Population Viability Analysis of Western Gulls on the Farallon Islands in Relation to Potential Mortality Due to House Mouse Eradication

Nadav Nur, Russell W. Bradley, Derek E. Lee,

Peter M. Warzybok, and Jaime Jahncke

PRBO Conservation Science

3820 Cypress Drive, Suite 11

Petaluma, CA 94951

**Summary**

Proposed house mouse eradication efforts on the Farallon National Wildlife Refuge have identified Western Gulls as a species at risk of non target mortality. Analyses of potential population level-impacts to the world’s largest colony of this species are crucial for evaluating the feasibility of this proposed project. Using PRBO’s long term datasets, we conducted a population viability analysis to model future trends for this population, assessing scenarios with and without eradication mortality, under varying environmental conditions. Scenarios were classified as: “optimistic” with high gull productivity; “realistic”, with average productivity; and “pessimistic”, with greater incidence of low productivity.

* Future population trends for Farallon Western Gulls, in the absence of any eradication-related mortality, will depend on likelihood of reoccurrence of years with especially low reproductive success, as was observed from 2009 to 2011, likely driven by environmental conditions.
* Under “optimistic” environmental conditions, the model predicts that this Western Gull population would grow by 10.6% after 20 years (median result; quartile range +41% to -14%).
* Under “realistic” environmental conditions, the model predicts that the population would decline by 8.7% after 20 years (median result; quartile range +18% to -29%).
* Alternatively, under “pessimistic” conditions, the model predicts that the population would decline by 27% after 20 years (median result; quartile range -4% to – 45%).
* We identify a mortality of 1700 gulls (of all age classes), as one that represents 95% overlap in expected outcomes under “realistic” scenarios.
* Under “realistic” conditions, mortality of 1700 gulls would cause the population to decline by an expected 12.7% after 20 years relative to initial conditions (median result, quartile range +4% to -47%).
* Given assumptions of the model and the high variability of parameters, mortality below that magnitude would not result in modeled outcomes that, after 20 years, are effectively distinguishable when comparing mortality and no-mortality scenarios.
* We conclude that a mortality of 1700 Western Gulls, given an overall population of 32,200 birds, represents an upper maximum of non target mortality to minimize the chances of pervasive long term population impacts.

**Introduction**

The Farallon Islands hold the world’s largest colony of Western Gulls. Proposed house mouse eradication efforts on the Farallon National Wildlife Refuge have identified Western Gulls as a species at risk of non target mortality, due to consumption of toxic rodenticide. While several mitigation measures are planned to minimize any mortality, analyses of potential population level impacts on Farallon Western Gulls is crucial for evaluating the feasibility of this proposed project. Our goals were to assess the future trajectory of this population, under varying environmental conditions, and to evaluate the impacts of any potential increased mortality on a twenty-year time scale.

**Scope of Study**

To meet our goals, we conducted a population viability analysis (PVA) of the Western Gull population on the Farallon Islands to contrast scenarios with additional mortality and scenarios without additional mortality. This study builds on data collection, compilation, previous demographic modeling, and analysis of demographic parameters of recent data for Farallon Western Gulls presented by Spear & Nur (1994), Nur et al. (1994), Pyle et al. (1997), and Lee (2011). The demographic modeling presented here relies on detailed observations and statistical analysis of the Farallon breeding population, covering the period 1986 to 2011, though the set of parameter values used focused on the latter half of the time series.

An important strength of Population Viability Analysis is that it incorporates stochasticity, the unpredictable variation in demographic parameters that reflects underlying environmental variability (Burgman et al. 1993, Beissinger 2002). This allows for a probabilistic assessment of future populations and evaluation of actions that may reduce or increase risk. (Nur & Sydeman 1999 and Akçakaya et al. 2004.)

Using information on the Western Gull population and how it may be impacted by additional mortality resulting from eradication efforts, we develop projections for the future using a time-frame of 20 years. We evaluate three scenarios that make different assumptions about future Western Gull productivity, likely a proxy for environmental conditions, and their impacts on the population dynamics of the Farallon population. For each scenario we contrast the “no additional mortality” scenario with a scenario of a specified level of mortality (which we call *C*). One goal of the study is to determine the value of *C* such that mortality below this level cannot effectively be distinguished from no mortality 20 years into the future, given assumptions regarding unpredictable variability in environmental and demographic parameters.

The three different scenarios all incorporate information on variation in demographic parameters observed during the recent time period (from 1999 to 2008 or 2009, depending on parameter), and vary in levels of reproductive success. Reproductive success in 2009, 2010, and 2011 was extremely low, less than 0.15 chicks fledged per pair in each of the three years. In the 23 years preceding, reproductive success had never been less than 0.30 chicks fledged per pair and was usually much higher. The cause of this near-failure in 2009-2011 has not been identified, and is likely linked to reduced prey availability for this species during the breeding season, as well as increased intra-specific predation on chicks. Thus, the three scenarios evaluated are:

1. Optimistic -“Near-failure” does not reoccur in the future. Reproductive success is variable but reflects observations made prior to 2009.
2. Realistic - “Near-failure” occurs at the historic frequency of 3 times per 26 years in the period analyzed 1986-2011.
3. Pessimistic - “Near-failure” occurs at the “recently observed frequency” of 3 times per 12 years.

For this exercise, we focus on modeling the Farallon population as observed during the recent time period, 1999 to 2011. We use population trend data for this period to derive a Leslie matrix population dynamic model that incorporates stochasticity (Nur & Sydeman 1999). We consider the recent time period to be most relevant for this exercise, as demographic data from the 1980’s and early 1990’s reflects a different population than exists at present – with the earlier part of the time series showing higher population numbers, lower recapture probability and survival, and higher reproductive success (Figures 1, 2, and 3) Therefore, we feel the more recent parameter data is more appropriate as a baseline for predicting future change.

Specific objectives addressed by this study are to:

1. Evaluate future population dynamics based on demographic parameter values and observed population trend, assuming no additional mortality, but considering different scenarios for future environmental conditions. This component of the study quantified the median (expected) behavior of the population as well as the risk of more extreme results (upper quartile and lower quartile of population results) under three different productivity scenarios.
2. Evaluate future population dynamics as in (1) but include impact of mortality of *C* gulls at the outset of the simulation. Part of this objective entailed determining the level of mortality (*C*) such that any mortality below this level, given the variability in parameters, cannot be effectively distinguished from the “no additional mortality” scenarios in this modeling exercise. For the purpose of this exercise, we considered the mortality scenarios to be effectively indistinguishable from each other if there was an overlap in the probability distribution for expected outcomes of at least 95%.

**Methods**

**Rationale of Our Approach**

The basis of the PVA is a Leslie matrix whose values (i.e., elements) are allowed to fluctuate in relation to variation in the future environment (Nur & Sydeman 1999a, Caswell 2001). Here we first briefly describe the demographic parameters being modeled: survival, reproductive success, and probability of breeding. Variation in demographic parameters with respect to age and environmental variability were simultaneously estimated.

1. **Survival** **of adults.** Annual survival was determined through capture/recapture analysis of banded gulls from 1986-2011, with respect to age and year-specific variation.
2. **Survival of juveniles and subadults.** This refers to annual survival of first-year, second-year, and third-year individuals. By the fourth year of life, evidence indicates that Western Gulls have reached adult levels of survival (Spear & Nur 1994, Pyle et al. 1997). Farallon Western Gulls generally disperse widely during the first one to three years of life (Spear & Nur 1994). Therefore it was not possible to derive accurate estimates of survival from capture/recapture using island-based observational data. Instead, we relied on empirical and statistical studies of age-specific survival of this population (Spear & Nur 1994, Pyle et al. 1997).
3. **Reproductive Success** is the number of young reared to fledging per breeding pair per breeding season. We used data from C,H, and K plots from 1986 to 2011. This estimate is conditional on an individual attempting to breed.
4. **Probability of Breeding** is a demographic component that reflects the likelihood that an individual that is resighted on the colony attempts to breed in that season. This parameter potentially varies with the age of the individual. Almost all adults were resighted only when attempting to breed; for that reason, recapture probability is used as an estimate of breeding probability. Note that, in terms of the demographic model, we partitioned probability of breeding into two components: the probability an individual is breeding for the first time and probability that an individual that has previously bred, is currently attempting to breed (see Nur & Sydeman 1999). We used demographic parameter estimates for both probabilities based on the capture/recapture analyses of individuals previously banded as well as observations of age of first-time breeders (see also Pyle et al. 1997).

We incorporate information on annual variation in these four demographic parameters based on observations made during the period 1986 to 2011, as described below, focusing on the most recent period, 1999 to 2011. An important feature of our study is that we calibrated the demographic parameter values used so that the model reproduced the observed population trend data during the recent time period, 1999 to 2011.

**Population Trend Data**

We used whole colony counts at peak incubation for the Farallon Islands for the period 1999 to 2011 and estimated the annual constant rate of change by conducting linear regression on ln-transformed counts (Nur et al. 1999). Results were very similar whether we considered the periods 1999 to 2011, 2000 to 2011, or 2001 to 2011. The observed trend over 1999 to 2011 was a modest growth of 0.74% per year (Figure 1). Therefore, our population model was calibrated to reproduce this growth rate.

**Estimation of Demographic Parameters in Relation to Annual Variation**

**Survival, Recapture Probability, and Reproductive Success:**

Annual survival (symbolized phi) and recapture probability (symbolized p) were estimated over the period 1986 to 2009, for both males and females (Figure 2). It was not possible to estimate year-specific survival beyond 2009 while simultaneously estimating year-specific recapture probability due to limitations of capture-recapture analysis (Cooch et al. 1996). For the initial parameter values in the population model we used mean survival estimates, averaged across the two sexes, based on the most recent 10 years, 1998/1999 to 2008/2009. We also assessed variation in survival and reproductive success across the entire time series (1986 to 2009, but found that the magnitude of variation differed between the two time series. The between year standard deviation (SD), was much greater for the longer time series, 15% greater for survival, 31 % greater for reproductive success. The between-year SD includes not only variation in underlying demographic parameters among years, but also variation due to sampling error (Gould & Nichols 1998). Recognizing that, we chose to use the smaller of the two between-year estimates of variance for modeling survival and reproductive success, that provided by the most recent time period. By using the smaller estimate, from the recent 10-year period rather than the 24-year period, we were reducing the effect of over-estimation due to inclusion of sampling error.

The between year SD in adult survival was determined from the year-specific analyses (above). For juvenile and subadult s survival, we scaled the between year SD relative to that of adults, given that survival is a binomially distributed random variable and its variance = phi (1-phi) (Mood et al. 1974). That is, the closer survival is to 0.50, the greater is its variance. See Table 1 for SD values used.

Reproductive success (the number of fledged young per breeding pair) was determined each year for our 3 study plots and then averaged across plots and years to determine a mean RS for the period from 1999 to 2008 (Figure 3). The poor reproductive success observed in 2009 to 2011 was modeled separately (see below).

We also quantified the mean annual capture probability (p), a measure of breeding probability, and the between year variation observed for this parameter. Capture probability, in this case, refers to the probability that an individual that has bred before breeds in a given year. However, we must also consider the probability that an individual that has never bred before, breeds in a given year (Nur & Sydeman 1999). While we were not able to explicitly estimate this latter parameter on a year by year basis over the 24 year time series, we were able to estimate how this probability varies with age, and used that in the modeling.

The demographic model also required estimation of variance in “net fecundity” where net fecundity is defined as the product of RS \* p \*0.5. We calculated variance in net fecundity based on the product of these individual parameters (Mood et al. 1974), assuming no covariance between RS and p. Thus, our estimate of variance in net fecundity is conservative because inclusion of positive covariance (likely the case: in “good” years both RS and p tend to be high and in “bad” years both tend to be low) would have increased the variance of net fecundity beyond what we were able to calculate.

**Poor Reproductive Success in Recent Years**

An important feature of the Farallon Western Gull population for the purposes of this modeling is that there has been unusually low reproductive success observed over the past three years (2009 to 2011). From 1986 to 2008, annual reproductive success ranged from 0.30 to 1.55 fledged young per pair (Figure 3). However in the most recent three years, an average of only 0.06 to 0.13 fledged young were produced per pair. Comparing the most recent 3 years to the 10 years previous to that (1999 to 2008), indicated a reduction of 86.2% in mean reproductive success (Figure 3). We believe that this recent “near-failure” could significantly impact population modeling if it were to continue over the coming years or repeat at some time in the future. Therefore, to model reproductive success we used the mean value over the recent period (1999 to 2008), but with between-year variability as described above (i.e., excluding 2009-2011) and added the probability of near-failure in reproduction occurring at three different frequencies.

**Age-specific Estimation of Parameters for the Population Matrix**

**Survival and Fecundity**

Survival by age was estimated using the program MARK (Cooch and White 2012), for individuals banded as chicks and subsequently captured or identified at the breeding colony. Age-specific estimates were then incorporated into the model as appropriate. For adults, age 4 and older, annual survival showed no clear pattern with respect to age, for either males or females (Lee 2011). Therefore the model assumed that all adults had the same survival value (see Table 1). Survival prior to age 4 could not be estimated from these capture-recapture analyses since a very small number of juvenile gulls return to the colony before breeding. Therefore, to estimate juvenile survival, we relied on prior analyses based on intensive field observations and statistical analysis by Spear & Nur (1994) and Pyle et al. (1997). We used mean values for males and females, for all ages, prior to calibration for the initial survival values in the model (Table 1).

The first component of fecundity, age-by-age reproductive success (RS) was directly estimated from females of known-age (Lee 2011). We assumed that patterns for males were similar to that of females (Pyle et al. 1997). RS appeared to differ with respect to age. RS increases with age up to age 7, then is fairly level through age 16, and then declines subsequently. On the basis of age by age estimates, we developed a simplified table, categorizing adults into four groups: Young adults (ages 4-5 yrs), transitional adults (age 6), prime-age adults (ages 7 to 16 yrs), and old adults (ages 17 and older) (Table 1).

Capture or resighting probability (p) was used to estimate breeding probability. Age-specific estimates were obtained as part of the survival modeling described above (see Lee 2011). Results indicated that p differed little with age for either sex and remained high throughout life (mean = 0.953 averaging across the two sexes; Lee 2011). Therefore we assumed that once an individual bred it did so with probability of 0.953 (see Table 1).

Age-specific breeding probability includes a second component, the probability an individual breeds for the first time. Capture-recapture analyses provided estimates of the transition from pre-breeder, never having bred before, to breeder (Lee 2011). The model assumed the earliest age of breeding is 4 years, with probability of breeding at age 4 being 19% (mean value for males and females). For 5 year olds, 52% attempt to breed, composed of individuals that bred the year before (as 4 year olds; 19%, see above) and an additional 33% that are breeding for the first time as 5-year olds. Similar calculations apply to age 6, at which age 81% are attempting to breed. By age 7, we assume that individuals reach the full-adult value of 95.3% breeding probability. Age-specific breeding probability is summarized in Table 1.

**Post breeding Census and Density Dependence**

The Leslie matrix population model can be implemented with respect to either a pre-breeding or post-breeding census (Caswell 2001, Akçakaya 2005). We chose the latter, primarily because it splits first-year survival into its own row, which can easily be manipulated. As a result, the youngest age class in the simulations refers to individuals who have just fledged. There is no evidence of density dependence for this population. Therefore we assumed population parameters to be independent of density.

**Calibration**

Estimates of survival, whether sub-adults or adults, will underestimate true survival due to permanent emigration of individuals from the study area (Clobert and Lebreton 1991). Such emigration could be from one part of the island to another, or off of the island altogether. The dispersal can be of pre-breeders or of individuals that have already bred.

To allow us to correct for this under-estimation, we calibrated the performance of the population model such that the set of demographic parameter values used produced a population whose median trajectory corresponded to the observed population behavior. From 1999 to 2011, the breeding population demonstrated an average (time-constant) increase of 0.74% per year (Figure 1). To replicate these conditions, we were required to increase survival by a small amount. For first-year survival, we increased the value from 0.582 to 0.610, but note that female survival was estimated by Spear & Nur (1994) at 0.61, so this simply means using the higher of the two sex-specific values, to allow for some emigration. For second-year survival , we increased the value from 0.794 to 0.810, but note that female survival was estimated by Spear & Nur (1994) at 0.81, so this, too, means simply using the higher of the two sex-specific values, to allow for some emigration. For third-year survival, we increased the value from 0.854 to 0.875, but note that female survival was estimated by Spear & Nur (1994) at 0.89, so this reflects a value that is in between the male and female estimates but shaded towards the female value. For survival in the fourth-year of life, we assumed the same value as adults (Pyle et al. 1997). For all individuals four years and older we adjusted survival from 0.885, this being the mean value for males and females, to 0.890, a very slight adjustment to allow for some emigration. Note that adult emigration is presumed less than that of juveniles and subadults, thus implying a smaller adjustment. We did not adjust fecundity values. All the simulations used the survival values adjusted through this calibration process. Survival and fecundity values used in the simulations, once the model was calibrated, are listed in Table 1.

**Details of the Stochastic Modeling**

The stochastic population modeling was carried out with RAMAS GIS version 5 (Akçakaya 2005). The primary outcome variable of the modeling was the number of individuals in each age class of the population in each year of the simulation, as a function of environmental variability and starting population size. The simulations depict results in which the demographic parameter values for survival and fecundity in a given year in a given simulation are randomly chosen from a distribution whose mean and variance were determined as described above.

In these analyses, we present results for a hypothetical 20-year simulation using the best data relative to the present state of the Farallon Western Gull population. Projections beyond 20 years would be excessively uncertain. In the output, years since the beginning of the simulation are shown as year 0, 1, 2, … 20.

**Starting Population Size, Mortality Scenarios, and Simulations**

The starting total population size for the simulations is 32,200 individuals of all age classes, in the absence of any additional mortality. This corresponds to a breeding population size of 17,400 individuals, the best recent estimate, from 2011 (Warzybok and Bradley 2011), assuming a stable age structure as determined by the Leslie matrix (Caswell 2001), and assuming average breeding propensity. In other words, our results indicate that given the calibrated demographic parameter values used and a breeding population size of 17,400 individuals, there are on average an additional 14,800 sub-adults and non-breeding adults. Note that the 3-year average for 2009-2011 is 17,100 breeding individuals, within 1.6% of the 2011-only value. Therefore, our results are robust to whether we use the most recent year or the 3-year average.

In scenarios with mortality, the starting population size was 32,200 – *C* gulls, where *C* was determined to be 1700 gulls. For these scenarios, we assumed that *C* gulls were removed in proportion to the age distribution of the total population. In other words, 5.3% (=1700/32200) of all age classes were removed at the start of the simulation.

This value of *C* was determined from an assessment of whether the set of outcomes under a “no-mortality” scenario is different from the set of outcomes under an “additional mortality” scenario – under “realistic” productivity values, as described above. We did this by assessing overlap of the modeled distributions for 20 years in the future. We defined two probability distributions to be different if their overlap was less than 95%. In other words, if the overlap was 95% or more between the “no-mortality” scenario and its companion “mortality” scenario, we considered the two distributions to be effectively indistinguishable even though statistically they may distinguishable (e.g., their medians may be statistically different).

To operationalize this definition we first identified the median of the no mortality distribution, call this mno. For example, this value might be 29,400. By definition, 50% of all outcomes were below this value, mno = 29,400. We then analyzed the distribution of outcomes under the same conditions except that *C* gulls were removed at the outset. We then identified the value of *C* such that with *C* gulls removed the distribution of outcomes had been shifted by 5%, i.e., 55% of outcomes were now below the original median. A displacement in the distribution of 5%, from 50% below mno to 55% of outcomes below mno, is equivalent to an overlap of 95% between two distributions, assuming the two distributions differ only in their location and they have the same shape and spread. Note that a displacement of 0% means an overlap of 100%, whereas a displacement of 50% entails an overlap of 50%. In the latter case, 100% of the new distribution lies below mno which in turn corresponds to the value below which 50% of the original distribution lies, i.e., the overlap is 50%: 50% of the original distribution lies above the maximum value observed for the new distribution.

To be clear, the value of *C* used in these modeling exercises was determined as the maximum level of mortality that produced ecologically indistinguishable differences in scenarios, defined here as 95% overlap, in the probability distributions of Western Gull population size 20 years in the future, for scenarios with and without mortality, under “realistic” productivity conditions, given our estimates of the total Farallon population. This level of mortality is completely independent from any assessment of predicted mortality based on gull attendance during a proposed eradication action, exposure to toxic rodenticide, or toxicity of rodenticide.

All scenarios depict results based on 10,000 simulations, the maximum for the RAMAS program. For the calculations of overlap of distributions we used 30,000 simulations, combining results of three different runs of 10,000 simulations. The simulations consider the 3 scenarios of Western Gull productivity: “Optimistic”, “Realistic”, and “Pessimistic” and 2 levels of mortality (i.e., no mortality or removal of *C* gulls)

**Results**

Results of the population viability analyses are summarized in Figures 4, 5, and 6, corresponding to Scenarios 1, 2, and 3 respectively. For each scenario we depict results with no additional mortality (starting population size is 32,200 individuals) and with removal of *C* gulls at the outset. By simulating results with different mortality levels, we determined that removal of 1700 gulls results in a shifting of the distributions by 5% and thus represents 95% overlap on a 20 year time horizon. This is the case assuming Scenario 2: “Realistic” environmental conditions where “near-failure” occurs at historic frequency (p = 0.1153 per year). The overlap in the two distributions under Scenario 2, with and without additional mortality is depicted graphically in Figure 7.

Figure 4 depicts results under the “Optimistic” no near-failure scenario. In the absence of additional mortality, the population is expected to grow by 10.6% after 20 years, to 35,600 individuals, using the median result of the modeling. However, there is a 25% probability of a decline of 14% or more after 20 years, and a 25% probability that the total increase will be 40% or more after 20 years. If the population incurs mortality in year 0, after 20 years it is expected to be at median value of 33,500, an increase of 4.0% compared to the pre-mortality population size of 32,200. Under the same set of assumptions, there is a 25% probability that there will be 26,100 individuals or fewer, which represents a population decline of 18.9% or greater compared to the pre-mortality population size. Thus, under this scenario, but not the other two, the population will have likely have increases in 20 years, even with additional mortality.. However, as in the other scenarios, there is also a substantial probability that the population will be at lower levels than it was prior to the mortality event in year 0.

Figure 5 depicts results under the scenario under “Realistic” conditions, of near-failure occurring at the historic frequency of 3 times per 26 years. In the absence of additional mortality, the population is expected to decline by 8.7% after 20 years, to a median outcome of 29,400 individuals. However there is a 25% probability of a decline of 29% or more after 20 years, and a 25% probability that the total increase will be 32% or more after 20 years. If the population incurs mortality in year 0, after 20 years it is expected (median value) to be at 28,100, a decline of 12.7% compared to the pre-mortality population size of 32,200. Under the same set of assumptions, there is a 25% probability that there will be 21,500 individuals or fewer, which represents a population decline of 33.2% or greater compared to the pre-mortality population size. That said, there is also a 25% probability that after 20 years, under this scenario, the population will have grown to 36,500 or more individuals, a 13.4% or greater increase compared to the pre-mortality size of 32,200, even though the population sustains a loss of 1700 gulls.

If near-failure occurs at the recent frequency of 3 times per 12 years, under the “Pessimistic” scenario, then we can expect population declines, at least by year 20 (Figure 6). In the absence of additional mortality, the population is expected to decline by 27% after 20 years, to a median outcome of 23,500 individuals. In addition, there is a 25% probability of a decline of 45% or more after 20 years, and a 25% probability that the decrease after 20 years will be 3.7% or less. In fact, under this scenario, and with no mortality, the probability of a net population increase of any magnitude after 20 years is 22%. If the population incurs mortality in year 0, after 20 years it is expected to be at a median value of 22,200, a decline of 31.1% compared to the pre-mortality population size of 32,200. Under the same set of assumptions, there is a 25% probability that there will be 17,900 individuals or fewer, which represents a population decline of 44.4% or greater compared to the pre-mortality population size. That said, there is also a 25% probability that after 20 years, under this scenario, the population will have declined to 29,300 or more individuals, that is the net decrease compared to the pre-mortality size of 32,200 is a decline of only 9.0%. Under this scenario, a loss of 1700 gulls would likely leave the population at a lower level than at the outset, prior to incurring additional mortality, with only the magnitude of the decline to be established.

**CONCLUSIONS**

Under “non mortality” scenarios, this Western Gull population will increase over the next twenty years, with “optimistic” productivity estimates, but decline with “realistic” productivity, and likely decline 3 times faster if incidence of recent near breeding failures were to increase.

In assessing mortality scenarios, we determined as the level of mortality which produced 95% overlap in the probability distributions of Western Gull population size 20 years in the future, for scenarios with and without mortality, under “realistic” productivity conditions, given our estimates of the total Farallon population. This value was 1,700 gulls, assuming a total Farallon population of 32,200 birds. These results are independent of any assessment of actual risk to this Western Gull population from rodenticide exposure. We fully support all efforts to mitigate and minimize any mortality associated with these proposed actions.

If the Western Gull population incurs a one-time loss of 1,700 individuals, this will have a noticeable effect on the population dynamics compared to no such mortality. For example, an expected 8.7% decline after 20 years could become, instead, after the one-time mortality event, a 12.7% decline under the “realistic” productivity scenario (Figure 5). Nevertheless, our results indicate that environmental variability due to “normal” variation in demographic parameters as well as the incidence of “near-failures” of reproductive success will have much greater impact than the effects of a mortality event such as loss of 1,700 gulls. Furthermore, the ability of the population to recover from the loss of 1,700 individuals will very much depend on the incidence of reproductive failures in the future, which is difficult to forecast.

Our analysis to assess the population viability of Farallon Western Gulls has been conducted using the best available demographic data for this species, in the population of interest, accounting for strong stochastic variability in parameters over a multi decadal time scale. This information should be strongly considered in assessments of population level impacts to this species for any future management actions.

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**Table 1.** Summary of compiled demographic parameters for Western Gull in relation to Age. Calibrated Survival and Net Fecundity values (and Standard Deviation) were used in the Population Dynamic Model Matrix. Excluding “Near-Failure” Years of 2009-2011. Data compiled from: Lee (2011), Spear & Nur (1995), Nur et al. (1994) and Pyle et al. (1997)

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| **Age** | | | **Repro**  **Success** | | | | **Breeding Probability** | | **Adult Survival** | **Calibrated**  **Survival** | **SD** | **Net**  **Fecundity** | **SD** |  |  |  |  |  |  |  |  |  | |  |
| **1** | | | 0 | | | | 0 | | 0.582 | 0.610 | 0.060 | 0 | 0 |  |  |  |  |  |  |  |  |  | |  |
| **2** | | | 0 | | | | 0 | | 0.794 | 0.810 | 0.049 | 0 | 0 |  |  |  |  |  |  |  |  |  | |  |
| **3** | | | 0 | | | | 0 | | 0.854 | 0.875 | 0.041 | 0 | 0 |  |  |  |  |  |  |  |  |  | |  |
| **4** | | | 0.436 | | | | 0.191 | | 0.885 | 0.890 | 0.039 | 0.0367 | 0.014 |  |  |  |  |  |  |  |  |  | |  |
| **5** | | | 0.436 | | | | 0.524 | | 0.885 | 0.890 | 0.039 | 0.101 | 0.039 |  |  |  |  |  |  |  |  |  | |  |
| **6** | | | 0.649 | | | | 0.810 | | 0.885 | 0.890 | 0.039 | 0.233 | 0.089 |  |  |  |  |  |  |  |  |  | |  |
| **7** | | | 0.882 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.372 | 0.143 |  |  |  |  |  |  |  |  |  | |  |
| **8** | | | 0.882 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.372 | 0.143 |  |  |  |  |  |  |  |  |  | |  |
| **9** | | | 0.882 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.372 | 0.143 |  |  |  |  |  |  |  |  |  | |  |
| **10** | | | 0.882 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.372 | 0.143 |  |  |  |  |  |  |  |  |  | |  |
| **11** | | | 0.882 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.372 | 0.143 |  |  |  |  |  |  |  |  |  | |  |
| **12** | | | 0.882 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.372 | 0.143 |  |  |  |  |  |  |  |  |  | |  |
| **13** | | | 0.882 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.372 | 0.143 |  |  |  |  |  |  |  |  |  | |  |
| **14** | | | 0.882 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.372 | 0.143 |  |  |  |  |  |  |  |  |  | |  |
| **15** | | | 0.882 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.372 | 0.143 |  |  |  |  |  |  |  |  |  | |  |
| **16** | | | 0.882 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.372 | 0.143 |  |  |  |  |  |  |  |  |  | |  |
| **17** | | | 0.718 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.303 | 0.116 |  |  |  |  |  |  |  |  |  | |  |
| **18** | | | 0.718 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.303 | 0.116 |  |  |  |  |  |  |  |  |  | |  |
| **19** | | | 0.718 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.303 | 0.116 |  |  |  |  |  |  |  |  |  | |  |
| **20** | | | 0.718 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.303 | 0.116 |  |  |  |  |  |  |  |  |  | |  |
| **21** | | | 0.535 | | | | 0.953 | | 0.885 | 0.890 | 0.039 | 0.226 | 0.087 |  |  |  |  |  |  |  |  |  | |  |

**Figures**

**Figure 1**. Western Gull breeding population trends for the Farallon Islands, 1986-2011.



**Figure 2.** Annual variation in recapture probability and survival (± SE) for Farallon Western Gulls from Long term Study Plots, 1986 to 2009 for both females and males. Missing vales for female recapture probability could not be estimated in program Mark.

Figure 2a. Female recapture probability

Figure 2b. Female Survival

Figure 2c. Male recapture probability

Figure 2d. Male survival

**Figure 3.** Annual estimates (± SE) for mean number of chicks fledged per female Western Gull breeding in C, H, or K plots of Southeast Farallon Island, California 1983-2011.

**Figure 4.** Population trajectory assuming “Optimistic” conditions: no re-occurrence of near-failure years, with (red) and without (black) eradication-associated mortality. Shown are the 25th percentile, 50th percentile (solid regression line and circles), and 75th percentile outcomes. Mortality scenario removes 1700 birds in year 0.

**Figure4.tif**

**Figure 5**. Population trajectory assuming “Realistic” conditions: re-occurrence of near-failure years at historic frequency (on average, 3 times per 26 years), with (red) and without eradication-associated mortality (black). Shown are the 25th percentile, 50th percentile (solid regression line and circles), and 75th percentile outcomes. Mortality scenario removes 1700 birds in year 0. Figure5.tif

**Figure 6.** Population trajectory assuming “Pessimistic” conditions: re-occurrence of near-failure years at recent frequency (on average, 3 times per 12 years), with (red) and without (black) eradication-associated mortality. Shown are the 25th percentile, 50th percentile (solid regression line and circles), and 75th percentile outcomes. Mortality scenario removes 1700 birds in year 0.

**Figure6.tif**

**Figure 7.** Probability distribution for “no mortality” scenario and “mortality of 1700” gulls, after 20 years, under “Realistic” Conditions: “historic” frequency of near-failure (results of 10,000 simulations for no mortality and 30,000 simulations for mortality of 1700 gulls). Note initial population size, with no mortality, is 32,200 individuals. Results binned into bins of 2,000 and then a polynomial (fourth-order) smoothing function was applied, except that the extreme tails are actual values. The two probability density functions overlap by approximately 95%.

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