between genetic and geographical distance was apparent when only within-stock comparisons were made (Asian stock, Mantel's $r = -0.098$, $n = 4$, $P = 0.491$, Western stock, Mantel's $r = 0.061$, $n = 12$,

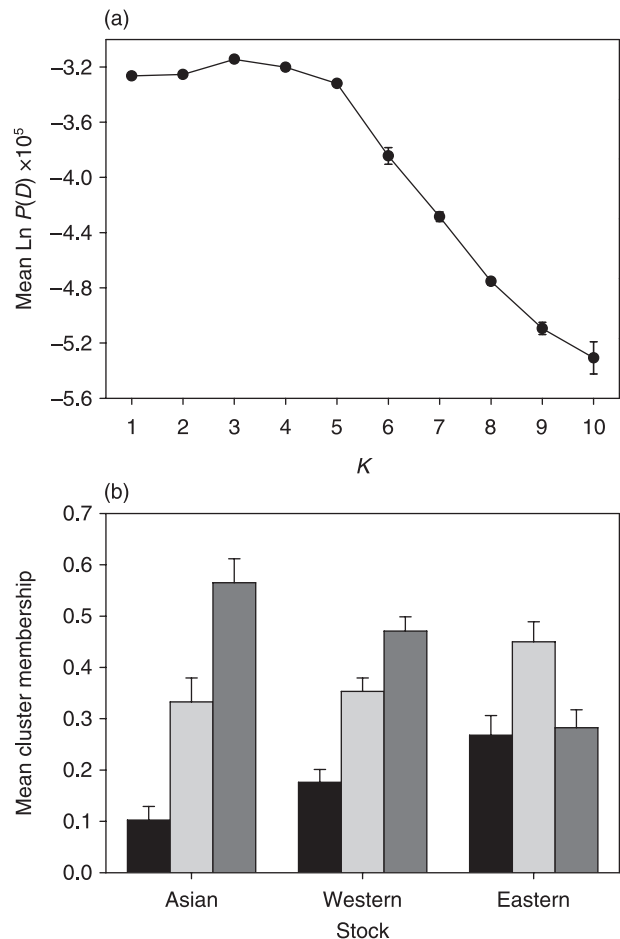


Fig. 3 Results of the Structure analysis of the AFLP data set. (a) Mean \pm SE $\ln P(D)$ values based on five replicates for each value of K ; (b) Mean \pm SE cluster membership coefficients for the three clusters (colour coded in black, light grey and dark grey, respectively) for each of the Steller's sea lion stocks.

$P = 0.348$, Eastern Stock, Mantel's $r = -0.027$, $n = 7$, $P = 0.486$) suggesting that the overall pattern is driven by among-stock comparisons. As expected, genetic distances calculated from the AFLP data matrix were positively correlated with equivalent values derived from both microsatellites and mtDNA (Mantel tests, $r = 0.371$, $n = 23$, $P = 0.002$ and $r = 0.326$, $n = 23$, $P < 0.001$, respectively), suggesting that all three of these markers provide concordant estimates of genetic differentiation. To further explore patterns of genetic divergence, we constructed a neighbour-joining tree at the regional level using 1000 bootstrapped F_{ST} matrices. The resulting topography was poorly resolved with the majority of nodes failing to gain 50% or greater bootstrap support, probably because of the small number of informative loci on which the genetic distances are calculated. Nevertheless, the regions of the eastern stock (SEA, BRC, ORE and NCA) form a distinct clade (data not shown), in

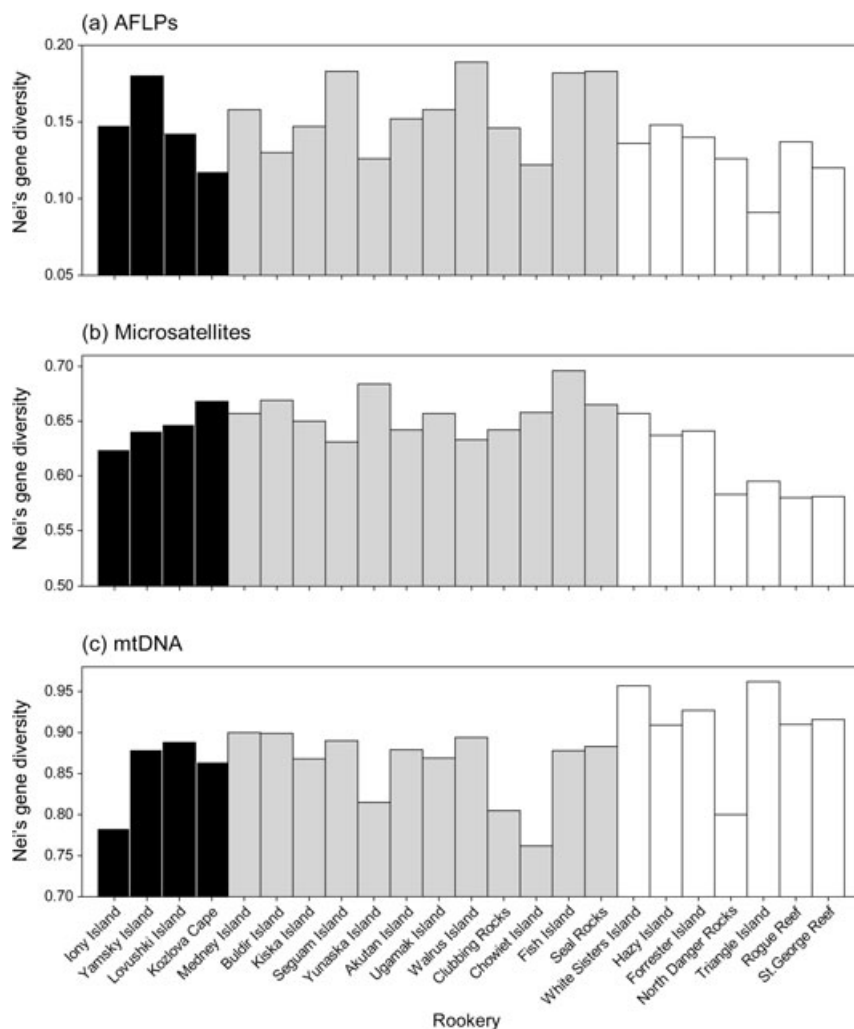


Fig. 4 Geographical variation in Nei's gene diversity across the current range of the Steller's sea lion calculated for (a) AFLPs, (b) microsatellites, and (c) mtDNA. Rookeries of the Asian, Western and Eastern stocks are denoted by black, grey and white-filled bars, respectively.

support of previous studies using both mtDNA and microsatellites (Baker *et al.* 2005; Hoffman *et al.* 2006).

Bayesian cluster analysis

We next implemented a Bayesian cluster analysis of the AFLP data set using the program Structure (Pritchard *et al.* 2000; Falush *et al.* 2007) in order to determine whether any genetic substructure could be detected without knowledge of the sampling locations of individuals. The resulting posterior probabilities were highly concordant among replicate runs, with the highest average value indicating the most likely number of population groups, K . Our data yielded a best estimate of $K = 3$ (Fig. 3a), which was also supported by a peak in Evanno *et al.*'s (2005) ΔK statistic. However, despite good support for $K = 3$, many individuals were poorly resolved in terms of group membership, probably because of the low resolution afforded by 37 unidominant markers. Consequently, we summarized the data by averaging the group membership coefficients

for all individuals in each of the three stocks (Fig. 3b). Average group membership coefficients were found to vary significantly among the stocks (in a two-way ANOVA fitting group, stock and the group:stock interaction, the interaction term was highly significant, $F_{(4,846)} = 11.4$, $P < 0.0001$). The first two clusters showed an increase in mean membership progressing from Asian through Western to the Eastern stock, while the third cluster shows the opposite, a pattern that is broadly consistent with isolation by distance.

Genetic diversity, population size and demography

To explore range-wide patterns of genetic diversity, we calculated Nei's gene diversity for each rookery using each of our three markers: AFLPs, microsatellites and mtDNA (Fig. 4). Significant variation was found among the three stocks for both AFLPs and microsatellites (one-way ANOVAs, $F_{2,20} = 3.60$, $P = 0.046$ and $F_{2,20} = 8.00$, $P = 0.003$, respectively) with the lowest Nei's gene diversity values

being found among the rookeries of the Eastern stock at both of these markers. In contrast, mtDNA diversity does not vary significantly among the three stocks (ANOVA, $F_{2,20} = 2.95$, $P = 0.075$). Moreover, for mtDNA the six highest gene diversity values all occur in the Eastern stock, the stock which has lowest nuclear diversity.

To relate genetic diversity at the three classes of marker to population size, we regressed gene diversity against log population size in 1960, chosen to reflect the best balance between a date before the declines begin yet where data coverage is still adequate and estimated by linear extrapolation from the available data. Nei's gene diversity was significantly correlated with the estimated population size in 1960 for microsatellites ($F_{1,21} = 8.14$; $P = 0.010$) but not for AFLPs or mtDNA ($F_{1,21} = 0.35$; $P = 0.562$ and $F_{1,21} = 0.01$; $P = 0.921$, respectively). However, when these regressions were repeated as GLMs of genetic diversity with population size fitted as a continuous variable and stock membership (e.g. Asian, Western or Eastern) fitted as a factor, neither of these terms were retained in the final model for AFLPs and mtDNA (although stock approached significance at $P = 0.055$ and 0.075 , respectively), and only stock was retained as a predictor of microsatellite diversity, explaining 45.0% of the null deviance ($F_{2,20} = 8.17$; $P = 0.003$).

Since the 1960s, the Western stock has undergone rapid decline, while the Eastern stock has increased. Consequently, we repeated the same analysis as above but this time fitting log population size in 2006 as a predictor of genetic diversity. Direct population count data were available for six rookeries of the Western Stock, and for the remaining rookeries, population size was obtained by linear extrapolation. AFLP and microsatellite diversity were both negatively associated with population size and mtDNA diversity was weakly but positively associated with population size, although none of these relationships were significant and only microsatellites approached significance ($F_{1,21} = 4.30$; $P = 0.050$). Again, when stock and population size in 2006 were fitted in full models of genetic diversity, no terms were retained for AFLPs and mtDNA, and only stock was retained in the GLM of microsatellite diversity.

Previous studies have found links between genetic diversity and population viability in a range of organisms (e.g. Saccheri *et al.* 1998; Rowe *et al.* 1999; Whiteman *et al.* 2006). Therefore, we sought to establish whether genetic diversity was linked to the rates at which different Steller's sea lion rookeries have declined. Microsatellite diversity explained a significant proportion of the variation in log population trend when fitted alone in a GLM ($F_{1,21} = 11.17$; $P = 0.003$), although the direction was in the reverse direction to that expected (e.g. growing colonies had lower gene diversity). In contrast, AFLPs and mtDNA did not explain significant variation when fitted alone

($F_{1,21} = 1.91$; $P = 0.180$ and $F_{1,21} = 0.53$; $P = 0.474$, respectively). Moreover, when full models of log population trend were constructed in which genetic diversity and stock were fitted together as predictors, only stock was retained ($F_{2,20} = 23.29$; $P < 0.0001$), explaining 70.0% of the total deviance.

To either confirm or refute the above trends, we ignored stock and instead used phylogenetic correction to allow for non-independence among populations. For this analysis, we explored the use of three alternative input phylogenies: (i) based on pooled data from AFLPs, mtDNA and microsatellites in which each pairwise distance was taken as the average of the three marker classes, normalized to force equal contribution from each marker class and with any negative distance values rounded to zero; (ii) what we consider subjectively the 'best' phylogeny, as judged by its ability to place neighbouring populations close to each other, based on the microsatellite F_{ST} values; and (iii) a non-genetic phylogeny based on great circle geographical distances. In each case and for each marker class, we tested for a correlation between genetic diversity and (i) population size in 1960, (ii) population size in 2006, (iii) the slope of population trend. Unfortunately, the results were highly variable. Using the geographical distance and microsatellite phylogenies, none of the tests were significant (ignoring one case of $P = 0.03$ that is best attributable to type I error). In contrast, using the full genetic phylogeny, we find that both AFLP and microsatellite diversity are correlated with population size in 1960 and 2006 ($P < 0.002$ in every case), while mtDNA diversity predicts the overall trend ($P < 0.001$).

Genetic bottleneck analyses

To determine whether low AFLP diversity in the Steller's sea lion could be at least partly due to a recent reduction in the effective population size, we next interrogated our microsatellite data set using the program Bottleneck (Piry *et al.* 1999). Significant heterozygosity excess was found in all three stocks using the IAM and within the Asian Stock using the SMM (Table 5). However, little evidence for a bottleneck was found using any of the probably more realistic TPM models. Moreover, none of the stocks deviated significantly from a normal L-shaped distribution of allele frequencies.

Patterns of genetic diversity across the Pinnipedia

Although $P_{5\%}$ appears low in the Steller's sea lion relative to most other vertebrates, there are problems in comparing different studies including the use of different restriction enzymes and selective primer combinations, variation in the geographical range of sampling and interobserver variation. Therefore, we sought to place the observed value

Table 5 Results of heterozygosity excess tests for the three Steller's sea lion stocks under a range of different mutational models using 13 polymorphic microsatellite loci (Hoffman *et al.* 2006)

Stock	Test		Mutation model					
			IAM	TPM 70	TPM 90	TPM 95	TPM 99	SMM
Asian	No. of loci with heterozygosity excess Wilcoxon test		11	10	7	6	5	4
		<i>P</i> value (one tail for heterozygosity deficiency)	0.999	0.905	0.658	0.368	0.658	0.047
		<i>P</i> value (one tail for heterozygosity excess)	0.002	0.108	0.368	0.658	0.368	0.960
	<i>P</i> value (two tails)	0.003	0.216	0.735	0.735	0.735	0.094	
Western	No. of loci with heterozygosity excess Wilcoxon test		12	9	8	7	6	6
		<i>P</i> value (one tail for heterozygosity deficiency)	0.998	0.960	0.632	0.393	0.170	0.095
		<i>P</i> value (one tail for heterozygosity excess)	0.003	0.047	0.393	0.632	0.847	0.916
	<i>P</i> value (two tails)	0.005	0.094	0.787	0.787	0.339	0.191	
Asian	No. of loci with heterozygosity excess Wilcoxon test		10	9	7	6	5	5
		<i>P</i> value (one tail for heterozygosity deficiency)	0.997	0.773	0.554	0.294	0.122	0.073
		<i>P</i> value (one tail for heterozygosity excess)	0.004	0.249	0.946	0.729	0.892	0.936
	<i>P</i> value (two tails)	0.009	0.497	0.946	0.588	0.244	0.146	

IAM, infinite alleles model; TPM, two-phase model (the number refers to the proportion of stepwise mutations); SMM, stepwise-mutation model. Significant *P* values are highlighted in bold.

into context by testing whether low AFLP diversity is a feature of pinnipeds in general by exploiting a data set comprising five samples each of 14 different pinniped species genotyped at eight AFLP loci and by collating data from microsatellite primer notes and papers containing mtDNA data (Table 4). Figure 5 shows that the Steller's sea lion has low diversity relative to most of the other pinniped species at all three markers. Moreover, a strong positive correlation was found between current population size and the proportion of polymorphic AFLP loci ($r^2 = 0.49$, $n = 14$, $P = 0.006$), suggesting that the low diversity at AFLP markers in this species may be a consequence of historically low population sizes. A similar but weaker pattern was obtained for mitochondrial DNA ($r^2 = 0.29$, $n = 19$, $P = 0.017$), but not for microsatellites ($r^2 = 0.14$, $n = 12$, $P = 0.231$). As with the population data, we also conducted tests using phylogenetic correction. For the phylogeny, we exploited the tree of Higdon *et al.* (2007), using the corrected divergence dates to produced branch lengths. When these trends were analysed using the program Continuous to correct for phylogenetic non-independence, the correlation between current population size and marker diversity was positive for all three markers and highly significant for AFLPs ($P < 0.0001$) but not for microsatellites ($P = 0.67$) or mtDNA ($P = 0.43$).

Discussion

Herein, we report a study using three different commonly used genetic markers, microsatellites, AFLPs and mitochondrial DNA to assess levels of genetic diversity and population structure in the Steller's sea lion, and then to place this species in the wider context of pinnipeds in

general. We find a significant pattern of isolation by distance and some evidence of a correlation between population size and genetic diversity. However, these patterns appear largely driven by differences among three different stocks with contrasting histories. Controlling for stock structure either by fitting stock as an extra predictor variable, or by implementing phylogenetic correction largely eliminates any population-specific effects. Among the pinnipeds, the Steller's sea lion seems to carry unexpectedly low levels of diversity, a pattern that is best reflected in AFLP markers which show the strongest correlation between diversity and current population size of the three markers we examined.

Expectations from different markers

Different markers are expected to reveal different aspects of a population's history, depending on their mode of inheritance and mutation rate. Thus, mitochondrial DNA will reveal patterns of maternally directed site fidelity while microsatellites, with their high mutation rates, will tend to recover high levels of variability following a population bottleneck faster than less mutable AFLP markers. Pinnipeds exhibit rather puzzling patterns of diversity that may be elucidated by the use of multiple marker types. Thus, elephant seals have extremely low diversity and were severely bottlenecked, but various fur seals were hunted just as hard and for longer yet carry the highest levels of microsatellite diversity seen among pinnipeds. Within a species, the Steller's sea lion has a broad geographical distribution and in parts of its range, populations are declining while elsewhere there appears to be expansion. These contrasting demographics may be

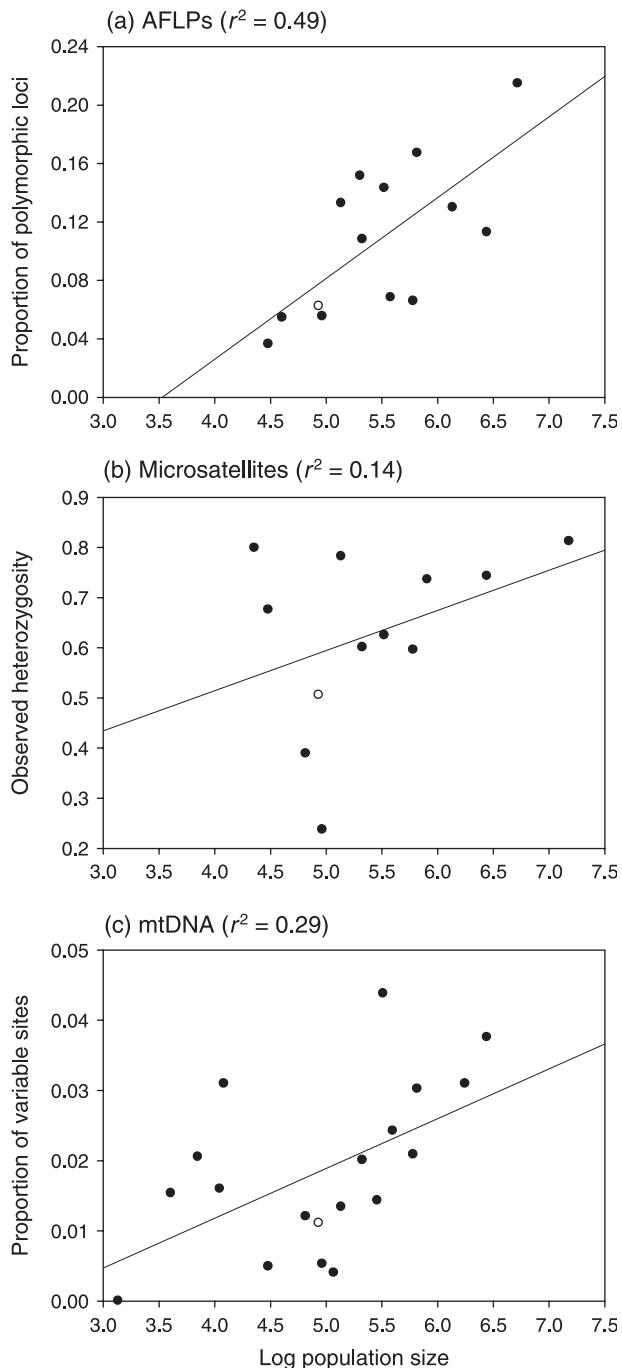


Fig. 5 Relationships between current estimated population size and genetic diversity at (a) AFLPs (b) microsatellites and (c) mtDNA across a range of pinniped species. White points indicate the Steller's sea lion. See Table 4 for details of sample sizes and literature references.

associated with the descendents of glacial refugia, which could in turn be reflected in distinct mitochondrial lineages. Finally, if widely reported trends linking microsatellite heterozygosity to individual fitness reflect a more general

tendency for greater genetic diversity to improve a population's health, we might expect that the contrasting population trends seen among modern populations of Steller's sea lions will correspond to each population's heterozygosity.

Genetic structure and links between diversity and population decline

A pattern of isolation by distance is likely to arise under a wide range of circumstances, even in potentially highly mobile aquatic species such as pinnipeds (e.g. Allen *et al.* 1995; Goodman 1998; Campbell *et al.* 2007). For example, in the grey seal, many adults show high levels of breeding site fidelity and dispersal occurs mainly to available neighbouring sites when an individual's natal colony has reached carrying capacity (Gaggiotti *et al.* 2002). One might expect the Steller's sea lion to be no exception, distributed as it is over a vast geographical range in a series of breeding rookeries along the Aleutian chain and beyond, and this is what a naïve analysis reveals. However, previous studies have suggested the existence of at least two and possibly three different stocks, possibly reflecting the existence of historical ice age refugia (Bickham *et al.* 1996; Baker *et al.* 2005; Harlin-Cognato *et al.* 2005). Indeed, more recent morphological analysis has led to the suggestion that the eastern and western stock might even be considered subspecies (Phillips *et al.* in press). Once this structure has been corrected for by fitting stock as an extra parameter, the pattern of isolation by distance becomes non-significant, suggesting that the main driver of the apparent isolation-by-distance pattern that we observe is the presence of rather dissimilar stocks. Use of Jost's D instead of F_{ST} in theory allowing for the reduced resolution of F_{ST} in highly polymorphic systems, if anything only weakened any pattern of isolation by distance.

The presence of different genetically distinct stocks is problematic for most of the genetic analyses one might wish to conduct. This is because the shared ancestry within a stock creates some degree of statistical non-independence. To take an extreme example, if 10 populations were sampled from each of two stocks, one with high diversity and one with low diversity, any regression of diversity on a trait linked to stock identity, such as recent demography or geographical location, would tend to be highly significant (20 data points and a clear trend), when in fact only two fully independent observations exist. Our data feature the same problem, with two main stocks, east and west, but separate demographic trends for each of many subpopulations (i.e. breeding colonies). When we ignore stock structure, we find interesting patterns, but when stock structure is properly controlled, either by fitting stock as a predictor variable or by using phylogenetic correction, most of these correlations are eliminated.

Our results using the program Continuous (Pagel 1997) also point to the need to assess critically whether the input phylogeny is valid. We obtained sharply contrasting results depending on the phylogeny we used. Based on data from all three markers pooled and normalized to ensure equal contribution from each, we obtained evidence of several perhaps suspiciously strong trends. However, when we used either a non-genetic tree based on geographical distances or the phylogeny that we believe most accurately reflects the likely true relationships among rookeries, these trends appear no longer significant. As yet, it is unclear which result is correct, although we prefer to be conservative and assume that the agreement between two contrasting but likely reliable trees provides the stronger evidence. Conversely, the fact that the combined marker phylogeny reveals trends not supported by two other trees suggests that while the combined tree maximizes the genetic information contributing to each genetic distance, tension between the different markers probably undermines the tree's reliability. Having said this, we feel this is an area where further work would be beneficial.

The general failure to uncover clear relationships between levels of genetic diversity and rookery size or population trend is perhaps not surprising. On the one hand, the levels of diversity that we find in this species appear rather low compared with other pinnipeds, arguably making it more difficult to resolve changes in diversity that might be linked to demography. At the same time, the main disjunction within the metapopulation is between the Asian/Western and the Eastern stocks, which are possibly even two subspecies (Phillips *et al.* in press), implying that within stocks there is appreciable gene flow among the different breeding colonies. Such gene flow will tend to mask or eliminate any possible differences in diversity that might otherwise result from the declines and expansions that have been documented over the last few decades. Moreover, the timescale of change is rather brief relative to the rate at which diversity is either lost or may accumulate, being of the order of only a few generations. Thus, when Bickham *et al.* (1998) used mtDNA to look for a loss in diversity between 1976 and 1978 and the 1990s in populations of the Central Gulf of Alaska, no significant differences were found, suggesting that reductions in population size over this period probably had a negligible effect on genetic diversity at this marker. However, the sample size used was small, including only 36 samples from the 1970s.

Levels of genetic diversity in the Steller's sea lion

In terms of AFLP diversity in particular, the Steller's sea lion seems to carry very low diversity. To place this observation in a broader context, we first attempted to compare our value with those reported for a range of

vertebrates by Milot *et al.* (2007). However, interspecific comparisons of AFLP diversity are not straightforward. First, relatively few studies give the proportion of the total number of AFLP loci amplified that are polymorphic. Second, biases may arise from the use of different restriction enzymes (*TaqI/EcoRI* produces more polymorphic banding patterns in mammals, Ajmone-Marsan *et al.* 1997) or selective primers (those containing a CG motif at their 3' end tend to amplify a higher proportion of polymorphic fragments, Bensch & Akesson 2005; Milot *et al.* 2007). Third, some studies sample over wider geographical ranges than others, potentially capturing greater genetic diversity, and sample sizes also vary. Inter-observer variation may also be important, although Bonin *et al.* (2004) showed that even with only very limited overlap in the specific bands scored by different observers, the underlying phylogenetic signal remains much the same. However, Milot's use of $P_{5\%}$ at least partly addresses these issues. For our data set, $P_{5\%}$ is 5.9%, which is low and comparable with the lowest values reported by Milot, those for the wandering (5.1%) and Amsterdam (2.1%) albatrosses which have undergone severe population bottlenecks. However, our measure of AFLP diversity may underestimate the true level of genetic diversity because we excluded 18 polymorphic loci that could not be scored reliably across all of the samples and which, if included, would raise our value to 21.5%. This represents another potential problem in comparing values of diversity from different studies. Nonetheless, an allozyme study of the Steller's sea lion showed an almost complete lack of genetic variability (Lidicker *et al.* 1981). Since protein electrophoretic studies largely utilize relatively conservative nuclear housekeeping genes, they are more likely to produce results comparable to an AFLP analysis than nuclear microsatellites or mtDNA.

An interesting observation is the contrasting pattern of diversity between the two main stocks for nuclear and mitochondrial markers, the eastern stock carrying the greatest mtDNA diversity but the least nuclear diversity. Several hypotheses can be advanced for why this might be so. First, an appreciable component of the modern patterns will relate to what happened during and immediately after the last ice age. If the eastern stock lies closest to whatever refugia existed, the general tendency for pinnipeds to exhibit maternally directed site fidelity might have caused the western stock to have been founded by only a subset of mitochondrial lineages, despite receiving most of the nuclear diversity due to higher levels of male-mediated gene flow. A more general version of this concept would be to state that while nuclear diversity tends to reflect total population size within a stock, mitochondrial diversity may instead be linked to the number and stability of breeding colonies. Consequently, any demographic changes that reduce population size but not colony number will tend to erode nuclear more than mitochondrial variability, while

local events that eliminate some colonies while allowing others to expand may have the converse effect. One further possibility that should not be discounted, particularly in a diving mammal where energy management is paramount, is that the mitochondrial genome could have at some point come under natural selection. Finally, our results are by no means unprecedented. For example, contrasting patterns of diversity were also found using AFLPs and mtDNA in the sonoma tree vole, with no population structure revealed by the former, but two distinct lineages uncovered using the latter (Blois & Arbogast 2006). Here, genetic diversity was much higher for the mtDNA, re-emphasizing the benefit of using multiple markers to guard against any one yielding an unexpected/misleading pattern.

Ours is one of rather few studies that have examined patterns of genetic diversity across the entire range of a widely distributed species. It is generally recognized that higher levels of genetic diversity usually occur towards the centre of a species' range (e.g. Arnaud-Haond *et al.* 2006 and Schwartz *et al.* 2003), a pattern also seen in our data. The reasons for such a pattern are multiple. Ficetola *et al.* (2007) found that distance from glacial refugia and geographical isolation together explain over 90% of variation in microsatellite diversity in the frog *Rana latastei* in northern Italy, suggesting a major impact of sequential bottlenecks and/or founder events. Similar patterns are seen even among modern humans, reflecting loss of diversity as we moved out of Africa to colonize the world (Manica *et al.* 2005; Prugnolle *et al.* 2005). Interestingly, song sparrows distributed along the Aleutian chain and hence overlapping with the Steller's sea lion distribution also reveal a stepwise loss of microsatellites diversity, apparently due to founder events as the species moved from island to island (Pruett & Winkler 2005).

Steller's sea lions appear to have unexpectedly low levels of genetic diversity, and one plausible explanation is a population bottleneck. However, applying the program Bottleneck, we failed to find any evidence of a recent severe reduction in population size. This largely supports other studies where although some species that have experienced a documented bottleneck such as the northern elephant seal have low diversity (Hoelzel *et al.* 1993), other species that were hunted to a similar or greater extent, such as many species of fur seal, currently have the highest levels of diversity seen among pinnipeds (e.g. Hoffman *et al.* 2003) and seem unaffected by sealing (e.g. Matthee *et al.* 2006). Indeed, one of the only species where a recent anthropogenic decline resulted in a detectable loss of diversity, verified by analysis of both pre- and postbottleneck samples, is the Mauritius kestrel, and this species declined to a single pair. Consequently, it seems likely that the patterns of diversity seen in modern populations will be dominated by longer-term demographic trends and have little to do with modern trends.

Interspecific comparisons

Even though short-term population trends appear to impact little on genetic diversity, the same may not be true of longer-term trends. Consequently, we examined a broad range of pinniped species to test whether current population size predicts diversity across the pinnipeds. Perhaps surprisingly, we find that while mtDNA and microsatellite diversity do not correlate significantly with population size after phylogenetic correction, AFLP markers do. The reason for the stronger relationship with AFLP markers is unclear but may relate to the relative rates of evolution of the three markers. Many of the largest changes in numbers have occurred recently and have been quite dramatic, with species of elephant seal, fur seal and sea lion having been exploited to near extinction and then rebounding. There are good reasons for believing that even these extreme histories will have reduced levels of diversity rather little, but any effects that are visible, both in terms of loss and regain of diversity will be most apparent in the fastest evolving markers which exhibit highest diversity, that is, mtDNA and microsatellites. Such markers are therefore more likely to be out of mutation-drift equilibrium, perhaps to some degree scrambling the relationship between diversity and current size. It would be interesting for future studies to ask whether stronger correlations could be obtained by using sighting data to reconstruct likely population histories for each species and then to allow for these in the estimation of current size. The strong result obtained for AFLP markers is also surprising because our sample sizes were small, at only five individuals per species. In terms only of assessing variability, the few individuals are in part compensated for by the scoring of large numbers of loci, although larger sample sizes would likely refine our estimates and, if anything, strengthen the AFLP result further. A bigger issue is likely to be whether five individuals can really represent a species across its entire range and possible population subdivisions. Our results surprisingly suggest it can, both from the strength of the regression of diversity on population size, and from the lack of structure seen within the Steller's sea lion. Clearly, this is an area where further study is warranted.

Comparing microsatellite and AFLP markers, we find a much stronger relationship with modern population size for the AFLPs. This is unexpected because many pinniped populations have in recent times experienced dramatic fluctuations due to hunting and habitat loss, and it seems logical that the faster evolving microsatellites would better track these changes. One possible explanation is that posthunting modern population sizes may have in many cases re-attained carrying capacity and hence approximate historical levels (e.g. Hodgson *et al.* 1998). If so, the slowly evolving AFLP markers may, through lack of response to rapid demographic change, exhibit a stronger correlation

than microsatellites, which suffered a larger displacement from equilibrium and are still catching up. An alternative explanation is suggested by a recent observation that, across diverse human populations, microsatellite length is strongly predicted by heterozygosity (Amos *et al.* 2008). Interpreted as support for a model in which heterozygote genotypes are more mutable than equivalent homozygotes, this study suggests that for microsatellites at least, a simple relationship between heterozygosity and population size may not exist. Such a model might also help to explain why otariids have high microsatellite diversity because in this group hybridization between sister species is not unusual. Under heterozygote instability, the large increase in heterozygosity caused by hybridization would feed back to increase microsatellite mutation rate and hence diversity.

Conclusion

Genetic diversity is widely accepted as an important component of fitness and rare alleles can easily be lost following population decline. We find that the Steller's sea lion has unusually low diversity even compared with related species, with potential management implications. However, despite rapid declines in population size, particularly in rookeries of the western stock, we failed to find significant trends between demography and genetic diversity. This does not mean that such trends are absent, but detection will require larger sample sizes collected over a longer time period. In the meantime, the best way to prevent further erosion of variability is probably through active measures of intervention designed to prevent disappearance of key rookeries such as in the western Aleutian Islands where populations seem to be highly vulnerable.

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This paper is one in a series resulting from a long-term study (since 1992) on the population genetics and systematics of Steller's sea lions sponsored mainly by NMFS. Joe Hoffman specialises in the molecular ecology of natural vertebrate populations and is also interested in the population genetics of a variety of Antarctic marine organisms. Kanchon Dasmahapatra is a molecular ecologist conducting research in speciation, phylogenetics and conservation genetics. Bill Amos runs the Molecular Ecology group and has a long-standing interest in understanding the distribution of variability in natural populations. His favorite wine of the moment is Chateau Musar 1991, outstanding! Caleb Phillips is interested in molecular evolution and phylogeography. Tom and his research group are responsible for investigating various research questions pertaining to the declining populations of Stellar's sea lions contribute to a wide assortment of projects involving animal foraging behaviour, demographics, and abundance estimation to collectively address these questions and provide fisheries management direction. John Bickham's research interests focus on genetic mutations and how they are produced and transmitted in individuals, populations, species and the evolutionary processes that affect genetic change. His current research projects include population genetics of Stellar's sea lions and bowhead whales, biodiversity studies in bats and ecotoxicological studies in contaminated environments in Azerbaijan.
