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ABUNDANCE OF HUMPBACK WHALES (*MEGAPTERA NOVAEANGLIAE*) WINTERING IN CENTRAL AMERICA AND SOUTHERN MEXICO FROM A ONE-DIMENSIONAL SPATIAL CAPTURE-RECAPTURE MODEL

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Executive Summary

Humpback whales (*Megaptera novaeangliae*) off the U.S. West Coast are a mixture of whales from different Distinct Population Segments (DPSs) under the U.S. Endangered Species Act (ESA), predominantly Central America and Mexico. Within DPSs, demographically independent populations (DIPs) of humpback whales are delineated as 'migratory herds' that share both wintering and feeding areas (Martien et al., 2020). The Central America DPS, composed of those whales that winter along the Pacific coast of Central America from Panama to Guatemala, corresponds to a single DIP that migrates almost exclusively to the U.S. West Coast. This DIP's wintering area is understood to extend into southern Mexico, and it is termed the CentAm/SMex-CA/OR/WA DIP for its wintering area and its feeding area off California, Oregon, and Washington (Fig. 1) (Taylor et al., 2021). The Mexico DPS includes multiple DIPs, with the DIP that migrates between northern mainland Mexico and the U.S. West Coast correspondingly termed the MMex-CA/OR/WA DIP (Martien et al., 2021). DIP-specific estimates of the number of whales using the U.S. West Coast Exclusive Economic Zone (EEZ), which are thus equivalent to estimates of the number of whales from each DPS using this feeding area, would be useful for management and decision-making involving these population units.

We estimated abundance for the CentAm/SMex-CA/OR/WA DIP from photo-identification data collected in their wintering area from 2019 to 2021. A randomization test suggested some fidelity in individual space use off Central America and Southern Mexico, implying that variable effort in time and space should be considered in capture-recapture estimates. We fitted a closed, one-dimensional spatial capture-recapture model to annual capture histories using a Bayesian framework. We accounted for uncertainty in the northern limit of the population and in the potential for movement of individuals across that limit by varying the northern population limit within the base model and exploring sensitivity to the northern boundary of the model domain in two alternate models.

We multiplied posterior distributions of abundance from the base and alternate models by a correction factor distribution, which was based on prior and new simulation work quantifying key expected sources of bias. The main sources of bias anticipated in estimating abundance for this dataset from a closed population model include births and deaths during the period of data collection, exclusion of first-year calves from the dataset, and sex heterogeneity in capture probability, which we assessed at a 3.4-fold (CV=0.463) greater chance of photo-identifying male individuals than females off Central America and Southern Mexico. The mean resulting correction factor is 1.35 (CV=0.143).

The base model produces a mean bias-corrected abundance estimate of 1,496 (CV=0.171), with a 20th percentile of 1,284. Alternate models with different northern model domain boundaries produce mean bias-corrected estimates of 1,313 (CV=0.167) and 1,601 (CV=0.166), corresponding to -12% and 7% differences in the 20th percentile from the base model. Comparison of the new abundance estimate for the

CentAm/SMex-CA/OR/WA DIP to one from 2004-06 that omits southern Mexico (Wade, 2021) suggests that the annual population growth rate is much lower than the 8.2% rate estimated for humpback whales off the U.S. West Coast as a whole (Calambokidis and Barlow, 2020). Population growth rate calculated directly from the current estimate including Southern Mexico and the Wade estimate is 4.8% per year (SD = 2.0%). Resummarizing our model results to exclude Southern Mexico animals results in a rate of 1.6% per year (SD = 2.0%).

We deduced the number of humpback whales from the MMex-CA/OR/WA DIP migrating to the U.S. West Coast EEZ by subtracting the new estimate for the CentAm/SMex-CA/OR/WA DIP from the most recent estimate of total abundance in the U.S. West Coast EEZ used in the draft 2021 stock assessment report (Calambokidis and Barlow, 2020; 86 FR 58887, October 25, 2021). The resulting mean estimate of abundance for humpback whales from the MMex-CA/OR/WA DIP using U.S. West Coast waters is 3,477 animals (CV=0.101).

Introduction

Humpback whales (*Megaptera novaeangliae*) have a circumglobal distribution, with most populations migrating between high-latitude feeding areas and low-latitude wintering areas (Kellogg, 1929). Individuals show fidelity to both their feeding and wintering areas. In the North Pacific, the relationships among humpback whale wintering and feeding areas are predominantly many-to-many, with animals from multiple wintering areas occurring in one feeding area, and vice versa (Calambokidis et al., 2001; Baker et al., 2013).

Under U.S. law, humpback whales are protected by the Endangered Species Act (ESA) and the Marine Mammal Protection Act (MMPA). While the two laws share similar goals of conservation and management of threats, the units for conservation differ. The unit of conservation under the ESA is a taxonomic species or a Distinct Population Segment (DPS) of a species. The unit of conservation under the MMPA is a “stock” of marine mammals, usually comprised of one or more demographically independent populations (DIPs).

Humpback whales in the North Pacific have been split into five DPSs under the ESA, based on genetic differences among wintering areas (Baker et al., 2013; 81 FR 62260, September 8, 2016). The vast majority of humpback whales occurring off the U.S. West Coast belong to the Central America DPS and the Mexico DPS (Calambokidis et al., 2000; Wade et al., 2016), which are listed as endangered and threatened, respectively, under the ESA. A much smaller number of whales from the Hawaii DPS have also been sighted or tagged in U.S. West Coast waters (Palacios et al., 2020; Calambokidis et al., 2000, 2001), and one animal seen near the border of Washington and British Columbia was even identified as being from the Western North Pacific DPS (Darling et al., 1996).

At the population level, individuals that share both feeding and wintering areas are considered to belong to the same “migratory herd”, a unit considered to be demographically independent from other migratory herds (Martien et al., 2020). Almost all whales in the Central America DPS migrate to the U.S. West Coast to feed, which corresponds to a single demographically independent population (DIP), known as the CentAm/SMex-CA/OR/WA DIP (Taylor et al., 2021). Whales from the Mexico DPS migrate either to the U.S. West Coast or to destinations further north off British Columbia, Alaska, the Aleutian Islands, and Russia, so they include multiple DIPs (Calambokidis et al., 2000; Urbán et al., 2000; Wade et al., 2016; Martien et al., 2021; Cheeseman, unpublished data). Those whales migrating from mainland Mexico to the U.S. West Coast are termed the MMex-CA/OR/WA DIP (Martien et al., 2021). Recent population estimates are available for total humpback whales in the U.S. West Coast EEZ (Becker et al., 2020; Calambokidis and Barlow, 2020), but not for abundance of whales from each of the two main contributing DIPs, which also would align with DPS-specific abundances.

The abundance of humpback whales in the North Pacific has been estimated at the basin scale and for individual feeding and wintering areas from photo-identification data collected in a coordinated effort called SPLASH (Structure of Populations, Levels of Abundance and Status of Humpback Whales in the North Pacific) (Barlow et al., 2011; Wade et al., 2016; Wade, 2021). SPLASH sampled all known wintering and feeding areas of humpback whales synoptically and repeatedly from 2004 to 2006 (Calambokidis et al., 2008). The wintering area abundance estimate from SPLASH for the Central America DPS is now 17 years old and also geographically out of date. Very few samples were obtained from Southern Mexico during SPLASH, and it was not included in either of the adjacent DPSs (Central America or Mexico) in analysis. The wintering area of the CentAm/SMex-CA/OR/WA DIP is now recognized as including southern Mexico based on recent photo-identification and genetic data, adding many animals to the population and rendering historic estimates inapplicable (Fig. 1; García Chávez et al., 2015; Ramírez Barragan et al., 2019; Martínez-Loustalot et al., 2020; Taylor et al., 2021). The abundance of humpback whales off the U.S. West Coast has been updated using mark-recapture analysis of photo-identification data collected off California and Oregon, with the most recent estimate ($4,973 \pm 239$ SE) based on data from 2015 to 2018 (Calambokidis and Barlow, 2020). A separate study using independent data and a different approach, habitat-based density estimation based on 2018 line-transect data along the entire U.S. West Coast, arrived at similar numbers, though with less precision ($4,784 \pm 1,469$ SE) (Becker et al., 2020). The former estimate was used in the draft 2021 stock assessment report for California/Oregon/Washington humpback whales (86 FR 58887, October 25, 2021).

Humpback whale photo-identification effort varies temporally and spatially off the Central American, Mexican, and U.S. West Coasts, which violates the assumptions of simple mark-recapture analysis methods. Moreover, humpback whales off the U.S. West Coast have been observed to exhibit spatial fidelity within their feeding area,

returning to similar locations each year (Calambokidis, unpublished data), which leads to individual heterogeneity in capture rates if effort is not spatiotemporally uniform. Similar individual fidelity to specific wintering locations is also likely. Spatial capture-recapture models, originally developed for terrestrial wildlife studies employing camera trap arrays, allow analysis of data with spatiotemporal heterogeneity in capture probability and individual variation in space use. These models have been adapted to one-dimensional systems, such as rivers, which closely approximate the narrow band of habitat along the coast that humpback whales inhabit (Royle et al., 2013).

We conducted a closed-population, mark-recapture analysis to estimate abundance of the CentAm/SMex-CA/OR/WA DIP from photo-identification data collected in its wintering area. First, we assessed divergence of individual space use from random for this population within its wintering area. We adapted a closed, one-dimensional, spatial capture-recapture framework to the context of spatially continuous photo-identification data collected in a narrow coastal band without effort information, and estimated population size from three years of spatial annual capture histories from 2019 to 2021. Sex heterogeneity in capture probability can also be an important source of bias in abundance estimates based on photo-identification data from wintering areas (Brown et al., 1995; Barlow et al., 2011). We estimated sex heterogeneity in capture probabilities in the CentAm/SMex-CA/OR/WA DIP's wintering area from capture histories of known-sex animals, quantified this source of anticipated bias in the abundance estimate using simulation, and combined the results with existing estimates of bias due to births, deaths, and exclusion of first-year whales from the North-Pacific-wide analysis to estimate a corrected abundance (Barlow et al., 2011). Finally, we calculated an estimate of the number of whales from the MMex-CA/OR/WA DIP using U.S. West Coast waters by subtracting the abundance estimate for the CentAm/SMex-CA/OR/WA DIP from the most recent U.S. West Coast capture-recapture-based abundance estimate.

Methods

We used sightings of humpback whales that were individually identified based on pigmentation, scarring, shape, and trailing edge serration in photographs of the ventral side of their flukes. These were collected both opportunistically and during dedicated research surveys, primarily from small vessels conducting daily trips from different locations in five Central American countries and southern Mexico (western Panama to Guerrero, Mexico) during the winter season. The winter season for a given year is defined as November of the preceding year to April, and also corresponds to the annual occasions we used for mark recapture analysis, so henceforth “year”, “annual”, or “occasion” all refer to this definition of the winter season. While our analysis focused on the data from the 2019 to 2021 seasons, we also used the entire available data series from 1988 to 2021 for some elements of the analysis as detailed below. These photographic identifications have been the basis of a number of past analyses using

subsets of these data (e.g., Rasmussen et al., 2011; Dobson et al., 2015; García Chávez et al., 2015; Steiger, et al., 2017; Ramírez Barragan et al., 2019; Ortega-Ortiz et al., In press). These include a coordinated effort and analyses from a collaborative North-Pacific-wide effort called SPLASH, conducted from 2004 to 2006, that included Central America and southern Mexico (Calambokidis et al. 2008, Barlow et al. 2011, Wade et al. 2016).

Photographic equipment and scoring and matching of identification photographs have changed through the years. Prior to 2005, photographs were generally taken with film SLR cameras with telephoto lenses, and after 2005, generally with Digital SLR cameras. For each individual humpback whale photographed in an encounter, the best photograph of the ventral side of the fluke was selected. Photographs were historically assigned to three quality levels based on five quality features (proportion visible, vertical angle, lateral angle, focus/sharpness, and exposure), using the same process developed previously (Calambokidis et al., 1997, 2000, 2008); more recently, a fourth, lower quality category has been added, as automated matching capabilities exceeded manual matching capabilities (see below) (Table 1). Photographs in the lower two quality levels were generally only retained for matching under special circumstances (e.g., photographs of individuals with a biopsy or known since birth). Photographs of individuals that matched the existing Cascadia Research Collective catalog of known individuals were assigned the corresponding identification number; those that did not match were assigned a new identification number and added to the catalog as new individuals. Through 2015, matching was performed manually by experienced matchers who compared photographs to similar ones based on general coloration patterns (Katona and Whitehead, 1981). From 2016 onward, increasing levels of automated matching were incorporated into the photo-identification matching process as automated matching algorithms developed and implemented through the Wildbook and Happywhale platforms. Initially, image recognition automation assisted with finding matches while manual review was retained to confirm new individuals (Flynn et al., 2017; Weideman et al., 2017; Weideman et al., 2020). In 2019, a sufficiently accurate algorithm was established that exceeded the accuracy of manual matching, allowing review of all previous identifications and more rapid management of new data (Cheeseman et al., 2021). All matches continue to be verified by humans.

We used data from the most recent three years (2019-2021), aggregated to annual occasions, to estimate abundance with a closed capture-recapture model. During this time, photographic identifications were not gathered in all locations in every year, especially in 2020 due to the start of the global pandemic in March, and identifications came from varying amounts of survey time in each region (Table 2). In 2021, a coordinated undertaking to obtain identifications in all countries/regions resulted in more dedicated surveys, more even effort among locations, and the largest sample of identifications compared to any previous year (Table 2). For the 2019-2021 period, information on survey effort was available only for 2021 at the time of the analysis.

Several violations of the assumptions of a closed-population model with constant capture probability among occasions and individuals (i.e., an M0 model) are likely for this dataset, including (1) variation in effort with time, (2) non-random sampling of the population, (3) errors in photo matching to the catalog, and (4) violation of the population closure assumption. These sources of bias were, where possible, assessed directly for this population, and addressed through data filtering, model specification, or estimation of bias correction factors from simulation.

Data filtering

Photographs in the lower two quality levels were omitted to minimize one cause of non-random sampling, individual heterogeneity in capture probability resulting from lower quality thresholds for individuals of interest. With use of Happywhale's matching algorithm, this quality filter, and exclusion of first-year whales (which may undergo substantial change in markings), bias due to missed matches is expected to be negligible in this data set (Cheeseman et al., 2021).

Model development: Assessment of potential biases

We further considered the following potential sources of bias: varying effort by occasion, varying effort with space and potential for associated non-random sampling, and sex heterogeneity in capture probability in the wintering area leading to non-random sampling. We did not consider individual heterogeneity, effectively assuming that given sex and individual space use, all whales are otherwise equally likely to fluke up, migrate to the wintering area, and migrate at a time overlapping with sampling in the wintering area.

Extensive variation in annual captures is evident in both time and space from 2019 to 2021 (Fig. 2). Spatial variation in capture probability is a concern if the population does not mix completely over its range among capture occasions. We used photo-identification data collected in the Central American and southern Mexico wintering area from 1988 to 2021 to test whether individuals are redistributed randomly or show consistency in spatial location among annual occasions (i.e., site fidelity at a finer scale than the extent of the population's wintering area). Data were subset to individuals captured during more than one occasion (totaling 268 individuals, or 31% of all identified whales). Distance along the relatively linear coast of Central America and southern Mexico was approximated by latitude. To account for spatiotemporal variation in effort, we compared mean individual differences in capture latitude among occasions from 1,000 replicates each of true and permuted datasets, preserving the distribution of effort for each occasion. For each replicate, a true dataset was drawn from a complete daily-resolution dataset of captures, randomly subsampling at most one daily mean capture location per occasion per individual. A corresponding "test" dataset was created

by permuting individual identifications among all capture locations within each occasion. For each dataset, mean distance among annual capture locations was calculated for each individual, then averaged across individuals for an overall mean inter-occasion distance per dataset (see Appendix A for code). The resulting distributions of true and permuted distances among captures on different occasions show that the individual distances among captures on different occasions are less than expected at random in the wintering area, indicating some level of site fidelity within the population’s wintering area (Fig. 3). Results were similar when distances were calculated in kilometers from latitude and longitude. Whales may be identified en route to a destination further south, and much of the effort since 2014 has been in the northern portion of the wintering area in Southern Mexico, so we also ran the test constrained to data from 1988 to 2013, with similar results.

To assess sex heterogeneity in capture probability in the wintering area for the CentAm/SMex-CA/OR/WA DIP, we compared the number of annual recaptures of males versus females over the full time series (1988-2021), using individuals that were genetically sexed from biopsies taken independently off the U.S. West Coast (Baker et al., 2013; Martien et al., 2020). Photo-identifications for quantifying annual captures were filtered for quality following the same procedure as for capture-recapture analysis (see “Data Filtering”). Genetic sex assignment followed either of two methods, using the *Sry* gene (Gilson et al., 1998) or the ZFX/ZFY 5’ exonuclease qPCR assay (Morin et al., 2005). Where assigned sex from more than one biopsy for the same animal disagreed, the assigned sex from the more accurate Morin et al. (2005) technique was used. Biopsies were filtered to those for which certainty of associated tail fluke identification was positive or probable, including all biopsies collected before this field started being assigned. Males were recaptured an average of 0.81 times (± 0.133 SE, $n=68$), and females an average of 0.24 times (± 0.104 SE, $n=34$). We approximated the ratio of male to female recaptures, and thus capture probabilities, as a lognormal distribution with the median equal to the ratio of the means, $0.81/0.24 = 3.4$, and coefficient of variation (CV) CV_h estimated analytically from the CVs of male and female recapture frequencies, CV_m and CV_f , as

$$CV_h \approx \sqrt{CV_m^2 + CV_f^2} = 0.46$$

Model development: Specification and data preparation

We specified a closed capture-recapture model using parameter-expanded data augmentation (Royle et al., 2007; Royle and Dorazio, 2012), and fitted it using a Bayesian analytical approach. Capture histories were aggregated to annual occasions, with or without a spatial dimension, as described below. The observed dataset was augmented by all-zero capture histories such that the total number of M potential individuals did not constrain the posterior for total existing individuals in preliminary

Markov Chain Monte Carlo (MCMC) simulations. A latent state of existence was estimated for each individual z_i with probability ψ as

$$z_i \sim \text{Bernoulli}(\psi),$$

using a vague prior for ψ ,

$$\psi \sim \text{Uniform}(0, 1).$$

In a non-spatial model, realized abundance N is equal to total existing individuals $\sum z_i$, and expected abundance is ψM .

Given evidence for some level of spatial fidelity and the spatially and temporally unbalanced sampling design (Fig. 2), we adapted a spatial capture-recapture (SCR) framework to account for individual space use (Efford 2004; Royle et al., 2013). Spatial capture-recapture models were originally developed for terrestrial wildlife studies employing camera trap arrays, and allow analysis of data with spatiotemporal heterogeneity in capture probability and individual variation in space use. We approximated space as one-dimensional, in degrees latitude, since the adjoining coast is roughly linear over the range of the population's wintering area (Fig. 1). A latent 'center of activity' covariate s_i , a latitude value, was estimated for each individual i , constrained to the model domain. Those individuals with an estimated s_i within specified population limits, which are potentially distinct from but contained within the model domain boundaries, were counted as part of the population (Fig. 4). The southern model domain boundary was defined as the accepted southern limit of the DIP, at the southern border of Panama ($\sim 7.25^\circ\text{N}$). The northern model domain boundary was defined as the northern border of the Mexican state of Colima ($\sim 19.2^\circ\text{N}$), halfway between the northern coastal border of Guerrero ($\sim 18^\circ\text{N}$) and the southern end of Bahía de Banderas, Jalisco ($\sim 20.4^\circ\text{N}$). We used a vague prior for s_i , corresponding to the model domain:

$$s_i \sim \text{Uniform}(7.25, 19.2).$$

We accommodated uncertainty in the northern population limit, which may overlap with the Mexico DPS between the states of Guerrero and Nayarit (Taylor et al., 2021), by uniformly varying it from 18°N to 19.2°N in summarizing abundance from the posteriors. We explored model uncertainty in terms of sensitivity to the northern model domain boundary, and thus to the range of geographic activity centers from which animals are "allowed" to be sighted in Guerrero, by fitting models with alternate northern model domains bounded to the north instead at 18.6°N (approximate northern border of Michoacán state) and 20.4°N . Both of these alternate scenarios used a northern population limit of 18.6°N , resulting in an envelope of reasonable minimum and maximum population estimates that capture the extent of geographic uncertainty in the population's distribution.

To create spatial annual capture histories that we could fit without effort information, we discretized capture locations into eight "traps" based on breaks in capture locations for

the full 1988 to 2021 time series. Seven of these traps contained captures for the 2019 to 2021 dataset (Fig. 2). Spatial captures within each trap were approximated as a point location, calculated as the mean latitude of all daily-level captures in that trap, so trap point locations correspond roughly to locations of focused field effort. Preliminary analyses showed abundance estimates were not sensitive to shifting cutoff points between traps or to increasing the number of traps. Individual capture probability by occasion and trap was calculated as

$$p_{i,j,k} = p0_{j,k} e^{-(s_i - X_j)^2 / (2\sigma^2)},$$

where X_j is the latitude of trap j , $p0$ is the capture probability intercept, which was allowed to vary by occasion k and trap j and given a uniform prior between zero and one,

$$p0_{j,k} \sim \text{Uniform}(0, 1),$$

and σ is the standard deviation of a univariate normal model of individual space usage, with vague prior

$$\sigma \sim \text{Uniform}(0, 50).$$

The model for the individual encounter events is

$$y_{i,j,k} \sim \text{Bernoulli}(z_i p_{i,j,k}),$$

where $y_{i,j,k}$ is a binary variable recording whether individual i was observed in trap j and occasion k .

Preliminary simulations indicated that this model tends to underestimate abundance in the parameter space of this analysis, with more accurate results returned by a model assuming a constant capture probability intercept across traps and occasions. However, preliminary abundance estimates from the two models were very similar, and the individual activity centers estimated in the model with constant capture probability were likely biased northward by the geographic gradient in effort (Table 2), so we used the model with space-and time-varying capture probability intercepts.

A non-spatial model with time-varying capture probabilities (i.e., an Mt model; Otis et al., 1978) with an otherwise similar specification using Bayesian parameter-expanded data augmentation was also fitted to the data for comparison to the results of the SCR model described above, providing additional information with respect to model uncertainty.

Preliminary simulation results based on maximum likelihood estimation (MLE) suggested that a model incorporating individual capture heterogeneity (i.e., an Mh model; Otis et al., 1978), such as that caused by sex heterogeneity, would not produce reliable results in the parameter space of this analysis. Mh models explored included Chao's, Poisson, and Darroch's models. Directly modeling sex heterogeneity was also not a viable option, because little or no information on sex heterogeneity in capture

probability was captured within the sparse 2019-2021 dataset. Instead, sex heterogeneity was addressed by estimating bias from simulations (see Bias estimation). The posterior distribution for abundance was then multiplied by the estimated distribution for bias correction to obtain a corrected abundance posterior.

Three MCMC chains were run for 22,500 iterations each after a burn-in of 2,500 iterations for the non-spatial Mt model, and for 55,000 iterations each after a burn-in of 5,000 iterations for the SCR model. MCMC chain mixing and stability and posterior sample size were evaluated by examining trace plots, running mean plots, and potential scale reduction factors (psrf) for each parameter. Trace plots of all parameters in each model showed good mixing of all three chains, running mean plots demonstrated stable results, and psrf was ≤ 1.01 for both point estimates and credible intervals.

Posterior-predictive goodness-of-fit checks were performed as described in Royle et al. (2013), including for deviation of spatial activity centers from a uniform distribution, deviation of expected from observed number of captures per individual per trap, deviation of expected from observed number of captures per trap, and deviation of expected from observed number of captures per individual. A posterior predictive check was also performed for the non-spatial Mt model results to evaluate deviation of expected from observed number of captures per individual.

Code for models and posterior predictive checks is provided in Appendix B.

Bias estimation

The magnitude of some important sources of bias had previously been estimated via simulation, and corresponding corrections estimated, for a three-year photo-identification data set of North-Pacific-wide humpback whales fitted with a closed capture-recapture model (Barlow et al., 2011). These sources of bias include not sampling animals in their first year (one source of non-random sampling) and violation of the population closure assumption.

To estimate the magnitude of bias expected from sex heterogeneity in capture probability, 5000 data sets were simulated based on abundance and occasion-specific capture probabilities estimated from the SCR and Mt models, respectively, as described above, an assumed sex ratio of 1:1, and a ratio of male to female capture probabilities set, in turn, to the 16th, 50th, and 84th percentiles of the lognormal distribution for the ratio estimate (see “Model development: Assessment of potential biases”). Male capture probabilities were set to those estimated from the Mt model. An Mt model was fitted to each dataset using MLE, and the mean ratio of estimated abundance to true abundance calculated. The three resulting estimates of bias were used to parameterize a normal distribution for estimation bias due to sex heterogeneity, with the mean equal to the bias estimated at the 50th percentile of the estimated distribution for sex heterogeneity, and

standard error calculated as the mean absolute difference between estimated bias at the 50th percentile and those at the 16th and 84th percentiles.

The biases considered here are approximately additive (Barlow et al., 2011). A distribution of correction factors was parameterized as

$$c \sim \frac{100\%}{100\% - \left(N(b_{open+calf}, \sigma_{b,open+calf}) + N(b_h, \sigma_{b,h}) \right)},$$

where $b_{open+calf}$ and $\sigma_{b,open+calf}$ (mean and standard error of combined bias due to births and deaths and excluding calves) were drawn from Barlow et al. (2011); and b_h and $\sigma_{b,h}$ (mean and standard error of estimated bias due to sex heterogeneity in capture probability) were estimated through simulation as described above. To obtain bias-corrected estimates, random samples were drawn from this distribution for multiplication by the values in the posterior distributions for abundance from the SCR and Mt models.

Data manipulation and filtering, visualization, simulations, and post-processing were conducted in R 3.6.2 (R Core Team, 2019), using the DBI, tidyverse, lubridate, coda, and mcmcplots packages (Plummer et al., 2006; Golemund and Wickham, 2011; Curtis, 2015; R Special Interest Group on Databases (R-SIG-DB) et al., 2019; Wickham et al. 2019). Model fitting was implemented with NIMBLE for Bayesian estimation, using the nimble and nimbleEcology packages (de Valpine et al., 2017; Goldstein et al., 2021; de Valpine et al., 2021), and with Rcapture for MLE (Rivest and Baillargeon, 2019).

Results and Discussion

A total of 430 distinct individuals were included in the 2019 to 2021 dataset off Central America and southern Mexico, of which 38 were captured during two distinct occasions and one in three occasions, and 59 were captured in more than one trap (i.e., spatial recaptures), in some cases within an occasion.

Two of the key abundance estimation biases anticipated in this analysis, along with associated uncertainty, were quantified by Barlow et al. (2011), with means of -10.5% for excluding animals in their first year and +5.2% for violating the population closure assumption. Barlow et al. (2011) based their simulations on sampling occasions separated by a maximum of 1.5 years, whereas the duration of our dataset is two years, which would lead to a greater positive bias due to births and deaths. If current vital rates in this DIP are markedly different from those simulated by Barlow et al. (2011) (0.96 annual adult survival rate, 11% birth rate, and 0.85 semi-annual survival rate for new calves), that could also affect the resulting bias, with lower adult survival increasing it and lower birth rates and calf survivals decreasing it. Similarly, the -10.5% estimated mean bias from excluding first-year animals would be an overprediction of the negative bias if the current population had lower birth rates than simulated in Barlow et al. (2011). Calambokidis and Barlow (2020) found a population growth rate for U.S. West Coast

humpbacks that is commensurate with that of the SPLASH years (Calambokidis et al., 2009), but we see evidence of lower growth rates for the CentAm/SMex-CA/OR/WA DIP (see below). Since the negative bias due to exclusion of first years is greater in magnitude than the opposing positive bias due to births and deaths, it would change faster with a change in birth rate. If the current population is growing more slowly due to a decrease in birth rate, the two combined biases would likely be less than the mean of -5.3% estimated in Barlow et al. (2011), making the current bias-corrected estimate a slight overestimate. Our simulations of unaccounted-for sex heterogeneity in capture probability, within the parameter space estimated from SCR and Mt models (1,100 total individuals; annual capture probabilities of 0.06, 0.06, and 0.23 in 2019, 2020, and 2021, respectively), resulted in a mean estimation bias of -19.0% (CV=0.521). Summing these biases and their associated uncertainties, and inverting the difference from unity, produces a mean overall correction factor of 1.35 (CV=0.143).

The base SCR model (northern model domain boundary at 19.2°N) resulted in a mean, bias-corrected, realized abundance estimate of 1,496 (CV=0.171) for the CentAm/SMex-CA/OR/WA DIP, with a 20th percentile of 1,284 (Fig. 5; Table 3). This estimate is somewhat lower than – though still above the 18th percentile of – that from the non-spatial Mt model, which resulted in a mean bias-corrected estimate of expected abundance of 1,804 (CV=0.191) (Table 3). Expected rather than realized abundance was calculated from the Mt model to take advantage of MCMC efficiencies in NIMBLE, but the two quantities were practically equivalent. The SCR estimate is only moderately sensitive to the location of the northern model domain boundary. To understand alternate model results, it is important to bear in mind that the dataset is the same throughout, limited to collections from Guerrero southward. Shifting the northern model domain boundary, and thus the limit of the potential activity center locations of animals in the dataset, southward to 18.6°N, at northern border of Michoacán state, increased the bias-corrected mean estimate slightly to 1,601 (CV=0.166), because it forced the spatial activity centers of observed animals to be concentrated in a smaller area, driving down the estimates of capture probability for the same number of captures. Shifting the northern model domain boundary northward to 20.4°N, at the transition to Bahía de Banderas, correspondingly drove up estimates of capture probability and lowered the bias-corrected mean abundance estimate to 1,313 (CV=0.167) (Table 3).

The mean estimate of space use σ in the SCR model is 3.0 degrees latitude (CV=0.131; Fig. 5). This estimate is similar to the mean individual difference in capture latitude among occasions found in the diagnostic randomization test using all years of data (Fig. 3). Note that this estimate cannot be translated to a distance in terms of kilometers, because it does not account for the corresponding change in longitude. Based on this estimate, traps are spaced at sufficient resolution (i.e., $\leq \sigma$; Fig. 2) to expose all individuals in the population to sampling and minimize bias and imprecision due to trap spacing alone (Sollman et al., 2012; Royle et al., 2013).

The posterior predictive checks for deviation of expected from observed number of captures per individual did not detect unexplained individual heterogeneity in either the Mt or SCR model (0.45 and 0.77 probabilities, respectively, of deviation from expected being less in the observed data than in the posterior predicted datasets). However, goodness-of-fit tests have extremely limited power to detect unexplained individual heterogeneity in relatively sparse captures histories (e.g., White and Cooch, 2017). Of the remaining posterior predictive checks – for deviation from a uniform distribution of spatial activity centers, deviation of expected from observed number of captures per trap, and deviation of expected from observed number of captures per individual per trap – none of the resulting statistics suggested a lack of fit (probabilities of deviation from expected being less in the observed data than in the posterior predicted datasets of 0.43, 0.38, and 0.84).

Given the goodness-of-fit results, the support for the importance of space in explaining capture probabilities of humpback whales in the CentAm/SMex-CA/OR/WA DIP in its wintering area, and the lower abundance estimate resulting from the SCR model, the estimate from the SCR model is the conservative choice. Sampling the full extent of the wintering area over multiple years, better characterizing the northern population limit for the wintering area, and obtaining effort data to be included as a covariate are crucial to obtaining a more robust abundance estimate for the CentAm/SMex-CA/OR/WA DIP.

Limited evidence is available to assess whether the CentAm/SMex-CA/OR/WA DIP, which corresponds to the endangered Central America DPS, has been growing at the same rate as observed for total humpback whales off the U.S. West Coast, estimated by Calambokidis and Barlow (2020) at 8.2% per year. The abundance estimate provided here for the CentAm/SMex-CA/OR/WA DIP is not comparable to a previous estimate for the Central American DPS for the years 2004-2006 (755, CV= 0.242) (Wade, 2021), because the DIP now includes animals off southern Mexico. The current abundance estimate is nearly double that obtained 15 years earlier, and would correspond to an increase of 4.8% per year (analytical SD = 2.0%) if it were purely due to population growth. Given the inclusion of southern Mexico whales in the current estimate, the true growth rate of this population is likely lower, and substantially lower than that observed for total humpback whales off the U.S. West Coast over the same time period. Alternatively, a population growth rate can be derived by resummarizing the results of the SCR model to include only whales with activity centers south of 14.5°N in the abundance estimate. Comparing this resummarized and bias-corrected result to the Wade estimate, for which the main sources of bias may nearly cancel each other out (Wade 2021), gives an increase of 1.6% per year (SD = 2.0%).

If the population growth rate for the CentAm/SMex-CA/OR/WA DIP is lower than that for U.S. West Coast humpbacks as a whole, we would expect a decrease over time in the proportion of Central America whales relative to Mexico whales in this feeding area. The current estimate for the CentAm/SMex-CA/OR/WA DIP (1,496, CV=0.171) represents 30% of the most recent mark-recapture-based estimate for the U.S. West Coast (4,973,

CV=0.048), which was based on data from 2015-2018 (Calambokidis and Barlow, 2020; 86 FR 58887, October 25, 2021). Previous estimates of this proportion, based on data from 2004-6, are indeed somewhat higher, at 42% in Wade (2021) and 55% (95% CI: 27-67%) in Lizewski et al. (2021), though still within the confidence interval of the latter. Although Martien et al. (2020) found no evidence of a change in genetic composition for the U.S. West Coast between 1988-89, 2004, and 2018, their statistical power was extremely low for detecting a change in proportions of this magnitude (K. Martien, pers. comm., February 2, 2022).

An estimate for the abundance of whales from the MMex-CA/OR/WA DIP in U.S. waters can be deduced from the remaining proportion of the U.S. West Coast estimate: $4,973 - 1,496 = 3,477$, with an analytical CV of 0.101. The 20th percentile of the corresponding lognormal distribution equals 3,179. The U.S. West Coast estimate was based on data from California and Oregon, but likely includes Washington animals due to interchange with that area (Calambokidis and Barlow, 2020).

The West Coast estimate is not bias-corrected for births and deaths or exclusion of calves from the dataset. The magnitude of the bias due to exclusion of calves would be expected to be similar to that of -10.5% estimated by Barlow et al (2011). But since the three-year (four annual occasions) timespan of the West Coast estimate is double that considered in Barlow et al. (2011), the positive bias due to births and deaths would be substantially greater than +5.2%, largely cancelling out the former.

Conclusions

We present the first population estimate for the CentAm/SMex-CA/OR/WA DIP, with a best estimate of 1,496 (CV=0.171) and a 20th percentile of 1,284. We also found evidence that the growth rate for this population is considerably lower than that found for total humpback whales along the U.S. West Coast. Our analysis provides strong support for spatial fidelity of the CentAm/SMex-CA/OR/WA DIP in their wintering area, with important implications for sampling needs to obtain accurate population estimates. We also provide an estimate of the magnitude of sex heterogeneity in capture probabilities in the wintering area for this population, with annual male capture probability a factor of 3.4 (CV=0.46) times higher than that for females. Finally, a recent estimate for humpback whales off the U.S. West Coast allows us to subtract the estimate for the CentAm/SMex-CA/OR/WA DIP for a first estimate of the number of humpback whales from the MMex-CA/OR/WA DIP in U.S. waters, at 3,477 (CV=0.101).

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Supplementary Information

Appendix A. R code for permutation test for individual spatial fidelity

Appendix B. NIMBLE and R code for models and posterior predictive checks

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Tables

Table 1. Descriptions of quality levels used by Cascadia Research Collective to categorize photographs for identification of humpback whales.

Quality level	Description
1	Best possible: sharp, well-lit, >75% view of target area, perpendicular to the photographer such that ridging is not distorted by angle or lighting
2	Pretty good: has some flaw, either in exposure/sharpness, slightly off angle, but still usable for identification
3	Fair: has flaws beyond a 2, exposure/sharpness compromised but still has parts in the photo that could be useful for identifying
4	Poor: deemed unsuitable for confident matching due to any combination of limited portion of fluke visible, angle, lack of focus, lighting, or pixelation

Table 2. Summaries by country or state (region) of photo-identifications for 2019 to 2021 winter seasons (November of preceding year through April) in Central America and southern Mexico. Regions do not always correspond to “traps” (Fig. 2). Summaries include date range of sightings and number of unique identifications. Summaries for 2021 also include number of survey days. Unique identifications (IDs) for all locations do not equal the total of the regions since some individuals were seen in multiple regions. Totals after filtering for photo quality are also provided by year and by location. Abbreviations used are: CRC = Cascadia Research Collective, UABCS = Universidad Autónoma de Baja California Sur, locs = locations.

Region	Organization	Principal collaborators	2019		2020		2021		2019-2021 Unique IDs (Filtered)	
			Date range	Unique IDs	Date range	Unique IDs	Date range	No. Survey Days		Unique IDs
Guerrero, Mexico	Whales of Guerrero	K. Audley, A. García Chávez, R. Ramírez Barragán	03-Jan-19 - 09-Mar-19	59	09-Jan-20 - 11-Mar-20	53	22-Dec-20 - 20-Mar-21	46	123	207
Oaxaca, Mexico	UABCS, Univ. del Mar	J. Urbán, P. Martínez, F. Villegas Zurita	03-Feb-19	1	27-Feb-20	1	10-Dec-20 - 26-Mar-21	38	100	83
Guatemala	CRC	E. Quintana-Rizzo	27-Feb-19 - 08-Mar-19	14	— ¹	—	08-Feb-21 - 16-Feb-21	8	17	24
El Salvador	Proy. Megaptera, Murdoch Univ.	N. Ransome ² , M. Castaneda	13-Jan-19 - 23-Feb-19	9	29-Jan-20 - 04-Mar-20	7	01-Feb-21 - 13-Mar-21	41	47	39
Nicaragua	ELI-Scientific	J. de Weerd	02-Mar-19	2	03-Jan-20 - 04-Mar-20	31	03-Jan-21 - 13-Apr-21	46	82	97
N and S Costa Rica	Panacetacea, CRC	F. Garita, J. Calambokidis, J. D. Palacios ³	—	—	16-Nov-19 - 05-Dec-19	2	28-Dec-20 - 28-Mar-21	57	47	41
Panama	Panacetacea	K. Rasmussen, B. Pérez	—	—	—	—	04-Feb-21 - 19-Feb-21	9	5	5
All Locs				84		90		244	361	
Filtered, All Locs				75		84			311	430

¹ Scheduled field work was cancelled due to the start of the global pandemic.

² 2019 IDS came from David Alfaro at Asociación Océano and Jose Baires.

³ 2020 IDs came from Sierra Goodman.

Table 3. Summaries of bias-corrected abundance estimates for the CentAm/SMex-CA/OR/WA DIP from capture-recapture models. All models are closed-population models fitted to capture histories from 2019-2021 off Central America and Southern Mexico. Base and alternate models are one-dimensional spatial capture-recapture models. Results are also shown for a non-spatial model with time-varying capture probabilities (Mt). Northern model domain boundary (MDB) and population limit(s) (PL) for each model are provided, along with the estimated mean abundance and coefficient of variation (CV). The base model estimate used for further inference is highlighted in bold.

Model	Northern MDB (Latitude)	Northern PL (Latitude)	Mean	CV (%)
Base	19.2	18°N - 19.2°N	1,496	17.1
Northern alternate MDB	20.4	18.6°N	1,313	16.7
Southern alternate MDB	18.6	18.6°N	1,601	16.6
Non-spatial Mt	–	–	1,804	19.1

Figures

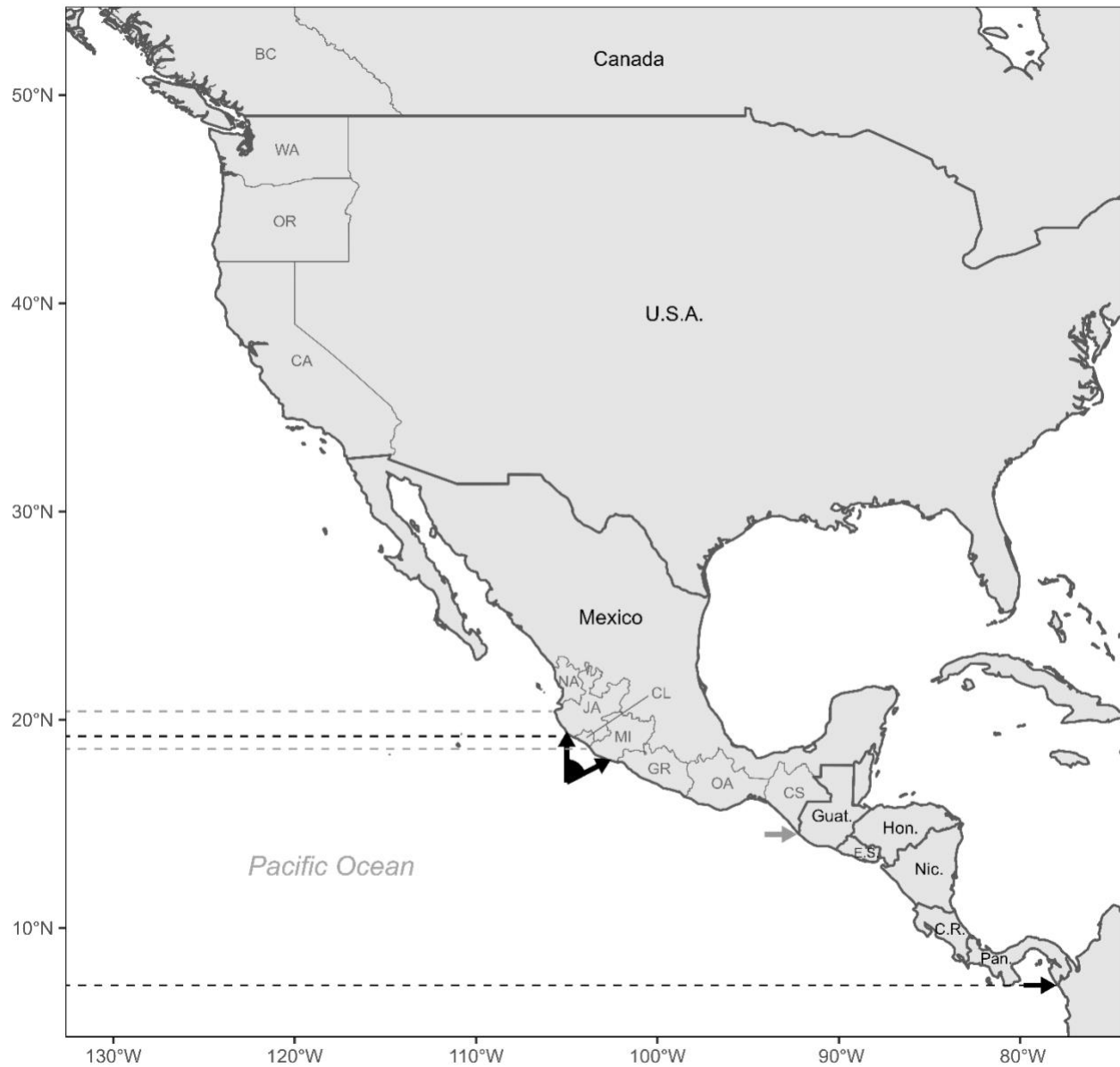


Figure 1. Map of Central American and southern Mexican Pacific coast, along which lies the wintering area of the CentAm/SMex-CA/OR/WA DIP of humpback whales. Black arrows indicate northern and southern population limits used in the base one-dimensional spatial capture-recapture model (i.e., limits for estimated activity centers of animals included in population), with the southern limit at 7.25°N and the range of northern population limits indicated as an arc from 18°N to 19.2°N. Gray arrow indicates originally designated northern limit of Central America DPS at 14.5°N (northern border of Guatemala). Black dashed lines at 7.25°N and 19.2°N indicate model domain boundaries for base model (i.e., within which estimated centers of activity of whales in the dataset, whether included in the population or not, are allowed to occur). Gray dashed lines show alternate northern model domain boundaries, at 18.6°N and 20.4°N. Abbreviations for Central American countries are Guat. = Guatemala, E.S. = El Salvador, Hon. = Honduras, Nic. = Nicaragua, C.R. = Costa Rica, Pan. = Panama. Abbreviations for subnational jurisdictions are BC = British Columbia, WA = Washington, OR = Oregon, CA = California, NA = Nayarit, JA = Jalisco, CL = Colima, MI = Michoacán, GR = Guerrero, OA = Oaxaca, CS = Chiapas.

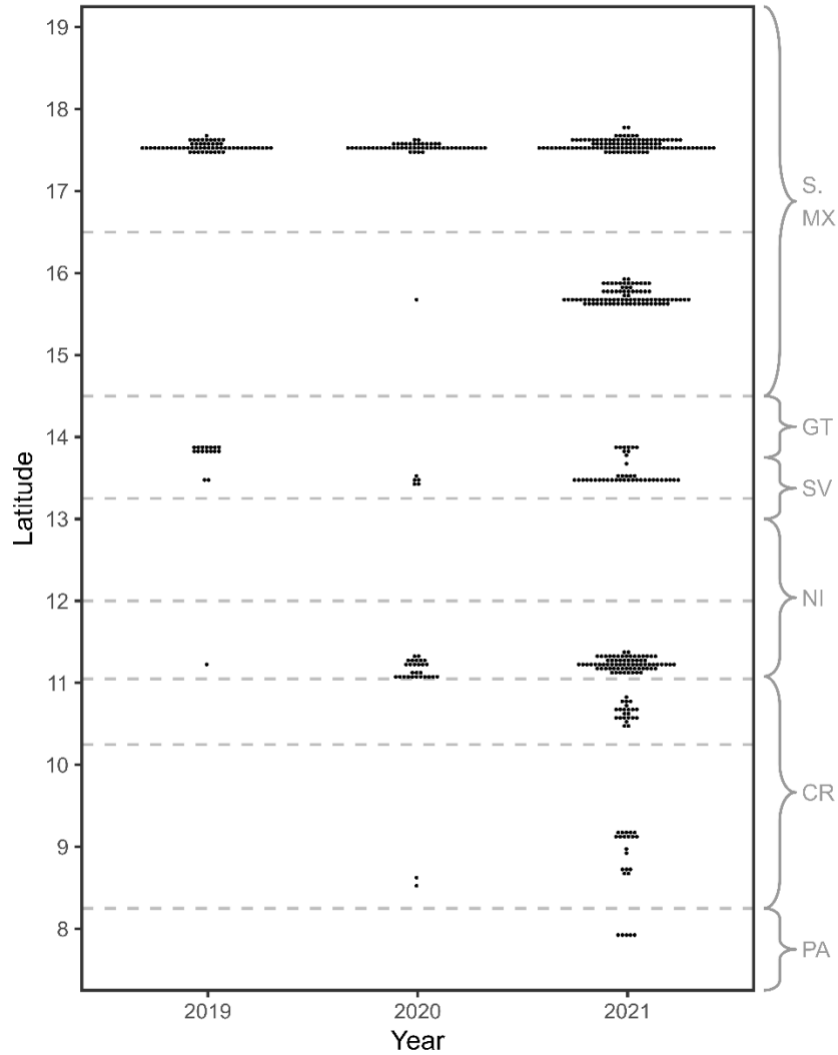


Figure 2. Spatiotemporal distribution of photo-identifications of humpback whales in Central America and southern Mexico from the 2019 to 2021 winter seasons. Each unique individual captured during an annual occasion and in the range of latitudes constituting a “trap” (denoted by dashed gray lines) is represented by a dot at the mean latitude of its captures within that occasion-trap, binned by 0.05 degrees latitude. Space is approximated by latitude, since the Pacific coast of Central America and southern Mexico is relatively straight. For reference, approximate latitudinal spans of Central American countries and southern Mexico are provided on the right-hand side in gray. Abbreviations are PA = Panama, CR = Costa Rica, NI = Nicaragua, SV = El Salvador, GT = Guatemala, S. MX = southern Mexico.

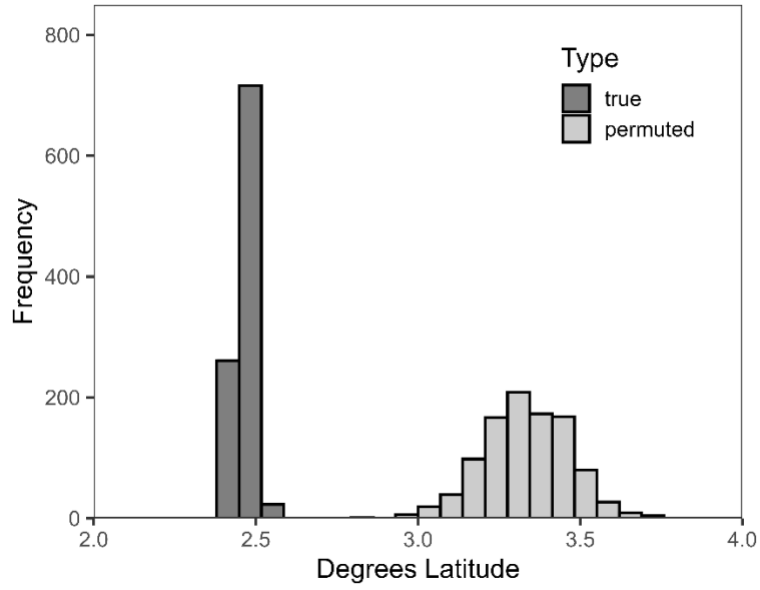


Figure 3. Results of permutation test for interannual spatial association within individuals. Histogram shows overall averaged mean distances per individual between annual capture locations (in degrees latitude), for true data and for permuted data, from 1000 replicates. For each replicate, a “true” dataset was drawn from a complete daily-resolution dataset of individual captures off Central America and southern Mexico from 1998 to 2021 by sampling one daily mean capture location per individual per occasion. A corresponding randomized dataset was created from this true dataset by permuting capture locations within occasion. For each true and each randomized dataset, mean distance among annual capture locations was calculated for each individual that was captured during more than one occasion, then averaged across individuals for the overall mean per dataset.

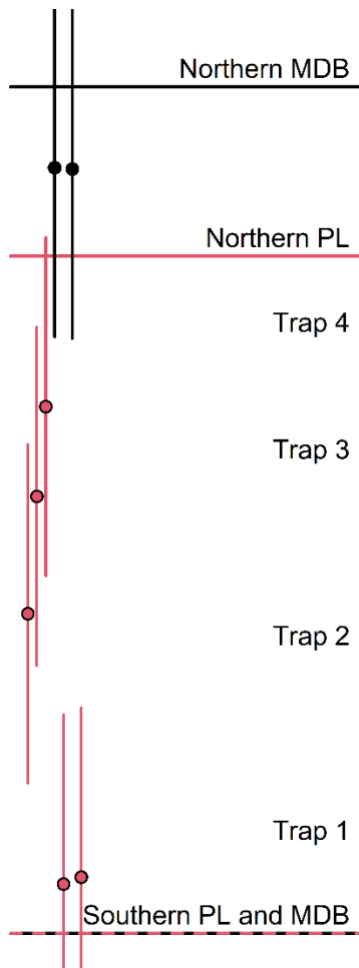


Figure 4. Conceptual diagram of model domain and population limits in one dimension (latitude). Model domain boundaries (MDB; black horizontal lines) must by definition include the population limits (PL; red horizontal lines) within them. For this analysis, the southern model domain boundary and southern population limit are the same, but the northern model domain boundary and northern population limit may differ. All estimated individual whale activity centers (dots) are restricted to the model domain. Those whales with activity centers within the population limits (activity centers indicated by red dots) are included in the abundance estimate. Probability of occurrence of an individual whale – and thus capture, if effort were constant in space – decreases exponentially with north-south distance from the activity center. If vertical whiskers to either side of each activity center represent the 95% probability density interval of occurrence locations for that individual, then that individual is most likely to be captured within “traps” (delineated by gray vertical bars to right) that overlap those whisker(s). Traps are ranges of latitudes within which the locations of all individual captures are approximated as the mean of those captures. An individual may be captured in more than one trap within an occasion.

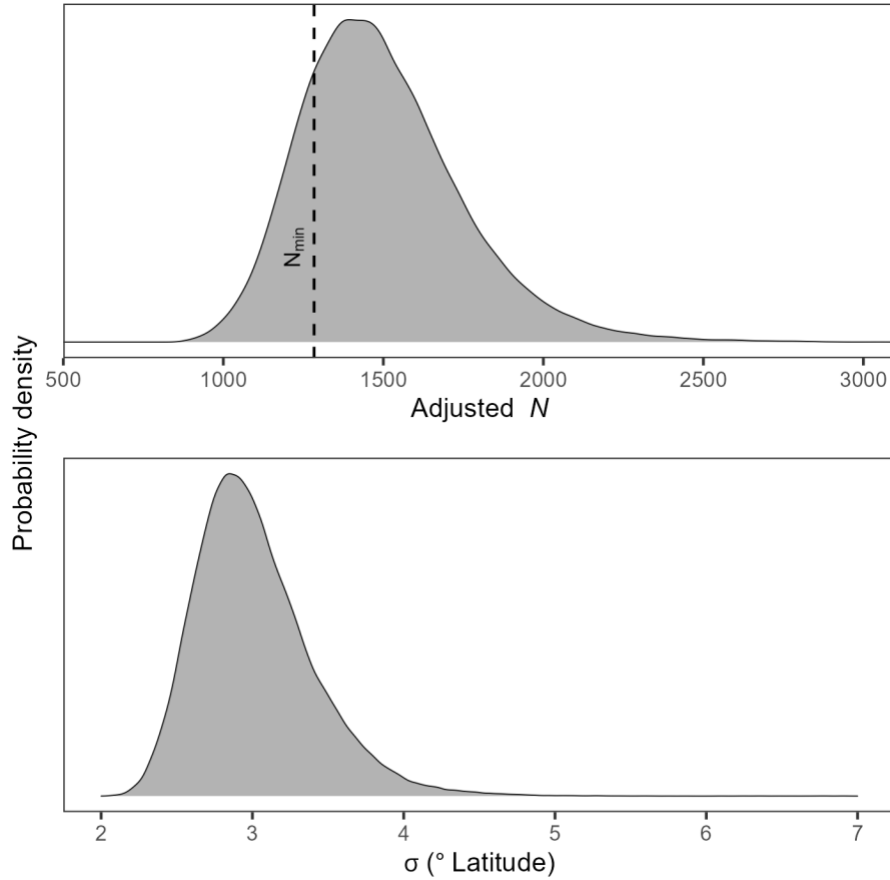


Figure 5. Posterior probability densities for abundance N (upper panel) and space use parameter σ (lower panel) for the CentAm/SMex-CA/OR/WA DIP of humpback whales. Parameters were estimated from a closed, one-dimensional, spatial capture-recapture model fitted to spatially discretized annual capture histories in the wintering area off Central America and southern Mexico from 2019 to 2021. The abundance estimate includes a correction factor to account for the combined bias, estimated through simulation, from birth and deaths, omission of first-years from the data set, and sex heterogeneity in capture probability. A vertical dashed line shows the 20th percentile of the abundance posterior, which would serve as the minimum population estimate N_{\min} for calculation of Potential Biological Removal of this population under the U.S. Marine Mammal Protection Act.

Appendix A. R code for permutation test for individual spatial fidelity

The following code performs a permutation test for non-randomness (fidelity) at the individual level in spatial location of captures among years. Test data sets are created by sampling a single record per individual per year. For each test data set, a permuted version is created by randomizing latitude within year. Then, for both permuted and unpermuted test data sets, the overall mean of mean distance per individual among years is calculated and the resulting distributions from 1000 replicates compared.

```
# function to calculate distance metric
# (overall mean of mean distance per individual among occasions)
dist.ind.mean <- function(x) {
  # filter multi-occasion individuals
  my.iy <- x %>% filter(ny>1)
  # calculate mean distance among within-individual captures
  d.ind <- my.iy %>%
    group_by(id) %>% tidyr::nest() %>%
    # # using kilometers
    # mutate(dists = purrr::map(data,
    #                           ~geosphere::distm(data.frame(lon=.x$lon,
    #                                                         lat=.x$lat))/1000),
    #        xdist = purrr::map(dists, function(x) mean(x[upper.tri(x)])))
    # using degrees latitude
    mutate(xdist = purrr::map(data, ~mean(dist(as.numeric(.x$lat)))))
  return(mean(unlist(d.ind$xdist)))
}

# df.day is a data frame containing capture records at daily resolution, with
# fields for individual identification number (id), annual occasion (occ),
# latitude (lat), longitude (lon), and number of annual occasions in which
# individual was captured (ny).

# permutation test
nrep <- 1000 # number of test replicates
truth <- as.numeric(rep(NA, nrep))
test <- as.numeric(rep(NA, nrep))
for (i in 1:nrep) {
  # create test data set by subsampling df.day
  all.iy <- df.day %>%
    group_by(id, occ) %>% sample_n(1) %>% ungroup() %>% arrange(occ, id)
  truth[i] <- dist.ind.mean(all.iy)
  # create permuted test data set
  perm <- all.iy %>%
    group_by(occ) %>%
    mutate(r=sample(1:n()), id=id[r], ny=ny[r]) %>%
    ungroup() %>% select(-r)
  test[i] <- dist.ind.mean(perm)
}
rm(all.iy, perm, i)

# calculate proportion of true test statistics that are less than their
# paired permuted counterparts
sum(truth<test)/nrep
```


Appendix B. NIMBLE and R code for models and posterior predictive checks

1) One-dimensional SCR model, where y is augmented with all-zero histories

Model code:

```
SCR0pjk.1D.code <- nimbleCode({
  for (j in 1:J) {
    for (k in 1:K) {
      p0[j,k] ~ dunif(0, 1)      # baseline encounter probability
    }
  }
  sigma ~ dunif(0, 50)        # scale parameter of encounter function
  psi ~ dunif(0, 1)          # DA parameter: E(N) = M*psi

  for(i in 1:M) {
    z[i] ~ dbern(psi)         # Is individual real?
    s[i] ~ dunif(ylim[1], ylim[2]) # 1D spatial coordinate

    for(j in 1:J) {
      # occasion-varying capture probabilities for individual i at trap j
      p[i,j,1:K] <- p0[j,1:K] * exp(-(s[i] - X[j])^2/(2*sigma^2))
      y[i,j,1:K] ~ dOcc_v(z[i], p[i,j,1:K], len = K)
    }
  }

  N <- sum(z[1:M])          # realized abundance
  EN <- psi*M
})
```

R code for goodness-of-fit (niter is total number of MCMC samples):

```
constants <- list(X = X, K = K, M = M, J = J, ylim = ylim)
m.scr01d <- nimbleModel(SCR0pjk.1D.code, constants = constants, data = list(y
= y))
cm.scr01d <- compileNimble(m.scr01d)
dataNodes <- m.scr01d$getNodeNames(dataOnly = TRUE, returnScalarComponents =
TRUE)
parentNodes <- m.scr01d$getParents(dataNodes, stochOnly = TRUE)
simNodes <- m.scr01d$getDependencies(parentNodes, self = FALSE)
vars <- parentNodes
nsbin <- 10
binlims <- ylim[1] + (0:nsbin)*((ylim[2]-ylim[1])/nsbin)
ppc <- data.frame(Ts = rep(NA, niter), Tsnew = rep(NA, niter),
                  Tij = rep(NA, niter), Tijnew = rep(NA, niter),
                  Ti = rep(NA, niter), Tinew = rep(NA, niter),
                  Tj = rep(NA, niter), Tjnew = rep(NA, niter))
for(ni in 1:niter) {
  values(cm.scr01d, vars) <- samples[ni, ]
  cm.scr01d$simulate(simNodes, includeData = TRUE)
  ysimflat <- values(cm.scr01d, dataNodes)
  ysim <- aperm(array(ysimflat, dim=c(K, J, M)), perm=c(3,2,1))
  pflat <- values(cm.scr01d, "p")
  p <- array(pflat, dim=c(M,J,K))
}
```

```

s <- values(cm.scrOld, "s")
z <- values(cm.scrOld, "z")
sran <- runif(M, ylim[1], ylim[2])
# Goodness of fit components
## spatial randomness
sbin <- rep(NA, nsbin)
snewbin <- rep(NA, nsbin)
for (b in 1:nsbin) {
  sbin[b] <- sum((z > 0) & (s > binlims[b]) & (s <= binlims[b+1]))
  snewbin[b] <- sum((z > 0) & (sran > binlims[b]) & (sran <= binlims[b+1]))
}
esbin <- sum(z)/nsbin
ppc$Ts[ni] <- sum(pow(sqrt(sbin[1:nsbin]) - sqrt(esbin), 2))
ppc$Tsnew[ni] <- sum(pow(sqrt(snewbin[1:nsbin]) - sqrt(esbin), 2))
## observations
esumyij <- matrix(NA, nrow=M, ncol=J)
sumyij <- matrix(NA, nrow=M, ncol=J)
sumynewij <- matrix(NA, nrow=M, ncol=J)
for (j in 1:J) {
  for (i in 1:M) {
    esumyij[i,j] <- sum(z[i]*p[i,j,])
    sumyij[i,j] <- sum(y[i,j,])
    sumynewij[i,j] <- sum(ysim[i,j,])
  }
}
esumyj <- colSums(esumyij)
sumyj <- colSums(sumyij)
sumynewj <- colSums(sumynewij)
esumyi <- rowSums(esumyij)
sumyi <- rowSums(sumyij)
sumynewi <- rowSums(sumynewij)
ppc$Tij[ni] <- sum(pow(sqrt(sumyij) - sqrt(esumyj), 2))
ppc$Tijnew[ni] <- sum(pow(sqrt(sumynewij) - sqrt(esumyj), 2))
ppc$Tj[ni] <- sum(pow(sqrt(sumyj) - sqrt(esumyj), 2))
ppc$Tjnew[ni] <- sum(pow(sqrt(sumynewj) - sqrt(esumyj), 2))
ppc$Ti[ni] <- sum(pow(sqrt(sumyi) - sqrt(esumyi), 2))
ppc$Tinew[ni] <- sum(pow(sqrt(sumynewi) - sqrt(esumyi), 2))
}
sum(ppc$Ts < ppc$Tsnew)/niter
sum(ppc$Tij < ppc$Tijnew)/niter
sum(ppc$Ti < ppc$Tinew)/niter
sum(ppc$Tj < ppc$Tjnew)/niter

```

2) Mt model (non-spatial closed model with time-varying capture probability), where y is augmented with all-zero histories

Model code:

```

nimbleCode({
  psi ~ dunif(0, 1)      # DA parameter: E(N) = M*psi

  for (k in 1:K) {
    p[k] ~ dunif(0, 1)  # baseline encounter probability
  }
}

```

```

for(i in 1:M) {
  y[i,1:K] ~ dOcc_v(psi, p[1:K], len = K)
  ynew[i,1:K] ~ dOcc_v(psi, p[1:K], len = K) # posterior predictive data
  sumynewi[i] <- sum(ynew[i,1:K])
  # constant sumyi calculated outside MCMC
}

# GOF metrics
## individual heterogeneity
Tinew <- var(sumynewi[1:M])/mean(sumynewi[1:M])
# constant Ti calculated outside mcmc
## temporal heterogeneity
for (k in 1:K) {
  sumyk[k] <- sum(y[1:M,k])
  sumynewk[k] <- sum(ynew[1:M,k])
  esumyk[k] <- p[k] * psi * M
}
Tk <- sum(pow(sqrt(sumyk[1:K]) - sqrt(esumyk[1:K]), 2))
Tknew <- sum(pow(sqrt(sumynewk[1:K]) - sqrt(esumyk[1:K]), 2))

EN <- psi*M
})

```

R code for goodness-of-fit:

```

sumyi <- c(rowSums(y), rep(0, M-dim(y)[1]))
Ti <- var(sumyi)/mean(sumyi)
# excess temporal heterogeneity?
sum(Tk > Tknew)/length(Tk)
# excess individual heterogeneity?
sum(Ti > Tinew)/length(Ti)

```