



# Determinants of genetic variation across eco-evolutionary scales in pinnipeds

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**The effective size of a population ( $N_e$ ), which determines its level of neutral variability, is a key evolutionary parameter.  $N_e$  can substantially depart from census sizes of present-day breeding populations ( $N_c$ ) as a result of past demographic changes, variation in life-history traits and selection at linked sites. Using genome-wide data we estimated the long-term coalescent  $N_e$  for 17 pinniped species represented by 36 population samples (total  $n = 458$  individuals).  $N_e$  estimates ranged from 8,936 to 91,178, were highly consistent within (sub)species and showed a strong positive correlation with  $N_c$  ( $R^2_{adj} = 0.59$ ;  $P = 0.0002$ ).  $N_e/N_c$  ratios were low (mean, 0.31; median, 0.13) and co-varied strongly with demographic history and, to a lesser degree, with species' ecological and life-history variables such as breeding habitat. Residual variation in  $N_e/N_c$ , after controlling for past demographic fluctuations, contained information about recent population size changes during the Anthropocene. Specifically, species of conservation concern typically had positive residuals indicative of a smaller contemporary  $N_c$  than would be expected from their long-term  $N_e$ . This study highlights the value of comparative population genomic analyses for gauging the evolutionary processes governing genetic variation in natural populations, and provides a framework for identifying populations deserving closer conservation attention.**

Biodiversity is declining at an alarming rate<sup>1</sup>. Intraspecific genetic variation provides the raw material for adaptive evolution and is an important component of biodiversity, as reflected by targets in the international treaty for the conservation of biodiversity<sup>2</sup>. Genetic variation is continuously generated by novel mutations entering a population, but is eroded by purifying selection and random genetic drift. The expected rate of loss of neutral or nearly neutral genetic variability is inversely related to the classic population genetic metric known as the effective population size, or  $N_e$  (refs. <sup>3,4</sup>).

In long-term stable populations,  $N_e$  is expected to scale proportionally with the observed number of potentially breeding individuals ( $N_c$ ) in the population<sup>5</sup>. Nevertheless, across taxonomic scales the empirical relationship between the level of genetic variability and census population size is often weak, and  $N_e$  tends to be orders of magnitude lower than  $N_c$  (ref. <sup>6</sup>). Several explanations for this observation, often referred to as Lewontin's paradox<sup>7</sup>, have been proposed<sup>8</sup>. One explanation involves the apparent inverse relationship

between mutation rate and population size<sup>9</sup>, resulting in a diminishing returns relation between genetic diversity and population size. Another explanation invokes selection at linked sites whereby both the spread of beneficial mutations and the removal of deleterious alleles reduce genetic variation of closely linked neutral polymorphisms<sup>10</sup>. Both forms of selection at linked sites, genetic hitch-hiking and background selection, have been shown to constrain neutral diversity, with the diversity-reducing effect being more pronounced the larger a population<sup>11,12</sup>. Another explanation for the common empirical disconnect between  $N_e$  and  $N_c$  can be sought in factors influencing the amount of genetic drift. In the absence of selection at linked sites, these include constitutive factors such as the species' mating system or other life-history traits introducing reproductive skew<sup>13</sup>, as well as demographic perturbations reflecting historical deviations from constant population sizes.

Estimates of  $N_e$  can be contrasted to  $N_c$ , with the ultimate goal of predicting one from the other<sup>14</sup> or identifying populations with a low  $N_e/N_c$  ratio, which, although a contentious issue<sup>15</sup>, has been

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related to a population's adaptive potential<sup>16,17</sup>.  $N_e/N_C$  can vary substantially among species with differences in life-history traits, such as the sex ratio. Factors increasing reproductive skew tend to result in lower  $N_e$  relative to current census sizes<sup>5,18,19</sup>. There are multiple approaches to estimating  $N_e$ , which differ in important aspects. Non-genetic methods based on demographic data<sup>5</sup> and pedigrees<sup>20</sup> generally provide a time-sensitive estimate over the course of a single or few generations. Methods using molecular data exploiting excess in heterozygosity over Hardy–Weinberg expectation, temporal sampling or linkage disequilibrium<sup>21</sup> integrate over slightly longer timescales, ranging from several dozen generations<sup>22</sup> to several thousand (although « $N_e$ » generations<sup>23</sup>).

In contrast, estimates of the coalescent effective population size capture the interaction of evolutionary forces shaping genetic variation over extended periods in the order of  $4N_e$  generations, which corresponds to the expected time to the most recent common ancestor of the alleles sampled from (diploid) current-day individuals. In vertebrates, this time frame spans tens of thousands to millions of years<sup>23–25</sup>. The coalescent  $N_e$  scales with the harmonic mean of a (rapidly fluctuating) sequence of values of effective sizes of each generation back to the most recent common ancestor. It thus encapsulates factors that have shaped the ancestral genetic variation that is still observed in a contemporary population sample<sup>26</sup>. Consequently, genomic data sampled from putatively neutral sites of remnant individuals provide a simple means for obtaining diversity ( $\theta_\pi$ )-based estimates of the coalescent  $N_e = \theta_\pi / 2c\mu$  if the mutation rate,  $\mu$ , and ploidy,  $c$ , are known. This relationship assumes that genetic diversity of the species' populations approximates an equilibrium between mutation and genetic drift, and holds within the limits of the infinite sites mutation model that is applicable for most species, except for 'hyper-diverse' organisms such as viruses<sup>27</sup>. It allows the investigation of processes affecting genetic diversity in the past and explores their relationship to current population sizes.

Investigation of the relationship between  $N_e$  and  $N_C$  has mostly been limited to single-species estimates or meta-analyses across large taxonomic scales<sup>6,11</sup>. Contrasting diversity patterns among closely related species, however, allows the inference of relevant factors shaping genetic diversity while circumventing phylogenetic inertia that may hamper the interpretation of the relevant life-history differences. Pinnipeds, comprising true seals, sea lions, fur seals and walruses, are a group of marine mammals that display considerable variation in life-history traits and breeding habitats<sup>28</sup>. Pinniped population abundances have fluctuated in response to changes in sea surface temperatures, as these are thought to impact the availability of prey and the location and extent of suitable breeding areas (for example, refs. <sup>29,30</sup>). Populations of many pinniped species were substantially reduced by commercial sealers from the late eighteenth to early twentieth century and which, coupled with climate change, has led to almost 50% of recognized taxa being considered threatened or endangered<sup>31</sup>.

Here we estimate coalescent  $N_e$  across pinniped species, assess the impact of ecological factors and population-specific demographic fluctuations and discuss the roles of drift and selection at linked sites in shaping diversity estimates. We compared genome-wide estimates of the coalescent  $N_e$  in 17 pinniped species to current  $N_C$  estimates, and inferred how this ratio is influenced by life-history traits and demographic perturbations preceding anthropogenic impact. The study has fundamental implications for understanding the influence of a species' life history and historical processes on contemporary genetic diversity and population size. It provides a basis to inform present-day conservation initiatives aimed at maintaining or restoring  $N_C$  and  $N_e$ .

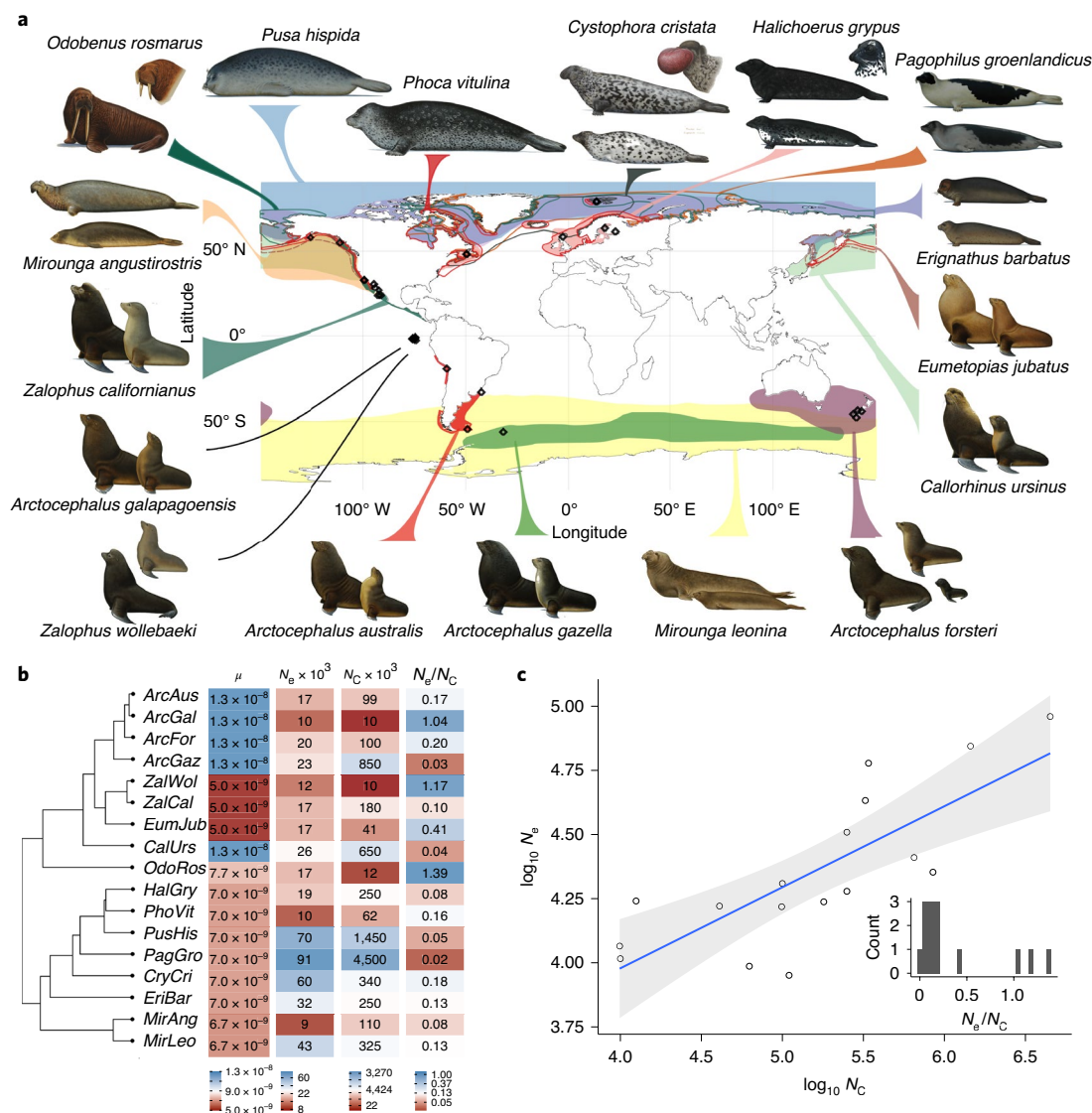
## Results and discussion

**Mutation rate  $\mu$ , genetic variation  $\theta_\pi$  and  $N_e/N_C$  ratio.** We generated double-digest restriction site-associated DNA (ddRAD) data for a total of 458 individuals sourced from 36 populations of 17 species

(20 taxa, including subspecies) distributed across the pinniped phylogeny (Fig. 1a,b). From an average of 31,956 (range 11,826–45,438) mapped loci per population remaining after rigorous filtering, we estimated the basic population genetic summary statistics Watterson's theta ( $\theta_w$ ), nucleotide diversity ( $\theta_\pi$ ) and Tajima's  $D$ . In addition, we inferred genome-wide mutation rates using 8,864 orthologous genes extracted from draft genome sequences of representatives for the three pinniped families: Otariidae (Antarctic fur seal, Californian sea lion), Odobenidae (walrus) and Phocidae (Weddell seal, Hawaiian monk seal) (Fig. 1b and Supplementary Table 1; see Methods for accession numbers). Mutation rate estimates of  $\mu$  per base pair (bp) and generation ranged from  $5.00 \times 10^{-9}$  to  $1.32 \times 10^{-8}$  ( $0.4$ – $1.21 \times 10^{-9}$  bp<sup>-1</sup> year<sup>-1</sup>), which is comparable to other mammalian lineages<sup>9</sup> including humans<sup>32</sup>. The similarity of these estimates, with no apparent effect of population size, precludes a strong influence of mutation rate on the variation in  $N_e$  across species.

Using the relationship  $N_e = \theta_\pi / 4\mu$ , we inferred  $N_e$  for each population separately (Supplementary Table 2). These resulting estimates ranged from 8,936 to 91,178 (median, 18,993) and were thus comparable to values for other large mammalian species, including humans<sup>33,34</sup>. We were interested in obtaining a global estimate of  $N_e$  for the entire (sub)species while facing the empirical limitation of access to local population sample(s). Under a broad set of migration models, theory suggests that the mean within-population diversity is most appropriate for comparison of properties across species<sup>4</sup>. In species with strong population structure, however, the mean within-population diversity may deviate markedly from the pooled population estimate, as quantified by  $F$ -statistics. A literature review suggests that the fixation index,  $F_{ST}$  is generally low for most of these species, findings supported in this study for species where multiple populations were available (Supplementary Table 3). Therefore, single-population samples are expected to contain robust information on genetic variation relevant at the level of the (sub)species. In line with this prediction,  $N_e$  estimates were consistent among population samples with little or no genetic differentiation (coefficient of variation (CV): New Zealand fur seal, *Arctocephalus forsteri* (*ArcFor*), 0.002; Galápagos sea lion, *Zalophus wollebaeki* (*ZalWol*), 0.002; Supplementary Table 3 and Extended Data Figs. 1–3). These estimates were also similar among moderately stratified populations (South American fur seal, *Arctocephalus australis* (*ArcAus*), 0.005; California sea lion, *Zalophus californianus* (*ZalCal*), 0.012), but could differ substantially among formally recognized subspecies with a high degree of evolutionary independence (ringed seal, *Pusa hispida* (*PusHis*), 0.029). We thus concluded that  $\theta_\pi$ -based  $N_e$  estimates approximated the amount of long-term genetic drift experienced by the (sub)species as a whole. Consequently, we extrapolated the 'species'  $N_e$  from the population best representing the core distribution of the nominate subspecies. We note that this estimate will differ from the mean value over all interconnected subpopulations<sup>35</sup> and will accordingly introduce additional variance in  $N_e/N_C$  estimates.

As a complement to the above approach, we calculated the harmonic mean of  $N_e$  estimates obtained as the rate of coalescence using multiple sequentially Markovian coalescent analyses for a single genome (PSMC)<sup>36</sup> where genome assemblies of sufficient quality were available. PSMC'-based values were similar to the ddRAD-derived estimates for the California sea lion, *ZalCal* (18,057 versus 16,398–22,766) and higher for the Antarctic fur seal, *Arctocephalus gazella*, *ArcGaz* (29,133 versus 22,550). Higher values for the PSMC'-based estimates are consistent with population structure decreasing the rate of coalescence, although this will depend on the timing and level of gene flow and population size changes<sup>37</sup>. For walrus, *Odobenus rosmarus* (*OdoRos*), PSMC' was conducted on an individual from the Pacific subspecies (*O. r. divergens*;  $N_C = 100,000$ ), whereas ddRAD estimates were based on a population sample from

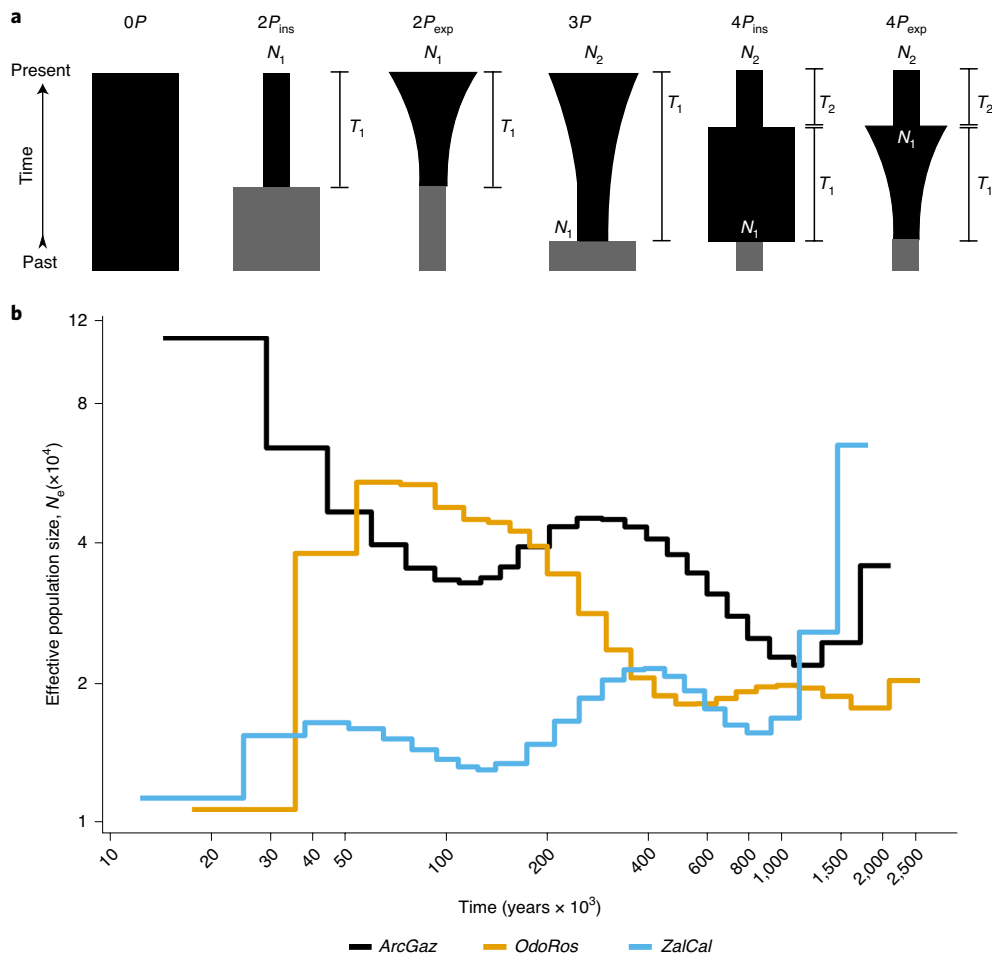


**Fig. 1 | Sampling set-up and core population parameters of the 17 pinniped species investigated in this study. a**, Sampling locations and geographic ranges of focal seal species and populations. **b**, Phylogenetic relationships among focal species following Higdon et al.<sup>48</sup>. Species abbreviations are the first three letters of both genus and species names. Mutation rate estimates,  $\mu$  ( $\text{bp}^{-1}$  per generation) derived from five genomes were used for all species within a clade. Population genetic summary statistics and estimates of census population sizes and  $N_e/N_c$  shown in the heatmap are based on the nominal subspecies and populations representing the core species range. **c**, Relationship between  $N_e$  and  $N_c$  shown on a logarithmic scale (base 10). The blue line depicts the regression line, the shaded area its 95% confidence intervals. The inset shows the frequency distribution of binned  $N_e/N_c$  ratios. Illustrations in **a** courtesy of Brett Jarrett; map retrieved from IUCN<sup>74</sup>.

the Atlantic subspecies (*O. r. rosmarus*;  $N_c = 12,500$ ). Despite the substantial difference in current-day  $N_c$ , estimates of  $N_e$  were similar across the two subspecies and methods (19,808 versus 15,491), which suggests that walrus in the Atlantic might still share substantial genetic variation with those in the Pacific, a finding consistent with mitochondrial results<sup>38</sup>. Overall, these results support the previous notion that the coalescent  $N_e$  contains information on the effects of long-term genetic drift, or selection at linked sites, shared among populations within species<sup>25</sup>.

Selection at linked sites probably affects  $N_e$  (see Main); however, in-depth investigation of its effects would require detailed knowledge of local mutation and recombination rates along the genomes of all species considered here. To assess the magnitude of the possible effect, we quantified the potential target size for selection (approximated by protein-coding genes) for walrus where both an annotated reference genome and ddRAD data were available. In total, 43.5% of

ddRAD loci in this species overlapped at least partially with annotated genes. Consistent with the diversity-reducing effects of selection both against amino acid substitutions and at linked sites, estimates of nucleotide diversity,  $\theta_\pi$ , within genic regions were lower than in intergenic regions (0.0005197 and 0.0005504, respectively; genome-wide mean, 0.0005371). This translates to a change in  $N_e$  from 16,984 in genic regions to 17,987 in intergenic regions (genome-wide, 17,551). The small magnitude of this difference is consistent with comparative analyses suggesting moderate effects of selection at linked sites for species with a relatively low  $N_e$ , such as pinnipeds<sup>11</sup>. It is also compatible with the proposition that, in the relatively large mammalian genomes, much intergenic sequence is far from selection targets. Within the range of  $N_e$  across species, changes that may be attributable to selection at linked sites are thus expected to be minor and are unlikely to affect the relationships with  $N_c$ , demographic change and the life-history variables investigated below.



**Fig. 2 | Historical changes in population size.** **a**, Demographic scenarios including (1) a null model of constant population size ( $0P$ ), (2) a single, instantaneous ( $2P_{ins}$ ) or gradual ( $2P_{exp}$ ) population size change, (3) an instantaneous population size change with subsequent gradual increase or reduction of population size ( $3P$ ) and (4) models incorporating two independent instantaneous ( $4P_{ins}$ ) or gradual ( $4P_{exp}$ ) population size changes. The magnitude of change is parameterized as  $N_1$  and  $N_2$  (size relative to ancestral population size  $N_{anc}$ , depicted in grey). The timepoints of change are accordingly parameterized as  $T_1$  and  $T_2$ . **b**, Changes in effective population size ( $N_e$ ) over time as inferred by PSMC' shown on a logarithmic scale.

Next, we compiled estimates for the number of breeding individuals,  $N_C$ , for each subspecies and explored their relationship to  $N_e$  (Fig. 1c, Extended Data Fig. 4 and Supplementary Table 2). Despite uncertainty in both estimates, the correlation was remarkably strong, explaining about 60% of the overall variance ( $P=0.0002$ ,  $R_{adj}^2=0.59$ ). This indicates that both independently derived variables share a substantial amount of information on species abundance. The  $N_e/N_C$  ratio was consistently below 1.0, with a median of 0.13 (mean, 0.31; exceptions: Galápagos sea lion (*ZalWol*), 1.17; Atlantic walrus (*OdoRos*), 1.39). The inferred  $N_e/N_C$  ratio was strikingly similar to published median estimates of 0.11 (ref. <sup>5</sup>) and 0.14 (ref. <sup>18</sup>) derived across a much larger taxonomic range and based on the temporal method, which capitalizes on the idea that variance in neutral allelic frequencies across generations is inversely proportional to  $N_e$ . Frankham<sup>5</sup> identified life-history parameters increasing the variance in reproductive success and population oscillation as the main factors responsible for low  $N_e/N_C$  ratios. Non-genetic approaches estimating  $N_e/N_C$  ratios on the basis of age-specific survival and fecundity rates have likewise identified life-history traits as the main contributor to reduced  $N_e$  estimates<sup>19</sup>. Despite the apparent similarities between  $N_e/N_C$  ratios obtained using these approaches and this study, they differ in terms of timescale and the information they carry.  $N_e$  estimates based on vital rates or allelic composition shifts integrate at most a handful of generations, and

thereby are insensitive to population oscillations across larger timescales. The estimate used in this study scales in coalescent units of  $4N_e$  generations which, in our dataset, translates to a median of approximately 1,584,549 years (range, 310,995–5,726,017 years). Assuming detailed information of dynamic changes in  $N_C$  across all generations in the past, the harmonic mean of  $N_C$  would similarly accommodate demographic change across a species' history<sup>39</sup>. Nevertheless, present-day estimates of  $N_C$  reflect only a snapshot of population history: we therefore expect a disproportionate role of past population oscillations on  $N_e/N_C$  through a reduction in  $N_e$ .

**Demographic changes.** To quantify past population size changes, we formulated a set of possible demographic models for each population, ranging from constant population size to the incorporation of two independent size changes (Fig. 2a). We then calculated their likelihood based on the diffusion approximation to the allele frequency spectrum<sup>40</sup>. Parameter estimates for the model with the best fit are reported in Fig. 3. Similar to estimates of  $N_e$ , model inference was highly consistent among interconnected populations with a shared evolutionary history. For example, for all eight populations of the Galápagos sea lion *ZalWol*, we inferred an exponential reduction to an average of 0.26 of the ancestral population size (range, 0.09–0.42; CV=0.39). For all four New Zealand fur seal *ArcFor* populations, the three-parameter model was preferred,

Species	Population	Ind.	Model	$N_1$	$N_2$	$T_1$	$T_2$	Taj's $D$
ArcAus	Peru	8	2Pins	0.002		0.000		0.334
	Brazil	11	3P	14.046	0.405	0.142		0.007
ArcGal	Isabela	12	3P	1.185	0.496	0.438		0.197
ArcFor	CFoulwind	8	3P	73.800	0.545	0.723		-0.026
	OBay	7	3P	54.878	0.609	0.568		-0.139
	OhauP	6	3P	42.943	0.470	0.456		-0.033
	VictoryB	6	3P	53.466	0.682	0.554		-0.178
ArcGaz	BirdIs	84	4Pins	22.785	0.305	6.154	0.017	-0.062
ZalWol	Baltra	10	2Pexp	0.330		0.516		0.311
	Espanola	10	2Pexp	0.107		0.039		0.312
	Fernandina	20	2Pins	0.088		0.007		0.413
	Genovesa	9	2Pexp	0.292		0.249		0.225
	IsabelaB	15	2Pexp	0.319		0.127		0.353
	IsabelaV	10	2Pexp	0.418		0.481		0.202
	Pinta	10	2Pexp	0.223		0.090		0.367
	SantaFe	9	2Pins	0.310		0.077		0.293
ZalCal	GCal1	12	3P	10.804	0.971	1.176		-0.010
	GCal2	13	2Pexp	2.238		0.171		-0.266
	GCal3	23	2Pins	12.613		0.073		-0.119
	GCal4	14	2Pexp	56.806		0.984		-0.110
	SMargarita	14	2Pins	0.012		0.000		0.203
	SMiguel	9	2Pins	4.023		0.234		-0.555
EumJub	GrassyIs	8	2Pins	0.308		0.352		0.018
CalUrs	SMiguel	8	2Pins	2.803		0.554		-0.576
OdoRos	Svalbard	12	3P	51.251	0.672	0.487		-0.208
HalGry	Scotland	3	2Pexp	0.008		0.138		-0.096
	StLawrence	6	2Pexp	0.067		0.346		0.158
PhoVit	Svalbard	13	4Pexp	1.237	1.292	0.180	0.035	-0.237
PusHis	Baltic	9	3P	41.192	0.808	0.412		-0.394
	Saimaa	10	2Pins	0.131		0.265		0.412
	Svalbard	14	3P	0.921	4.437	0.546		-0.664
PagGro	StLawrence	9	3P	32.197	1.653	0.419		-0.652
CysCri	Svalbard	8	3P	59.262	1.187	0.588		-0.511
EriBar	Svalbard	12	2Pins	0.858		0.399		-0.029
MirAng	NAtlantic	15	3P	0.039	0.534	0.155		0.066
MirLeo	SAtlantic	11	2Pexp	5.083		0.266		-0.299

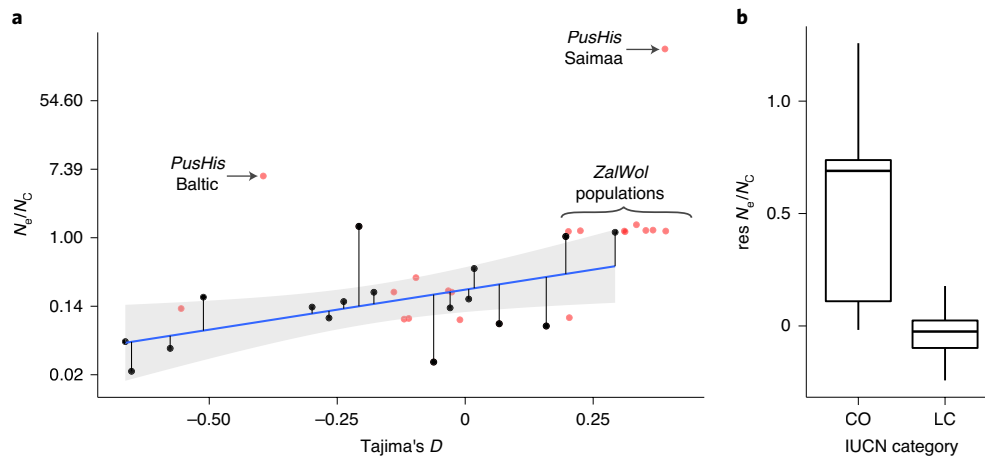
**Fig. 3 | Demographic parameter estimates.** Summary of parameter estimates for demographic population histories according to the model best fitting the data for each pinniped species and population. For parameter nomenclature, see Fig. 2a.  $N_1$ ,  $N_2$ , Tajima's  $D$ : blue background colour indicates population growth, red population decline. Ind represents the number of samples used for the analysis, with populations used for subsequent model analysis (see Fig. 2a) shown in grey.

portraying an initial 52-fold increase in population size ( $N_1$  range, 42–74;  $CV=0.22$ ) followed by a bottleneck ( $N_2$  range, 0.47–0.68;  $CV=0.15$ ). Demographic reconstructions were also similar in more stratified populations: subspecies of the ringed seal *PusHis* showed signatures of a large population reduction in both the Saimaa ringed seal (*P. h. saimensis*,  $N_1=0.13$ ) that colonized Lake Saimaa at least 9,500 years ago<sup>41</sup> and the Baltic sea population (*P. h. botnica*,  $N_1=41.19$ ,  $N_2=0.81$ ), although in the Baltic this was preceded by a large population increase.

Single-population estimates of the demographic history are probably an oversimplification because they ignore the confounding effect of population structure and gene flow<sup>42</sup> (Extended Data Figs. 1–3). Nevertheless, in the absence of comprehensive sampling of all possible populations for each species, this is the only operational way forward. Moreover, PSMC' analyses, based on individual genome assemblies that assume panmixia, were broadly consistent with the demographic scenarios inferred from ddRAD population data (Fig. 2b). In walrus *OdoRos*, both analyses showed a population increase followed by a large decline to below the ancestral population size (PSMC' time estimate  $\approx 500,000$ –40,000 years ago, estimate

from  $\delta a \delta i$  software  $T_1=0.49$ ), despite using data from distinct subspecies (Fig. 2b and Fig. 1). In the Antarctic fur seal *ArcGaz*, both methods recovered an increase in population size followed by a more recent decrease (PSMC' time estimate  $\approx 1,000,000$ –100,000 years ago,  $\delta a \delta i$   $T_1=6.1$ ). PSMC' then recovered a successive population increase. For the California sea lion *ZalCal*, the demographic histories inferred from the ddRAD data covered different timescales ( $T_1$ , Fig. 1), with some populations showing greater population growth than the PSMC' (Fig. 2b). A multi-population model incorporating migration might ultimately improve demographic estimates for the species and provide a better comparison to the PSMC'.

Tajima's  $D$  is a summary statistic that contains information on the demographic history of a population by comparing two measures of genetic diversity,  $\theta_\pi$  and  $\theta_w$ . The latter is more sensitive to the presence of rare alleles, which accumulate following population expansion. Assuming neutrality, negative values of Tajima's  $D$  are consistent with a scenario of recent population expansion whereas positive values indicate a recent period of demographic contraction. Tajima's  $D$  co-varied significantly with our estimates concerning recent population size changes  $N_1$  and  $N_2$  ( $P=0.0002$ ,  $R_{adj}^2=0.59$ )



**Fig. 4 | The relationship between  $N_e/N_c$  ratio and Tajima's  $D$ .** **a**, Relationship between  $N_e/N_c$  and Tajima's  $D$  reflecting the demographic history of species. Note that the regression line (blue) refers only to the depicted relationship and does not consider the influence of other predictor variables. The grey shaded area depicts 95% confidence intervals of the regression line. Black points, populations used for model inference; red points, additional subspecies and populations. The y axis follows a natural logarithmic scale. **b**, Relationship between the residual variance from **a** ( $\text{res } N_e/N_c$  on y axis) and IUCN Red List category. Residual values per species (black points in **a**) were grouped into categories of concern (CO,  $n=5$ ) or least concern (LC,  $n=12$ ).

and, hence, captured basic information of the explicit demographic models. Using Tajima's  $D$  as a proxy for demographic history, we investigated whether demographic processes were shared among species with similar ecological niches and life histories. Variables analysed included breeding habitat (ice or land), species range, length of breeding season and reproductive skew based on harem size (Supplementary Table 2). Stoffel et al.<sup>22</sup> suggested that recent human-induced bottlenecks are related to the mating system and breeding habitat in pinnipeds, with bottlenecks linked to dense agglomerations in polygynous species breeding on land. In our models, breeding habitat likewise received statistical support (cumulative weight of evidence ( $wAIC_c$ )=0.59; Supplementary Table 4). This similarity with Stoffel et al.<sup>22</sup> is surprising considering the different timescales investigated. Stoffel et al.<sup>22</sup> based their inferences on summary statistics from microsatellite data exploring recent demographic changes in the range of hundreds of years, as opposed to the scale of several hundred thousand years considered in this study. In contrast to Stoffel et al.<sup>22</sup>, mating system had a relatively small impact as expressed in Akaike's weights (mating system  $wAIC_c=0.14$ ). Models including a species' geographic range size were more strongly supported on average ( $wAIC_c=0.85$ ; Supplementary Table 4); Tajima's  $D$  was negatively correlated with species range ( $P=0.008$ ,  $R^2_{\text{adj}}=0.34$ ), implying that species with large current distributions were more likely to have experienced a population expansion in the past. Overall, these results motivate two key interpretations. First, a species' ecological condition and life history affects its demographic stability through evolutionary time. Second, present distribution ranges contain information on past demographic processes—just as current species abundance contains information on long-term population sizes in the order of  $4N_e$  generations (see Fig. 1c). On a cautionary note, our estimate of long-term effective population size,  $N_e$ , using the relationship  $N_e=\theta_\pi/4\mu$ , is based on the expectation of stationary allele frequency distributions; hence, it assumes that populations have been approximately stable or had sufficient time for mutations to accumulate and reach equilibrium. In pinnipeds, recent human-induced bottlenecks<sup>22</sup> and demographic changes (this study) challenge this assumption. Despite the potential deviation from the equilibrium assumption,  $N_e$  estimates were correlated with estimates of population size ( $N_1$  or  $N_2$ ) from the demographic models ( $P<0.05$ , with phylogenetic correction  $P<0.001$ ), supporting the notion that  $N_e$

estimates based on  $\theta_\pi$  constitute a simple, but meaningful, measure of long-term population size.

**The effect of demographic change and life-history variables on  $N_e/N_c$ .** We next explored the relationship between  $N_e/N_c$ , demographic change as inferred from Tajima's  $D$  and life-history trait variables, including breeding habitat (ice or land), breeding latitude, length of breeding season and reproductive skew as inferred by harem size. The best-supported model, explaining 30% of the variance in  $N_e/N_c$ , included only Tajima's  $D$  ( $P=0.02$ ), which was also the best-supported predictor variable overall ( $wAIC_c=0.60$ ; Supplementary Table 5). While models including life-history parameters such as the length of the breeding season ( $wAIC_c=0.42$ ) and breeding habitat ( $wAIC_c=0.19$ ) also received some support, demographic change was the major variable explaining variation in  $N_e/N_c$  (Fig. 4a and Extended Data Fig. 5). Species inferred to have experienced recent expansions, such as the harp seal, *Pagophilus groenlandicus* (*PagGro*), had the lowest  $N_e/N_c$  ratio of 0.02, whereas both Galápagos species (*A. galapagoensis* and *Z. wollebaeki* (*ArcGal* and *ZalWol*), with genetic signatures indicative of a bottleneck, had values around 1.0. This relationship appears to run counter to the common interpretation of the  $N_e/N_c$  ratio based on short-term measures of  $N_c$ : a population with small  $N_e/N_c$  will lose genetic diversity and fix deleterious mutations faster than an equally sized population with a higher ratio. Accordingly, a low  $N_e/N_c$  value is usually interpreted as a signal for increased relative genetic risk of that population. The interpretation based on the coalescent  $N_e$  estimate used in this study is necessarily different, as  $N_e$  and  $N_c$  react on different timescales. The diversity-based measure of  $N_e=\theta_\pi/4\mu$  reflects the mean coalescent times for a sample of two (that is,  $\theta_\pi$ ). It is therefore less sensitive to very recent and rapid changes in population size than the actual size of a breeding population,  $N_c$ , which can change substantially in the course of a few generations. Hence, a  $N_e/N_c$  ratio close to 1.0, as seen for the Galápagos pinnipeds, is entirely consistent with a recent bottleneck scenario strongly reducing current-day  $N_c$  while not affecting long-term  $N_e$  (Extended Data Fig. 6).

Following this logic, we can expect that recent anthropogenic activity, such as active harvest or habitat degradation<sup>22,31</sup>, will have a strong impact on  $N_c$  but less so on the coalescent  $N_e$ . Substantial human-associated population size changes are expected to be in the range of hundreds of years, which corresponds to  $<10^{-3}N_e$

generations. Even heavy exploitation during such a time frame is not expected to leave visible traces in estimates of the coalescent  $N_e$ . Very recent changes in  $N_e$  should introduce residual variation in  $N_e/N_C$  not accounted for by the long-term demographic history of the population and by life-history characteristics of the species. To quantify this potential anthropogenic component of changes in  $N_e$ , we subtracted  $N_e/N_C$  estimates from the values predicted by the model that controlled for demographic change (the residuals from Fig. 4a,  $\text{res}N_e/N_C$ ). We reason that species with positive residuals ( $\text{res}N_e/N_C > 0$ ) have reduced  $N_e$  relative to the expectation of the model and may therefore deserve specific conservation attention. For example,  $\text{res}N_e/N_C$  of the isolated ringed seal *PusHis* subspecies reached 6.0 for the Baltic Sea and  $>220$  for Lake Saimaa. These populations have been separated for only a relatively short period from the core species range and thus harbour more (ancestral) genetic variation than would be expected by their current-day  $N_C$ . A similar pattern is seen in the Galápagos sea lion *ZalWol*, which has a disproportionately high amount of genetic variation given its  $N_C$ . The large  $\text{res}N_e/N_C$  for the small population of Atlantic walrus, *OdoRos*, similarly indicates that its relatively high genetic diversity may be reminiscent of a large ancestral population or continued cohesion with the large Pacific stock. In line with this assumption, species classified by the International Union for Conservation of Nature (IUCN) as of ‘concern’ had significantly higher  $\text{res}N_e/N_C$  values than species classified as ‘least concern’ ( $P = 0.001$ ,  $R^2_{\text{adj}} = 0.47$ ; Fig. 4b).

## Conclusions

Genetic diversity is the raw material of evolution. The coalescent  $N_e$  reflects the rate at which neutral or nearly neutral genetic variation is lost, and thus constitutes a useful measure for intraspecific biodiversity, which may contribute to adaptive potential in changing environments. Here, we demonstrated for a mammalian clade that the long-term dynamics of  $N_e$  co-vary strongly with the actual number of extant breeding individuals,  $N_C$  (Fig. 1). The strong correlation between  $N_e$  and  $N_C$  contributes to the debate on the relative roles of selection at linked sites and population size in shaping diversity (for example, refs. <sup>11,12,43</sup>), and implies a strong role for classical genetic drift in species with small to medium effective population sizes.

Both  $N_e$  and  $N_C$  were affected by historical processes, though at largely different timescales (Fig. 2 and Fig. 1). Considering absolute values,  $N_C$  exceeded  $N_e$  by one order of magnitude and showed considerably higher variation across taxa: estimates of  $N_C$  varied by up to 450-fold, whereas those of  $N_e$  were remarkably consistent, differing at most tenfold. This result fits with predictions of population genetic theory:  $N_e$  is expected to scale approximately with the harmonic mean of single-generation population size ‘snapshots’. Mean genetic diversity is thus most strongly influenced by historical periods of small population size with below-average numbers of breeding individuals. As a corollary, mean genetic diversity has reduced variance relative to  $N_C$  and is expected to be less sensitive to recent, short-term population changes during the Anthropocene<sup>44,45</sup>. Strong, recent contractions of  $N_C$  will, however, cause losses or rare variants<sup>22</sup> which, in species with moderate population sizes, may constitute an important component of adaptive potential.

When controlling for the influence of demographic oscillation, which interacted with a species’ life history, residual variation of the close relationship between  $N_e$  and  $N_C$  contained information on population vulnerability (Fig. 4). This result may serve as proof of concept in helping to identify species at risk. A low  $N_e/N_C$  ratio considered per se without invoking information on long-term population dynamics and estimates for related species is not an immediate cause for concern. On the contrary, it may point towards current-day population levels exceeding numbers of effectively breeding individuals as inscribed in the long-term genetic record of the species. However, residuals in  $N_e/N_C$  relative to other species

might flag up unaccounted population stratification or admixture<sup>46</sup> and human-mediated population crashes.

The approach used herein is readily applicable to other clades, and requires only three sets of data: estimates of mutation rate, genome-wide genetic diversity and current-day population size estimates. Mutation rate estimates, if not inferred from the target group, can generally be extrapolated from closely related species. Neither should estimation of the genetic parameters be a limitation, as it requires estimation of only two central population genetic parameters,  $\theta_\pi$  and Tajima’s  $D$ , the latter a composite of  $\theta_w$  and  $\theta_\pi$ . Accurate estimates of  $N_C$  are arguably the most difficult to obtain. Nevertheless, extracting data from the IUCN database, which is noisy and with high levels of uncertainty<sup>47</sup>, still yielded a remarkably high correlation with  $N_e$ .

## Methods

**Species and population sampling.** A total of 17 globally distributed pinniped species were collected from across the phylogenetic range<sup>48</sup> (Fig. 1). Samples were chosen to cover the two major pinniped families, Otariidae ( $n = 8$ ) and Phocidae ( $n = 8$ ), in even proportion, and to include the walrus (*O. rosmarus*), which is the only extant member of the family Odobenidae ( $n = 1$ ). Several subspecies or populations were sampled for six of the species. To attribute taxonomic and conservation status, we followed the current classification from IUCN v.2018-1. For the South American fur seal, a population sample from the subspecies *A. australis australis* from Brazil was included, and a population from Peru with contentious subspecific status<sup>49</sup>. The Steller sea lion was represented by the subspecies Loughlin’s Steller Sea Lion (*Eumetopias jubatus monteriensis*), and samples from the walrus come from the Atlantic stock *O. rosmarus rosmarus*. Samples from the grey seal were obtained from the subspecies *Halichoerus grypus grypus* in the Western Atlantic, and from *H. g. macrorhynchus* from the United Kingdom. For the ringed seal, three different subspecies from Svalbard (*P. hispida hispida*), the Baltic Sea (*P. h. botnica*) and Lake Saimaa (*P. h. saimensis*) were included. Multiple populations were sampled for *A. forsteri* (four populations) and *Z. californianus* (six populations). In addition, the *Z. wolfebaeki* dataset from Shafer et al.<sup>50</sup>, consisting of 94 individuals from eight rookeries (93 after filtering in this study), was used, with each rookery analysed separately. See Supplementary Table 6 for details.

**Reduced-representation library preparation and data processing.** Samples consisted of frozen blood, flipper or muscle tissue extracted using either an UltraClean Blood DNA Isolation Kit (MoBio) or a DNeasy Blood and Tissue Kit (QIAGEN). We generated ddRAD libraries using a modified protocol from Brelford et al.<sup>51</sup> and described in Shafer et al.<sup>50</sup> using *MseI* and *SbfI* restriction enzymes followed by size selection targeting 200–430-bp insert size. Samples were distributed across a total of ten lanes and were 125-bp paired-end sequenced on an Illumina HiSeq2500 machine. After trimming adaptors using cutadapt v.1.8 (ref. <sup>52</sup>), reads were de-multiplexed using the process\_radtags module of STACKS<sup>53</sup>. Overlapping reads were merged using PEAR 0.9.6 (ref. <sup>54</sup>) with default parameters, and all merged read-pairs of length  $>50$  bp were retained. Previous work has shown that mapping of restriction site-associated read data to genomes of closely related species yields more reliable genotypes than de-novo-based approaches<sup>55</sup>, particularly with reference to demographic inference<sup>50</sup>. At the beginning of our study, four pinniped reference genomes were available to us: walrus (*O. rosmarus*, Odobenidae, genome v.1.0, GCF\_000321225.1)<sup>56</sup>, Antarctic fur seal (*A. gazella*, Otariidae, genome v.1.4, GCA\_900500725)<sup>57</sup>, Weddell seal (*Leptonychotes weddellii*, Phocidae, genome v.1.0 GCF\_000349705.1) and a draft assembly of the California sea lion (*Z. californianus*, genome v.1.2). Reads were mapped to the closest available reference genome using STAMPY 1.0.23 (ref. <sup>58</sup>), with the substitution-rate parameter set based on the distance between each species and its closest relative with a reference genome in the phylogeny of Higdon et al.<sup>48</sup>. Because the northern fur seal, *Callorhinus ursinus*, is equally distant in this phylogeny to both the Antarctic fur seal and the California sea lion, for this species the analysis was repeated using both reference genomes. Results were highly congruent, so only the statistics based on the Antarctic fur seal genome are reported.

Sequence alignment files (.bam) resulting from the mapping procedure were filtered with a wrapper developed for this study, ClustRAD, described in Supplementary Information (see also Extended Data Fig. 7). The resulting sequence clusters were distributed across the entire genome assembly, and thus represent genome-wide genetic variation (Extended Data Fig. 8). To reduce genotype errors mimicking rare alleles with a strong impact on demographic inference, we generated folded-site frequency spectra (SFS) incorporating genotype uncertainty and up to 20% missing data using ANGSD 0.921 (ref. <sup>59</sup>). Due to the difficulty of inferring genotypes in hemizygous sex chromosomes, partially missing sex information and expected discordance in demographic histories between the X chromosome and autosomes, loci mapping to X-chromosomal scaffolds were excluded. X-linked scaffolds were inferred by

synteny to the dog (*Canis familiaris*) genome. Lastz v.1.04 (ref. 60) was used to align each pinniped reference genome to the dog genome (CanFam3.1) using matches of at least 10 kb (parameters  $M=254$ ,  $K=4,500$ ,  $L=3,000$ ,  $Y=15,000$ ,  $C=2$ ,  $T=2$ , ambiguous = iupac, matchcount = 10,000). Any scaffolds with matches to the dog X chromosome were classified as putatively X-linked and excluded from further analysis. In addition, scaffolds <400 kb, for which power to establish synteny was low, were likewise excluded. We investigated population structure by performing a principal component analysis for each species with NGScovar in NGStools<sup>61,62</sup> using genotype probabilities estimated with ANGSD 0.921 (ref. 59). Only putative polymorphic ( $P < 0.001$ ), bi-allelic sites with a maximum of 20% missing data, a minimum site quality and mapping quality of 20 were included in the analysis.

**Estimation of summary statistics.** For each population, we derived estimates of nucleotide diversity  $\theta_x$  (ref. 63), Watterson's theta  $\theta_w$  (ref. 64) and Tajima's  $D$  (ref. 65) from the folded-site frequency spectra as inferred by ANGSD. Because the ddRAD clusters in this dataset are small, often containing no segregating sites, we estimated parameters in two ways. First, we calculated the mean for each scaffold excluding scaffolds with <3,000 cumulative sites located in ddRAD clusters. This approach incorporates variation across the genome. Second, we obtained a single, genome-wide estimate concatenating all ddRAD clusters and using the sum of the per-site estimates of  $\theta_x$  and  $\theta_w$  to calculate Tajima's  $D$ . The results of both approaches were highly correlated ( $\theta_x$ ,  $R_{adj}^2 = 0.997$ ;  $\theta_w$ ,  $R_{adj}^2 = 0.998$ ; Tajima's  $D$ ,  $R_{adj}^2 = 0.939$ ), and we proceeded with the first approach.  $F_{ST}$  estimates were calculated using the approach used in ref. 66 for species with multiple populations sampled that were sequenced in the same ddRAD libraries.

Under the simplifying assumption of mutation-drift equilibrium,  $N_e$  can be inferred from the relationship  $N_e = \theta_x / 4\mu$ . Because mutation rates of pinnipeds are unknown, they were inferred from branch-specific substitution rates of putatively neutrally evolving, synonymous sites in coding sequences. We first compiled a set of 11,256 orthologues for the two most closely related outgroup species, dog and panda (*Ailuropoda melanoleuca*), available from the OrthoMam v.9 database<sup>67</sup>. We then screened GenBank for corresponding orthologues in the three species of pinnipeds for which annotations were available: Weddell seal, Hawaiian monk seal (*Neomonachus schauinslandi*) and walrus. In addition, we extracted 1/1 orthologues for the Antarctic fur seal genome v.1.4 (ref. 57) and the California sea lion genome v.1.2. To minimize the risk of including paralogs, for each species of pinniped we next performed a reciprocal TBLASTN to the dog reference and retained only sequences with unique hits. We then aligned the obtained sequences using MACSE<sup>68</sup> (with default parameters except for  $-\text{gap op} = 3$  and  $-\text{ext gap ratio} = 0.75$ ) and filtered the amino acid alignments using HMMcleaner<sup>69</sup> to remove misaligned regions in sequences, which can heavily impact estimation of substitution rates. We applied a very conservative threshold of a maximum of five consecutive positions differing from the correspondent profiles. Sites filtered in this step were then masked in the nucleotide sequences and, within each alignment, only sequences where >50% of the original sites remained unmasked were retained. We then kept only those alignments in which at most one taxon was missing, which left us with a total of 8,864 alignments.

Separate branch-specific substitution rates were calculated for the phocid clade, represented by the Hawaiian monk seal and Weddell seal, estimated to have diverged 18.2 million years ago (Ma); the otariid clade, represented by the California sea lion and Antarctic fur seal, which diverged from each other around 9.2 Ma; and the odobenid clade, represented by walrus, which diverged from otariids 22 Ma. The divergence between phocids and the clade including the other three pinnipeds has been estimated to have occurred around 27 Ma (Extended Data Fig. 9, modified from ref. 48). To obtain estimates of neutral substitution rates of synonymous sites, we used a substitution mapping approach<sup>70</sup>. This approach first performed a maximum likelihood optimization of branch lengths and then mapped substitutions with the inferred model parameters using the *bppml* and *mapnh* programmes available in the Program Suite (BppSuite) and testnh libraries of the Bio++ libraries<sup>71</sup>. We additionally performed a correction to take into account the opportunity for mutation using the *kappa* parameter estimated in the previous step<sup>72,73</sup>. Results for all genes were averaged for each species and then reported as substitution rate per site and per year following divergence dates in ref. 48. For the estimation of  $N_e$ , the substitution rate of the branch leading to the closest relative of the target species was chosen, and mutation rate was approximated as the product of the estimated substitution rate per site and per year and the harmonic mean of generation times of all species sampled in the clade (see Supplementary Table 2 and ref. 74).

Both mutation rate estimates,  $\mu$ , and estimates of  $\theta_x$  are subject to sampling variance and systematic variation across the genome. In the absence of positional information of the loci and recombination rate governing the correlation between parameters and neighbouring loci, sensible confidence intervals around  $N_e$  cannot be provided. We therefore report only genome-wide averages of  $N_e$  based on the mean of  $\theta_x$  estimates and the point estimate of  $\mu$ , as described above.

**Demographic inference.** Inference of demographic history was based on the folded SFS for each species derived as described above, and modelled

using a composite likelihood approach implemented in *δaδi*<sup>40</sup>. Previous works simulating RAD-like genotyping-by-sequencing data demonstrated that demographic parameters are well estimated for simple models using this approach<sup>75</sup> and *δaδi* can use the SFS directly from ANGSD, avoiding the need for genotyping and reducing bias<sup>76</sup>. Several demographic models were considered (see Fig. 2a):

- (1) Constant population size (null model (model 0P)).
- (2) Two-parameter models (2P) characterizing a single-population size change at time  $T_1$  (in units of  $2N_{anc}$  generations). Population size change was modelled either as (a) an instantaneous change ( $(2P_{ins})$ ) resulting in rapid population expansion or bottleneck or (b) gradual exponential change in population size ( $(2P_{exp})$ , gradual growth or contraction). The parameter characterizing population size change was expressed as relative population size after the change  $N_t$  (relative to the ancestral population size  $N_{anc}$ ). A reduction in population size was indicated if  $N_t < 1$ , and expansion if  $N_t > 1$ . In the case of gradual population size change, change was characterized by an exponential growth parameter.
- (3) A three-parameter model including one rapid population change of size  $N_t$  at time  $T_1$ , followed by gradual change to the present population size  $N_2$  (3P).
- (4) Four-parameter models incorporating two independent population size changes, again considering (a) instantaneous change ( $4P_{ins}$ ) or (b) gradual change following an exponential distribution ( $4P_{exp}$ ). Accordingly, a total of four parameters were required to describe the timing ( $T_1$  and  $T_2$ ) of population size changes  $N_1$  and  $N_2$  (both relative to the ancestral population size,  $N_{anc}$ ). Each model was run independently four times, with confidence intervals based on 100 bootstrap replicates. Only the replicate with the highest mean log-likelihood for each model was considered for comparison between models. A statistical test of the goodness of fit between models was performed using likelihood-ratio tests with the R package extRemes v.2.0.8, using a significance level of  $P = 0.05$  and degrees of freedom equal to the difference in the number of model parameters. Nested models were tested in pairs, starting with the null model (0P) followed by pairs of models with increasing numbers of parameters. If models had the same number of parameters, the model with the highest mean log-likelihood was selected. Parameter estimates are reported for the model with the best fit.

As a complement to the above-mentioned approach, we used PSMC' (ref. 36) to estimate changes in effective population size over time for all species where both ddRAD and whole-genome sequencing data were available. PSMC' was run on the whole-genome shotgun sequencing data generated for each of the pinniped reference genome assemblies, including the Antarctic fur seal (SRR2658532–SRR2658561), California sea lion (10.5281/zenodo.3741488) and walrus (SRR575502–SRR575518). We mapped the short-read data to their respective reference genome (versions as above for ddRAD analyses) using bwa<sup>77</sup> and called single-nucleotide polymorphisms with Samtools pileup/bcftools v.1.3 (ref. 78), with a minimum quality score of 20. Scaffolds inferred to be X-linked were excluded. Regions with low mapping success were masked using a mask generated with SNPable (<http://lh3lh3.users.sourceforge.net/snpable.shtml>), as were areas with excessively low or high coverage when the input files were formatted using scripts from the MSMCtools repository (<https://github.com/stschiff/msmc-tools>). Each analysis was performed for a single individual using unphased data (PSMC' approach). The substitution rates calculated above and generation times were used to rescale both time and effective population sizes. Long-term effective population size was calculated as the harmonic mean of population sizes inferred at each coalescent event.

**Statistical inference.** We were interested in assessing the relationship between the genetic variation and other biological traits of the species. We limited our considerations to explanatory variables for which estimates were available for all species, and compiled the following information.

- (1) Estimates of the current census population size ( $N_c$ ) of species and subspecies were extracted from the IUCN website (<https://www.iucn.org/> accessed 15 January 2019)<sup>74</sup>. These numbers mostly represent the number of mature individuals. Where ranges were presented, we chose the midpoint. IUCN population estimates were not available for *Erignathus barbatus*, but past estimates put the total species at 700,000, and *E. b. barbatus*, the subspecies that we have sampled, at 250,000 individuals<sup>79</sup>.
- (2) Similarly, for the assessment of conservation status we followed the classification of the IUCN<sup>74</sup>. Following Stoffel et al.<sup>22</sup>, we grouped species listed as 'near threatened', 'vulnerable' or 'endangered' into a 'concern' category and contrasted them to species of 'least concern'.
- (3) Data on species-specific life-history traits were extracted from ref. 28. Variables with right-skewed distributions were transformed logarithmically. Several of the predictor variables were strongly intercorrelated (Extended Data Fig. 10), which is expected to lead to suppression effects and make problematic the interpretation of regression coefficients of linear models. Moreover, the number of predictor variables was high relative to sample size, given by the number of populations or species. We therefore eliminated intercorrelated predictor variables with high Pearson regression coefficients



a priori to reduce the effect of collinearity and to avoid overfitting models. We kept the variables with presumed lower measurement error and more immediate biological relevance. This led to the following four predictor variables:

- (a) Harem size rather than mean sexual size dimorphism ( $r=0.91$ ) as a more direct measure of the mating system
- (b) Length of reproductive season rather than breeding latitude ( $r=-0.61$ ) and length of lactation period ( $r=0.47$ )
- (c) Species distribution range rather than breeding latitude ( $r=0.77$ ); because the former was significantly associated with Tajima's  $D$ , it was replaced by the latter in models in which Tajima's  $D$  was used as a predictor variable
- (d) Breeding habitat (ice or land), which has been shown to be an important predictor of recent human-induced population size changes<sup>22</sup>.

Linear models exploring the relationship between genetic and predictor variables were fitted using  $R^{80}$ . We fitted all possible models with a maximum of two predictor variables. Model selection was based on Akaike's information criterion for small sample sizes ( $AIC_c$ ), which provides information about the goodness of fit of the respective model to the observed data. Following common convention, models with  $\Delta AIC_c < 2$  were regarded as having statistical support. We further provide for each model information-theory based model selection statistics ( $AIC_c$ ,  $\Delta AIC_c$  and  $wAIC_c$ ) and  $R^2$  adjusted for the number of parameters. The sum of Akaike weights ( $\Sigma wAIC_c$ , ranging from 0 to 1) was used to estimate and compare the relative contribution of explanatory parameters. Different numbers of populations were sampled for each species and, in some cases, populations or subspecies from the same species were sequenced in different ddRAD libraries. To mitigate the effects of this, only the population that best represents the core species range was used for model fitting (Extended Data Fig. 5).

The statistical model employed assumes independence of data points. To test for possible effects of genealogical non-independence, we also conducted all correlations in a phylogenetically explicit framework using the package nlme v.3.1–131.1. (refs. <sup>80,81</sup>). This did not qualitatively change the results.

**Reporting Summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

## Data availability

All data generated for this study are archived in the sequence read archive under bioproject no. PRJEB37019 at the National Centre of Biotechnology Information ([www.ncbi.nlm.nih.gov/sra](http://www.ncbi.nlm.nih.gov/sra)). For individual accession numbers see also Supplementary Table 6. All code used for the analyses and the alignments used to infer substitution rates are available at 10.5281/zenodo.3741488.

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## Author contributions

Study design was performed by J.B.W.W., A.B.A.S. and C.R.P. Laboratory work was done by A.B.A.S. and C.-C.W. Data analysis was carried out by C.R.P., S.T., S.D.P., E.B.-C., J.B.W.W. and A.B.A.S. Samples were provided by A.B.B., A.J.O., C.L., D.A.-G., F.G., J.W.B., J.F., J.L.H., J.B.W.W., K.M.K., L.R.O., M.K., M.V., N.J.G., S.S. and T.N. The initial manuscript draft was written by C.R.P., A.B.A.S. and J.B.W.W. All authors edited the manuscript, contributed to interpretation of results and approved the final version of the manuscript.

## Competing interests

The authors declare no competing interests. The views and conclusions contained in this document are those of the authors and should not be interpreted as representing the opinions or policies of the US Government, nor does mention of trade names or commercial products constitute endorsement or recommendation for use. Additionally, the findings and conclusions in the paper are those of the author(s) and do not necessarily represent the views of the National Marine Fisheries Service, NOAA.

## Additional information

**Extended data** is available for this paper at <https://doi.org/10.1038/s41559-020-1215-5>.

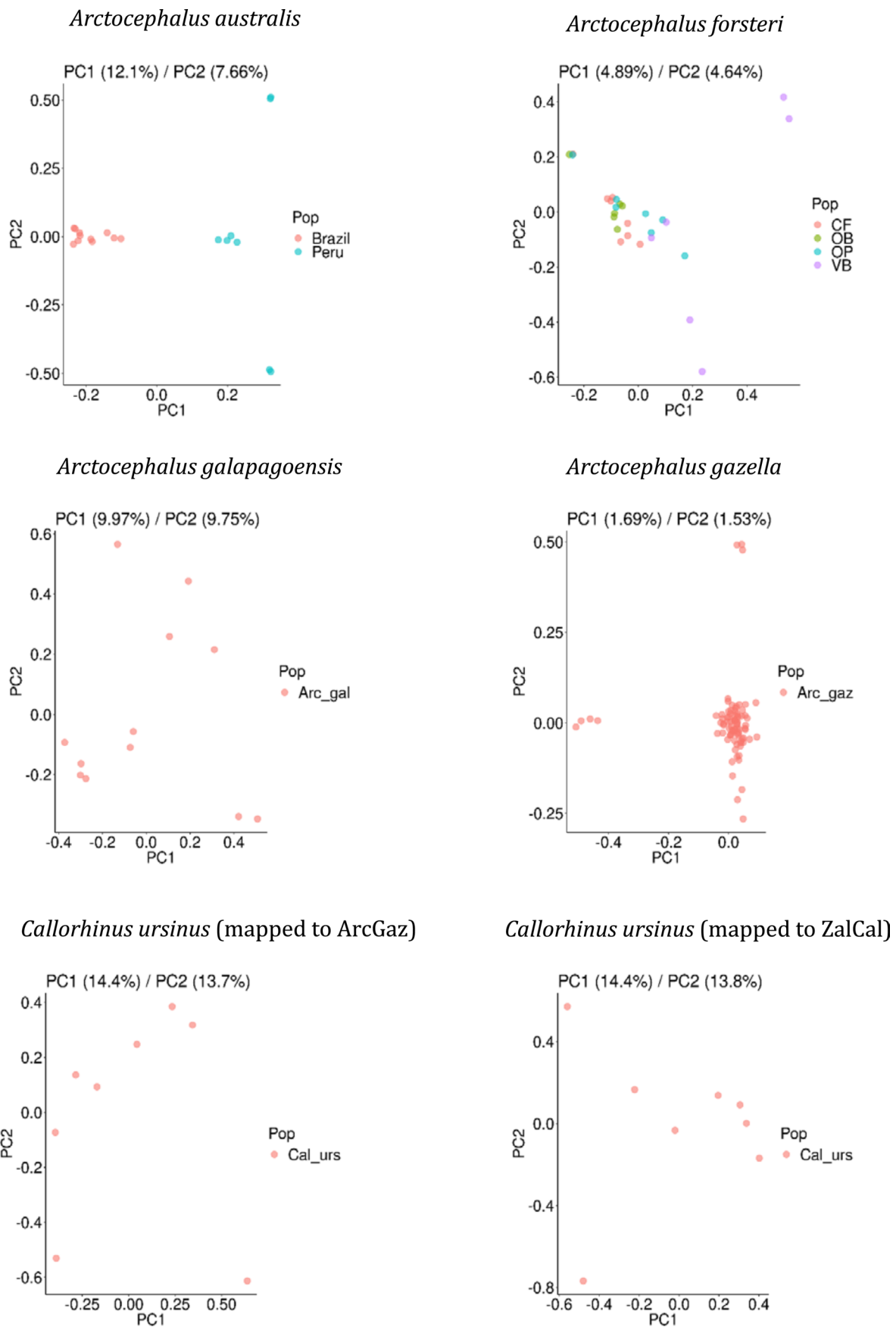
**Supplementary information** is available for this paper at <https://doi.org/10.1038/s41559-020-1215-5>.

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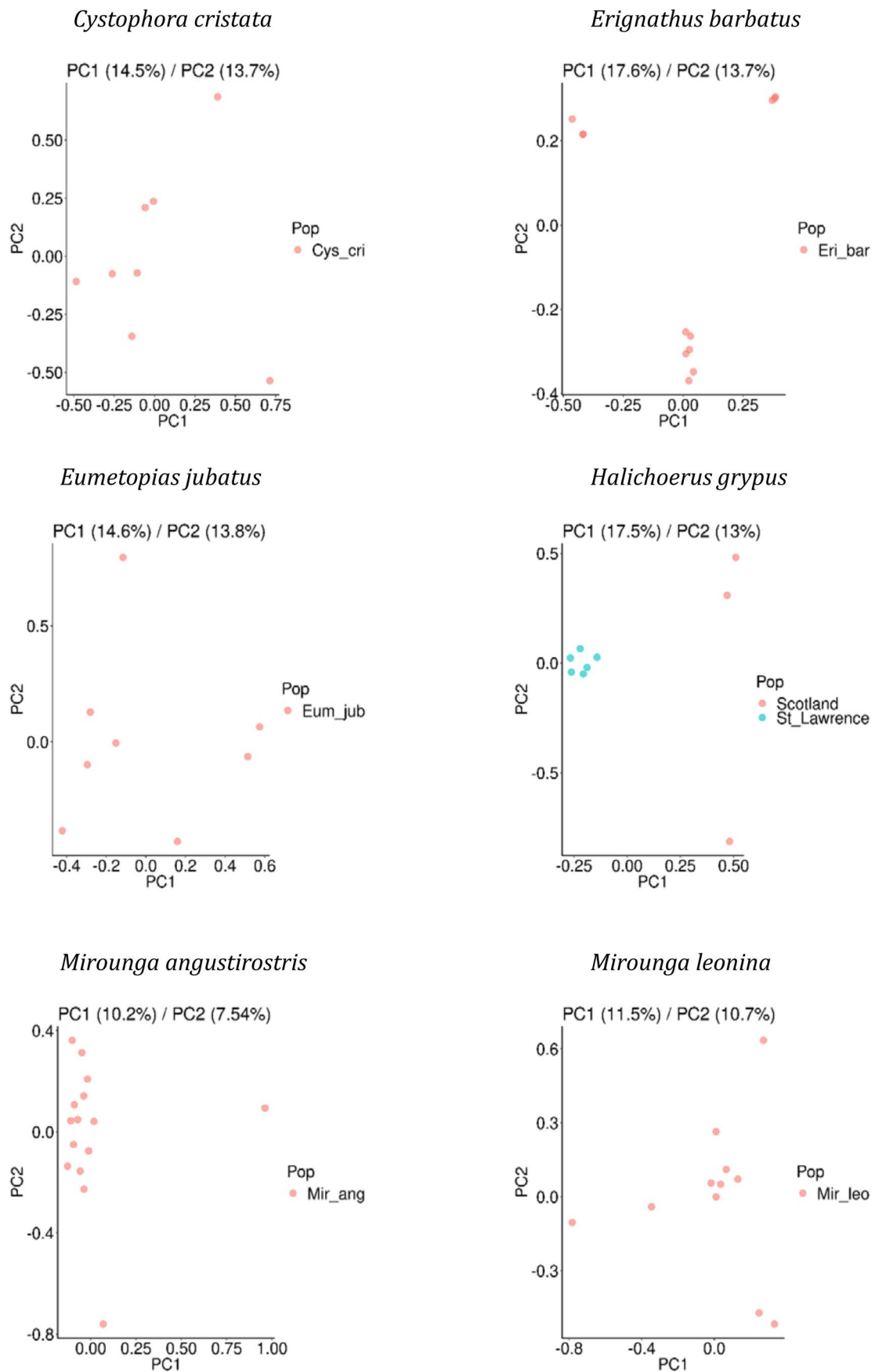
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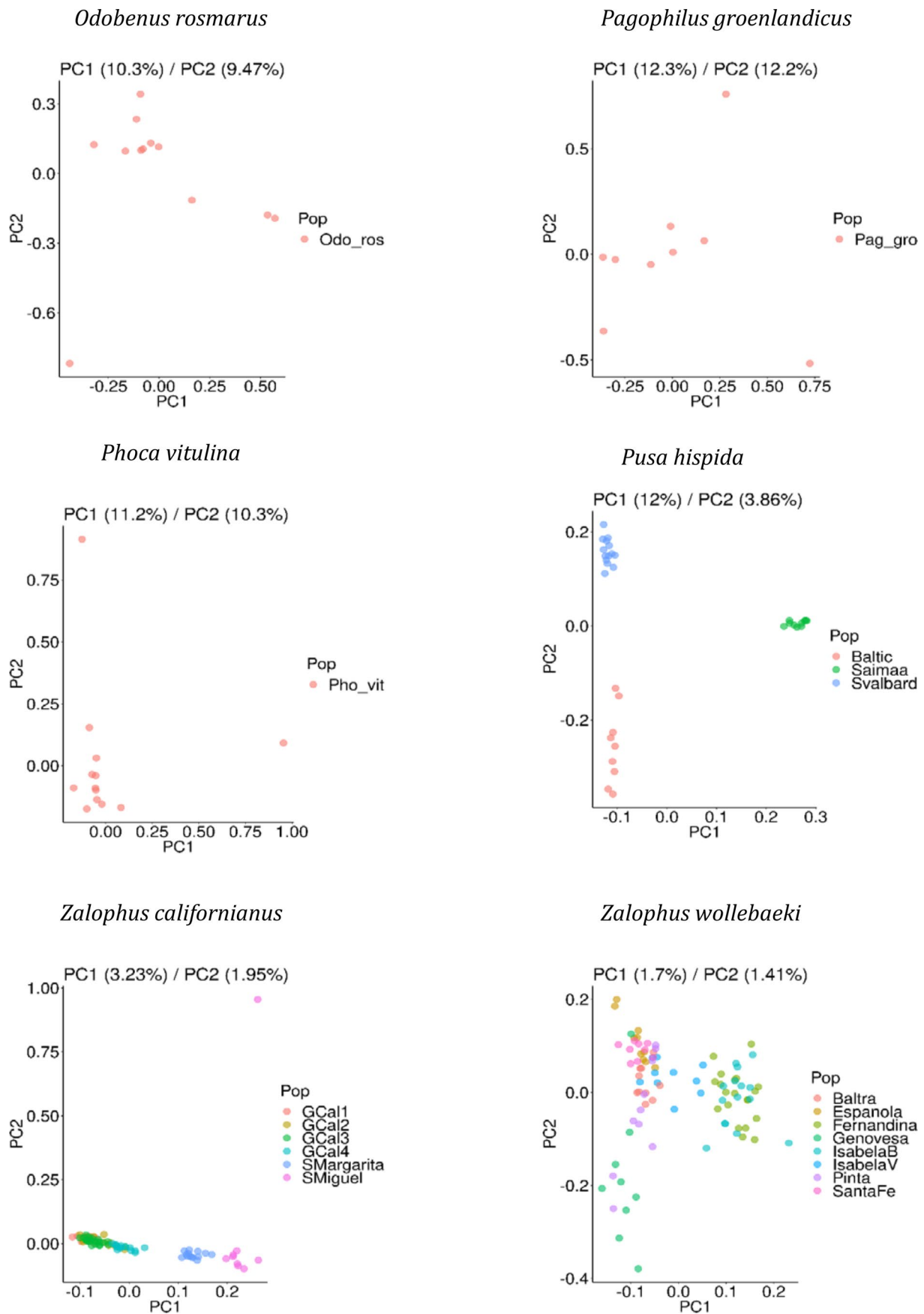
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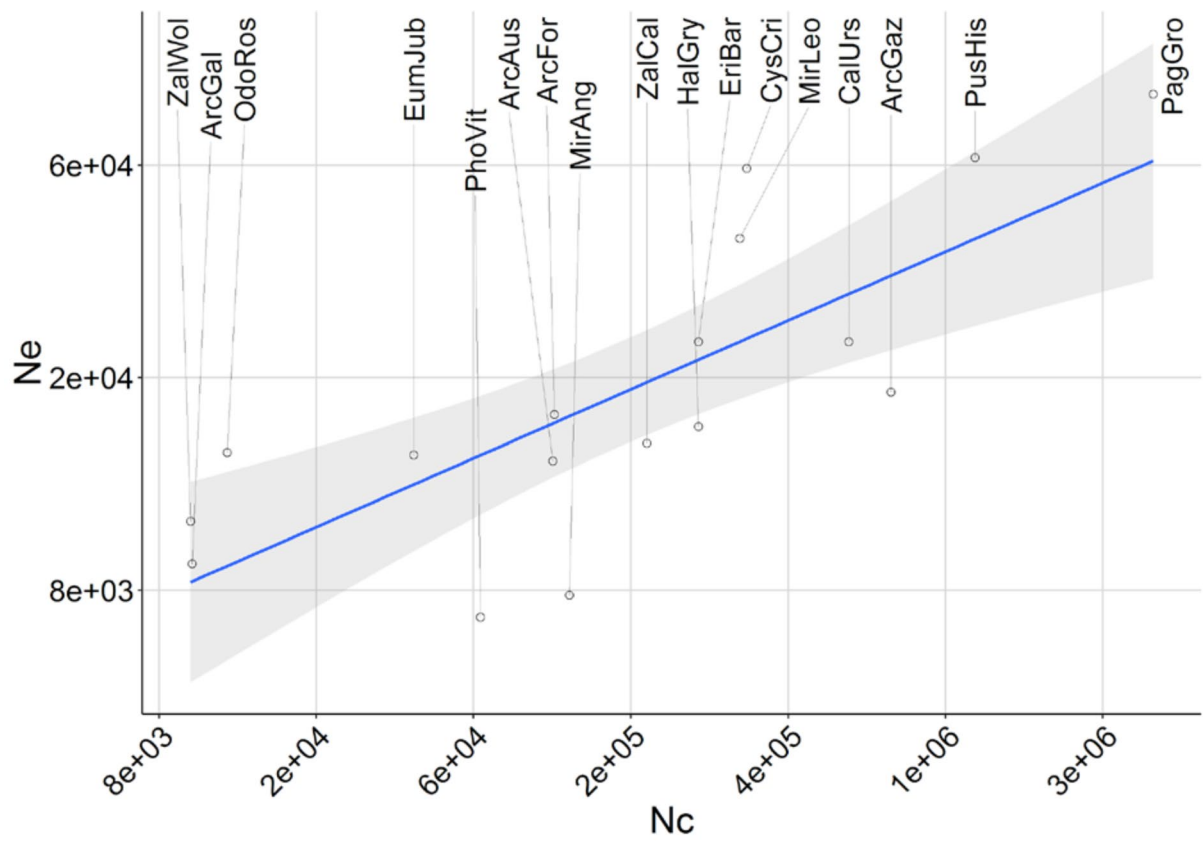
**Extended Data Fig. 1 | Principal component analysis (part1).** Principal component analysis (first two axes) for *Arctocephalus australis*, *Arctocephalus forsteri*, *Arctocephalus galapagoensis*, *Arctocephalus gazella* and *Callorhinus ursinus*.



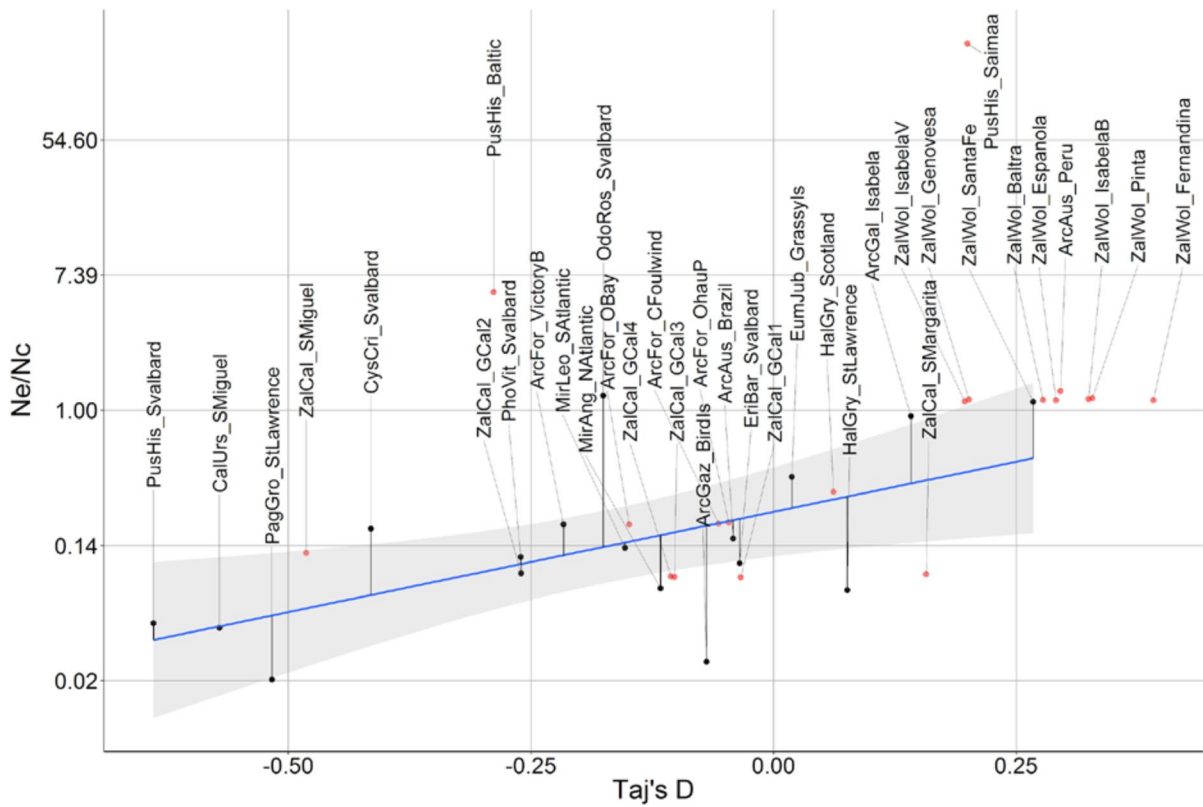
**Extended Data Fig. 2 | Principal component analysis (part2).** Principal component analysis (first two axes) for *Cystophora cristata*, *Erignathus barbatus*, *Eumetopias jubatus*, *Halichoerus grypus*, *Mirounga angustirostris* and *Mirounga leonina*.



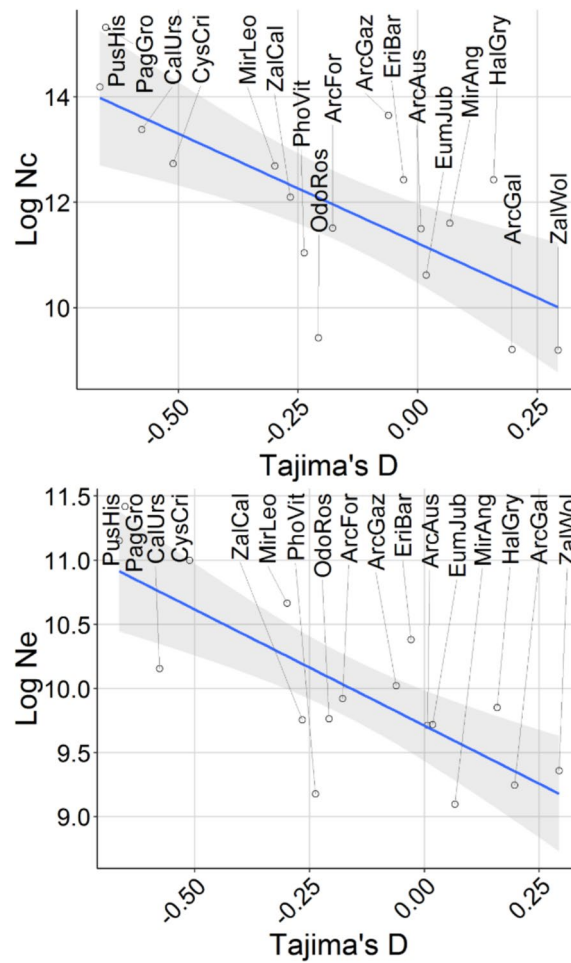
**Extended Data Fig. 3 | Principal component analysis (part3).** Principal component analysis (first two axes) for *Odobenus rosmarus*, *Pagophilus groenlandicus*, *Phoca vitulina*, *Pusa hispida*, *Zalophus californianus* and *Zalophus wollebaeki*.



**Extended Data Fig. 4 | Relationship between effective population size and census population size.** Relationship between effective population size ( $N_e$ ) and census population size ( $N_c$ ) per species as in Fig. 1c (main text) including species labels. Blue line: lineal regression line; shade: 95% confidence interval of regression. The y-axis follows a natural logarithmic scale.

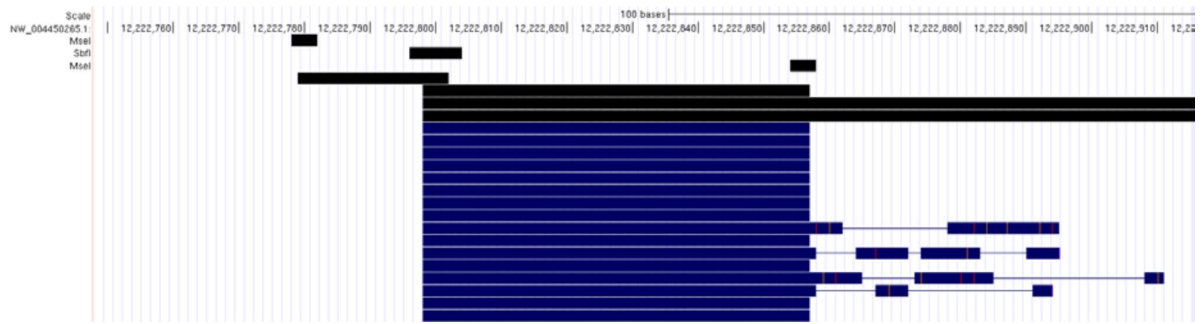


**Extended Data Fig. 5 | The ratio of effective population size and census population size in relation to Tajima's D.** The ratio of effective population size and census population size in relation to Tajima's D representing the species' demographic history; as in Fig. 4a (main text) including species and population labels. Note that the lineal regression (blue line) only refers to the population chosen to represent the species (black points) and the depicted relationship without considering the influence of other predictor variables. 95% confidence interval of the lineal regression is shown in shade. The y-axis follows a natural logarithmic scale.

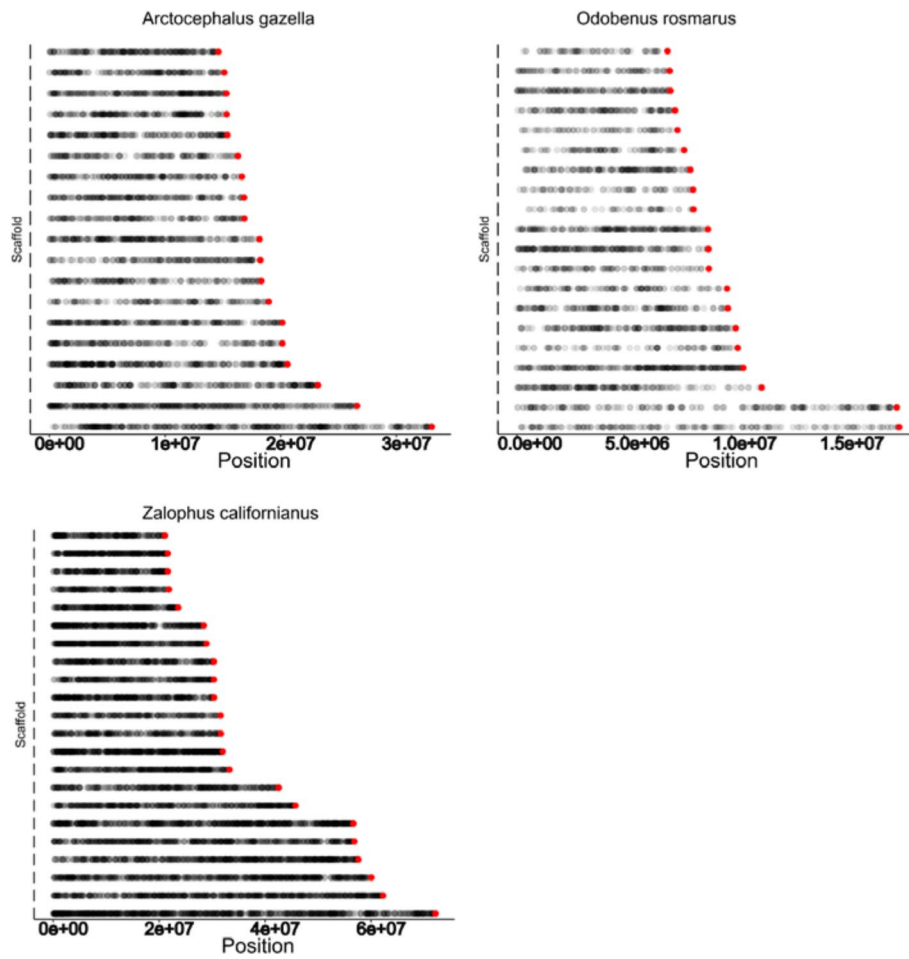


**Extended Data Fig. 6 | Census population size ( $N_c$ ) and population size ( $N_e$ ) in relation to Tajima's  $D$  per species.** Census population size ( $N_c$ ) and population size ( $N_e$ ) in relation to Tajima's  $D$  per species (top and bottom panels respectively). The natural logarithm has been used for transformation of  $N_e$  and  $N_c$  values. Blue line: lineal regression line; shade: 95% confidence interval of regression. Note that the Galápagos seals (ArcGal, ZalWol) lie close to the expected relationship between Tajima's  $D$  and  $N_e$ , whereas they show strong negative residuals for the relationship with  $N_c$ . This is consistent with a very recent reduction in census population size contributing to the high  $N_e/N_c$  ratio above one.

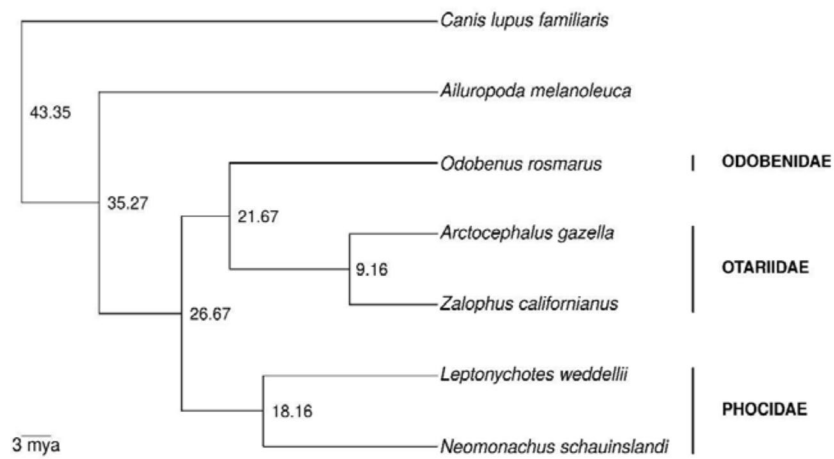




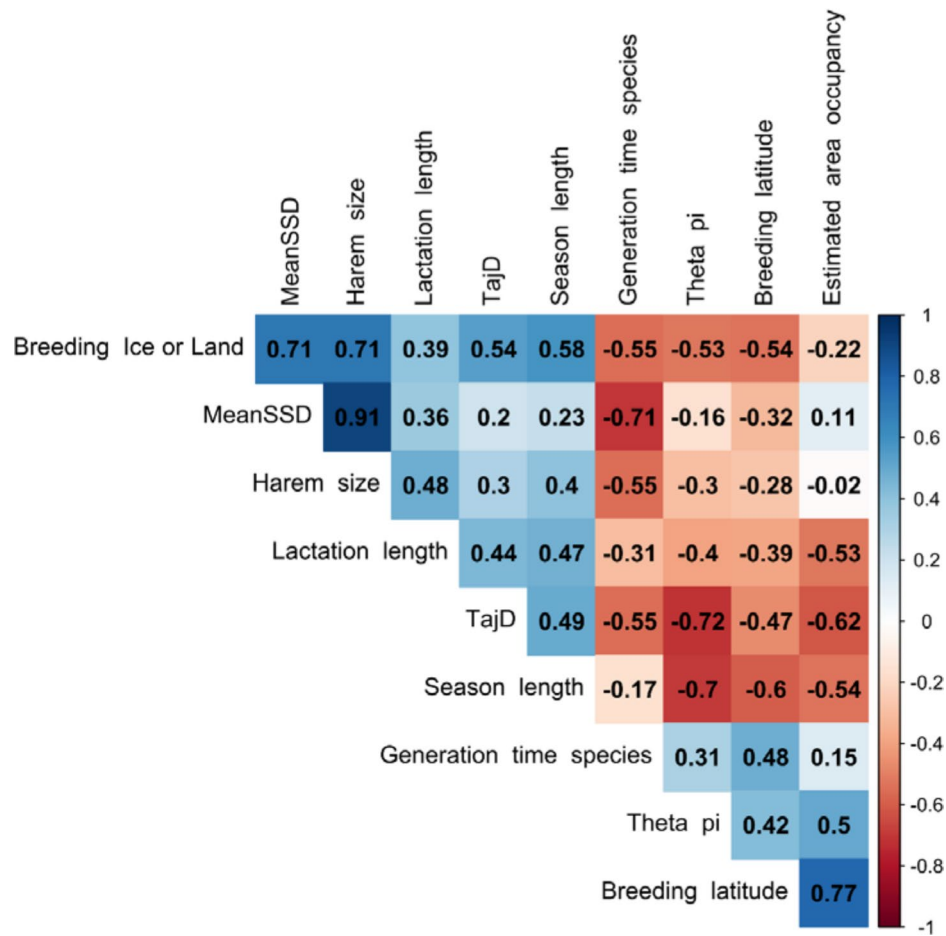
**Extended Data Fig. 7 | Examples of problems identified from visual inspection of the mapped ddRAD reads.** Screenshot examples of mapped reads displayed in the UCSC browser resulting from a ddRAD library constructed using Msel and SbfI. Black bars joining the SbfI to Msel cut sites are potential clusters. Blue bars show aligned reads. The extension of reads beyond the Msel cut site are the 'ghost' PCR extensions with higher visible mismatches and indels.



**Extended Data Fig. 8 | Distribution of ddRAD loci across the reference genomes.** Distribution of ddRAD loci (black points) across the largest 20 scaffolds in species with both reference genome and ddRAD data. Red points correspond to the end of scaffold. For *Zalophus californianus* data correspond to population SMargarita.



**Extended Data Fig. 9 | Phylogeny of species used to estimate  $\mu$ .** Phylogenetic relationships of five pinniped species (*Arctocephalus gazella*, *Leptonychotes weddellii*, *Neomonachus schauinslandii*, *Odobenus rosmarus*, *Zalophus californianus*) and the outgroups dog (*Canis lupus familiaris*) and panda (*Ailuropoda melanoleuca*). The phylogeny was obtained by pruning the tree of Higdon et al.<sup>48</sup> to include only the five species above. Branch lengths are proportional to divergence times, numbers next to nodes show estimated node ages.



**Extended Data Fig. 10** | The collinearity of life-history traits extracted from ref. <sup>28</sup>. Correlogram of the explanatory variables illustrating the degree of collinearity.

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Study description	Estimation of genetic variation and its relationship to species abundance, life history parameters, and conservation status as described in detail in the methods section.
Research sample	Tissue samples from individuals of pinniped species from across the globe. All samples were available previously and no additional field work was done for the purpose of this study.
Sampling strategy	We aimed for a minimum of 10 chromosomes to obtain a reliable estimate of genetic variation in the population for 17 different species (as e.g. used for simulations in Carling & Brumfield (2007)). Where possible, several populations were included to test for robustness of 'species' genetic variation estimates.  1. M. D. Carling, R. T. Brumfield, Gene Sampling Strategies for Multi-Locus Population Estimates of Genetic Diversity ( $\theta$ ). PLoS ONE. 2, e160 (2007).
Data collection	From each sample, we extracted DNA and generated ddRAD data as made available in the SRA archive of NCBI. Published and unpublished genomes were acquired from identified sources.
Timing and spatial scale	As specified above, the samples have been obtained as part as individual studies from a large group of researchers. They were obtained in different batches and ddRAD data was generated accordingly. Batch effects have been explicitly considered. Once our goal of >15 species was reached we started analyzing the genetic data altogether.
Data exclusions	Data were excluded according to data quality as specified in detail in the methods section.
Reproducibility	We explicitly included several populations for each species to test for reproducibility of estimates among populations with shallow differentiation. As a precursor study for this manuscript, we (i.e. Shafer et al.) ran a series of tests to assess robustness for population genetic estimates from ddRAD data.  1. A. B. A. Shafer et al., Bioinformatic processing of RAD-seq data dramatically impacts downstream population genetic inference. Methods Ecol Evol. 8, 907–917 (2017).
Randomization	All statistical analyses were also performed controlling for phylogenetic relationships. Demographic analyses were subjected to bootstrapping.
Blinding	Library preparation were done by lab staff who had no insight into the study setup. For the analyses standardized bioinformatic pipelines have been used on all samples excluding human bias.
Did the study involve field work?	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No

## Field work, collection and transport

Field conditions	Samples were obtained as part of independent research programs unrelated to this study. This being a global collection of samples conditions varied across species.
Location	see Table S6.
Access and import/export	Sampling and sample import was conducted in accordance with local legal requirements.
Disturbance	Disturbance includes stress to the animals during capture and biometric measurements, and possible pain during tissue sampling. Restraining was reduced to a time minimum, and tissue samples were reduced in size allowing for sufficient DNA extraction to reduce pain to a minimum.

## Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

## Materials &amp; experimental systems

n/a	Involvement
<input checked="" type="checkbox"/>	<input type="checkbox"/> Antibodies
<input checked="" type="checkbox"/>	<input type="checkbox"/> Eukaryotic cell lines
<input checked="" type="checkbox"/>	<input type="checkbox"/> Palaeontology
<input type="checkbox"/>	<input checked="" type="checkbox"/> Animals and other organisms
<input checked="" type="checkbox"/>	<input type="checkbox"/> Human research participants
<input checked="" type="checkbox"/>	<input type="checkbox"/> Clinical data

## Methods

n/a	Involvement
<input checked="" type="checkbox"/>	<input type="checkbox"/> ChIP-seq
<input checked="" type="checkbox"/>	<input type="checkbox"/> Flow cytometry
<input checked="" type="checkbox"/>	<input type="checkbox"/> MRI-based neuroimaging

## Animals and other organisms

Policy information about [studies involving animals](#); [ARRIVE guidelines](#) recommended for reporting animal research

Laboratory animals

n.a.

Wild animals

see Supplementary Table 6.

Field-collected samples

Tissue samples were cryopreserved prior to extraction of DNA.

Ethics oversight

Ethical approval only relates to local sampling which was conducted prior to this study. Sampling permissions are specified in Supplementary Table 2.

Note that full information on the approval of the study protocol must also be provided in the manuscript.