Evaluating anthropogenic threats to endangered killer whales to inform effective recovery plans

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

Robert C. Lacy, 1✉
Email rlacy@ix.netcom.com

Rob Williams, 2

Erin Ashe, 2

Kenneth C. Balcomb III, 3

Lauren J. N. Brent, 4

Christopher W. Clark, 5

Darren P. Croft, 4

Deborah A. Giles, 3

Misty MacDuffee, 6

Paul C. Paquet, 6,7

1 Chicago Zoological Society, Brookfield, IL 60513 USA

2 Oceans Initiative, Seattle, WA 98102 USA

3 Center for Whale Research, Friday Harbor, WA, 98250 USA
Abstract

Understanding cumulative effects of multiple threats is key to guiding effective management to conserve endangered species. The critically endangered, Southern Resident killer whale population of the northeastern Pacific Ocean provides a data-rich case to explore anthropogenic threats on population viability. Primary threats include: limitation of preferred prey, Chinook salmon; anthropogenic noise and disturbance, which reduce foraging efficiency; and high levels of stored contaminants, including PCBs. We constructed a population viability analysis to explore possible demographic trajectories and the relative importance of anthropogenic stressors. The population is fragile, with no growth projected under current conditions, and decline expected if new or increased threats are imposed. Improvements in fecundity and calf survival are needed to reach a conservation objective of 2.3% annual population growth. Prey limitation is the most important factor affecting population growth. However, to meet recovery targets through prey management alone, Chinook abundance would have to be sustained near the highest levels since the 1970s. The most optimistic mitigation of noise and contaminants would make the difference between a declining and increasing population, but would be insufficient to reach recovery targets. Reducing acoustic disturbance by 50% combined with increasing Chinook by 15% would allow the population to reach 2.3% growth.

Introduction

Conservation science is tasked with quantifying the relative importance of multiple anthropogenic threats to species, both to determine if cumulative impacts exceed
sustainable levels and to guide effective recovery plans[1, 2, 3, 4]. However, cumulative human impacts are often poorly understood and inadequately addressed in conservation and management[5]. Fundamental research is still needed to integrate information on qualitatively different stressors into comprehensive models that reveal the cumulative impacts on measures of population growth, stability, and resilience[6]. Such work is needed, in part, because threats vary widely in their amenity to mitigation. When regulators require users to forego economic opportunities, it is important to have confidence that management actions will achieve the desired effect[7]. One way to accomplish this is to conduct “population viability analyses” (PVA) that use models of population dynamics to evaluate the relative importance of multiple anthropogenic stressors, singly and in combination, so that conservation can be directed toward efforts most likely to promote species recovery[8]. PVA can be a powerful tool for informing management and conservation decisions. However, the detailed population models used in PVA depend on: availability of estimates for demographic rates (both fecundity and survival and the variability in such rates); confidence that observed past rates are predictors of ongoing demography, or that trends can be foreseen; data for quantifying effects of threats on demographic rates; and a population model that adequately captures the key demographic, social, genetic, and environmental processes that drive the dynamics of the population of concern. Nevertheless, even when data on certain aspects of the population or its threats are not available, we can use PVA models to explore possible outcomes across a plausible range of values, and thereby identify which factors might be important and the target of additional research.

AQ1

The Southern Resident killer whale (Orcinus orca, SRKW) population in the northeastern Pacific Ocean is one of the most critically endangered populations of marine mammals in the USA[9] and Canada[10]. The USA and Canada have listed this transboundary population as Endangered, citing three primary risk factors: lack of the whales’ preferred prey, Chinook salmon (Oncorhynchus tshawytscha); chronic and acute underwater noise and physical disturbance (e.g., from ferries, commercial ships, whale-watching boats, fishing boats, and recreational traffic); and high levels of contaminants, including polychlorinated biphenyls (PCBs)[10, 11]. A recent Status Review[12] highlighted also the potential risk to this small, localized population from catastrophic events such as an oil spill. Governments and non-governmental organizations are currently seeking effective conservation measures for this high-profile population. Fortunately, the biological and environmental data available for SRKWs are rich by the standards of any marine
mammal population. Long-term annual censuses, with continuous monitoring since 1976, coupled with the specialized diet, have allowed inference of quantitative relationships between prey and various metrics of fecundity and survival[13, 14]. Thus, the prerequisites for a robust PVA suitable for guiding conservation are met.

PVA uses demographic models to assess risk to wildlife populations and evaluate the likely efficacy of protection measures, recovery targets, and restoration options[15, 16]. We used the Vortex PVA model to examine the dynamics of SRKWs. Vortex[17, 18, 19] is a flexible, individual-based simulation that is freely available. Vortex has been used to set recovery goals and guide actions for many threatened species, including the Mexican wolf (*Canis lupus baileyi*)[20], Florida panther (*Puma concolor coryi*)[21], and Florida manatee (*Trichechus manatus latirostris*)[22]. Several recent PVAs on the SRKWs have shown how variability in demography[23] or inter annual variability in Chinook salmon abundance[12, 24, 25] could affect the population. We extend those approaches to consider also the sub-lethal effects of contaminants and acoustic disturbance, and the cumulative impacts of threats and interactions among them.

We first parameterized a Baseline model with demographic rates observed over 1976 through 2014, and tested the sensitivity of population growth to each demographic parameter. We then constructed one model that quantifies the population consequences of all three anthropogenic threats to SRKWs identified in Canadian[10] and USA[11] recovery plans. We compared the relative importance of each threat by projecting the population growth across the possible range of each threat. Finally, we used the PVA to explore the degree to which threats would have to be mitigated, alone or in combination, to reach a quantitative USA recovery target of sustained 2.3% growth over 28 years[11].

## Results

Five sets of population models and the scenarios examined in each are listed in Table 1. The Baseline model projects mean population growth over the next 100 years of \( r = -0.002 \), with variation across years of \( SD = 0.045 \) (Fig. 1). These projections match very closely to the rate of \( r = 0.002 \), with \( SD = 0.042 \), observed over 1976 to 2014. The marginally lower growth in the model can be accounted for by future accumulation of low levels of inbreeding. After 100 years, the projected mean inbreeding coefficient is 0.067, about the same as results from mating between first-cousins. When inbreeding depression was eliminated from the Baseline model, the projected growth was \( r = 0.002 \), with \( SD = 0.043 \) – nearly identical growth and variation in growth to the trend in recent decades, and thereby
confirming that the model replicates accurately the recent dynamics of the population.

Table 1

Models of viability of the SRKW population for assessing current viability, sensitivity to anthropogenic threats, and responses to management.

| Set                        | Scenario                  | Parameters varied                  | Population growth (r) |
|---------------------------|---------------------------|-----------------------------------|-----------------------|
| Baseline                  | Baseline                  | Rates as observed 1976–2015        | −0.002                |
|                           | Sensitivity Tests         | See Supplementary Information (S.I.) | See S.I.              |
| Individual Threats        | Current                   | Chinook = 1.0; Noise = 85%; PCB = 2 ppm/y | −0.001                |
|                           | Chinook                   | 0.6 to 1.3 x baseline              | −0.038 to +0.025      |
|                           | Noise                     | 0 to 100% of time                  | +0.017 to −0.004      |
|                           | PCB                       | 0 to 5 ppm/y                       | +0.003 to −0.008      |
| Cumulative Threats        | No Anthropogenic Threats  | baseline Chinook; no noise, no PCB; no oil spills; no ship strikes | +0.019                |
|                           | Low Development           | 25% decline in Chinook; 92.5% noise; low frequency oil spills and ship strikes (see Table 2) | −0.008                |
|                           | High Development          | 50% decline in Chinook; 100% noise; higher frequency oil spills and ship strikes (see Table 2) | −0.017                |
| Demographic Management    | Fecundity                 | 1 to 1.5 x baseline                | +0.016                |
### Set | Scenario | Parameters varied | Population growth (r)
--- | --- | --- | ---
Adult Mortality | 1 to 0.5 x baseline | + 0.009
Calf Mortality | 1 to 0.5 x baseline | + 0.004

**Threat Management**

Chinook | 1 to 1.3 x baseline | + 0.025
Noise | 85% to 0% | + 0.017
PCB | 2 to 0 ppm/y | + 0.004
Chinook & Noise | 1 to 1.3 x Chinook; 42.5% Noise | + 0.036

Population growth rates are mean $r$ for Baseline, ranges for tests of Individual Threats, means for Cumulative Threat scenarios, and maxima for ranges tested in Demographic Management and Threat Management scenarios.

---

**Figure 1**

The distribution of 10,000 simulated trajectories with means and SD of the population size for northeastern Pacific Ocean SRKWs projected for 100 years, based on demographic rates observed from 1976 through 2014, applied to a starting population as it existed in 2015.

![Population Size vs Year Graph](image)

Sensitivity tests of the influence of each demographic rate in the baseline PVA (Supplementary Information) show that, across the ranges of values tested, variation in fecundity (defined for the model as the mean proportion of adult...
females giving birth per year) accounts for most (77%) of the uncertainty in population growth rate. Annual adult mortality has some influence on the population trajectories (6%), but because mortality is already close to 0, there is comparatively less opportunity to improve the value of this parameter. Calf (first year) and juvenile (1 y to 10 y) mortality each accounted for about 3% of variation in population growth. Individual variation in reproductive success and temporal fluctuations (EV) in demographic rates had almost no effect on long-term population growth, as would be expected for a very long-lived species in which short-term fluctuations average out over time. Therefore, although our estimates of annual variation in rates are uncertain, refining the estimates would not change any conclusions about the effects of threats on the viability of the population. Given the small population size, inbreeding depression might cause sufficient adverse impact on population viability (6% of the total variance explained) such that it should not be ignored in assessments of long-term population viability. The impact of inbreeding was exacerbated slightly when we did not include avoidance of very close inbreeding (Supplemental Information).

**Individual Threats**

The set of models that includes estimates for the threats identified in the recovery plans – Chinook prey availability, noise and disturbance, and contaminants – was calibrated so that in the Current Threats scenario the demographic rates at existing threat levels reflect the mean demographic rates observed from 1976 through 2014. Thus, the Current Threats scenario mirrored the simpler Baseline scenario, except that rounding error in estimating effects of threats led to very slight deviation from the Baseline. The levels of these threats were then varied across broad ranges of values to determine which threat would have the greatest impact on population growth. Over the ranges tested, the effects of Chinook prey abundance on fecundity and survival had a greater effect on the population growth rate than did the other two factors (Fig. 2). Noise disturbance acts through decreased feeding efficiency in our model, but has a lesser effect than prey abundance because the maximum impact of boat noise 100% of the time would be to reduce foraging by about 20%. PCB accumulation rates that we tested result in mean levels in adult females of 0 to 132 ppm. Across this range, calf mortality is predicted to rise from about 7% to 50% (see Methods), and this impact shifts population growth from slightly positive to negative.

**Figure 2**
Effect of Chinook prey abundance (index varied from 0.60 to 1.30), noise and disturbance (boats present from 0% to 100% of time), and PCB contaminants (accumulation rate from 0 to 5 ppm/year) on mean population growth, while holding the other two factors at their baseline levels (1.0 prey index, 85% noise, and 2 ppm/y PCB accumulation). The x-axis is standardized to the range tested for each variable.

Cumulative Threats

Threats may interact, such that cumulative effects differ from those projected based on the summation of individual impacts. Full exploration of all of the possible interactions among the threats to the SRKW is not warranted at this time because individual threats are not yet well quantified. As more data on the above threats and other threats are acquired, management authorities can use the PVA framework to examine specific interactions of interest or full statistical analysis of all possible interactions[26]. To illustrate how cumulative threats can be assessed within the PVA model, we examined combinations of threat levels that represent the cumulative impacts of multiple threats for a few sample scenarios. We compared the Current Threats to a scenario with no anthropogenic threats and to scenarios with an increase in current threats and the addition of new threats. Figure 3 compares the population trajectory for the Current Threats with a scenario in which noise and PCB contamination were set to 0, and with two scenarios that describe levels of threat that could occur with proposed further industrial development and climate change. Table 2 shows the mean growth rates, probabilities of decline below 30 animals, and probabilities of extinction within 100 years under these scenarios.

Figure 3
Mean projected SRKW population sizes for scenarios with (from top to bottom): no anthropogenic noise or contaminants; current Chinook abundance, noise, and PCBs; reduced Chinook, increased noise, and additional threats of oil spills and ship strikes as estimated for low level impacts of future industrial development; and these increased and additional threats with higher level impacts of development.

![Graph showing population sizes over years for different scenarios]

**Table 2**

Measures of viability of the SRKW population over 100 years under scenarios of minimal threats, current threats, and two levels of increased threats due to development.

| Scenario                          | Threats modelled | Population projec |
|----------------------------------|------------------|-------------------|
|                                  | Chinook trend    | Noise  | PCB (ppm/y) | Oil spill (big; small) | Ship strikes | Population growth (r) | Probability extinct |
| No anthropogenic threats         | constant         | 0      | 0            | 0                      | 0           | 0.019               | 0                   |
| Current threats                  | constant         | 85%    | 2            | 0                      | 0           | −0.001              | 0                   |
| Low increase                     | −25% in 100 y    | 92.5%  | 2            | 0.21%; 1.08%           | 1 per 10 y | −0.008              | 5%                  |
| Higher increase                  | −50% in 100 y    | 100%   | 2            | 0.42%; 2.16%           | 2 per 10 y | −0.017              | 25%                 |

See text for explanation of threats modelled.
The population could show robust growth if all anthropogenic threats were removed, but has no growth under current threat levels (Fig. 3). The combination of increased and additional threats expected under planned further industrial development in the habitat of the SRKW would cause population decline.

Demographic Management

The potential benefits of improvements in the primary demographic rates were examined in a set of Demographic Management scenarios. The demographic analyses indicate that reaching the SRKW recovery target of 2.3% growth is impossible by improving any single rate by a plausible amount, although increased fecundity would have the greatest positive influence on population growth (Fig. 4). To reach the recovery target, sustained mitigation of threats will be necessary to promote both increased fecundity and reduced mortality.

Figure 4

Mean population growth for SRKW achieved by improvements in demographic rates. Fecundity was increased from baseline to 1.5x baseline; mortality rates were decreased from baseline to 0.5x baseline. Dashed lines indicate a stated recovery target (2.3% growth) and r = 0.

Threat Management

Improvements in demographic rates would need to be achieved by management actions that reduce threats or otherwise enhance the environment for SRKW. We therefore examined how population growth would respond to reductions in the levels of current threats. To achieve the recovery goal by increasing Chinook abundance alone would require a return to nearly the highest rates of Chinook abundance observed since 1979 (Fig. 5). If eliminating acoustic disturbance while
maintaining current levels of Chinook abundance were possible, annual population growth could reach 1.7%. Removal of PCBs from the habitat would result in marginally positive (0.3%) growth, but the effect is much smaller than the impact of reduced noise and disturbance or increased Chinook abundance. Complete removal of both acoustic disturbance and PCBs is predicted to result in 1.9% growth. Therefore, reaching the recovery target without increasing Chinook salmon numbers is likely impossible. Reducing acoustic disturbance by 50% and simultaneously increasing Chinook by more than 1.15x would allow the population to reach the 2.3% growth target. Other combinations of mitigation should be explored by management authorities as conservation options are identified.

**Figure 5**
Mean population growth for SRKW achieved by mitigation of anthropogenic threats. Threat reductions are scaled on the x-axis from no reduction to the maximum reductions tested: Chinook abundance increased up to 1.3x the long-term mean; noise disturbance during feeding was reduced from 85% to 0; and PCBs were reduced from accumulation rates of 2 ppm/y to 0. The top line shows growth rates under a combination of varying levels of improved Chinook abundance plus mitigation of noise to half the current level.

![Graph showing mean population growth for SRKW](image)

**Discussion**
The SRKW population has experienced almost no population growth during the past four decades, and it declined in the last two decades. Intensive monitoring of the population since 1976 provides the information for construction of a detailed PVA model that closely replicates the observed population dynamics, and thereby provides a basis for projections under scenarios of increased anthropogenic threats.
or, conversely, increased mitigation actions. Models projecting population changes based on average demographic rates and fluctuations in those rates project that under the status quo the population will most likely remain near its current size. However, our use of baseline demographic rates averaged across 38 years of monitoring might give an overly optimistic projection for the SRKW if rates have deteriorated in recent years. A population projection based on demographic rates observed through 2011 projected a 1% annual mean growth[25], but a recent Status Review[12] projects a decline of 0.65% per year if demographic rates (such as recently lower fecundity) remain as they have been during 2011-2016. If ongoing monitoring indicates that these are not just short-term fluctuations in rates, then assessments of current viability, vulnerability to new or increased threats, and measures needed to achieve recovery will need to be revised.

When examined over ranges that encompass plausible improvements, the demographic parameter that presents the better opportunity for a large benefit to population growth is fecundity, rather than mortality. This finding is similar to a study of two bottlenose dolphin (Tursiops aduncus) populations off Australia, which found that variability in reproduction was more important than variability in mortality in driving differences between the populations[27]. There is simply more potential for improving reproduction than for improving adult survival when survival is already close to 1. Even complete elimination of adult mortality in the SRKW (not a biological possibility) would result in a population growth rate of 1.8%, still below the recovery goal of 2.3% growth. Although recovery cannot be achieved solely by improving adult survival, any decline in adult survival caused by new or exacerbated threats could have serious consequences for the population.

The PVA was useful for exploring scenarios representing the three main anthropogenic threats – prey limitation, acoustic and physical disturbance, and PCBs – that might worsen with increased development, or could be mitigated through management. Across the ranges of threat levels that we examined, reduction of the prey base was the single factor projected to have the largest effect on depressing population size and possibly leading to extinction, although either higher levels of noise and disturbance or higher levels of PCB contamination are sufficient to push the population from slow positive growth into decline. If additional threats from proposed and approved shipping developments (such as catastrophic and chronic oil spills, ship strikes, and increased vessel noise) combine with the predicted decline of Chinook due to climate change[28], then the population could decline by as much as 1.7% annually, have a 70% probability of
declining to fewer than 30 animals, and have a 25% chance of complete extirpation within 100 years.

Mitigating multiple anthropogenic threats sufficiently to reach the recovery target will be difficult. The PVA is a useful way for managers to identify priorities for future research, and to focus conversations with ocean users and other special interests about the most pragmatic ways to promote recovery of endangered species. Those discussions must be integrated with considerations of feasibility, cost, societal impact, and timeframe for effective implementation. If a threat cannot be mitigated in a timescale relevant to conservation, or if costs are so high that they are prohibitive, thinking of those intractable problems as “fixed costs” in a cumulative impact management framework[4] might be useful. For example, our model results show that eliminating PCBs would provide less benefit to SRKWs than improving salmon returns or reducing anthropogenic noise and disturbance. This is fortuitous because imagining a way to eliminate PCBs that are persistent in the ecosystem is problematic[29], even though levels in tissues of SRKWs have been slowly declining in recent decades[30]. Identifying fixed costs that are difficult or impossible to mitigate allows a practical discussion about how to rank recovery actions among the anthropogenic factors that can be managed.

Of the three threats we considered, across wide but plausible ranges of each, salmon abundance is the greatest factor affecting SRKW population dynamics. Previously reported correlations of demographic rates with Chinook abundance[13, 14, 24] were used to parameterize our model, and Wasser et al.[31] recently offered insights into a mechanism that could cause the effect on fecundity: hormone levels indicate that SRKWs experience nutritional stress related to periods of lower abundance of Chinook prey and that this stress results in fewer successful pregnancies. Our PVA model estimated that SRKW recovery cannot be achieved without reaching the highest levels of salmon abundance observed since 1979, which was 30% higher Chinook salmon abundance than the long-term average between 1979 and 2008. This model result allows managers to focus discussions on whether achieving such a high sustained level of salmon abundance is attainable, and if so, how to achieve it. For example, removal of a hydroelectric dam on the Elwha River in the state of Washington is expected to increase spawning habitat for all five wild Pacific salmon species in the Salish Sea, but discussions about dam removal began in the 1960s[32] and the cost was in the hundreds of millions of US dollars. Restoration of spawning and rearing habitat could improve growth and survival of wild, juvenile salmon, but this takes political will, time, and money[33]. Improvement of marine survival of juvenile salmon might be possible by better
management of net-pen salmon aquaculture sites that host and amplify viruses and parasites that have the potential to reduce survival of wild salmon[34, 35]. Reducing Chinook harvest could provide an interim and strategic opportunity to rebuild depressed wild Chinook salmon runs and increase the number of Chinook available to whales in terminal areas like the Salish Sea[36]. Harvest reductions without longer term rebuilding plans might be an incomplete measure in places where Chinook harvests are already low due to abundance concerns or other constraints[37].

The SRKW population could be adversely affected by any new threats and further intensified impacts of the anthropogenic threats that we did assess. For example, pollutants other than PCBs might affect the population, and PCBs are known to have adverse effects beyond just reduced infant survival – such as reduced immune function[38]. However, other than calf survival, sufficient data are not yet available on the impacts of PCBs on demographic rates to allow incorporation of those threats in the population model. Moreover, threats to the population likely interact, perhaps in non-linear ways. For example, cetaceans that are food-limited might mobilize more lipids, and this will change the accumulated loads and harmful effects of PCBs and other organic pollutants. Similarly, reduction in foraging success because of boat noise might be of little consequence if prey is abundant, but could be critical if killer whales have difficulty procuring enough prey. If we can obtain data on additional threats and the interactions among threats, such effects could be included in the PVA models. At present, given that only estimates of approximate average effects of some threats are included in the model, inclusion of higher level interactions is premature.

While acknowledging that we examined only the identified primary threats to the SRKWs and that we cannot yet fully assess possible complex interactions among those threats, an important finding from our PVA is that reaching the recovery target will likely require mitigation of multiple threats. For example, the PVA projects that a 50% noise reduction plus a 15% increase in Chinook would allow the population to reach the 2.3% growth target. Noise is a particularly attractive issue to address in a management context, because it is amenable to several possible mitigation scenarios[39, 40]. With respect to noise from commercial shipping, preliminary calculations suggest that the distribution of source levels of individual ships follows a power law, implying that quieting the noisiest ships will reduce overall noise levels by a disproportionate amount[41]. Identifying the noisiest ships operating in SRKW critical habitat[42] and creating incentives to reduce their noise outputs through speed restrictions and maintenance might generate considerable
reductions in noise levels. The International Maritime Organization and the
International Whaling Commission have urged nations to reduce the contribution of
shipping to ocean ambient noise, with some countries adopting a pledge to reduce
anthropogenic noise levels by 50% in the next decade[43]. However, from the
perspective of a foraging killer whale that emits high-frequency (18-32 kHz)
echolocation clicks to detect and capture salmon, high-frequency noise from small,
outboard vessels that follow whales might cause a greater reduction in a killer
whale’s foraging success than low-frequency (<1 kHz) background noise from
commercial shipping[44].

Clearly, even without new or increased external threats, the SRKW population has
no scope to withstand additional pressures. The current situation for SRKWs gives
little cause for optimism. This is likely to worsen, given the energy-related project
proposals already approved for the region[45], which will increase broadband ocean
noise levels and the risk of ship strikes and oil spills[46]. Our models of the
additional threats expected with a proposed increase in oil shipping show that these
threats will push a fragile population into steady decline. Obviously, countering
such additional threats sufficiently to achieve SRKW population recovery would
require even more aggressive mitigation actions than if there were no such
increasing threats to the population.

The case study we present offers an unusual opportunity to examine multiple
anthropogenic threats in a wildlife population that is extremely data-rich by the
standard of any marine ecology study[5]. One threat (the impact of prey abundance
through the prey-demography link) has been well studied for decades. Another
(acoustic disturbance) is relatively well appreciated in that there are documented
relationships between higher noise level and reduction in foraging success.
However, a conceptual step is required to convert the reduction in foraging to a
reduction in prey acquisition. Full consideration of noise impacts would need to
include complex interactions among reduced foraging time, reduced detection
space, and reductions in prey availability. The third kind of threat (population
consequences of PCBs and other persistent pollutants) relies on very few data
points to calibrate the effect of the PCBs only on whale calf survival, which
underestimates the total population consequences of contaminants in two ways.
Lack of concentration-response studies on compounds other than PCBs hinder our
ability to model population consequences of PBDEs or other contaminants.
Similarly, existing studies do not allow us to predict effects of contaminants on
pregnancy rate or adult mortality. This spectrum of data-rich to data-poor steps in
predicting population consequences of multiple stressors is ubiquitous in
conservation and ecological studies[2, 47]. The funding to fill knowledge gaps with empirical data may be lacking, or in the case of critically endangered species, time to wait for science to fill data gaps may be insufficient[48]. Some authors use expert elicitation[49, 50] to fill data gaps. Expert opinion or examination of hypothetical, but plausible scenarios should be used to augment rather than replace the available data.

The case study presented here illustrates the use of PVA as a method to inform difficult conservation decisions, by simulating across plausible ranges of uncertainty. For example, sensitivity analyses revealed that some factors (e.g., individual variability in breeding success) have no effect, and such knowledge gaps should not be a barrier to management action. Given our inability to manage some insidious threats, such as persistent organic pollutants that are already in the environment, it is reassuring that the model predicts that this stressor has the smallest adverse impact on the population, at least via the pathway of reduced calf survival. The PVA can focus priority research on questions that make a practical difference. Studies of foraging efficiency under varying levels of anthropogenic disturbance are needed only because the population is prey-limited. If doubling Chinook salmon numbers were possible, and returning them to levels seen in the 1920s[51], consideration of other anthropogenic impacts on the whales’ foraging efficiency might not be necessary. Alas, this is not a realistic scenario, and the model therefore points to the importance of including both improvement in prey abundance and reduction in noise as the more effective mitigation pathway.

Unfortunately, focus on only the immediate, tractable threats is all too common in conservation. For example, conservation of grizzly bears (*Ursus arctos horribilis*) in the continental United States focuses on roads and development activities, but the primary concern is that the species has been absent from most of its range since the 1800s[52]. Similarly, the current small size of the SRKW population was not caused by lack of salmon. The whales’ depleted status is due in large part to the legacy of an unsustainable live-capture fishery for display in aquariums[53]. Salmon, noise, and contaminants are important factors that can prevent recovery. Many policies, including the US National Environmental Policy Act, require regulators to consider the effect of a proposed activity “which results from the incremental impact of the action when added to other past, present, and reasonably foreseeable future actions regardless of what agency (Federal or non-Federal) or person undertakes such other actions (40 CFR § 1508.7).” Allocating impacts among multiple ocean user sectors may be difficult, but in the case study we
present, the population is sufficiently imperilled that it has little or no scope for tolerating additional stressors.

**Methods**

The SRKW population is closed to immigration and emigration, every individual in the population is known, and the population has been censused annually for decades[11]. Individuals were identified by their unique fin shapes, saddle patches, and the presence of any nicks or scratches, and sexed using distinctive pigmentation patterns around the genital slits. Male and female offspring remain within the natal, matrilineal unit, although mating occurs within and between these pods. The term “resident” refers to their residency in inshore waters of southern British Columbia (Canada) and Washington state (USA) in the summer months, when they feed almost exclusively on Chinook salmon[13, 14, 54, 55]. Given that there is no dispersal from the population[56], mortality was recorded if an individual’s matriline was observed in the population within a year but the individual did not appear.

We used values of demographic parameters calculated from the census data to build the population model in the Vortex PVA program[17, 18]. We included temporal variation in demographic rates (“environmental variation”), based on inter-annual variability in parameters observed since 1976, and we included individual variation in age of maturity and probability of reproductive success. The Vortex simulation model of possible future population trajectories includes demographic stochasticity (binomial variation in individual fates); random assignment of sex and a bi-sexual mating system, resulting in fluctuations in sex ratio and mate availability that can affect small populations; and projections of loss of genetic diversity, allowing for inclusion of inbreeding depression. We quantified population growth as the mean exponential rate of increase (r = ln[N_{t+1}/N_t]).

Modelling was conducted in stages. First, a “Baseline” model was developed to represent the population trajectories if demographic rates remain the same as have been observed in recent decades. We confirmed this Baseline model by comparing simulated dynamics with recent population trends. Secondly, we conducted sensitivity tests on uncertain demographic rates in the model to determine which parameters had large effects on the projected population growth. Thirdly, we used a set of models of Individual Threats that tested ranges of values for the primary threats identified in the recovery plans to determine which would have the greatest effects on population projections. Fourthly, we examined Cumulative Threats scenarios to project the fate of the population if further industrial development
increases existing threats and adds new ones. A set of Demographic Management scenarios was then examined to determine the population growth that could be achieved by improvements in demographic rates. Finally, we explored Threat Management scenarios to assess the plausibility of reaching sustained annual population growth of 2.3% given various options for increasing salmon abundance, reducing ocean noise levels, or reducing contaminant levels. The following section describes key parameter estimates used in the model. More detailed description of the modelling methods is presented in Supplementary Information. The input files for the Vortex project are available at http://www.vortex10.org/SRKW.zip and from the Dryad Digital Repository at https://dx.doi.org/10.5061/dryad.46vq7xxxx.

Baseline PVA

We started the simulations with the ages, sexes, and pod membership of the killer whales living in 2015. We specified the mother of each animal, where known (for 76 of 80 living animals)\[57\]. Based on previous genetic data on paternity\[58\], we specified in the simulation that females would not mate with their father, a son, or a maternal half-sibling. What effect lower levels of inbreeding or the inevitable accumulated inbreeding in a closed population will have on any cetacean is unknown. We modelled inbreeding depression as being caused by recessive lethal alleles, with 6.29 “lethal equivalents” (the negative of the slope of log(recruitment) against the inbreeding coefficient), the mean combined effect of inbreeding on fecundity and first-year survival in a survey of impacts on wild species\[59\].

Demographic rates were calculated from individual animal histories compiled by the Center for Whale Research\[57\], using data collected from 1976 through 2014. The time series begins when the population was depleted by live-captures for display in aquariums\[60\]. The time series therefore includes periods of moderate population growth (1976 to 1993), subsequent decline, and approximate stability. Demographic rates were estimated for the age-class groupings used in recent models\[24, 61\], except that we set an upper limit for female breeding at 45 y rather than 50 y, because no females in the population have been documented to produce calves at older ages. Thus, we calculated survival and (for adult females) fecundity rates for calves (first year), juveniles (defined as from 1 year through 9 years of age), young mature females (10–30 years), older reproductive females (31–45 y), post-reproductive females (46 y and older), young mature males (10–21 y), and older males (22 y and older). Killer whales can survive many years after reproductive senescence, but estimating maximum longevity is difficult in such a long-lived species\[62\]. We set an upper limit of age to 90 y in our models, although
only about 2% of females would be expected to reach this age, and only about 2% of males (with higher mortality) would be expected to exceed 50 y. Females stop breeding long before the maximum age, so the long-term population growth would not be affected by the upper age limit unless post-reproductive females benefit the pod in ways other than through their own reproduction.

Mortality for each age-sex class was averaged across the 39 years of data to obtain mean annual rates. We did not try to partition observed mortality into presumed causes of death. The use of these historic data for our Baseline model makes the implicit assumption that the frequency of deaths due to the various causes remains the same as has been observed across recent decades. The variation in mortality observed across years has two components: 1) environmental variation (fluctuations in the probability of survival), and 2) demographic stochasticity (binomial variation in individual fates). To determine how much of the observed variation was due to environmental variation, the variance due to demographic stochasticity can be calculated from the expectation for a binomial process, and then subtracted from the total variation across years. Calculated annual mortality rates (and environmental variation) ranged from a low of 0.97% (SD = 0) for young adult females to 17.48% (SD = 17.96) for calves. Although the lack of evidence for annual variation in the mortality adult females beyond that expected from random sampling of a constant probability might seem optimistic, for long-lived species a low level of annual variation in rates would have negligible effect on long-term population trajectories. We confirmed through sensitivity tests (Supplementary Information) that the environmental variation entered into the population model has no effect on our results.

The breeding system is polygamous, with some males able to obtain multiple mates, females mating with different males over their lifetimes, and mating between and within pods. Males become sexually mature (actively breeding, which may occur several years after they are physiologically capable of breeding) from 12 to 18 years of age. Thus, in the model, each male was assigned an age of sexual maturity by randomly selecting a value from 12 to 18. Variance in reproductive success among individual females and males will cause genetic diversity to be depleted faster and inbreeding to accumulate faster than would occur if mating was assumed random. Information is available on male mating success[51], and we incorporated variation in male and female reproductive success in the model (Supplementary Information). Our models project an effective population size that is 37% of the total size, close to an estimate obtained from genetic data[58].
Breeding rates, expressed as the proportion of the females of an age class that produce a calf each year, were calculated from annual census data. Rates ranged from 0% for post-reproductive females (age >45 y), to 7.88% (SD = 4.15) for older adult females (age 31–45 y), to 12.04% (SD = 3.54) for young adult females (age 10–30 y).

The upper limit on population size was set to 300, so that carrying capacity (K) would not restrict future population growth except under the best conditions tested. In the projections of current or expected conditions, the SRKW populations never reached this limiting size, and rarely exceeded 150 animals in any of the independent iterations of each simulation. Population recovery was assessed by the mean growth rate each year calculated before any carrying capacity truncation. Thus, the growth rate reflects the demographic potential and is not affected by the limit on population size in the model.

The SRKW population was projected for 100 years. For the initial exploration of parameter uncertainty, the simulation was repeated in 10,000 independent iterations to obtain high precision in mean and variance estimations. For comparisons among alternative management scenarios, less iteration is needed to obtain the relative influence of input values, and tests were run with 1,000 iterations. Sensitivity tests were conducted by varying each basic demographic rate (life table values for fecundity and mortality) over a range of ± 10% around the baseline value. For several model variables that describe other aspects of the population dynamics and are also very uncertain, a wider range of values was tested (see Supplementary Information).

Individual Threats

We explored the effects of three threats identified in the recovery strategies. For each of prey abundance, noise disturbance, and PCB contaminants, we scaled impacts such that the estimated current level of the threat resulted in the mean demographic rates reported over recent decades. Effects of prey limitation were modelled using published relationships linking inter-annual variability in Chinook salmon to inter-annual variability in calf and adult mortality[63] and fecundity[13, 61]. A prey index was calculated by dividing the total salmon abundance in each year by its average abundance over the 1979–2008 period[63]. The relationship of mortality to prey abundance was modelled with a multiplier of baseline mortality that is a linear function scaled to 1 when salmon abundance was at the mean observed level over period of observation: 

\[
\text{MortalityFactor} = 3.0412 - 2.0412 \times \text{PreyIndex}.
\]
PreyIndex. The relationship of birth rate to prey was modelled with logistic functions, with the intercept scaled to yield the observed birth rates for young females (12.04%) and older females (7.88%) when PreyIndex = 1. For relationships of form BirthRate = \exp(A + B\times PreyIndex)/[1 + \exp(A + B\times PreyIndex)], the function parameters were $A = -3.0$ and $B = 1.0$ for young females, and $A = -3.46$ and $B = 1.0$ for older females. (See Supplementary Information for more details on these relationships.) To explore the impacts of prey abundance across a range of plausible values, we varied the prey index from approximately the lowest level (0.60) reported since 1978 to approximately the highest level (1.30).

Effects of noise on demography were modelled using the approach outlined in previous analyses of loss of acoustic communication space[4, 64]. We used summertime observations to estimate the proportion of time boats were present (during daylight hours) while the whales were foraging and the reduction in foraging expected with that amount of acoustic disturbance. We calibrated the model of noise impacts so that the mean Baseline demographic rates are obtained at the reported level of disturbance. We then simulated the relative change in foraging time and consequently demographic rates across the spectrum from no noise impact at all, to the upper limit expected if boat disturbance increased from current, already high, levels to 100% of time. We do not have data on the amount of acoustic disturbance in the winter feeding areas, but the modelling based on observed summertime disturbance provides a means to project a range of population consequences if changes in disturbance overall mirror those that are possible in the summertime habitat. Land-based observations have shown that SRKWs reduce their time spent feeding in the presence of boats by 25%[65]. Vessels are present 85% of the daytime, and SRKWs are foraging in the presence of vessels an estimated 78% of that time. Thus, for the 85% current (baseline) exposure to vessels, feeding is expected to be reduced by 16.6% ($= 85\% \times 78\% \times 25\%$) due to disturbance by boats. To translate the reduction in feeding into its demographic consequences, we multiplied the prey index by a factor of $(1 - 0.195 * \text{Noise})/(1 - 0.166)$ to obtain the proportional availability of prey. This proportion is thus 1 in the current, baseline conditions (Noise = 0.85), 0.965 when vessels are always present (Noise = 1.00), and 1.20 assuming no disturbance from vessels. The noise-modified index of prey availability was then used to determine the consequent mortality and fecundity rates. We recognize that anthropogenic noise can also have less direct effects on wildlife, including disruption of social behaviours and even impeding responsiveness to other sensory modalities[66].
Our model of accumulation, depuration, and impact on calf survival of PCBs was based on the approach described by Hall et al.[67, 68] with modifications in rates for SRKW[69]. Calves obtain their initial load of contaminants from their dams through gestation and lactation, and females producing calves thereby depurate an estimated 77% of their contaminants[67]. Otherwise, males and non-breeding females accumulate PCBs in the blubber at a rate that we varied from 0 to 5 ppm/year in our tests. Few data are available on PCBs in the SRKW population with which to calibrate the model of PCB bioaccumulation, and the levels of PCBs reported in SRKW might have been dropping slowly in recent years. Reported levels in adult female SRKW range from 55 ± 19 ppm sampled in 1993–1996, 37 ± 42 ppm sampled in 2004–2007, and 30 ± 31 ppm sampled in 2008–2013[30]. Our population model generates a mean 28, 55, and 81 ppm PCBs in adult females when bioaccumulation rate is 1, 2, and 3 ppm/year, respectively. Effects of maternal PCB load on calf mortality were modelled using a logistic response function (survival = \( \exp(2.65 - 0.02 \times \text{PCB}) / [1 + \exp(2.65 - 0.02 \times \text{PCB})] \)), fitted to the two observed data points for SRKW (survival = 0.8252) and the nearby northern resident killer whales (survival = 0.9218)[24], with the mean PCB levels (55.4 ppm and 9.3 ppm, respectively)[70] reported from the time period in the middle of the span over which mortality rates were calculated. If we use the more recent, lower estimates of PCB loads in SRKW to estimate the impacts, our response function would have a steeper slope. There are not yet sufficient data on effects of PCBs on other demographic rates to allow inclusion of any other effects of PCBs (or other contaminants) in our PVA model.

Cumulative Threats

We modelled two scenarios to represent the cumulative impacts of possible increases in threats, based in part on a recent environmental impact assessment submitted to Canada National Energy Board[45] evaluating effects of a proposed oil pipeline and associated tanker traffic. For the purposes of this PVA, projected increases in anthropogenic threats are not meant to mimic any one industrial development, but rather a general process of industrialization reflecting the number of port expansions, pipeline proposals, and liquefied natural gas terminal proposals pending for the BC coast[4]. For a low level scenario, we used the catastrophe option in Vortex to add the possibility of large (>16,500 m³) and smaller (>8,250 m³) oil spills. The frequencies of a big spill (0.21% chance per year) and a smaller spill (1.08%) were based on an industry projection of the likelihood of such spills caused by proposed increase in tanker traffic[71]. Based on the percent overlap of oil coverage and critical habitat, we estimate that if a large oil spill were to occur,
about 50% of the SRKWs would be killed due to direct exposure to the oil. We estimate that 12.5% of the SRKWs would be killed by exposure to oil from a smaller spill. For a scenario with higher level impacts of development, we doubled the frequency of oil spills.

These energy development scenarios also included an increase in vessel noise and disturbance of feeding, with the current vessel presence of 85% of time increased to 92.5% in the low level scenario and to 100% in the high level scenario. We also included a probability of additional deaths of killer whales due to ship strikes, with one death per decade in the low level and two deaths per decade in the high level scenario. Although some persistent organic pollutants might increase under increased industrial activity in the SRKW habitat, PCBs have been phased out of production and are in decline in at least some fish species in low-development basins[72]. Lacking data on likely long-term trends in the contaminant loads of SRKW prey, we did not include any change in such pollutants in these scenarios.

Climate change is projected to cause a decline in Chinook abundance[28], and we modelled this possibility with a projected 25% (low scenario) or 50% (high scenario) decrease in Chinook over the next 100 years.

**Demographic Management and Threat Management scenarios**

We used the PVA to simulate how much improvement in demographic parameters or how much reduction in anthropogenic threats, singly or in combination, would be required to reach a stated recovery objective of sustained annual population growth of 2.3% for 28 years[11]. In calculating the growth for these models, we started the tally 20 years into the simulation to avoid short-term demographic fluctuations as the age structure adjusts to new demographic rates, and growth was tallied over the subsequent 28 years. For the set of Demographic Management scenarios, we assessed the relationship between improved demography and population growth. Birth rate was incremented by 1.1x, 1.2x, 1.3x, 1.4x, and 1.5x, whereas calf mortality and adult mortality were decreased by 0.9x, 0.8x, 0.7x, 0.6x, and 0.5x. Next, in Threat Management scenarios, we modelled the effects of reduced threats, with the consequences resulting from the functional relationships to demography. We increased salmon abundance (up to the highest level of the Chinook index observed between 1979 and 2008, namely 1.3 times the long-term average). We simulated the improved demography if acoustic disturbance were reduced or eliminated. We considered the population consequences of improved calf survival resulting from reduction of PCBs, testing rates of future accumulation in SRKW from the estimated current 2 ppm/y to down to 0 ppm/y. Finally, we
tested scenarios that both reduced acoustic disturbance by half and increased salmon abundance up to 1.3x.

Electronic supplementary material

**Supplementary information** accompanies this paper at https://doi.org/10.1038/s41598-017-14471-0.

**Publisher's note:** Springer Nature remains neutral with regard to jurisdicational claims in published maps and institutional affiliations.

**Acknowledgements**

We thank the team members for the Center for Whale Research for providing the complete databases of censuses and demographic trends from 1976 through 2014. We thank Ailsa Hall, Sea Mammal Research Unit, University of St. Andrews, for providing the model of bioaccumulation of PCBs in cetaceans.

**Author Contributions**

P.P. coordinated the project. K.C.B. and D.A.G. provide the core census and demographic database. R.W., E.A., L.J.N.B., C.W.C., D.P.C., P.P., and M.M. provided data on threats. R.C.L. built the population model and conducted the simulations. R.W., R.C.L. and P.P. led the writing. All authors contributed to and reviewed the manuscript.

**Competing Interests** The authors declare that they have no competing interests.

Electronic supplementary material

**Supplementary Information**

**References**

1. Wright, A. J. & Kyhn, L. A. Practical management of cumulative anthropogenic impacts with working marine examples. *Conserv. Biol.* **29**, 333–340 (2014).

2. Côté, I. M., Darling, E. S. & Brown, C. J. Interactions among ecosystem stressors and their importance in conservation. *Proc. Biol. Sci.* **283**, 20152592,
https://doi.org/10.1098/rspb.2015.2592 (2016).

3. National Academy of Sciences. *Approaches to Understanding the Cumulative Effects of Stressors on Marine Mammals* (The National Academies Press, Washington, DC, 2016) https://doi.org/10.17226/23479.

4. Williams, R., Thomas, L., Ashe, E., Clark, C. W. & Hammond, P. S. Gauging allowable harm limits to cumulative, sub-lethal effects of human activities on wildlife: A case-study approach using two whale populations. *Mar. Policy* **70**, 58–64 (2016).

5. Maxwell, S. M. *et al.* Cumulative human impacts on marine predators. *Nat. Commun.* **4**, 2688, https://doi.org/10.1038/ncomms3688 (2013).

6. Lacy, R. C. *et al.* Metamodels for transdisciplinary analysis of population dynamics. *PLoS ONE* **8**, e84211, https://doi.org/10.1371/journal.pone.0084211 (2013).

7. Agardy, T., Di Sciara, G. N. & Christie, P. Mind the gap: addressing the shortcomings of marine protected areas through large scale marine spatial planning. *Mar. Policy* **35**, 226–232 (2011).

8. Boyce, M. S. Population viability analysis. *Annu. Rev. Ecol. Syst.* **23**, 481–506 (1992).

9. Reynolds, J. E. III, Marsh, H. & Ragen, T. J. Marine mammal conservation. *Endang. Species Res.* **7**, 23–28 (2009).

10. Fisheries and Oceans Canada. *Action Plan for the Northern and Southern Resident Killer Whale (Orcinus orca) in Canada*. Species at Risk Act Action Plan Series. (Fisheries and Oceans Canada, Ottawa, 2017).

11. National Marine Fisheries Service. *Recovery Plan for Southern Resident Killer Whales (Orcinus orca)* (National Marine Fisheries Service Northwest Region, Seattle, 2008).

12. National Marine Fisheries Service. *Southern Resident Killer Whales (Orcinus orca) 5-Year Review: Summary and Evaluation*. (National Marine Fisheries Service West Coast Region, Seattle, 2016)
13. Ward, E. J., Holmes, E. E. & Balcomb, K. C. Quantifying the effects of prey abundance on killer whale reproduction. *J. Appl. Ecol.* **46**, 632–640 (2009).

14. Ford, J. K. B., Ellis, G. M., Olesiuk, P. F. & Balcomb, K. C. Linking killer whale survival and prey abundance: food limitation in the oceans’ apex predator? *Biol. Lett.* **6**, 139–142 (2010).

15. Shaffer, M. L. Population viability analysis. *Conserv. Biol.* **4**, 39–40 (1990).

16. Beissinger, S.R. & McCullough, D.R. (eds). *Population Viability Analysis* (University of Chicago Press, Chicago, 2002).

17. Lacy, R. C. Structure of the VORTEX simulation model for population viability analysis. *Ecol. Bull.* **48**, 191–203 (2000).

18. Lacy, R.C. & Pollak, J.P. *Vortex: A Stochastic Simulation of the Extinction Process*. Version 10.0. (Chicago Zoological Society, Brookfield, Illinois, 2014); www.vortex10.org/Vortex10.aspx.

19. Lacy, R.C., Miller, P.S. & Traylor-Holzer, K. *Vortex10 User’s Manual* (IUCN SSC Conservation Breeding Specialist Group, and Chicago Zoological Society, Apple Valley, Minnesota, 2014).

20. Carroll, C., Fredrickson, R. J. & Lacy, R. C. Developing metapopulation connectivity criteria from genetic and habitat data to recover the endangered Mexican wolf. *Conserv. Biol.* **28**, 76–86 (2014).

21. Maehr, D.S., Lacy, R.C., Land, E.D., Bass, O.L. Jr. & Hoctor, T.S. Evolution of population viability assessments for the Florida panther: a multiperspective approach. In *Population Viability Analysis* (eds. Beissinger, S. R. & McCullough, D.R.) 284-311 (University of Chicago Press, Chicago, 2002).

22. Marmontel, M., Humphrey, S. R. & O’Shea, T. J. Population viability analysis of the Florida manatee (*Trichechus manatus latirostris*), 1976–1991. *Conserv. Biol.* **11**, 467–481 (1997).
23. Krahn, M.M. et al. 2004 Status review of Southern Resident killer whales (Orcinus orca) under the Endangered Species Act. NOAA Tech. Memo. NMFS-NWFSC-62 (U.S. Dept. Commerce, Washington, DC, 2013) https://www.nwwsc.noaa.gov/assets/25/6377_02102005_172234_krahnstatusrevtm62fi

24. Vélez-Espino, L. A. et al. Comparative demography and viability of northeastern Pacific resident killer whale populations at risk. Canadian Technical Report of Fisheries and Aquatic Sciences 2014, 3084 (2014).

25. Ward, E.J. et al. Estimating the impacts of Chinook salmon abundance and prey removal by ocean fishing on Southern Resident killer whale population dynamics. NOAA Tech. Memo. NMFS-NWFSC-123. (U.S. Dept. Commerce, Washington, DC, https://www.nwwsc.noaa.gov/assets/25/4647_08132013_113012_ImpactsOnSRKWsTM2013).

26. Prowse, T. A. A. et al. An efficient protocol for the sensitivity analysis of complex ecological models. Ecosphere 7, e01238, https://doi.org/10.1002/ecs2.1238 (2016).

27. Manlik, O. et al. The relative importance of reproduction and survival for the conservation of two dolphin populations. Ecol. Evol. 6, 3496–3512 (2016).

28. Muñoz, N. J., Farrell, A. P., Heath, J. W. & Neff, B. N. Adaptive potential of a Pacific salmon challenged by climate change. Nat. Clim. Change 5, 163–166 (2014).

29. Hickie, B. E., Ross, P. S., Macdonald, R. W. & Ford, J. K. B. Killer whales (Orcinus orca) face protracted health risks associated with lifetime exposure to PCBs. Environ. Sci. Technol. 41, 6613–6619 (2007).

30. Mongillo, T.M. et al. Exposure to a Mixture of Toxic Chemicals: Implications for the Health of Endangered Southern Resident Killer Whales. NOAA Tech. Memo. NMFS-NWFSC-135 (U.S. Dept. Commerce, Washington, DC, 2016) https://doi.org/10.7289/V5/TM-NWFSC-135.

31. Wasser, S. K. et al. Population growth is limited by nutritional impacts on pregnancy success in endangered Southern Resident killer whales (Orcinus orca). PLoS One 12, e0179824 (2017).
32. Witze, A. Let the river run: Removing two dams from the Elwha is a can’t-miss restoration experiment. *Science News* **187**, 22–26 (2015).

33. Sommer, T. R., Nobriga, M. L., Harrell, W. C., Batham, W. & Kimmerer, W. Floodplain rearing of juvenile Chinook salmon: evidence of enhanced growth and survival. *Can. J. Fish. Aquat. Sci.* **58**, 325–333 (2001).

34. Krkošek, M. *et al.* Declining wild salmon populations in relation to parasites from farm salmon. *Science* **318**, 1772–1775 (2007).

35. Miller, K. M. *et al.* Genomic signatures predict migration and spawning failure in wild Canadian salmon. *Science* **331**, 214–217 (2011).

36. Williams, R. *et al.* Competing conservation objectives for predators and prey: estimating killer whale prey requirements for Chinook salmon. *PLoS ONE* **6**, e26738, https://doi.org/10.1371/journal.pone.0026738 (2011).

37. Hilborn, R. *et al.* The Effects of Salmon Fisheries on Southern Resident Killer Whales: Final Report of the Independent Science Panel (ESSA Technologies, Vancouver, BC, 2012).

38. Desforges, J.-P. W. *et al.* Immunotoxic effects of environmental pollutants in marine mammals. *Environment International* **86**, 126–139 (2016).

39. Williams, R., Erbe, C., Ashe, E., Beerman, A. S. & Smith, J. C. Severity of killer whale behavioral responses to ship noise: A dose-response study. *Mar. Pollut. Bull.* **79**, 254–60 (2014).

40. Williams, R., Erbe, C., Ashe, E. & Clark, C. W. Quiet(er) marine protected areas. *Mar. Pollut. Bull.* **100**, 154–161 (2015).

41. Leaper, R. & Renilson, M.R. A review of practical methods for reducing underwater noise pollution from large commercial vessels. *Trans. RINA* **154** (A2) (2012).

42. Veirs, S., Veirs, V. & Wood, J. D. Ship noise in an urban estuary extends to frequencies used for echolocation by endangered killer whales. *PeerJ* **4**, e1657, https://doi.org/10.7717/peerj.1657 (2016).
43. Malakoff, D. A push for quieter ships. *Science* 328, 1502–1503 (2010).

44. Silber, G. K., Vanderlaan, A. S. M., Arceredillo, A. T. & Johnson, L. L. The role of the International Maritime Organization in reducing vessel threat to whales: process, options, action and effectiveness. *Mar. Policy* 36, 1221–1233 (2012).

45. Canada National Energy Board. *National Energy Board Report. Trans Mountain Expansion Project* (National Energy Board, Calgary, Alberta.). https://docs.neb-one.gc.ca/ll-eng/lisisapi.dll/2969681/National_Energy_Board_Report_-_OH-001-2014_-_A5A9H1.pdf 2016

46. Gaydos, J. K., Thixton, S. & Donatuto, J. Evaluating threats in multinational marine ecosystems: a Coast Salish First Nations and tribal perspective. *PLoS ONE* 10, e0144861, https://doi.org/10.1371/journal.pone.0144861 (2015).

47. King, S. L., Schick, R. S. & Donovan, C. An interim framework for assessing the population consequences of disturbance. *Method. Ecol. Evol.* 6, 1150–1158 (2015).

48. González-Suárez, M., Lucas, P. M. & Revilla, E. Biases in comparative analyses of extinction risk: mind the gap. *J. Anim. Ecol.* 81, 1211–1222 (2012).

49. Donlan, C., Wingfield, D. K., Crowder, L. B. & Wilcox, C. Using expert opinion surveys to rank threats to endangered species: a case study with sea turtles. *Conserv. Biol.* 24, 1586–1595 (2010).

50. Teck, S. J., Halpern, B. S. & Kappel, C. V. Using expert judgment to estimate marine ecosystem vulnerability in the California Current. *Ecol. Appl.* 20, 1402–1416 (2010).

51. Irvine, J. R. & Fukuwaka, M. A. Pacific salmon abundance trends and climate change. *ICES J. Mar. Sci.* 68, 1122–1130 (2011).

52. Noss, R. F., Quigley, H. B., Hornocker, M. G., Merrill, T. & Paquet, P. C. Conservation biology and carnivore conservation in the Rocky Mountains. *Conserv. Biol.* 10, 949–963 (1996).
53. Bigg, M. A. & Wolman, A. A. Live-capture killer whale (Orcinus orca) fishery, British Columbia and Washington, 1962-73. J. Fish. Res. Board Can. 32, 1213–1221 (1975).

54. Hanson, M. B. et al. Species and stock identification of prey consumed by endangered southern resident killer whales in their summer range. Endang. Species Res. 11, 69–82 (2010).

55. Ford, M. J. et al. Estimation of a killer whale (Orcinus orca) population’s diet using sequencing analysis of DNA from feces. PLoS ONE 11, e0144956, https://doi.org/10.1371/journal.pone.0144956 (2016).

56. Olesiuk, P. F., Bigg, M. A. & Ellis, G. M. Life history and population dynamics of resident killer whales (Orcinus orca) in coastal waters of British Columbia and Washington State. Rep. Int. Whal. Comm. 12, 209–243 (1990).

57. Center for Whale Research. Southern Resident Orca Survey. Review Copy. 20150320 update. (Center for WhaleResearch, Friday Harbor, Washington, 2015).

58. Ford, M. J. et al. Inferred paternity and male reproductive success in a killer whale (Orcinus orca) population. J. Hered. 102, 537–553 (2011).

59. O’Grady, J. J. et al. Realistic levels of inbreeding depression strongly affect extinction risk in wild populations. Biol. Conserv. 133, 42–51 (2006).

60. Williams, R. & Lusseau, D. A killer whale social network is vulnerable to targeted removals. Biol. Lett. 2, 497–500 (2006).

61. Vélez-Espino, L. A. et al. Relative importance of Chinook salmon abundance on resident killer whale population growth and viability. Aquat. Conserv. 25, 756–780 (2015).

62. Ward, E. J., Parsons, K., Holmes, E. E., Balcomb, K. C. I. I. I. & Ford, J. K. B. The role of menopause and reproductive senescence in a long-lived social mammal. Front. Zool. 6, 4, https://doi.org/10.1186/1742-9994-6-4 (2009).

63. Ford, J.K.B., Wright, B.M., Ellis, G.M., & Candy, J.R. Chinook Salmon Predation by Resident Killer Whales: Seasonal and Regional Selectivity, Stock
Identity of Prey, and Consumption Rates. (Fisheries and Oceans Canada, Nanaimo BC, 2009).

64. Williams, R., Clark, C. W., Ponirakis, D. & Ashe, E. Acoustic quality of critical habitats for three threatened whale populations. *Anim. Conserv.* **17**, 174–185 (2014).

65. Lusseau, D., Bain, D. E., Williams, R. & Smith, J. C. Vessel traffic disrupts the foraging behavior of southern resident killer whales *Orcinus orca*. *Endang. Species Res.* **6**, 211–221 (2009).

66. Morris-Drake, A., Kern, J. M. & Radford, A. N. Cross-modal impacts of anthropogenic noise on information use. *Curr. Biol.* **26**, 2697–2840 (2016).

67. Hall, A. J. *et al.* Population consequences of polychlorinated biphenyl exposure in bottlenose dolphins – an individual based model approach. *Environ. Health Perspect.* **114**(Suppl. 1), 60–64 (2006).

68. Hall, A.J. *et al.* Assessing the population consequences of pollutant exposure to cetaceans using an individual based modelling framework. Tech. Rep. of International Whaling Commission, Tromsø, Norway, SC63–E5 (2011).

69. Hall, A.J. & Williams, R. The potential effect of PCBs on Killer whales – using the ‘SPOC’ individual based pollution model approach to estimate impacts on population growth. IWC SC meeting document SC/66a/E/2 (2015).

70. Ross, P. S., Ellis, G. M. & Ikonomou, M. G. High PCB concentrations in free-ranging Pacific killer whales, *Orcinus orca*: Effects of age, sex and dietary preference. *Mar. Pollut. Bull.* **40**, 504–515 (2000).

71. Det Norske Veritas (USA). *Termpol 3.15 – General Risk Analysis and Intended Methods of Reducing Risks. Trans Mountain Expansion Project.* https://apps.neb-one.gc.ca/REGDOCS/File/Download/2393696 and https://apps.neb-one.gc.ca/REGDOCS/File/Download/2393360 (2013).

72. West, J. E., O’Neill, S. M. & Ylitalo, G. M. Time trends of persistent organic pollutants in benthic and pelagic indicator fishes from Puget Sound, Washington, USA. *Arch. Environ. Contam. Toxicol.* **73**, 207–229 (2017).
Supplementary Information -- Evaluating anthropogenic threats to endangered killer whales to inform effective recovery plans

Robert C. Lacy, Rob Williams, Erin Ashe, Kenneth C. Balcomb III, Lauren J. N. Brent, Christopher W. Clark, Darren P. Croft, Deborah A. Giles, Misty MacDuffee, Paul C. Paquet

Expanded modelling methods

Primary data source for baseline demographic rates

Demographic rates were calculated from the database on individual animal histories compiled by the Center for Whale Research\(^5\). Data from 1976 through 2014 were used. The SRKW population experienced moderate population growth from 1976 to 1993, and experienced a moderate decline subsequently. This may indicate that some aspects of the habitat declined in quality, or the change in trajectory may have been due to random fluctuations in reproduction and survival, rather than indicating any long-term trend. It might also be that the population peaked when it reached an ecological carrying capacity – the number of animals that can be supported by the available resource base. By including data back to 1976 for calculation of demographic rates, therefore, we may be providing an optimistic snapshot of current conditions. However, to restrict analyses to only the demography experienced in more recent years could under-estimate the capacity of the population to grow.

Vortex is an individual-based model, with the fate of each animal tracked through its lifetime. In contrast to population-based models or analytical calculations of average population growth, individual-based models can include individual variation in reproductive success, loss of genetic diversity, inbreeding, inbreeding avoidance, and any demographic consequences of these processes. We started the projections of population dynamics with the actual ages and sexes of the killer whales currently living in 2015. We also specified the mother of each animal, when it was known (for 76 of 80 living animals)\(^5\). These designations of maternity are important, because we specified in the simulation model that females would not mate with their father, son, or maternal half-
sibling. For the oldest animals in the population, we do not know mothers or other close relatives (other than their own offspring). Thus, the analyses will underestimate the accumulation of inbreeding, because living animals may be related in ways not documented.

Because the sample sizes are necessarily small, the data are not sufficient to determine accurate estimates of demographic rates for each annual age class. We therefore used the same age class groupings as were used in recent models\(^{34,61}\), except that we set an upper limit for female breeding at 45y rather than 50y. Not all SRKW calves and juveniles have been sexed, and the data do not indicate a significant difference in survival between males and females in these younger age classes, so survival was calculated with both sexes combined for calves and for juveniles. Survival rates of young adult females and young adult males have been better than older females and males, so it is appropriate to consider these age classes separately in the analyses. Killer whales can survive many years after reproduce senescence\(^{62}\). Although it is not known what age they can reach, and estimates of birth dates for the older animals in the population are uncertain, one female has been estimated to be 104 years old. The next oldest females are estimated to be 87y and 50y. We set an upper limit of age to 90y in our models. Although age estimates for the oldest animals are highly uncertain, the upper limit that we place on age in the model will have no impact on long-term population growth in our model. Post-reproductive females might provide benefits to the pod other than through reproduction, but we do not have quantitative data on such effects that would allow us to include them in the population model.

**Reproductive system**

The breeding system is polygamous, with some males able to obtain multiple mates, and females mating with different males over their lifetimes. There is evidence that there is variation among males and among females in reproductive success. Genetic evidence indicates that the effective population size is about 1/3 of the total size\(^{58}\), as would occur if some animals are more successful breeders than others. To represent this phenomenon in the PVA model, we assigned each male a score from 0 to 1 as his likelihood of being a
capable breeder (i.e., with full access to breeding females) each year. The mean was set at 0.50, with a SD of 0.20 in a beta-distributed distribution. This leads to about 75% of offspring being sired by 15% of the males born in our Baseline model. Similarly, we assigned an individual reproductive success score to each female, used to represent the factor by which the female’s probability of breeding each year was below (score <1) or above (score >1) the overall mean for females of her age. These scores for variation in female reproductive success had mean = 1.0 and SD = 0.2, created by sampling from a beta distribution (mean = 0.50, SD = 0.1) and doubling those values. The beta distribution was chosen over a normal distribution to ensure that values were bounded between 0 and 1.

Genetic data indicate that females usually mate with males from the other pods, but occasionally do mate within their own pod⁵⁸. However, no cases of father-daughter, mother-son, or maternal half-sibling matings were observed, even though the within-pod matings would make such inbreeding statistically likely (given the small population size) if mating were random. To include this inbreeding avoidance in the model, we specified that females would not mate with their fathers, sons, or brothers or half-brothers from the same mother. Matings between paternal half-siblings and more distant relatives were allowed in the model. The avoidance of very close inbreeding will reduce the extent of damage caused by inbreeding in the closed SRKW population, but could also lead to a situation in which some females might be unable to find a suitable mate if the population becomes small enough so that no unrelated males are available.

**Reproductive rates**

Breeding rates – expressed as the proportion of the females of an age class that produce a calf each year – were calculated to be:

12.04% for young adult females (age 10-30), with SD = 3.54

7.88% for older females (age 31-45), with SD = 4.15

0% for post-reproductive females (age > 45)
The above standard deviations expressing variation in rates across years were adjusted by removing the part of the observed variation that would be expected to be due to the demographic stochasticity (chance fluctuations in numbers breeding even when probabilities of breeding are constant), following methods in Lacy et al.\textsuperscript{19}. Our estimated fecundities are a little higher than those estimated from the 1987-2011 data\textsuperscript{24}. Our estimates of annual variation are lower because we removed the demographic stochasticity.

**Mortality**

Mortality rates were calculated from the tallies of deaths each year from 1976 through 2014. The mortality rate for each age-sex class was averaged across the 39 years of data to obtain the mean annual rates. Calculated annual mortality rates and environmental variation (represented as a standard deviation) were as follows:

- **Calves:** 17.48\% (SD = 17.96)
- **Juveniles:** 2.15\% (SD = 2.42)
- **Young adult females:** 0.97\% (SD = 0; i.e., the variation in mortality across years was no greater than expected based on the demographic stochasticity if the probability of death is constant over time)
- **Older adult females:** 2.25\% (SD = 0)
- **Post-reproductive females:** 6.29\% (SD = 7.77)
- **Young adult males:** 3.03\% (SD = 2.53)
- **Older adult males:** 9.80\% (SD = 8.68)

The above mortality rates are all a little better (i.e., lower) than previously published rates\textsuperscript{24}, which is due to their use of data from 1987-2011, which omits a prior decade that had better demographic rates. The estimates of annual variation above are all
considerably lower than previous estimates because of our removal of the expected demographic stochasticity from the annual variation in observed death rates.

**Impacts of inbreeding**

Long-term pedigree information are not available from which the effect of inbreeding could be determined for killer whales or for any cetacean species. The reported avoidance of mating between some closely related pod members, however, indicates that the species has reduced fitness when inbred. Lacking data on the severity of inbreeding depression in cetaceans, we assumed in the model that the effect of inbreeding on reproductive success (birth rate and calf survival) is the same as the mean value reported (6.29 “lethal equivalents”) in a survey of impacts on wild species. Inbreeding might also affect other components of fitness – such as longevity – but data on such effects are too few to allow estimation of typical impacts, and we therefore did not include those effects in our model. We used the option in Vortex to model inbreeding depression as being caused by the presence of recessive lethal alleles. Each initial animal in the model was assigned an average of 6.29 unique lethal alleles across its genome, distributed among founders according to a Poisson distribution. Vortex simulates the transmission of alleles at multiple loci through the pedigree, with offspring receiving an allele from each parent at each locus. Inbreeding depression is modeled by the death of any offspring that are homozygous for one of the recessive lethal alleles. See the Vortex manual for further description of how genetics and inbreeding depression are modelled.

**Sensitivity tests**

To determine the sensitivity of the model projections to the uncertainty in 11 demographic input variables, we ran 100 iterations each of 1000 combinations of input parameters sampled from ranges. The sampled parameters were evenly spaced across the range set for each, according to a “Latin hyperspace” design. This method of sampling provides high statistical power for determination of the effect of each parameter. The ranges sampled for each variable were:

Inbreeding effects: Lethal equivalents = 3.145 to 9.435 (± 50% around the baseline value)
Male variance in Reproductive Success: \( SD = 0 \text{ to } 0.4 \)

Female variance in Reproductive Success: \( SD = 0 \text{ to } 0.4 \)

Birth rate: \( \pm 10\% \) around the baseline values

Annual variation in birth rate: \( \pm 10\% \) around the baseline values

Calf mortality rate: \( \pm 10\% \) around the baseline values

Annual variation in calf mortality: \( \pm 10\% \) around the baseline values

Juvenile mortality rate: \( \pm 10\% \) around the baseline values

Annual variation in juvenile mortality: \( \pm 10\% \) around the baseline values

Adult mortality rate: \( \pm 10\% \) around the baseline values

Annual variation in adult mortality: \( \pm 10\% \) around the baseline values

The above ranges for sensitivity tests are different from a common practice of examining the amount of change in the population growth for a constant proportional change in each variable. However, the apparent standardization of sensitivities of growth to changes in variables is arbitrary. It is arbitrary because the same proportional change in different demographic rates might not be likely or achievable for the population of interest. For other model variables, such as the inbreeding depression, it is not clear that a 10% change in the value has the same meaning as a 10% shift in other demographic rates measured on very different and not always linear scales. Moreover, different results can be obtained if ages are binned into more or fewer age class categories for the calculations. Although strict comparison of the effect of proportional changes in model parameter values is often not meaningful, the sensitivity tests can be useful guides as to which model parameters are the most important determinants of population trajectories, over the range of values that are chosen for testing. Thus, it is often useful to vary parameter values by amounts that reflect our uncertainty in each input value.
To test the effect of inbreeding avoidance on population growth, we compared the baseline model to one in which there was no avoidance of close inbreeding.

Modelling of impacts of Chinook salmon abundance

A large body of evidence has shown a strong relationship between abundance of Chinook salmon and demographic rates for killer whales\textsuperscript{13,14,24}. In the previous studies, details of the data used to quantify Chinook abundance, the co-factors considered, and the statistical methodologies differ in some key respects. One study examined fecundity for young and old females separately and examined impacts on survival rates of different age classes\textsuperscript{24}, whereas others\textsuperscript{13,14} assessed relationships with pooled demographic data. Each derived an index of relative prey abundance, but the salmon stocks included and the period over which the indices were averaged differed among studies. Studies have examined the relationships with various time lags (0 to 2 years), although for projecting long-term population viability in simulation models, it is not important if changes in fecundity and survival due to changing Chinook abundance occur in the year of the Chinook index or with a lag of a year or two.

All three studies above identified large effects of Chinook salmon abundance on SRKW demographic rates. Two regression models are available that link the relationship between salmon abundance and logit(birth rate)\textsuperscript{13,24}. We rescaled the regressions provided by both studies so that Chinook abundance index was set to 1.0 for the average over the span of data (1976-2014 for our demographic analyses), and so that the regression predicts the observed long-term average birth rate when Chinook index = 1. After that rescaling, the relationship for overall fecundity and for fecundity of the younger (more productive) females are very similar to the previous studies\textsuperscript{13,24} and are both approximately:

\[
\text{logit(birth rate)} = -3.0 + \text{Chinook}
\]

in which Chinook is the abundance relative to the long-term average.

This relationship leads to a predicted shift from a birth rate of 12% to 18.5% if the Chinook abundance increases by 50%, and a shift down to 7.6% if prey abundance
decreases by 50%. A linear relationship derived by Ford et al.\textsuperscript{14} would lead to smaller but still significant shifts (up to 13.3% and down to 9.2%, respectively) in the birth rate.

There is evidence for a stronger effect on fecundity of older females in relation to the Puget Sound terminal run, but a similar slope for Fraser + Puget abundance\textsuperscript{24}. If the intercept in equation (1) is adjusted so that the fecundity of the older breeding females is correctly predicted at the baseline of Chinook = 1, then the relationship for that age class becomes approximately:

\[ \text{logit}(\text{birth rate}) = -3.46 + \text{Chinook} \] \hspace{1cm} (2)

Ford et al.\textsuperscript{14} provide the relationship of mortality to Chinook abundance in a form that is easily transferred to the Vortex model, namely:

\[ \text{Mortality index (relative to the expected value)} = 4.0066 - 2.6504 \times \text{Chinook (relative abundance index)} \] \hspace{1cm} (3)

Rescaling this relationship so that the mortality index is 1 when Chinook = 1 gives:

\[ \text{Mortality index (relative to the baseline)} = 3.6504 - 2.6504 \times \text{Chinook} \] \hspace{1cm} (4)

This relationship translates to a 26% increase in mortality for each 10% decrease in Chinook abundance. We used equation (4) to relate Chinook salmon abundance to SRKW mortality in our PVA.

The rescaling of the relationships above was confirmed to produce the same population growth in the Vortex model ($r = -0.002$, with SD = 0.045) as was projected from the baseline model. These general relationships can then be used for projecting the primary impacts of changing Chinook abundance on population viability. We note that other regressions might be more appropriate for identifying the consequences to SRKWs of losing individual salmon stocks (“runs”), or the benefit of restoring runs in particular streams. The above-mentioned equations are appropriate for looking at broad-scale changes in SRKW habitat.

**Sensitivity testing of model parameters**
We ran simulations with higher and lower values of each demographic parameter to determine the importance of each to population growth. For each model parameter that was tested, Table S1 gives the value used in the baseline model (birth rate calculated separately for younger and older females), the minimum and maximum values of the range that was tested, the population growth rates that resulted from the min value (r-min) and max value (r-max) for that parameter while all other parameters were held at their baseline, and the % of variance in population growth (as a percent of the total variation accounted for by the varied parameters) due to variation in that parameter when all other parameters were simultaneously and independently varied across their ranges of uncertainty.

Table S1. Values for each parameter in the sensitivity analysis, their effect on projected population growth, and proportion of variation explained by variation in that parameter. The residual variance is that due to non-linearities of and interactions among effects.

| Parameter                  | Baseline | Min tested | Max tested | r-min   | r-max   | %variance |
|----------------------------|----------|------------|------------|---------|---------|-----------|
| Inbreeding depression      | 6.29     | 3.145      | 9.435      | -0.0005 | -0.0029 | 6.1       |
| Variance in male RS        | 0.20     | 0.00       | 0.40       | -0.0017 | -0.0023 | 0.4       |
| Variance in female RS      | 0.20     | 0.00       | 0.40       | -0.0017 | -0.0019 | 0.0       |
| Mean birth rate            | 0.1204   | 0.1084     | 0.1324     | -0.0062 | 0.0019  | 77.2      |
| SD in birth rate           | 0.0354   | 0.0319     | 0.0389     | -0.0020 | -0.0018 | 0.0       |
| Calf mortality             | 0.1748   | 0.1573     | 0.1923     | -0.0012 | -0.0029 | 3.2       |
| SD in calf mortality       | 0.1796   | 0.1616     | 0.1976     | -0.0017 | -0.0019 | 0.0       |
| Juvenile mortality         | 0.0215   | 0.0194     | 0.0237     | -0.0011 | -0.0028 | 3.1       |
| SD in juv. Mortality       | 0.0242   | 0.0218     | 0.0266     | -0.0015 | -0.0019 | 0.0       |
When inbreeding avoidance was dropped from the Baseline model, mean population growth was reduced very slightly, from $r = -0.0020$ to $r = -0.0024$.

**References**

All references cited in this Supplementary Information are given in the main manuscript.