



# **Integrated population model of Antipodean albatross for simulating management scenarios**

Technical report prepared for Department of Conservation – June 2021

**Authors:**  
Yvan Richard



PO Box 27535, Wellington 6141  
New Zealand  
[dragonfly.co.nz](http://dragonfly.co.nz)

### Cover Notes

To be cited as:

Richard, Yvan (2021). Integrated population model of Antipodean albatross for simulating management scenarios, 31 pages. Technical report prepared for Department of Conservation – June 2021.

Cover image:



<https://www.flickr.com/photos/angrysunbird/2549595283>

## EXECUTIVE SUMMARY

Antipodean albatross *Diomedea antipodensis antipodensis* are endemic to New Zealand, with the quasi-totality of the population nesting on Antipodes Island. The species is classified as Nationally Critical due to a potential demographic decline. Threats to the population include incidental mortality in fisheries (in New Zealand and in international waters) and climate change.

The objective of this project was to provide a tool that allows stakeholders to explore the potential impact of threats and the demographic outcomes of management strategies. Using the tool, simulations of the demographic impact of different scenarios may be carried out so that management strategies can be assessed and prioritised.

A small subset of the population of Antipodean albatross has been studied since 1994, and these field data were used to perform the simulations. A Bayesian integrated population model was developed to estimate the main demographic parameters of the population. The model considered detectability of individuals, inter-annual variability, and movements in and out of the study area; it was fitted using the software Stan.

From the model, the annual survival rate for females was estimated to decline from 0.947 (95% c.i.: 0.914 – 0.974) in the period from 1994 to 2004, to 0.882 (95% c.i.: 0.814 – 0.94) after 2005. Estimated survival for males was higher, at 0.946 (95% c.i.: 0.913 – 0.972) and 0.927 (95% c.i.: 0.887 – 0.961) for the two periods. Breeding success also declined between the two periods, from 72.4% (95% c.i.: 65.8% – 78.6%) from 1994 to 2004 to 63.7% (95% c.i.: 53.4% – 73%) subsequently.

Under the current scenario, simulations suggest a significant decline of the population, with an annual growth rate of -4.84% (95% c.i.: -6.07% – -3.65%). Limitations in the data and in the model assumptions may cause the decline to be slightly overestimated; however, the trend remains of concern.

The simulation tool is aimed to assist conservation managers with the prioritisation of management strategies to mitigate threats to the Antipodean albatross population and to guarantee the persistence of this species. The tool can be accessed at <https://dragonfly-science.shinyapps.io/antipodean-albatross-simulations>.

## 1. INTRODUCTION

The seabird species Antipodean albatross (*Diomedea antipodensis antipodensis*) is endemic to New Zealand and consists of two subspecies, Antipodean albatross (*D. a. antipodensis*) and Gibson's albatross (*D. a. gibsoni*). The subspecies Antipodean albatross breeds almost exclusively on Antipodes Island, with a few pairs breeding on Chatham and Campbell islands, whereas Gibson's albatross breeds on Auckland Island. The species is classified as Endangered by the International Union for Conservation of Nature (BirdLife International 2018), and each subspecies is classified individually as Nationally Critical in New Zealand (Robertson et al. 2017).

The population of Antipodean albatross is exposed to a number of at-sea threats. They are caught incidentally in surface-longline fisheries in New Zealand waters and globally (Richard & Abraham 2020). Climate change may also impact the population indirectly, increasing heat stress to chicks and affecting the distribution or abundance of prey species.

On Antipodes Island, the species' population has been monitored within a 29-ha (0.29-square kilometre) area every year since 1994, except in 2006. Field data from this area (Elliott & Walker 2020) and quantitative modelling (Edwards et al. 2017) suggest a population decline since 2007, via a decrease in female survival and in breeding success, and an increase in recruitment age. Tracking data of individual at-sea movements also suggest a potential change in the foraging grounds over time (Elliott & Walker 2020), which may have increased their overlap with fisheries (Bose & Debski 2020). A number of mitigation techniques exist to reduce the level of incidental captures in fisheries and are already in place in a number of fisheries, in New Zealand and worldwide (Løkkeborg 2011).

The main objective of this project was to develop an online tool to facilitate the prioritisation of management strategies around population threats. The online tool allows the running of simulations of the fate of the population under different scenarios, leading to the identification of strategies with the highest positive impact on the population. The simulations rely on estimates of the main demographic parameters of this subspecies. A Bayesian integrated population model was developed for this purpose, based on individual capture-recapture, breeding, and nest success data that have been collected in the study area on Antipodes Island since 1994.

## 2. METHODS

The Antipodean albatross subspecies breeds almost exclusively on Antipodes Island (Agreement on the Conservation of Albatrosses and Petrels 2009). When breeding, a single egg is laid on a nest consisting of a low pedestal build of soil and vegetation, often re-used between breeding attempts. It takes a year for an egg to produce a fledgling. For this reason, adults can only breed every second year when successful. Fledglings spend at least three years at sea before returning to the colony, and subsequently spend another year or more before breeding for the first time (Agreement on the Conservation of Albatrosses and Petrels 2009).

Since 1994, the 29-ha (0.29-square kilometre) area on Antipodes Island has been surveyed every year, except in 2006; the most recent survey was in 2021. Survey visits to the island

were generally conducted in January, so that the outcome of the previous year's breeding attempts could be observed, while simultaneously recording new breeding attempts. Each visit was on average for a month to allow sufficient time to survey the birds present and to band any new birds in the study area. Due to the remoteness of the island and its limited accessibility, logistic constraints led to variation in the exact timing and length of visits between years.

The data collected in the field consist of the date and location of detected banded individuals at the site, their breeding status and stage, and their sex when identifiable.

Additionally, a buffer around the study area was frequently visited, next to two other blocks on the island. In these areas, the sightings and breeding status of banded individuals were also recorded, and identified as being outside the study area. A description of the field data is presented in Edwards et al. 2017

The data were aggregated to create individually- and annually-based capture histories, representing the state of individuals each year between 1994 and 2021. Individuals were categorised into three age classes: juvenile (between fledging and first return to the colony), pre-breeder (from first return to first breeding at the colony), and adult (after first breeding). Eight observed states were represented:

1. adult breeding inside the study area;
2. adult non-breeding inside the study area;
3. adult outside the study area (breeding or not);
4. pre-breeder inside the study area;
5. pre-breeder outside the study area;
6. juvenile;
7. dead;
8. not seen.

Adults sighted both inside and outside the study area one year were considered inside the study area. Adults only sighted outside the study area were not split between breeders and non-breeders as their breeding status cannot be identified precisely (especially for birds seen early in the season). Because surveys of the study area overlapped between the end of the previous breeding season and the beginning of the next one, the aggregated data were prepared to represent the status of the population just before breeding occurs; i.e., chicks of the current breeding year first appear in the prepared data the following year after fledging (if successful). Only birds banded within the study area were included in the final dataset, representing a total of 3,176 individuals, 1,730 of which were banded as chicks.

Nest success was recorded at the nest level, as the nesting individuals might not necessarily be seen, and nests were considered successful if they produced a fledgling. A successful nests could either have a chick being very close to fledging at the last observation, or empty but showing indications of recent breeding activity without showing any sign of failure (e.g.; broken shells, dead body parts).

## **2.1 Integrated population model**

To estimate the main demographic parameters of the population of Antipodean albatross, a multi-state Bayesian capture-recapture model was developed. This type of model aims to alleviate the main biases in the data, which are common to most population survey data.

The state of an individual can be unknown, and an individual may be undetected but still alive. Individuals may be undetected in a given year for several reasons. They could be at sea, such as juveniles, adults previously breeding successfully or on a “sabbatical” year, or breeding adults on a foraging trip may not be detected during short visits to the island. Undetected individuals could also be present at the colony, but outside the study area.

For these reasons, the “actual” state of individuals was considered as a latent variable in the model, with year-to-year transitions between the states determined by explicit biological rules. For example, an adult cannot become a juvenile, or an adult breeding successfully cannot breed again the following year. In addition, an observation process was considered, linking the latent state to the observed state, and determined by both the survey effort and the birds’ behaviour.

### **2.1.1 Latent states**

A total of eight latent states were considered in the model, different from the observed states:

1. adult breeding inside the study area;
2. adult breeding outside the study area;
3. adult non-breeding inside the study area;
4. adult non-breeding outside the study area;
5. pre-breeder inside the study area;
6. pre-breeder outside the study area;
7. juvenile;
8. dead.

The transition matrix between the eight latent states required specifying the probability of being in each latent state given the previous one, representing 64 transition probabilities.

For juveniles (J), pre-breeders (PB), breeding adults (B), and non-breeding adults (NB), the probabilities of changing to a different live state given the previous state were:

$$P(\text{PB}_t | \text{J}_{t-1}) = R_a \phi_{\text{J}}, \quad (1)$$

$$P(\text{B}_t | \text{PB}_{t-1}) = B_a \phi_{\text{PB}}, \quad (2)$$

$$P(\text{B}_t | \text{NB}_{t-1}) = P(\text{breed} | \text{non-breeder}) \phi_s, \quad (3)$$

$$P(\text{NB}_t | \text{B}_{t-1}) = \begin{cases} 1 & \text{after a successful breeding attempt,} \\ (1 - P(\text{breed} | \text{fail})) \phi_s & \text{after a failed breeding attempt,} \end{cases} \quad (4)$$

where  $t$  is the year,  $\phi_{\{\text{J}, \text{PB}, s\}}$  the annual survival rate of juveniles, pre-breeders, and adults of sex  $s$ , respectively,  $R_a$  the probability of a juvenile of age  $a$  returning to the colony,  $B_a$  the probability of a pre-breeder of age  $a$  breeding for the first time,  $P(\text{breed} | \text{fail})$  the probability of an adult breeding in a particular year, given it was an unsuccessful breeder the previous year,  $P(\text{breed} | \text{non-breeder})$  the probability of an adult breeding in a particular year, given it was a non-breeding adult the previous year.

When the sex was unknown, conditional probabilities were used; e.g., the annual survival rate of an individual of unknown sex was  $P(\varnothing) \phi_{\varnothing} + (1 - P(\varnothing)) \phi_{\sigma}$ , where  $P(\varnothing)$  is the probability that an individual in the study area is a female.

The probabilities of remaining in the same live state from one year to the next were:

$$P(\text{J}_t | \text{J}_{t-1}) = (1 - R_a) \phi_{\text{J}}, \quad (5)$$

$$P(\text{PB}_t | \text{PB}_{t-1}) = (1 - B_a) \phi_{\text{PB}}, \quad (6)$$

$$P(\text{NB}_t | \text{NB}_{t-1}) = (1 - P(\text{breed} | \text{non-breeder})) \phi_s, \quad (7)$$

$$P(\text{B}_t | \text{B}_{t-1}) = \begin{cases} 0 & \text{after a successful breeding attempt,} \\ P(\text{breed} | \text{fail}) \phi_s & \text{after a failed breeding attempt,} \\ (1 - P(\text{success})) P(\text{breed} | \text{fail}) \phi_s & \text{after an unknown outcome.} \end{cases} \quad (8)$$

In addition, the transition probabilities were multiplied by the probability of moving inside or outside the study area, depending on the state:

$$P(\text{Out}_t | \text{Int}_{t-1}) = E_s, \quad (9)$$

$$P(\text{In}_t | \text{Out}_{t-1}) = I_s, \quad (10)$$

$$P(\text{Out}_t | \text{Out}_{t-1}) = 1 - I_s, \quad (11)$$

$$P(\text{In}_t | \text{In}_{t-1}) = 1 - E_s, \quad (12)$$

where  $E_s$  is the probability of an individual of sex  $s$  moving out of the study area (emigrate), and  $I_s$  the probability of an individual of sex  $s$  moving into the study area (immigrate).

The probabilities of being dead ( $D$ ) in a particular year were:

$$P(\text{D}_t | \text{J}_{t-1}) = 1 - \phi_{\text{J}}, \quad (13)$$

$$P(\text{D}_t | \text{PB}_{t-1}) = 1 - \phi_{\text{PB}}, \quad (14)$$

$$P(\text{D}_t | \text{B}_{t-1}) = 1 - \phi_s, \quad (15)$$

$$P(\text{D}_t | \text{NB}_{t-1}) = 1 - \phi_s, \quad (16)$$

$$P(\text{D}_t | \text{D}_{t-1}) = 1. \quad (17)$$

The probability of impossible transitions—e.g., from adult to juvenile or to pre-breeder, from pre-breeder to juvenile, and from dead to alive—were fixed to zero.

The adult annual survival rate was estimated independently for females and males, and was allowed to vary randomly between years, with the survival rate  $\phi_{s,t}$  for sex  $s$  at year  $t$  being defined on the logit scale as:

$$\text{logit}(\phi_{s,t}) = \text{logit}(\bar{\phi}_s) + \epsilon_{s,t} s_s, \quad (18)$$

where  $\bar{\phi}_s$  is the mean survival rate across years for sex  $s$ ,  $\epsilon_{s,t}$  is the normally-distributed random effect for each sex and year, and  $s_s$  is the sex-specific variability of the random effect among years.

The annual survival rate of juveniles and pre-breeders was assumed to be constant over time, and the same between males and females in the model.

Breeding success, i.e., the probability that a nest produces a fledgling, was also modelled as a random effect over time.

The probability  $R_a$  of a juvenile of age  $a$  returning to the colony and becoming a pre-breeder was set to 0 at ages below the minimum observed age at first return (3 years), and set to 1 for birds of age 9 and above, as all birds are expected to have returned to the colony by age 9 (G. Elliott, pers. comm.). The age-specific probability of return for birds aged 3 to 8 was modelled as a random effect.

Similarly, the probability  $B_a$  of a pre-breeder of age  $a$  to become a breeder for the first time was set to 0 for birds under 7 years old, the minimum recorded breeding age. The age-specific probability of first breeding for birds aged 7 to 20 was modelled as a random effect. The probability for birds aged 21 and above was set to be constant to represent the long tail in the distribution of age at first breeding (i.e., some birds take a long time to breed or do not breed)

Both  $R_a$  and  $B_a$  were dependent on age, but assumed not to vary with year.

### 2.1.2 Observation process

In the model, latent states are related to observed states via an observation matrix, representing the probability of recording any of the eight observed states given a latent state (one of 8 latent states, different from the observed states).

The probability of detection was estimated separately in the model for:

- breeding adults inside the study area,
- non-breeding adults inside the study area that previously bred successfully,
- other non-breeding adults inside the study area,
- pre-breeders inside the study area,
- adults and pre-breeders outside the study area,
- juveniles (outside the study area by definition),

- dead individuals.

There were only a few recorded observations of juveniles and dead individuals, with all juveniles and most deaths being recorded at sea. For this reason, their detection probability was assumed to be constant among years.

Because year-to-year variations are most likely to reflect the timing and amount of observations on the island, the other detection probabilities were allowed to vary among years, but with the same annual variability among them; they were defined as:

$$\text{logit}(\gamma_{x,t}) = \text{logit}(\gamma_x) + \epsilon_t s, \quad (19)$$

where  $\gamma_{x,t}$  is the detection probability of birds of category  $x$  at year  $t$ ,  $\text{logit}(\gamma_x)$  the average detection probability for category  $x$  among years,  $\epsilon_t$  the random annual effect of year  $t$  for all categories, and  $s$  the variability among years for all categories.

For 2006, when the population was not surveyed, all detection probabilities were fixed to zero.

### 2.1.3 Model fitting

The model was written in the Stan language and fitted in the R statistical package (R Core Team 2019) using the *rstan* library (Stan Development Team 2020).

Stan was chosen over alternatives such as Bugs or JAGS as it implements the no-U-turn sampler (NUTS; Hoffman & Gelman 2014) which improves model convergence and allows fitting times to be reduced by an order of magnitude (from days to hours).

One disadvantage of Stan is that it does not support the direct sampling of discrete parameters. Nevertheless, multi-state models can still be fitted by marginalising discrete latent states, i.e., summing at each time step the likelihood of the observed state over all possible latent states, iteratively over each individual capture history (Yackulic et al. 2020).

The model was fitted using Markov chain Monte Carlo (MCMC) methods, using four chains, for 6,000 iterations, after a burn-in period of 5,000 iterations.

The code of the Stan model is provided in Appendix A.

## 2.2 Population simulations

The main aim of this project was to provide stakeholders with a tool to simulate the fate of the Antipodean albatross population under different scenarios. For this purpose, an interactive online application written in R and using the Shiny framework was developed.

Because the demographic model does not provide the latent state of individuals at each time step directly due to the marginalisation of discrete latent variables, the initial population structure for the simulations was derived separately. For this purpose, the latent state at each time step for each individual was drawn randomly from the previous state and the observed state. Using Bayes' theorem, the probability of an individual to be

in the latent state  $\Pi_i$  given the observed state  $O$  is:

$$P(\Pi_i|O) = \frac{P(O|\Pi_i)P(\Pi_i)}{P(O)}, \quad (20)$$

where  $P(O|\Pi_i)$  is the probability of the observed state  $O$  given the latent state, which is the detection probability of that state, as estimated by the model.  $P(\Pi_i)$  is the probability of state  $\Pi_i$  and is the transition probability from the previous latent state, as estimated by the model.  $P(O)$  is the probability of the observed state, and is the sum of observing  $O$  given all possible latent states, i.e.,  $\sum_k P(O|\Pi_k)P(\Pi_k)$ . In addition, the probability of a dead individual at a given time step was set to zero when the individual was subsequently detected alive. The process was repeated for each of the 6 000 MCMC samples from the model, and the resulting population structure in 2021—and its uncertainty—was taken as the initial population for the simulations. Pre-breeders and adults outside the study area were not included, to simulate only the population inside the study area and the juveniles that fledged from there.

The population size from the simulations was scaled up by the ratio of the total number of breeding pairs on the island to the number of breeding pairs inside the study area. The total number of breeding pairs was estimated from extensive surveys of the whole island in 1994, 1995, and 1996. The scaling of the studied population size to the whole island, therefore, assumes that the ratio did not change over time. The proportion of the number of breeding pairs that were inside the study area was estimated to be 2.7332% averaged across the three censuses (Elliott & Walker 2020), and the inverse of this value (36.58715) was used to scale up the simulation population size to the whole island.

The population simulations consisted of predicting the fate of each individual in the initial 2021 population, and of new fledglings produced each year, every year for 30 years, based on the demographic parameters estimated in the model. For each simulated year, an actual year between 2008 and 2020 was first drawn randomly to represent the interannual variability estimated in the model, while considering only the most recent years. The drawn year defined the value of survival rates and breeding success. Surviving individuals were drawn following a Bernoulli process with a probability equal to the survival rate of the drawn year and of the individual class (juvenile, pre-breeder, adult female, or adult male). Juveniles and pre-breeders either remained in their age class or moved to the next one depending on the age-specific transition probabilities. Adults breeding that year were then drawn according to the probability of breeding, depending on whether they bred successfully (or not) the previous year. The success of breeding adults was then drawn randomly from the probability of success of that year. Among successful breeders, the number of fledglings produced was taken as the minimum number of female or male adults, and new individuals of age 0 were created, with a sex assigned randomly with a probability of 0.5. This process was then repeated iteratively for the 30 simulated years, and for each iteration of the MCMC methods.

In the online tool, scenarios are specified in terms of direct impacts, affecting specific demographic parameters. Threats can impact the annual survival rate of juveniles, pre-breeders, adult males, and adult females separately, or can also impact breeding probability or breeding success. The threats can be defined as being either already present, in which case the impact is removed from the population in the simulations, or potential, with the impact added to the population. For example, to assess the potential effect of introducing new mitigation measures in fisheries, the impact would need to be

specified as already present, and the incidental mortalities would be removed from the population in the simulations.

Impacts may be specified as an absolute change in the demographic parameter, or as a number of individuals for survival rates. When using individuals, the impact is converted to the absolute change in survival rate,  $\Delta$ , based on the total number of individuals in the affected category:

$$\Delta = S' - S = 1 - \frac{(1 - \Phi)N - I}{N} - \Phi, \quad (21)$$

where  $S'$  is the new survival rate,  $\Phi$  the survival rate of the population category (juvenile, pre-breeder, adult female, or adult male),  $N$  the scaled-up number of individuals in the category, and  $I$  the number of mortalities caused by the threat. The conversion of impacts from individuals to a change in demographic rates assumes that the impact of threats is consistently proportional to the population size.

Multiple threats and impacts may be specified for a given scenario. In this case, the overall change in demographic parameters is calculated by summing the absolute changes across threats and impacts within each demographic parameter.

Upon completion of the simulations, the mean and 95% credible interval of the population size, of the number of annual breeding pairs, and population mean annual growth rate, and the mean population structure are calculated and reported, in tables and figures.

To illustrate how the removal of threats (i.e., their mitigation) may affect population growth, two hypothetical scenarios were simulated, representing two existing threats. Each threat resulted in the death of 500 individuals. Under one scenario, all deaths were of juveniles, whereas under the second scenario, all deaths were of adults (male and female).

### 3. RESULTS

#### 3.1 Model parameters

The MCMC traces indicated that the model converged reasonably well, as the four chains were well mixed and did not show significant autocorrelation (see Appendix B for the MCMC traces and values of each demographic parameter estimated by the model). One exception was the parameter related to the detection probability, which converged but showed marked autocorrelation, but this would not impact the results.

The estimated adult annual survival rate between 1994 and 2020 showed changes over time (Figure 1). Before 2005, the estimated survival rate was similar between sexes, with an annual mean of 0.947 (95% c.i.: 0.914 – 0.973). From 2005, however, estimated female survival declined to a mean of 0.882 (95% c.i.: 0.814 – 0.94); female survival was lowest in 2013, estimated at 0.821 (95% c.i.: 0.752 – 0.883). In contrast, male survival only slightly declined to a mean of 0.927 (95% c.i.: 0.887 – 0.961), with a minimum around 0.90 in 2007.

The estimated survival in the three most recent years (2018 to 2020) suggested a possible increase to levels similar to estimates before 2004, with female adult survival reaching 0.929 (95% c.i.: 0.861 – 0.976) in 2020, and adult male survival at 0.971 (95% c.i.:

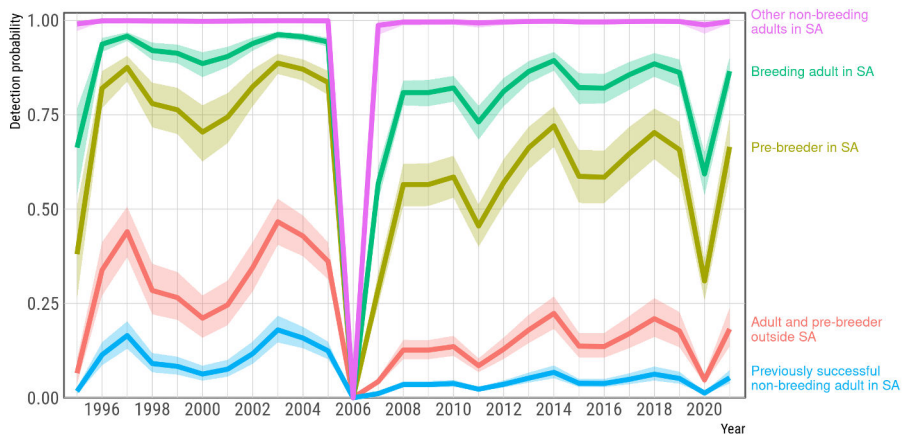


**Figure 1:** Adult annual survival for female and male Antipodean albatross between 1994 and 2020, estimated from the demographic model. Lines indicate the mean, shading the 95% credible interval.

0.943 – 0.991).

The annual survival rate of juveniles and pre-breeders, assumed to be constant among years, was estimated at 0.879 (95% c.i.: 0.869 – 0.888) and 0.922 (95% c.i.: 0.913 – 0.931), respectively.

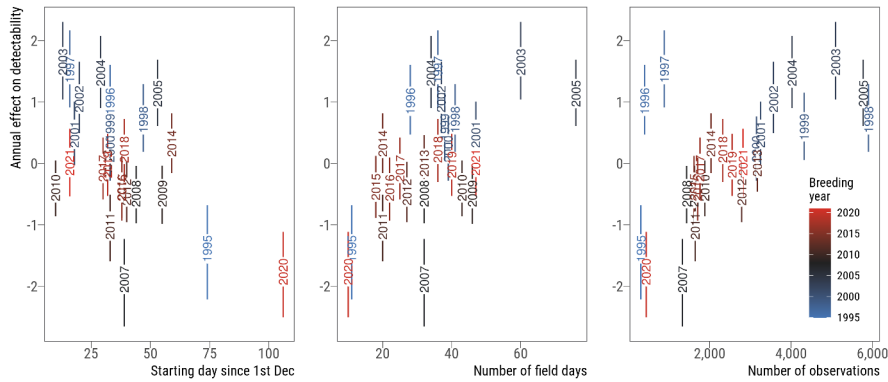
The change of adult survival rates over time was considerable even though the interannual variability in the probability of detection was controlled in the model. The detection probability also showed a decrease over time, i.e., after 2006 (Figure 2).



**Figure 2:** Detection probability of individuals inside the study area (SA) for breeding adults, non-breeders that were previously successful breeders, other non-breeders, and pre-breeders, and for adults and pre-breeders combined outside the study area. Lines indicate the mean, shading the 95% credible interval.

The interannual change in detectability, applied to all individuals present on the island, was related to both the timing and length of the field seasons on the island (Figure 3). Estimates of detectability were highest when the field season started early

(early December) and when the survey effort was high, both in the number of days with recorded field observations, and in the total number of recorded observations in the season.



**Figure 3:** Relation between the interannual variability of the probability of detection and the timing and effort of population surveys. The timing of surveys was measured here as the number of days between the 1 December preceding the breeding season and the first day of recorded observations. Observation effort is in the number of days with observations, and the total number of observations recorded during the breeding season. The annual effect on detectability is shown as the 95% credible interval of the annual random effect as estimated in the model, and the label showing the year of the field season is centred on the mean estimate.

Amongst the years with the lowest detectability, 1995 and 2020 were characterised by a low number of field days and observations, and started late in the season (mid-February and mid-March, respectively). In contrast, the highest estimated detectability was in 2003, when the field season was both the second earliest (mid-December) and the second longest (60 days of observations).

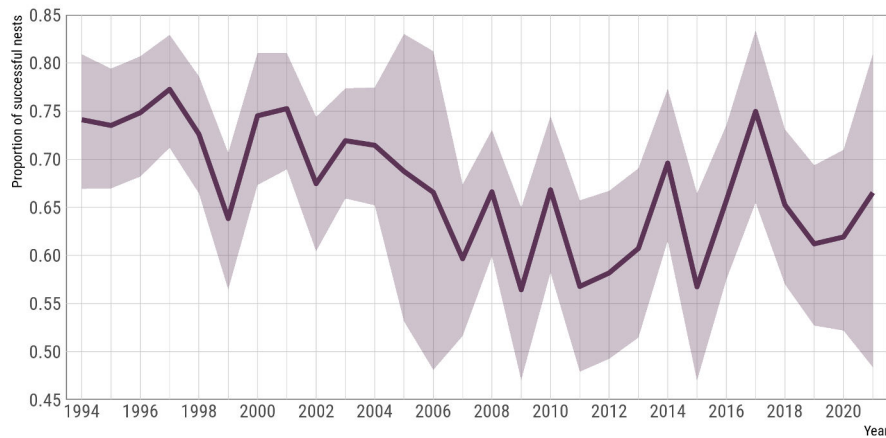
The estimated probability of detection varied significantly between individuals considered in the model (Table 1 and Figure 2). This probability was around 5.2% for non-breeding adults that were successful breeders the previous years, 18.0% for adults and pre-breeders outside the study area, 66.1% for pre-breeders inside the study area, 86.4% for adults breeding inside the study area, and 99.7% for non-breeding adults that were not successful breeders in the previous year. Additionally, the detectability was estimated close to zero for both juveniles and dead individuals, with a mean of 0.019% (95% c.i.: 0% – 0.073%) and 0.083% (95% c.i.: 0.054% – 0.118%), respectively.

The probability of breeding was estimated in the model depending on the previous breeding status, and was assumed to be constant among years. For adults that were failed breeders the previous year, the probability of breeding was estimated at 70.5% (95% c.i.: 68.6% – 72.3%). The probability was considerably lower for previous non-breeders, at 64.1% (95% c.i.: 62.8% – 65.4%).

As for survival, breeding success was also allowed to vary among years in the model. Modelled as the probability that a nest successfully produces a fledgling, breeding success also declined between the period 1994–2004 and 2005–2021 (Figure 4). Prior to 2005, the mean breeding success was estimated at 72.4% (95% c.i.: 65.8% – 78.6%), but at 63.7% (95% c.i.: 53.4% – 73%) after 2005.

**Table 1:** Mean estimates (and credible interval, c.i.) of the probability of detection among the different individual types in the Antipodean albatross population considered in the demographic model (SA, study area).

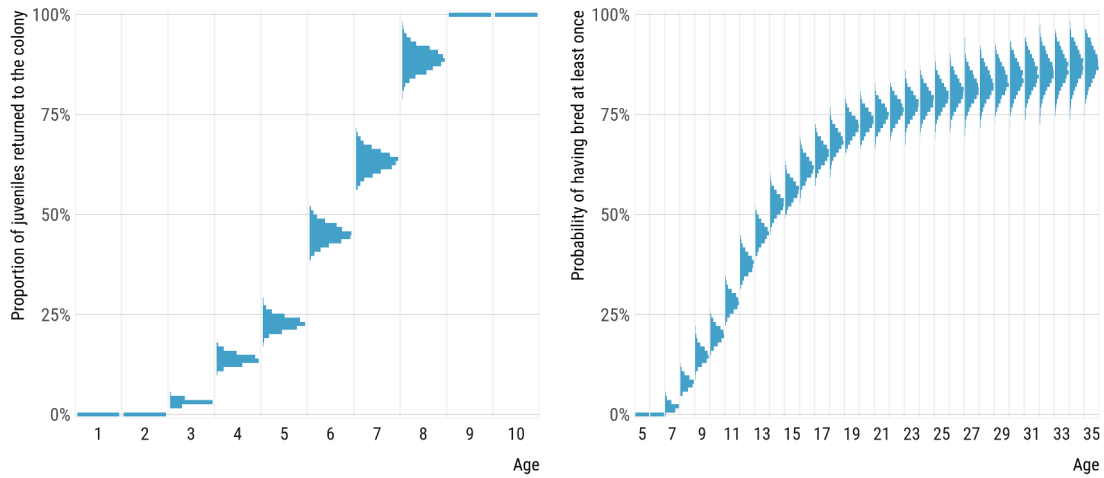
Type	Mean	95% c.i.
Breeding adult in SA	0.864	0.816 – 0.900
Previously successful non-breeding adult in SA	0.052	0.036 – 0.072
Other non-breeding adults in SA	0.997	0.992 – 1.000
Pre-breeder in SA	0.661	0.575 – 0.736
Adult and pre-breeder outside SA	0.180	0.132 – 0.234



**Figure 4:** Breeding success by year for Antipodean albatross between 1994 and 2021, measured as the proportion of nests producing a fledgling. Line indicates the mean, shading the 95% credible interval.

To take into account bird movements in and out of the study area for the estimation of survival rates, the probability of individuals that were inside the study area leaving the area, and conversely the probability of individuals that were outside the study area returning to it, were estimated for females and males independently, and assumed to be constant among years. These probabilities suggest that females are less faithful to their area than males, as females had a 9% (95% c.i.: 8.1% – 10%) probability of leaving the study area, compared with 4% (95% c.i.: 3.5% – 4.6%) for males. Similarly, females had an estimated probability of 17.7% (95% c.i.: 15.2% – 20.3%) to return to the colony after leaving it, compared with 25.4% (95% c.i.: 21.9% – 29.1%) for males.

The ages at first return and at first breeding were also estimated in the model (Figure 5). The age at first return varied between 3 and 9 years, with an average at 6.26 years. The minimum age at first breeding was 7 years, and by age 13, half of the individuals had bred at least once, although some individuals did not breed at all.



**Figure 5:** Proportion of individuals that returned to the colony (left) and proportion of individuals that bred at least once as function of age (right). For each age, a histogram of the Markov chain Monte Carlo values is shown as estimated by the model.

### 3.2 Online simulation tool

Based on the demographic parameters obtained from the model, an online application was developed to simulate the population dynamics of Antipodean albatross under different scenarios (see a screenshot of the online simulation tool in Figure 6). The application can be accessed at <https://dragonfly-science.shinyapps.io/antipodean-albatross-simulations>.

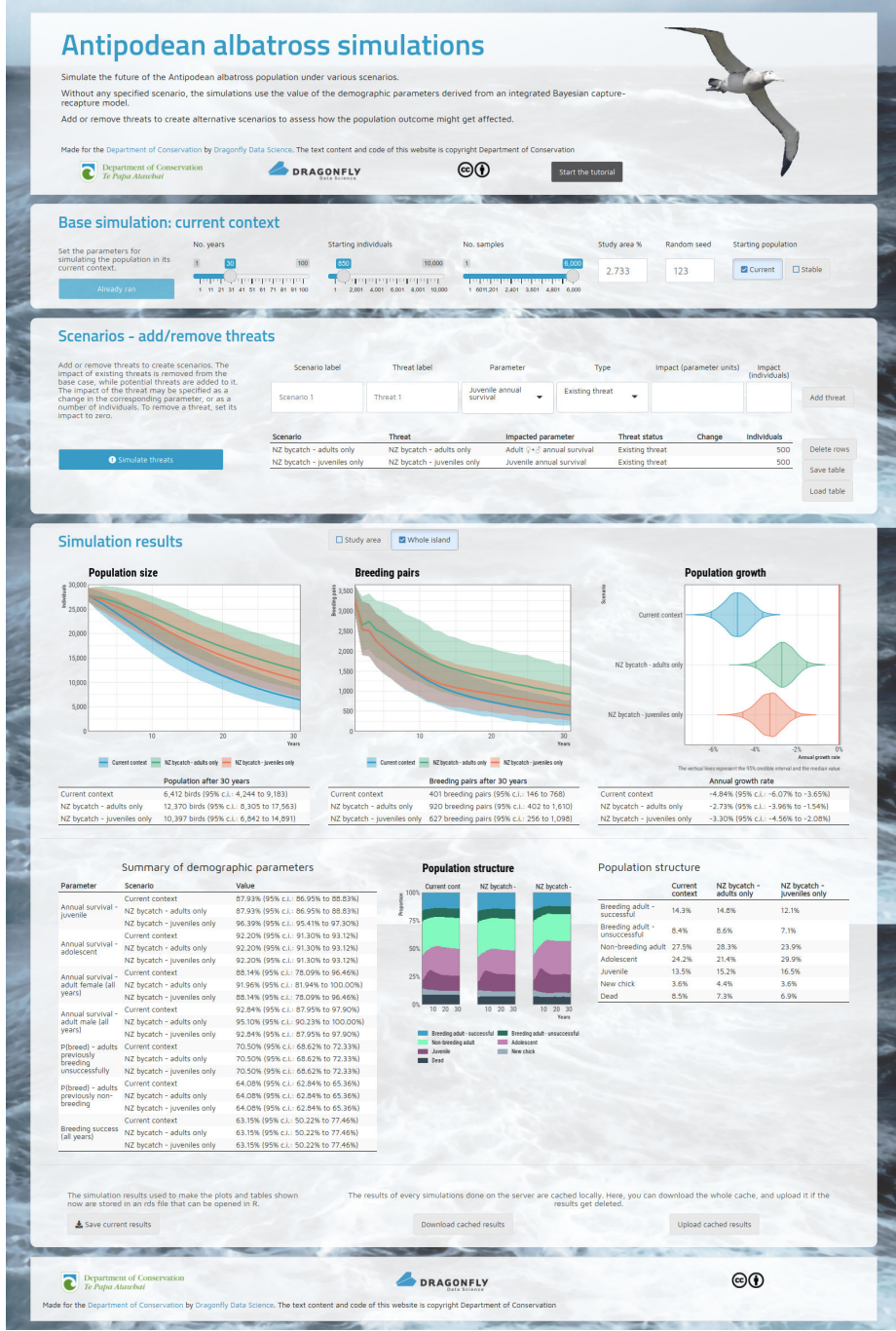
The structure of the population in 2021 was used for the initialisation of the simulations, and was obtained from drawing iteratively the latent state of each individual in the study area each year when the state was unknown (examples of the predictions of individual state are shown in Figure 7).

The number of breeding pairs inside the study area from on-site surveys was similar to the estimates derived from the model (Figure 8). Nevertheless, the model estimate was higher overall. This difference was due to the model estimate including the individuals that are not detected during surveys.

The population in 2021 used to initialise the simulations was estimated inside the study area at 90 (95% c.i.: 81 – 100) breeding pairs, and 762 (95% c.i.: 726 – 801) total individuals. Scaling up to the entire island, these estimates represent a total of 3,292 (95% c.i.: 2,964 – 3,659) breeding pairs and 27,893 (95% c.i.: 26,562 – 29,306) total individuals.

On average, the population consisted of 15.7% juvenile, 21.3% pre-breeders, 37.5% non-breeding adults, 17% successful breeding adults, and 8.6% unsuccessful breeding adults.

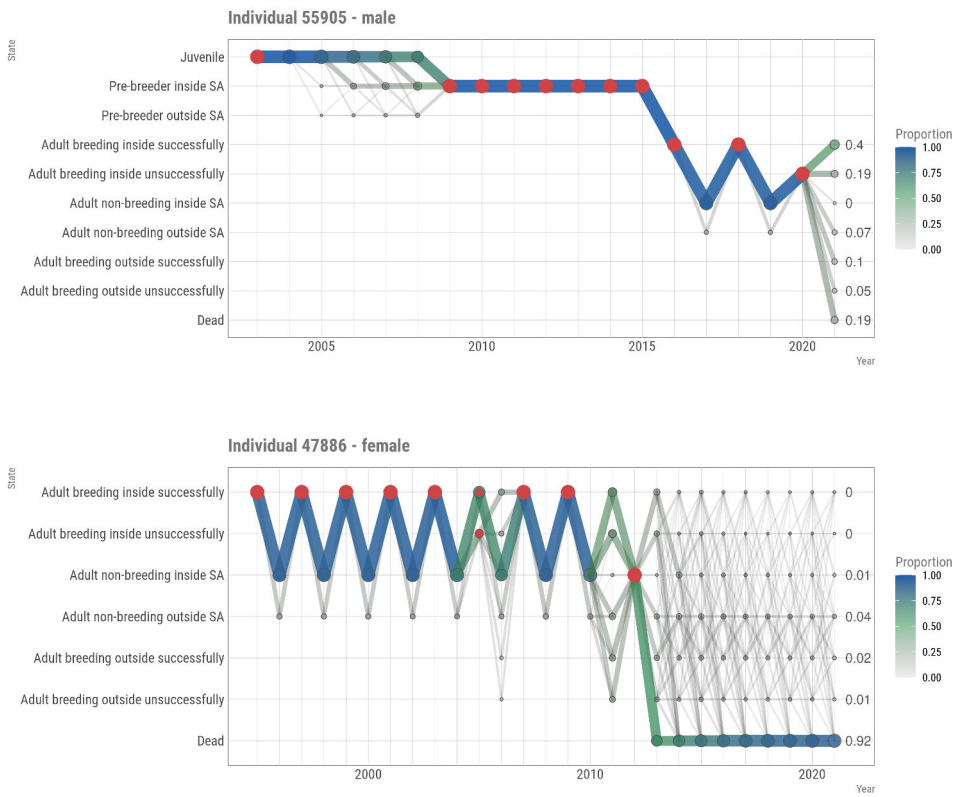
In the current context, i.e., without specifying any management scenario, simulations predicted a population decline of 4.84% (95% c.i.: 3.65% – 6.07%) with the total annual number of breeding pairs in the study area decreasing from 90 (95% c.i.: 81 – 100) to 11 (95% c.i.: 4 – 21) after 30 years (“Current context” in Figure 9). Scaling up the study area population to the entire island, this estimate corresponded to a decline from 3,292 (95%



**Figure 6:** Screenshot of the online application tool to run predictions of the Antipodean albatross population in the future under different scenarios.

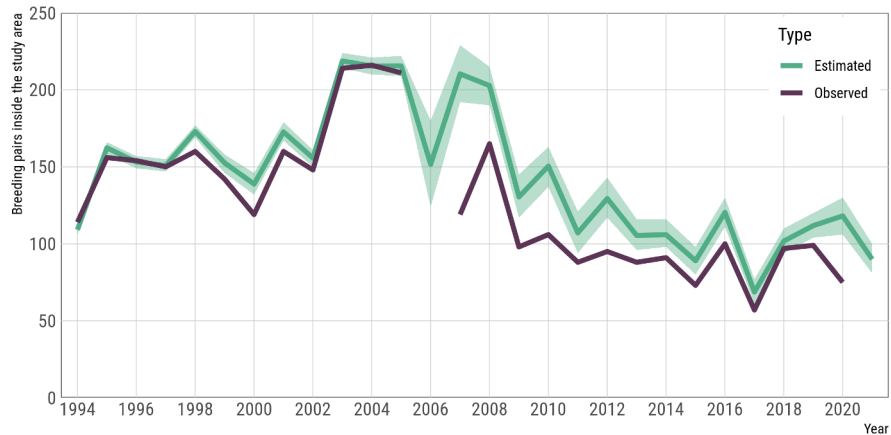
c.i.: 2,964 – 3,659) breeding pairs to 401 (95% c.i.: 146 – 768), or for the whole population, from 27,893 (95% c.i.: 26,562 – 29,306) birds to 6,412 (95% c.i.: 4,244 – 9,183).

When simulating a hypothetical scenario of mitigating an existing threat causing the death of 500 juveniles, the rate of decline decreased to 3.3% (95% c.i.: 2.1% – 4.6%); when

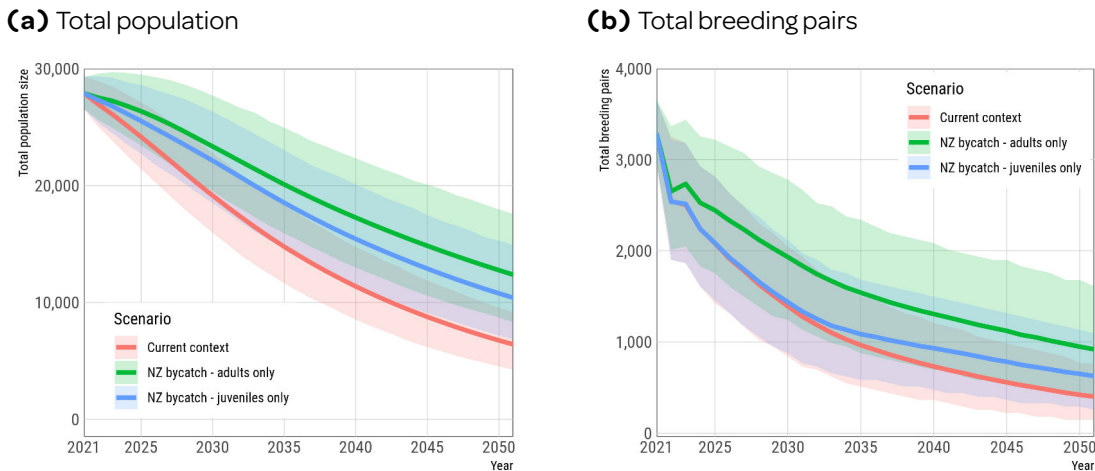


**Figure 7:** Examples of drawing the latent state of individuals from their observed state. Red dots represent the latent states that are possible given the observed state of an individual that was detected. The size and colour of segments indicate the probability of transition between two successive states. Numbers indicate the probability of each state in 2021, used to draw the initial population structure for population projections.

the mortalities only affected adults, the rate further decreased to 2.7% (95% c.i.: 1.5% – 4%) (Figure 9).



**Figure 8:** Comparison of the annual number of breeding pairs when recorded during field surveys (Observed) and when estimated from the model (Estimated). Lines indicate the mean, shading the 95% credible interval for the estimate.



**Figure 9:** Simulation of the population size (a) and of the number of annual breeding pairs (b) of Antipodean albatross over the next 30 years on Antipodes Island. The simulation is based on the demographic parameters estimated in the model, only keeping the time-varying values between 2008 and 2020. The mean and 95% credible interval are shown.

## 4. DISCUSSION

The aim of this project was to provide an online simulation tool for predicting the outcome of management strategies on the demography of Antipodean albatross. As for any model, the accuracy of the prediction depends on the input field data, the complexity of the factors affecting the demography, and the change over time in the threats to the species.

Although movements in and out of the study area were included in the model, any permanent emigration from the study area was more likely to be considered as local mortality, and may underestimate annual survival rate. The area around the study

site has been visited regularly, and sightings recorded there were used in the model to estimate the rate of movements between areas. It is a relatively small area compared with the rest of the island; some individuals may not be seen again once they relocate permanently, making their emigration indistinguishable from death. Nevertheless, the observations of the researchers when moving across the island suggest that permanent emigration by a significant number of individuals is unlikely (G. Elliott, pers. comm.).

The current model specification was designed to provide a basis for the simulations, requiring the balancing of realism and simplicity. For example, a number of parameters were not dependent on years, such as the probability of breeding or the survival rate of pre-breeders, and the model presented here may not be the closest representation of reality. The model results and absolute projections into the future need to be viewed in this context. Nevertheless, model limitations do not detract from the overall trajectory of a declining Antipodean albatross population, and the simulation tool allows comparison of the relative impact of alternative management strategies under different scenarios.

The recent increase in survival rates since 2018 may be a probabilistic coincidence, but could also indicate an alleviation of the threats affecting females predominantly. For example, fisheries may operate in different areas over time, or the areas where individuals forage may also vary, resulting in a change in the overlap between the species and fishery threats. The next few years of field data will inform whether this trend continues.

#### 4.1 Recommendations

Regular updates of the demographic model after each field season would ensure that the best estimates are used for the simulations. In particular, additional years would reduce the sampling effect of drawing each simulated year from a limited set of annual estimates, and would provide a more complete representation of inter-annual variability.

The conversion of the impact of threats from individuals to parameter units in the simulations relies on the proportion of the number of breeding pairs that were inside the study area. This scalar was obtained from surveys between 1994 and 1996, and it is possible that it no longer represents current conditions. For instance, a number of landslides during an extreme weather event in January 2014 (Chilvers & Hiscock 2019) may have changed the spatial distribution of some colonies. An updated and complete population survey of the island would provide a new estimate of this critical parameter.

An island survey would also be an opportunity to record all sightings of birds that were banded inside the study area. This record would allow reviewing the estimates of movements in and out the study area, which may improve the estimate of survival rates.

Further developments to the model may improve the accuracy of the population predictions. For example, no inter-annual variability of the probability of breeding was considered, and may be beneficial to future updates of the model.

## 5. ACKNOWLEDGMENTS

I am sincerely thankful to Graeme Elliott and Kath Walker for providing the data they have tenaciously collected over decades, and for sharing their experience of the species

and field work.

The assistance of Philipp Neubauer in the development and fitting of the Bayesian model in Stan, and of Katrin Berkenbusch in the editing of this report, were much appreciated.

This work was funded by Department of Conservation (through project BCBC2020-09). I am grateful to Johannes Fischer and Igor Debski for the constructive talks during the project.

Data preparation and statistical analyses were carried out using R (including the libraries `data.table` and `rstan`) and Stan, writing scripts using Emacs, containerised using Docker, and this document was produced using L<sup>A</sup>T<sub>E</sub>X. I am grateful to the many people who contribute to these key open source software projects and make them available.

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## APPENDIX A STAN MODEL CODE

```

1  functions{
2
3    matrix trans_probs(int nstates, real s_ad, real s_prebr, real s_juv,
4                      real p_mv_out, real p_mv_in, int succ, vector p_breed,
5                      real p_rec, real p_bead, real p_succ) {
6
7      /** TRANSITIONS and SURVIVAL **/
8
9      // 1: adults breeding inside SA
10     // 2: adults breeding outside SA
11     // 3: adults non-breeding inside SA
12     // 4: adults non-breeding outside SA
13     // 5: pre-breeders inside SA
14     // 6: pre-breeders outside SA
15     // 7: juvs
16     // 8: deads
17
18     matrix[nstates, nstates] tmat;
19
20     /** ADULTS PREVIOUSLY BREEDING WITHIN STUDY AREA **/
21     // re-breeding in SA (SA = study area)
22     tmat[1, 1] = succ == 2 ?
23         0 :
24         (succ == 1 ?
25             p_breed[1] * s_ad * (1-p_mv_out) :
26             (1-p_succ) * p_breed[1] * s_ad * (1-p_mv_out));
27     // re-breeding outside SA
28     tmat[1, 2] = succ == 2 ?
29         0 :
30         (succ == 1 ?
31             p_breed[1] * s_ad * p_mv_out :
32             (1-p_succ) * p_breed[1] * s_ad * p_mv_out);
33     // non-breeding in SA
34     tmat[1, 3] = succ == 2 ?
35         s_ad * (1-p_mv_out) :
36         (succ == 1 ?
37             (1-p_breed[1]) * s_ad * (1-p_mv_out) :
38             (1-p_succ) * (1-p_breed[1]) * s_ad * (1-p_mv_out) +
39             p_succ * s_ad * (1-p_mv_out));
40     // non-breeding outside SA
41     tmat[1, 4] = succ == 2 ?
42         s_ad * p_mv_out :
43         (succ == 1 ?
44             (1-p_breed[1]) * s_ad * p_mv_out :
45             (1-p_succ) * (1-p_breed[1]) * s_ad * p_mv_out +
46             p_succ * s_ad * p_mv_out);
47     tmat[1, 5] = 0;           // pre-breeders inside SA
48     tmat[1, 6] = 0;           // pre-breeders outside SA
49     tmat[1, 7] = 0;           // juvs
50     tmat[1, 8] = 1-s_ad;     // dead
51
52     /** ADULTS PREVIOUSLY BREEDING OUTSIDE STUDY AREA **/
53     // re-breeding in SA (SA = study area)
54     tmat[2, 1] = succ == 2 ?
55         0 :
56         (succ == 1 ?
57             p_breed[1] * s_ad * p_mv_in :
58             (1-p_succ) * p_breed[1] * s_ad * p_mv_in);
59     // re-breeding outside SA
60     tmat[2, 2] = succ == 2 ?
61         0 :
62         (succ == 1 ?
63             p_breed[1] * s_ad * (1-p_mv_in) :
64             (1-p_succ) * p_breed[1] * s_ad * (1-p_mv_in));
65     // non-breeding in SA
66     tmat[2, 3] = succ == 2 ?
67         s_ad * p_mv_in :
68         (succ == 1 ?
69             (1-p_breed[1]) * s_ad * p_mv_in :
70             (1-p_succ) * (1-p_breed[1]) * s_ad * p_mv_in + p_succ * s_ad * p_mv_in);
71     // non-breeding outside SA
72     tmat[2, 4] = succ == 2 ?
73         s_ad * (1-p_mv_in) :
74         (succ == 1 ?
75             (1-p_breed[1]) * s_ad * (1-p_mv_in) :
76             (1-p_succ) * (1-p_breed[1]) * s_ad * (1-p_mv_in) + p_succ * s_ad * (1-p_mv_in));
77     tmat[2, 5] = 0;           // pre-breeders inside SA
78     tmat[2, 6] = 0;           // pre-breeders outside SA
79     tmat[2, 7] = 0;           // juvs
80     tmat[2, 8] = 1-s_ad;     // dead
81
82     /** ADULTS PREVIOUSLY NOT BREEDING WITHIN STUDY AREA **/
83     tmat[3, 1] = p_breed[2] * s_ad * (1-p_mv_out); // breeding in SA (SA = study area)
84     tmat[3, 2] = p_breed[2] * s_ad * p_mv_out; // breeding outside SA
85     tmat[3, 3] = (1-p_breed[2]) * s_ad * (1-p_mv_out); // non-breeding in SA
86     tmat[3, 4] = (1-p_breed[2]) * s_ad * p_mv_out; // non-breeding outside SA
87     tmat[3, 5] = 0;           // pre-breeders inside SA
88     tmat[3, 6] = 0;           // pre-breeders outside SA
89     tmat[3, 7] = 0;           // juvs

```

```

90  tmat[3, 8] = 1-s_ad;                                // dead
91
92  /* ADULTS PREVIOUSLY NOT BREEDING OUTSIDE THE STUDY AREA */
93  tmat[4, 1] = p_breed[2] * s_ad * p_mv_in;           // breeding in SA (SA = study area)
94  tmat[4, 2] = p_breed[2] * s_ad * (1-p_mv_in);       // breeding outside SA
95  tmat[4, 3] = (1-p_breed[2]) * s_ad * p_mv_in;     // non-breeding in SA
96  tmat[4, 4] = (1-p_breed[2]) * s_ad * (1-p_mv_in); // non-breeding outside SA
97  tmat[4, 5] = 0;                                     // pre-breeders inside SA
98  tmat[4, 6] = 0;                                     // pre-breeders outside SA
99  tmat[4, 7] = 0;                                     // juvs
100 tmat[4, 8] = 1-s_ad;                                // dead
101
102 /* PRE-BREEDERS INSIDE THE STUDY AREA */
103 tmat[5, 1] = s_prebr * p_beard * (1-p_mv_out);    // breeding in SA (SA = study area)
104 tmat[5, 2] = s_prebr * p_beard * p_mv_out;         // breeding outside SA
105 tmat[5, 3] = 0;                                     // non-breeding in SA
106 tmat[5, 4] = 0;                                     // non-breeding outside SA
107 tmat[5, 5] = s_prebr * (1-p_beard) * (1-p_mv_out); // pre-breeders inside SA
108 tmat[5, 6] = s_prebr * (1-p_beard) * p_mv_out;     // pre-breeders outside SA
109 tmat[5, 7] = 0;                                     // juvs
110 tmat[5, 8] = 1-s_prebr;                            // dead
111
112 /* PRE-BREEDERS OUTSIDE THE STUDY AREA */
113 tmat[6, 1] = s_prebr * p_beard * p_mv_in;          // breeding in SA (SA = study area)
114 tmat[6, 2] = s_prebr * p_beard * (1-p_mv_in);      // breeding outside SA
115 tmat[6, 3] = 0;                                     // non-breeding in SA
116 tmat[6, 4] = 0;                                     // non-breeding outside SA
117 tmat[6, 5] = s_prebr * (1-p_beard) * p_mv_in;     // pre-breeders inside SA
118 tmat[6, 6] = s_prebr * (1-p_beard) * (1-p_mv_in); // pre-breeders outside SA
119 tmat[6, 7] = 0;                                     // juvs
120 tmat[6, 8] = 1-s_prebr;                            // dead
121
122 /* JUVENILES */
123 tmat[7, 1] = 0;                                     // breeding in SA (SA = study area)
124 tmat[7, 2] = 0;                                     // breeding outside SA
125 tmat[7, 3] = 0;                                     // non-breeding in SA
126 tmat[7, 4] = 0;                                     // non-breeding outside SA
127 tmat[7, 5] = s_juv * p_rec * (1-p_mv_out);        // pre-breeders inside SA
128 tmat[7, 6] = s_juv * p_rec * p_mv_out;            // pre-breeders outside SA
129 tmat[7, 7] = s_juv * (1-p_rec);                   // juvs
130 tmat[7, 8] = 1-s_juv;                             // dead
131
132 /* DEADS */
133 tmat[8, 1] = 0;                                     // breeding in SA (SA = study area)
134 tmat[8, 2] = 0;                                     // breeding outside SA
135 tmat[8, 3] = 0;                                     // non-breeding in SA
136 tmat[8, 4] = 0;                                     // non-breeding outside SA
137 tmat[8, 5] = 0;                                     // pre-breeders inside SA
138 tmat[8, 6] = 0;                                     // pre-breeders outside SA
139 tmat[8, 7] = 0;                                     // juvs
140 tmat[8, 8] = 1;                                     // dead
141
142 return tmat;
143 }
144
145
146 matrix obs_probs(int n_obs_states, real[] p_obs, real p_detect_juv, real p_detect_dead,
147                   real p_female, real p_succ, int succ, int no_visit) {
148
149  /* OBSERVED STATES */
150
151  // 1: adults breeding in SA
152  // 2: adults non-breeding in SA
153  // 3: adults outside SA
154  // 4: pre-breeders inside SA
155  // 5: pre-breeders outside SA
156  // 6: juvs
157  // 7: dead
158  // 8: not seen
159
160  matrix[n_obs_states, n_obs_states] pmat;
161
162  /* ADULTS BREEDING WITHIN STUDY AREA */
163  pmat[1, 1] = no_visit == 1 ? 0 : p_obs[1]; // ad breeding in SA (SA = study area)
164  pmat[1, 2] = 0;                           // ad non-breeding in SA
165  pmat[1, 3] = 0;                           // ad outside SA
166  pmat[1, 4] = 0;                           // pre-breeders inside SA
167  pmat[1, 5] = 0;                           // pre-breeders outside SA
168  pmat[1, 6] = 0;                           // juvs
169  pmat[1, 7] = 0;                           // dead
170  pmat[1, 8] = 1 - pmat[1, 1];             // not seen
171
172  /* ADULTS BREEDING OUTSIDE STUDY AREA */
173  pmat[2, 1] = 0;                           // ad breeding in SA (SA = study area)
174  pmat[2, 2] = 0;                           // ad non-breeding in SA
175  pmat[2, 3] = no_visit == 1 ? 0 : p_obs[5]; // ad outside SA
176  pmat[2, 4] = 0;                           // pre-breeders inside SA
177  pmat[2, 5] = 0;                           // pre-breeders outside SA
178  pmat[2, 6] = 0;                           // juvs
179  pmat[2, 7] = 0;                           // dead
180  pmat[2, 8] = 1 - pmat[2, 3];             // not seen

```

```

181  /* ADULTS NON-BREEDING INSIDE STUDY AREA */
182  pmat[3, 1] = 0;                      // ad breeding in SA (SA = study area)
183  pmat[3, 2] = no_visit == 1 ?          // ad non-breeding in SA
184  0 :
185  (succ == 2 ?
186  p_obs[2] :
187  (succ == 1 ?
188  p_obs[3] :
189  p_succ * p_obs[2] + (1-p_succ) * p_obs[3]));
190  pmat[3, 3] = 0;                      // ad outside SA
191  pmat[3, 4] = 0;                      // pre-breeders inside SA
192  pmat[3, 5] = 0;                      // pre-breeders outside SA
193  pmat[3, 6] = 0;                      // juvs
194  pmat[3, 7] = 0;                      // dead
195  pmat[3, 8] = 1 - pmat[3, 2];        // not seen
196
197  /* ADULTS NON-BREEDING OUTSIDE STUDY AREA */
198  pmat[4, 1] = 0;                      // ad breeding in SA (SA = study area)
199  pmat[4, 2] = 0;                      // ad non-breeding in SA
200  pmat[4, 3] = no_visit == 1 ? 0 : p_obs[5]; // ad outside SA
201  pmat[4, 4] = 0;                      // pre-breeders inside SA
202  pmat[4, 5] = 0;                      // pre-breeders outside SA
203  pmat[4, 6] = 0;                      // juvs
204  pmat[4, 7] = 0;                      // dead
205  pmat[4, 8] = 1 - pmat[4, 3];        // not seen
206
207  /* PRE-BREEDERS INSIDE STUDY AREA */
208  pmat[5, 1] = 0;                      // ad breeding in SA (SA = study area)
209  pmat[5, 2] = 0;                      // ad non-breeding in SA
210  pmat[5, 3] = 0;                      // ad outside SA
211  pmat[5, 4] = no_visit == 1 ? 0 : p_obs[4]; // pre-breeders inside SA
212  pmat[5, 5] = 0;                      // pre-breeders outside SA
213  pmat[5, 6] = 0;                      // juvs
214  pmat[5, 7] = 0;                      // dead
215  pmat[5, 8] = 1 - pmat[5, 4];        // not seen
216
217  /* PRE-BREEDERS OUTSIDE STUDY AREA */
218  pmat[6, 1] = 0;                      // ad breeding in SA (SA = study area)
219  pmat[6, 2] = 0;                      // ad non-breeding in SA
220  pmat[6, 3] = 0;                      // ad outside SA
221  pmat[6, 4] = 0;                      // pre-breeders inside SA
222  pmat[6, 5] = no_visit == 1 ? 0 : p_obs[5]; // pre-breeders outside SA
223  pmat[6, 6] = 0;                      // juvs
224  pmat[6, 7] = 0;                      // dead
225  pmat[6, 8] = 1 - pmat[6, 5];        // not seen
226
227  /* JUVENILES */
228  pmat[7, 1] = 0;                      // ad breeding in SA (SA = study area)
229  pmat[7, 2] = 0;                      // ad non-breeding in SA
230  pmat[7, 3] = 0;                      // ad outside SA
231  pmat[7, 4] = 0;                      // pre-breeders inside SA
232  pmat[7, 5] = 0;                      // pre-breeders outside SA
233  pmat[7, 6] = no_visit == 1 ? 0 : p_detect_juv; // juvs
234  pmat[7, 7] = 0;                      // dead
235  pmat[7, 8] = 1 - pmat[7, 6];        // not seen
236
237  /* DEADS */
238  pmat[8, 1] = 0;                      // ad breeding in SA (SA = study area)
239  pmat[8, 2] = 0;                      // ad non-breeding in SA
240  pmat[8, 3] = 0;                      // ad outside SA
241  pmat[8, 4] = 0;                      // pre-breeders inside SA
242  pmat[8, 5] = 0;                      // pre-breeders outside SA
243  pmat[8, 6] = 0;                      // juvs
244  pmat[8, 7] = no_visit == 1 ? 0 : p_detect_dead; // dead
245  pmat[8, 8] = 1 - pmat[8, 7];        // not seen
246
247  return pmat;
248 }
249
250
251
252  real log_sum_one_indiv (int N_STATES, int sex, int[] age, int MAX_T, int first_cap, int last_cap,
253  int[] c_hist, real[] s_ad, real s_prebr, real s_juv, real[] p_moveout,
254  real[] p_movein, int[] b_success, vector p_breed, vector p_recruit,
255  vector p_beadult, real[] p_success, int N_STATES_P, real[,] p_obs,
256  real p_detect_juv, real p_detect_dead, real p_female, int[] NO_VISIT,
257  int first_state) {
258
259  matrix[N_STATES, N_STATES] tmat;
260  matrix[N_STATES_P, N_STATES_P] pmat;
261  vector[N_STATES] pz[MAX_T];
262  real temp[N_STATES];
263  real lsum;
264
265  for (j in 1:N_STATES) {
266    pz[first_cap, j] = (j == first_state);
267  }
268  for (t in (first_cap+1):last_cap) {
269    tmat = trans_probs(N_STATES, s_ad[sex+1, t-1], s_prebr, s_juv,
270                      p_moveout[sex+1], p_movein[sex+1], b_success[t-1],
271                      p_breed, p_recruit[age[t]], p_beadult[age[t]]),
272

```

```

272         p_success[t-1]);
273     pmat = obs_probs(N_STATES_P, p_obs[t-1], p_detect_juv, p_detect_dead, p_female,
274                      p_success[t-1], b_success[t-1], NO_VISIT[t]);
275     for (i in 1:N_STATES) {
276         for (j in 1:N_STATES) {
277             temp[j] = pz[t-1, j] * tmat[j, i] * pmat[i, c_hist[t]];
278         }
279         pz[t, i] = sum(temp);
280     }
281 }
282 lsum = log(sum(pz[last_cap]));
283
284 return lsum;
285
286 }
287
288
289 real calc_log_sum_multi (int[] INDS, int start, int end, int N_STATES, int[,] AGE,
290                           int MAX_T, int[] FIRST_CAP, int[] LAST_CAP, int[,] C_HIST,
291                           real[,] s_ad, real s_prebr, real s_juv,
292                           real[] p_moveout, real[] p_movein, int[,] B_SUCCESS, vector p_breed,
293                           vector p_recruit, vector p_beadult, real[] p_success,
294                           int N_STATES_P, real[,] p_obs, real p_detect_juv, real p_detect_dead, real p_female,
295                           int[] NO_VISIT, int[] FIRST_STATE) {
296     real lsum;
297
298     lsum = 0.0;
299     for (ind in start:end) {
300
301         lsum += log_sum_one_indiv(N_STATES, SEX[ind], AGE[ind], MAX_T, FIRST_CAP[ind], LAST_CAP[ind], C_HIST[ind],
302                                   s_ad, s_prebr, s_juv, p_moveout, p_movein,
303                                   B_SUCCESS[ind], p_breed,
304                                   p_recruit, p_beadult, p_success,
305                                   N_STATES_P, p_obs, p_detect_juv, p_detect_dead, p_female,
306                                   NO_VISIT, FIRST_STATE[ind]); //, ind);
307     }
308
309     return lsum;
310 }
311
312 }
313
314 }
315
316 data {
317
318     int<lower=1> N_INDS;
319     int<lower=1> INDS [N_INDS];
320     int<lower=1> FIRST_STATE [N_INDS];
321
322     int<lower=0, upper=2> SEX [N_INDS];
323     int<lower=1> N_SEXED;
324     int<lower=0, upper=1> IS_FEMALE [N_SEXED];
325
326     int<lower=1> N_NESTS;
327     int<lower=0, upper=1> NEST_SUCCESS [N_NESTS];
328     int<lower=1> NEST_YEAR [N_NESTS];
329
330     int<lower=1> FIRST_CAP [N_INDS];
331     int<lower=1> LAST_CAP [N_INDS];
332
333     int<lower=1> MAX_T;
334     int<lower=1> MAX_AGE;
335     int<lower=1> AGE[N_INDS, MAX_T];
336
337     int<lower=1, upper=MAX_AGE> MIN_R_AGE;
338     int<lower=MIN_R_AGE, upper=MAX_AGE> MAX_R_AGE;
339     int<lower=MIN_R_AGE, upper=MAX_AGE> MIN_B_AGE;
340     int<lower=MIN_B_AGE, upper=MAX_AGE> MIN_B_AGE2;
341
342     int<lower=0, upper=1> NO_VISIT[MAX_T];
343
344     int<lower=1> N_STATES;
345     int<lower=1> N_STATES_P;
346     int<lower=1> N_PDETECTS;
347     int<lower=1, upper=N_STATES_P> C_HIST[N_INDS, MAX_T];
348
349     int<lower=0, upper=2> B_SUCCESS[N_INDS, MAX_T];
350
351 }
352
353
354 transformed data {
355     int<lower=1> grainsize=1;
356 }
357
358
359 parameters{
360
361     real<lower=0, upper=1> p_female;
362

```

```

363 real<lower=0, upper=1> p_rec [MAX_R_AGE - MIN_R_AGE + 1];
364
365 real<lower=0, upper=1> p_br [MIN_B_AGE2 - MIN_B_AGE];
366 real<lower=0, upper=1> p_br_post;
367
368 vector<lower=0, upper=1>[2] p_breed; // 1: previously unsuccessful breeders; 2: other non-breeders
369
370 real<lower=0> sigma_re_bsucc;
371 real bsucc_lg_re [MAX_T];
372 real bsucc_lg_mean;
373
374 real<lower=0, upper=1> s_prebr;
375 real<lower=0, upper=1> s_juv;
376
377 /* Random effect on recruitment */
378 real<lower=0> sigma_re_rec;
379 real rec_lg_re [MAX_R_AGE - MIN_R_AGE + 1];
380 real rec_lg_mean;
381
382 /* Random effect on becoming adult */
383 real<lower=0> sigma_re_bead;
384 real bead_lg_re [MIN_B_AGE2 - MIN_B_AGE];
385 real bead_lg_mean;
386
387 /* Random effect on adult survival */
388 real<lower=0> sigma_re_ad_s;
389 real surv_ad_lg_re [2, MAX_T-1];
390 real surv_ad_lg_mean [2];
391
392 /* Random effect on detectability */
393 real<lower=0> sigma_re_p;
394 real p_detect_lg_re [MAX_T-1];
395 real p_detect_lg_mean [N_PDETECTS];
396 real<lower=0, upper=1> p_detect_juv;
397 real<lower=0, upper=1> p_detect_dead;
398
399 real<lower=0, upper=1> p_leave[2];
400 real<lower=0, upper=1> p_back[2];
401
402 }
403
404 transformed parameters {
405
406     vector<lower=0, upper=1>[MAX_AGE] p_recruit;
407     vector<lower=0, upper=1>[MAX_AGE] p_beadult;
408
409     real<lower=0, upper=1> s_adult [2, MAX_T-1];
410     real<lower=0, upper=1> s_ad[3, MAX_T-1];
411
412     real<lower=0, upper=1> p_detect [N_PDETECTS, MAX_T-1]; // 1: breeding ad (inside sa); 2: non-breeding ad previously
413     successful (inside sa); 3: other non-breeders (inside sa); 4: prebr inside SA; 5: ad or prebr outside SA
414
415     real<lower=0, upper=1> p_success [MAX_T];
416
417     real<lower=0, upper=1> p_moveout [3];
418     real<lower=0, upper=1> p_movein [3];
419
420     real<lower=0, upper=1> p_obs [MAX_T-1, N_PDETECTS];
421
422     /* Juvs becoming pre-breeders (recruitment to the colony) */
423     for (a in 1:(MIN_R_AGE-1)) {
424         p_recruit[a] = 0;
425     }
426     for (a in MIN_R_AGE:MAX_R_AGE) {
427         p_recruit[a] = inv_logit(rec_lg_mean + rec_lg_re[a - MIN_R_AGE + 1] * sigma_re_rec);
428     }
429     for (a in (MAX_R_AGE+1):MAX_AGE) {
430         p_recruit[a] = 1;
431     }
432
433     /* Pre-Breeders becoming adults (start breeding) */
434     for (a in 1:(MIN_B_AGE-1)) {
435         p_beadult[a] = 0;
436     }
437     for (a in MIN_B_AGE:(MIN_B_AGE2-1)) {
438         p_beadult[a] = inv_logit(bead_lg_mean + bead_lg_re[a - MIN_B_AGE + 1] * sigma_re_bead);
439     }
440     for (a in MIN_B_AGE2:MAX_AGE) {
441         p_beadult[a] = p_br_post;
442     }
443
444     for (t in 1:(MAX_T-1)) {
445         for (s in 1:N_PDETECTS) {
446             p_detect[s, t] = inv_logit(p_detect_lg_mean[s] + p_detect_lg_re[t] * sigma_re_p);
447         }
448         for (sex in 1:2) {
449             s_adult[sex, t] = inv_logit(surv_ad_lg_mean[sex] + surv_ad_lg_re[sex, t] * sigma_re_ad_s);
450         }
451     }
452 }
```

```

453   for (t in 1:MAX_T) {
454     p_success[t] = inv_logit(bsucc_lg_mean + bsucc_lg_re[t] * sigma_re_bsucc);
455   }
456
457   for (sex in 0:2) {
458     p_moveout[sex+1] = sex != 0 ? p_leave[sex] : p_female * p_leave[1] + (1-p_female) * p_leave[2];
459     p_movein[sex+1] = sex != 0 ? p_back[sex] : p_female * p_back[1] + (1-p_female) * p_back[2];
460     for (t in 1:(MAX_T-1)) {
461       s_ad[sex+1, t] = sex != 0 ? s_adult[sex, t] : p_female * s_adult[1, t] + (1-p_female) * s_adult[2, t];
462     }
463   }
464
465   for (s in 1:N_PDETECTS) {
466     for (t in 1:(MAX_T-1)) {
467       for (sex in 0:2) {
468         p_obs[t, s] = p_detect[s, t];
469       }
470     }
471   }
472
473
474 }
475
476
477 model {
478
479   matrix[N_STATES, N_STATES] tmat;
480   matrix[N_STATES_P, N_STATES_P] pmat;
481
482   real temp[N_STATES];
483
484   p_female ~ beta(1, 1);
485   IS_FEMALE ~ bernoulli(p_female);
486
487   /* Return to colony */
488   p_rec ~ beta(1, 1);
489
490   /* Becoming an adult (breeding for the first time) */
491   p_br ~ beta(1, 1);
492   p_br_post ~ beta(1, 1);
493
494   /* Probability of adult to breed (1: failed breeders; 2: non-breeders) */
495   p_breed ~ beta(1, 1);
496
497   /* Survival */
498   s_juv ~ beta(1, 1);
499   s_prebr ~ beta(1, 1);
500
501   /* Recruitment to colony */
502   sigma_re_rec ~ cauchy(0, 2);
503   rec_lg_re ~ normal(0, 1);
504   rec_lg_mean ~ normal(0, 2);
505
506   /* Becoming adult */
507   sigma_re_beard ~ cauchy(0, 2);
508   bead_lg_re ~ normal(0, 1);
509   bead_lg_mean ~ normal(0, 2);
510
511   /* P(successful breeding) */
512   for (n in 1:N_NESTS) {
513     NEST_SUCCESS[n] ~ bernoulli(p_success[NEST_YEAR[n]]);
514   }
515   sigma_re_bsucc ~ cauchy(0, 2);
516   bsucc_lg_re ~ normal(0, 1);
517   bsucc_lg_mean ~ normal(0, 2);
518
519   /* P(leaving/returning the study area) */
520   p_leave ~ beta(1, 1);
521   p_back ~ beta(1, 1);
522
523   /* Survival */
524   for (sex in 1:2) {
525     surv_ad_lg_mean[sex] ~ normal(0, 2);
526     for (t in 1:(MAX_T-1)) {
527       surv_ad_lg_re[sex, t] ~ normal(0, 1); // Time effect varies by sex
528     }
529   }
530   sigma_re_ad_s ~ cauchy(0, 2);
531
532   /* Detectability */
533   for (s in 1:N_PDETECTS) {
534     p_detect_lg_mean[s] ~ normal(0, 2);
535   }
536   p_detect_juv ~ beta(1, 1);
537   p_detect_dead ~ beta(1, 1);
538
539   // Same time effect for all classes and sexes (reflects changes in surveys)
540   for (t in 1:(MAX_T-1)) {
541     p_detect_lg_re[t] ~ normal(0, 1);
542   }
543   sigma_re_p ~ cauchy(0, 2);

```

```
544     /* Likelihood */
545     target += reduce_sum(calc_log_sum_multi, INDS, grainsize, N_STATES, SEX, AGE, MAX_T,
546                         FIRST_CAP, LAST_CAP, C_HIST,
547                         s_ad, s_prebr, s_juv, p_moveout, p_movein,
548                         B_SUCCESS, p_breed,
549                         p_recruit, p_beadult, p_success,
550                         N_STATES_P, p_obs, p_detect_juv, p_detect_dead, p_female,
551                         NO_VISIT, FIRST_STATE);
552
553 }
```

## APPENDIX B MODEL ESTIMATES

**Table B-1:** Annual survival rate of adults by year and sex, and of pre-breeders and juveniles. Shown are the mean, 95% credible interval (c.i.), and the MCMC trace of the parameter.

Year	Females			Males		
	Mean	95% c.i.	Trace	Mean	95% c.i.	Trace
1994	0.961	0.924 – 0.987		0.959	0.919 – 0.986	
1995	0.970	0.944 – 0.989		0.963	0.933 – 0.986	
1996	0.951	0.917 – 0.977		0.936	0.901 – 0.966	
1997	0.950	0.916 – 0.975		0.950	0.916 – 0.978	
1998	0.924	0.884 – 0.959		0.936	0.900 – 0.965	
1999	0.923	0.881 – 0.959		0.917	0.879 – 0.949	
2000	0.941	0.901 – 0.972		0.941	0.908 – 0.968	
2001	0.935	0.898 – 0.965		0.951	0.924 – 0.975	
2002	0.951	0.925 – 0.973		0.949	0.921 – 0.973	
2003	0.975	0.954 – 0.990		0.965	0.942 – 0.983	
2004	0.941	0.906 – 0.971		0.936	0.907 – 0.960	
2005	0.898	0.806 – 0.969		0.941	0.889 – 0.980	
2006	0.878	0.789 – 0.963		0.933	0.883 – 0.978	
2007	0.883	0.822 – 0.936		0.897	0.853 – 0.935	
2008	0.845	0.776 – 0.911		0.920	0.879 – 0.954	
2009	0.894	0.832 – 0.947		0.914	0.874 – 0.949	
2010	0.906	0.840 – 0.960		0.926	0.889 – 0.958	
2011	0.841	0.770 – 0.906		0.937	0.901 – 0.969	
2012	0.829	0.760 – 0.891		0.911	0.872 – 0.943	
2013	0.821	0.752 – 0.883		0.936	0.902 – 0.963	
2014	0.929	0.876 – 0.970		0.950	0.918 – 0.977	
2015	0.848	0.781 – 0.909		0.914	0.873 – 0.949	
2016	0.871	0.803 – 0.930		0.913	0.871 – 0.949	
2017	0.937	0.884 – 0.977		0.924	0.879 – 0.959	
2018	0.901	0.832 – 0.956		0.916	0.871 – 0.952	
2019	0.908	0.839 – 0.963		0.938	0.894 – 0.972	
2020	0.929	0.861 – 0.976		0.971	0.943 – 0.991	

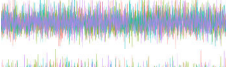
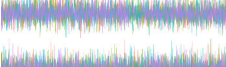
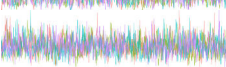
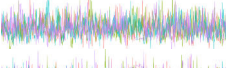
  

Age class	Mean	95% c.i.	Trace
Pre-breeders	0.922	0.913 – 0.931	
Juveniles	0.879	0.869 – 0.888	

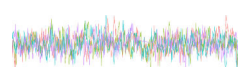
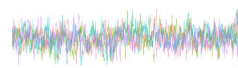
**Table B-2:** Probabilities of successful breeding by year. Shown are the mean, 95% credible interval (c.i.), and the MCMC trace of the parameter.

Parameter	Year	Mean	95% c.i.	Trace
P(successful breeding)	1994	0.74	0.67 – 0.81	
	1995	0.74	0.67 – 0.79	
	1996	0.75	0.68 – 0.81	
	1997	0.77	0.71 – 0.83	
	1998	0.73	0.66 – 0.79	
	1999	0.64	0.56 – 0.71	
	2000	0.75	0.67 – 0.81	
	2001	0.75	0.69 – 0.81	
	2002	0.67	0.60 – 0.74	
	2003	0.72	0.66 – 0.77	
	2004	0.71	0.65 – 0.77	
	2005	0.69	0.53 – 0.83	
	2006	0.67	0.48 – 0.81	
	2007	0.60	0.52 – 0.67	
	2008	0.67	0.60 – 0.73	
	2009	0.56	0.47 – 0.65	
	2010	0.67	0.58 – 0.74	
	2011	0.57	0.48 – 0.66	
	2012	0.58	0.49 – 0.67	
	2013	0.61	0.51 – 0.69	
	2014	0.70	0.62 – 0.77	
	2015	0.57	0.47 – 0.66	
	2016	0.66	0.57 – 0.74	
	2017	0.75	0.65 – 0.83	
	2018	0.65	0.57 – 0.73	
	2019	0.61	0.53 – 0.69	
	2020	0.62	0.52 – 0.71	
	2021	0.67	0.48 – 0.81	

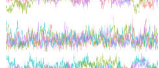
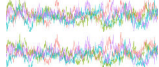
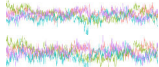
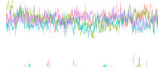
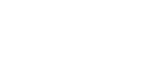
**Table B-3:** Probabilities of returning to the colony and to breed for the first time, as function of age. Shown are the mean, 95% credible interval (c.i.), and the MCMC trace of the parameter.

Parameter	Age	Mean	95% c.i.	Trace
P(return to colony)	3	0.03	0.02 – 0.04	
	4	0.11	0.09 – 0.14	
	5	0.10	0.07 – 0.14	
	6	0.29	0.24 – 0.34	
	7	0.33	0.25 – 0.42	
	8	0.69	0.56 – 0.83	
P(breed for first time)	7	0.02	0.01 – 0.04	
	8	0.06	0.04 – 0.09	
	9	0.07	0.05 – 0.10	
	10	0.06	0.04 – 0.09	
	11	0.10	0.07 – 0.14	
	12	0.13	0.09 – 0.18	
	13	0.12	0.08 – 0.17	
	14	0.13	0.09 – 0.18	
	15	0.07	0.04 – 0.12	
	16	0.13	0.08 – 0.19	
	17	0.10	0.05 – 0.16	
	18	0.09	0.04 – 0.15	
	19	0.12	0.06 – 0.20	
	20	0.05	0.02 – 0.10	

**Table B-4:** Probabilities of adults breeding, that an individual is female, that a bird inside the study area move outside it, and probability that a bird outside the study area returns inside. Shown are the mean, 95% credible interval (c.i.), and the MCMC trace of the parameter.

Parameter	Category	Mean	95% c.i.	Trace
P(breeding)	Previously unsuccessful breeders	0.70	0.69 – 0.72	
	Other non-breeders	0.64	0.63 – 0.65	
P(female)		0.51	0.49 – 0.53	
P(leave the study area)	Female	0.09	0.08 – 0.10	
	Male	0.04	0.04 – 0.05	
P(return to the study area)	Female	0.18	0.15 – 0.20	
	Male	0.25	0.22 – 0.29	

**Table B-5:** Detection probabilities: annual averages, year effect, and interannual variability, as well as the time-invariant detection probabilities of juveniles and dead birds. Shown are the mean, 95% credible interval (c.i.), and the MCMC trace of the parameter.

Parameter	Category	Mean	95% c.i.	Trace
P(detection) - overall	Breeding adult (inside SA)	0.864	0.816 – 0.900	
	Non-breeding adult (inside SA)	0.052	0.036 – 0.072	
	Other non-breeders (inside SA)	0.997	0.992 – 1.000	
	Pre-breeders (inside SA)	0.661	0.575 – 0.736	
	Adults and pre-breeders outside SA	0.180	0.132 – 0.234	
Year effect (logit scale)	1995	-1.388	-2.214 – -0.679	
	1996	1.014	0.471 – 1.605	
	1997	1.529	0.915 – 2.168	
	1998	0.711	0.189 – 1.294	
	1999	0.596	0.058 – 1.152	
	2000	0.238	-0.282 – 0.766	
	2001	0.475	-0.028 – 1.007	
	2002	1.052	0.494 – 1.657	
	2003	1.656	1.044 – 2.305	
	2004	1.475	0.901 – 2.078	
	2005	1.139	0.612 – 1.692	
	2006	0.017	-1.792 – 1.849	
	2007	-1.887	-2.654 – -1.229	
	2008	-0.491	-0.973 – -0.039	
	2009	-0.492	-0.982 – -0.034	
	2010	-0.394	-0.859 – -0.050	
	2011	-1.019	-1.593 – -0.507	
	2012	-0.464	-0.955 – -0.027	
	2013	0.004	-0.457 – -0.462	
	2014	0.329	-0.156 – -0.816	
	2015	-0.383	-0.885 – -0.124	
	2016	-0.395	-0.931 – -0.100	
	2017	-0.079	-0.585 – -0.424	
	2018	0.229	-0.302 – -0.726	
	2019	-0.021	-0.527 – -0.484	
	2020	-1.763	-2.506 – -1.113	
	2021	0.018	-0.536 – -0.565	
Inter-annual variability		0.858	0.638 – 1.169	
Parameter		Mean	95% c.i.	Trace
P(detection) - Juveniles		0.0002	0.0000 – 0.0007	
P(detection) - Dead birds		0.0008	0.0005 – 0.0012	