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Managing the long-term persistence of a rare cockatoo under climate change

J. Berton C. Harris¹*, Damien A. Fordham¹, Patricia A. Mooney², Lynn P. Pedler², Miguel B. Araújo^{3,4}, David C. Paton¹, Michael G. Stead¹, Michael J. Watts¹, H. Reşit Akçakaya⁵ and Barry W. Brook¹

¹School of Earth and Environmental Sciences, University of Adelaide, SA 5005, Australia; ²Glossy Black-Cockatoo Recovery Program, Department for Environment and Heritage, Kingscote, SA 5223, Australia; ³Department of Biodiversity and Evolutionary Biology, National Museum of Natural Sciences, CSIC, C/José Gutierrez Abascal, 2, Madrid 28006, Spain; ⁴Rui Nabeiro Biodiversity Chair, CIBIO, University of Évora, Largo dos Colegiais, 7000 Évora, Portugal; and ⁵Department of Ecology and Evolution, Stony Brook University, Stony Brook, NY 11794, USA

Summary

1. Linked demographic-bioclimatic models are emerging tools for forecasting climate change impacts on well-studied species, but these methods have been used in few management applications, and species interactions have not been incorporated. We combined population and bioclimatic envelope models to estimate future risks to the viability of a cockatoo population posed by climate change, increased fire frequency, beak-and-feather disease and reduced management.

2. The South Australian glossy black-cockatoo *Calyptorhynchus lathami halmaturinus* is restricted to Kangaroo Island, Australia, where it numbers 350 birds and is managed intensively. The cockatoo may be at particular risk from climate change because of its insular geographic constraints and specialised diet on a single plant species, *Allocasuarina verticillata*. The cockatoo population model was parameterised with mark-resight-derived estimates of survival and fecundity from 13 years of demographic data. Species interactions were incorporated by using a climate-change-driven bioclimatic model of *Allocasuarina verticillata* as a dynamic driver of habitat suitability. A novel application of Latin Hypercube sampling was used to assess the model's sensitivity to input parameters.

3. Results suggest that unmitigated climate change is likely to be a substantial threat for the cockatoo: all high-CO₂-concentration scenarios had expected minimum abundances of < 160 birds. Extinction was virtually certain if management of nest-predating brush-tail possums *Trichosurus vulpecula* was stopped, or adult survival reduced by as little as 5%. In contrast, the population is predicted to increase under low-emissions scenarios.

4. Disease outbreak, increased fire frequency and reductions in revegetation and management of competitive little corellas *Cacatua sanguinea*, were all predicted to exacerbate decline, but these effects were buffered by the cockatoo population's high fecundity.

5. Spatial correlates of extinction risk, such as range area and total habitat suitability, were nonlinearly related to projected population size in the high- CO_2 -concentration scenario.

6. *Synthesis and applications.* Mechanistic demographic-bioclimatic simulations that incorporate species interactions can provide more detailed viability analyses than traditional bioclimatic models and be used to rank the cost-effectiveness of management interventions. Our results highlight the importance of managing possum predation and maintaining high adult cockatoo survival. In contrast, corella and revegetation management could be experimentally reduced to save resources.

Key-words: beak-and-feather disease, bioclimatic envelope, *Calyptorhynchus lathami*, climate change, glossy black-cockatoo, management, population viability analysis, revegeta-tion, wildfire

*Correspondence author. E-mails: aramidopsis@gmail.com; bert. harris@adelaide.edu.au

Introduction

Climate change may be one of the most potent extinction drivers in the future, especially because it can exacerbate existing threats, and there is an urgent need for conservation science to improve tools to predict species' vulnerability to climate change (Sekercioglu et al. 2008). One popular approach is the use of bioclimatic envelope models (BEMs), also known as species distribution models. These models use associations of present-day distributions with climate to forecast changes in species' bioclimatic envelopes (Pearson & Dawson 2003). BEMs have, in some cases, been used to assess extinction risk for thousands of species under climate change scenarios (e.g. Sekercioglu et al. 2008). However, predictions from these models are of constrained value because they (i) are correlative and yet typically require extrapolation to environmental space that is beyond the bounds of the statistical fitting (Thuiller et al. 2004); (ii) use range area type estimates to infer extinction risk rather than measuring threat to population persistence (Fordham et al. 2011); (iii) suffer from model selection uncertainty (Araújo & Rahbek 2006); and (iv) do not consider biotic interactions (e.g. Araújo & Luoto 2007).

Spatially explicit population-modelling techniques that link demographic models with BEMs are being used to add ecological realism to correlative BEM forecasts (Huntley *et al.* 2010). Combining quantitative population models and BEMs provides a more mechanistic and probabilistic approach compared to modelling distribution alone, because it links demographic parameters to climate and other explanatory variables and explores a range of uncertain outcomes using stochastic simulation (Brook *et al.* 2009). Several studies have combined habitat and population models to assess population viability (e.g. Akçakaya *et al.* 2004), but few analyses have coupled population and bioclimatic models to estimate extinction risk in the context of climate change (Keith *et al.* 2008; Anderson *et al.* 2009; Fordham *et al.* in press), and this methodology has rarely been used in birds (but see Aiello-Lammens *et al.* 2011). Ideal case-study species for this approach are those with long-term estimates of vital rates (and their variance), representative occurrence data over their geographic range and detailed knowledge of the environmental drivers influencing range and abundance.

The South Australian glossy black-cockatoo Calvptorhynchus lathami halmaturinus Temminck (GBC) formerly inhabited mainland South Australia, but now survives only on Kangaroo Island (located off the southern coast of central Australia) and is considered 'endangered' by the Australian government (DEH 2000; Fig. 1). When the GBC recovery program began in 1995, the cockatoo population comprised c. 200 individuals. From 1998 to the present, the intensively managed population has increased gradually to the current estimate of c. 350 birds (Pedler & Sobey 2008). The GBC's specialised habitat requirements and slow life history make it inherently vulnerable to decline (Cameron 2006), and its small population size and insular geographic constraints (single location) put it at high risk from population-wide catastrophes such as fire and disease (Pepper 1997). High-quality Allocasuarina verticillata Lam. L.A.S. Johnson, drooping

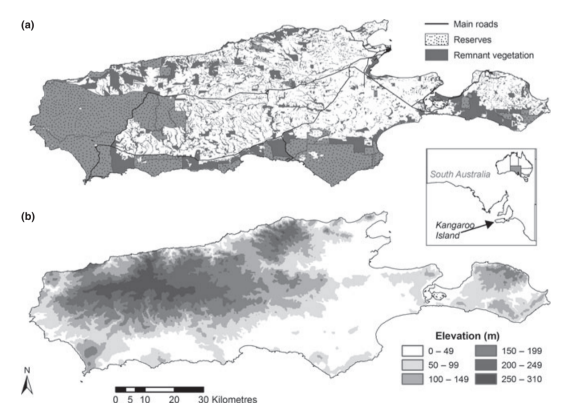


Fig. 1. The South Australian glossy black-cockatoo *Calyptorhynchus lathami halmaturinus* is restricted to Kangaroo Island, South Australia. Maps showing (a) remnant native vegetation and protected areas and (b) elevation.

The GBC faces an interacting set of current and future threats including nest competition and predation, wildfire, climate change and disease (Mooney & Pedler 2005). GBC recruitment can be severely impaired by nest predation from arboreal brush-tail possums Trichosurus vulpecula Kerr. Protecting nest trees from possum predation by fitting metal collars and pruning adjacent tree crowns increased nest success from 23 to 42% (Garnett, Pedler & Crowley 1999). Approximately, 45% of nests are now placed in artificial hollows fitted by managers. Little corellas Cacatua sanguinea Gould and honeybees Apis mellifera L. are nest competitors that are also managed (Mooney & Pedler 2005). Wildfires are another threat that can kill nestlings and destroy large areas of habitat (Sobey & Pedler 2008). Kangaroo Island is expected to warm by 0.3-1.5 °C and receive 0-20% less rainfall by 2050 compared to 1990 levels, under a mid-range greenhouse-gas emissions scenario (CSIRO 2007). Climate change is likely to threaten the GBC by causing A. verticillata's climatic niche to shift and compress southwards towards the southern ocean boundary (Stead 2008), causing heat- and drought-induced mortality (Cameron 2008) and an increased frequency of extreme events, such as fire and drought (Dunlop & Brown 2008). In addition, A. verticillata cone production may decrease as conditions become warmer and drier (D. C. Paton personal observations), limiting the GBC's food supply. Lastly, psittacine beak-and-feather disease, although not yet reported in Kangaroo Island GBCs, could potentially cause substantial declines in the population if an outbreak occurred (DEH 2005; Appendix S5).

Here, we develop a detailed spatial population viability model for the GBC by building a demographic model, linking the demographic model to landscape and climate variables and testing scenarios in a population viability analysis. The analysis is based on a comprehensive location-specific data set and incorporates climate change and its interaction with fire, disease and management. Two earlier attempts at modelling the GBC used non-spatial simulations to investigate extinction risk (Pepper 1996; Southgate 2002), but both were limited in scope and made simplifying assumptions. For instance, in contrast to known population increases, Pepper (1996) predicted a rapid decline to extinction, and Southgate (2002) suggested the population would decline by 10% annually (Appendix S1). These studies were hampered by the limited data available when the analyses were performed and did not consider fire, disease, climate change or the positive influence of management. By contrast, we use a detailed data set collected by the GBC recovery program since 1995, consisting of 13 years of mark-resight and reproductive data and extensive documentation of catastrophes and

management intervention, to parameterise our models. Few parrots have such complete demographic data available (Snyder *et al.* 2004).

Our approach incorporates a critical biotic interaction between the GBC and its primary food source, A. verticillata, by incorporating projected changes in the plant's range in the spatially explicit cockatoo model to provide direct measures of extinction threat (e.g. expected minimum abundance) as well as implied measures calculated from changes in habitat suitability and range size (Fordham et al. 2011). Similar approximations of species interactions have been used with BEMs (e.g. Araújo & Luoto 2007; Barbet-Massin & Jiguet 2011), but never in combination with a demographic model. Specifically, we sought to (i) model the population trajectory and extinction risk of the GBC up to the year 2100; (ii) determine the possible future effects of current and emerging threats to the subspecies; (iii) assess the impact of choosing different management strategies on GBC population trends; and (iv) evaluate the relative importance of demography and anthropogenic extinction drivers on the GBC's population viability.

Materials and methods

POPULATION MODEL

For the demographic component of the model, we used 13 years of mark-resight surveys to estimate survival rates using program MARK v.5.1 (Cooch & White 2008). Birds are marked with numbered bands as nestlings at several sites across the island (some areas are better sampled than others), and telescopes are used to resight marked birds during the annual post-breeding census. The markresight analysis was used to test the importance of management and environmental variables on survival rates of juvenile (<1 year old) and sub-adult/adult GBCs (Table S1). Fecundity was calculated as the number of fledglings of each sex produced per female of breeding age from 1996 to 2008 (see Appendix S2 for details on the markresight analysis, fecundity calculations and standard deviations used in the population model). Survival and fecundity estimates were combined with other life-history information, such as age of first breeding, to build a stage- and sex-structured, stochastic population model of the GBC (Table 1). We used RAMAS GIS (Akçakaya & Root 2005) to create a spatially explicit metapopulation model that links the subspecies' demography to landscape data, comprising dynamic bioclimatic maps for Allocasuarina verticillata (the GBC's primary food source), and raster layers of native vegetation, substrate and slope (see below).

BIOCLIMATIC SUITABILITY MAPS FOR ALLOCASUARINA VERTICILLATA

Climate change was incorporated by modelling the potential distribution of *Allocasuarina verticillata*, as a function of three key climate variables that influence the species' distribution (annual rainfall, January temperature and July temperature; Stead 2008). We used meteorological data to estimate long-term average annual rainfall and mean monthly January and July temperature (1980–1999) for Australia (Fordham, Wigley & Brook 2012). We used thin-plate splines and a digital elevation model to interpolate between weather stations (Hutchinson 1995; Appendix S3). An annual time series of climate change layers was generated for each climate variable based

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	Age 0	Age 1	Age 2+		SAD(%)	
Female						
Age 0	0	0	0.2324 (0.09	951)	7.3	
Age 1	0.612 (0.0951)	0	0	,	4.3	
Age 2+	0	0.913 (0.0951)	0.913 (0.714	18)	32.4	
	Age 0	Age 1	Age 2	Age 3	Age 4+	SAD(%)
Male						
Age 0	0	0	0	0	0*	9.3
Age 1	0.612	0	0	0	0	5.5
Age 2	0	0.913	0	0	0	4.9
Age 3	0	0	0.913	0	0	4.3
Age 4+	0	0	0	0.913	0.913	32.0

Table 1. Stage matrices used in the model with stable age distribution (SAD) of each age class

The top row in each matrix represents fecundities, and the subdiagonal and diagonal in the bottom right elements represent survival rates. The first stage (age 0) for both sexes is the sub-adult stage. The final stages (female, age 2+; male, age 4+) are the adult stages. The intermediate stages are pre-breeding sub-adult stages. The proportional sensitivities of the finite rate of increase to small changes in each of the non-zero elements of the female matrix (elasticities) are in parentheses.

*In RAMAS, we specified fecundity values of 0.2324 and 0.296 for females and males, respectively (Appendix S2).

on two emission scenarios: a high-CO₂-concentration stabilisation reference scenario, WRE750, and a strong greenhouse gas mitigation policy scenario, LEV1 (Wigley *et al.* 2009). WRE750 assumes that atmospheric CO₂ will stabilize at about 750 parts per million (ppm), whilst under the LEV1 intervention scenario CO₂ concentration stabilizes at about 450 ppm. Future climate layers were created by first generating climate anomalies from an ensemble of nine general circulation models and then downscaling the anomalies to an ecologically relevant scale (*c.* 1 km² grid cells) (Fordham, Wigley & Brook 2012; Fordham *et al.* 2012; Appendix S3). Averages from multiple climate models tend to agree better with observed climate compared to single climate models, at least at global scales (Fordham, Wigley & Brook 2012).

Occurrence records for A. verticillata (n = 572) came from cleaned records from the South Australian biological survey. An equal number of pseudoabsences were generated randomly within the study region (Appendix S3). Although our focus was on Kangaroo Island, we modelled the distribution of the species across South Australia (325,608 grid cells) to better capture its regional niche (see Barbet-Massin, Thuiller & Jiguet 2010). We modelled the potential current and future climatic suitability of the landscape for A. verticillata with an ensemble of seven bioclimatic modelling techniques, including simple surface-range envelope models and more complex machine learning approaches, in BIOENSEMBLES software (Diniz-Filho et al. 2009; Appendix S3). Ensemble modelling generates consensus projections that circumvent some of the problems of relying on single-model projections of climate change impacts on species' potential distributions (Araújo & New 2007). We used BIOENSEMBLES models to forecast annually for 90 years (i.e. climate suitability maps for each year were created from 2010 to 2100). Nonetheless, our model assumed that the A. verticillata-GBC relationship would remain strong and we were unable to consider other species interactions.

INTEGRATING THE POPULATION MODEL AND SPATIAL INFORMATION

Binomial generalised linear models (GLMs) were used to relate GBC occurrence records to *A. verticillata* present-day climate suitability

(above) and three landscape variables that are known to influence the distributions of the GBC and *A. verticillata*: substrate (Raymond & Retter 2010), native vegetation cover (http://www.environment.gov.au/erin/nvis/index.html), and slope (http://www.ga.gov.au/meta/ANZCW0703011541.html; Appendix S4). Verified GBC occurrence records (n = 349) consist of presences only. Pseudoabsences were generated by down-weighting cells close to a known sighting (Appendix S4). The analysis was performed with package MuMIn (Bartoń 2012) in R (v. 2.12.1; R Development Core Team, http://www.R-project.org). The best model (determined by AIC_c) from this analysis was used to parameterise the habitat-suitability function in RAMAS (Appendix S4).

RAMAS uses the habitat-suitability function to assign a habitatsuitability value to each grid cell of the study area based on values of the input rasters (in this case *A. verticillata* climatic suitability, substrate, native vegetation and slope). Every grid cell above the habitatsuitability threshold is considered suitable, and suitable cells are aggregated based on neighbourhood distance (the spatial distance at which the species can be assumed to be panmictic; Akçakaya & Root 2005). The habitat-suitability threshold (0.83) and neighbourhood distance (four cells) values were derived iteratively to match the wellknown current extent of suitable habitat for the GBC on the island (Mooney & Pedler 2005).

The initial population size in all scenarios was 350 birds, in accordance with recent estimates (Pedler & Sobey 2008). The island's current carrying capacity was estimated at 653 birds by combining feeding habitat requirements (Chapman & Paton 2002) with data on A. verticillata area (Appendix S4). Dispersal estimates came from data on movements of marked individuals (Fig. S1). A ceiling model of density dependence was used to approximate the GBC's intraspecific competition for nest hollows and feeding habitat (Mooney & Pedler 2005). Population dynamics were linked to habitat via the density dependence function: habitat determines carrying capacity that conditions demographic rates (survival and fecundity) in each year, as a function of population size and carrying capacity in that year (Akçakaya & Root 2005). Each simulation incorporated environmental and demographic stochasticity and was run 10 000 times (Akçakaya et al. 2004).

Our main measures of population viability were expected minimum abundance (EMA) and mean final population size of persisting runs. EMA, which is equivalent to the area under the quasi-extinction risk curve (McCarthy 1996), provides a better (continuous, unbounded) representation of extinction risk than probability of extinction or quasi-extinction (McCarthy & Thompson 2001). We calculated EMA by taking the smallest population size observed in each iteration and averaging these minima.

We also calculated three spatial measures that are commonly used to infer extinction likelihood: change in total habitat suitability (from RAMAS), occupied range area (area of cells greater than habitatsuitability threshold) and average cockatoo density (see Fordham *et al.* 2011 for details). Density was calculated by relating the population size at each time step to habitat-suitability values per grid cell in suitable patches.

MODEL SCENARIOS

We generated RAMAS models for three climate scenarios: WRE750, LEV1 and a control scenario with no climate change. For each climate scenario, we assessed GBC population viability given changes in fire frequency, disease outbreak and changes in management from funding constraints. We modelled severe fires as reducing GBC fecundity by 10% and adult and sub-adult survival by 3%, based on responses measured in 2007 (Sobey & Pedler 2008; P.A. Mooney personal communication). Wildfire frequency was modelled as increasing with building fuel loads. Baseline scenarios include an annual probability of severe fire of 6.8% (Appendix S5). We modelled 5%, 25% and 220% (i.e. 2·2-fold) increases in fire frequency under climate change (Lucas et al. 2007). It was not realistic to model any fire increases for the no climate change scenario or the 25% or 220% increase for the mitigation LEV1 scenario (Appendix S5). Psittacine beak-andfeather-disease outbreaks were modelled as reducing sub-adult survival by 50%, with an annual probability of an outbreak of 5% (DEH 2005; Appendix S5). We modelled ending brush-tail possum, little corella and revegetation management as causing 44%, 7% and 3% reductions in fecundity, respectively (Mooney & Pedler 2005).

SENSITIVITY ANALYSIS

We used a Latin Hypercube sensitivity analysis to assess the impact of varying the values of six key input parameters (adult survival, varied by $\pm 5\%$; sub-adult survival, $\pm 10\%$; fecundity, $\pm 10\%$; carrying capacity, $\pm 20\%$; and proportion of population dispersing annually, $\pm 20\%$) on GBC mean final population size (Iman, Helson & Campbell 1981). Latin Hypercube sampling, which simultaneously varies the values of the input parameters and then estimates sensitivity by fitting a spline regression model, is arguably preferable to other Monte Carlo techniques because it requires many fewer iterations to sample the parameter space whilst allowing for co-variation in parameter choices (McKay, Beckman & Conover 1979). We fit a Poisson GLM with all six predictors (a segmented linear model was used for adult survival; segmented package in R; Appendix S5) and calculated standardised regression coefficients (fitted slopes divided by their standard errors) to rank the importance of the input parameters (Conroy & Brook 2003). We also tested the model's sensitivity to parameterisation of disease outbreaks by doubling the frequency of simulated outbreaks, increasing the impact to a 75% reduction in survival and combining these parameterisations.

Results

DEMOGRAPHY

The best-supported mark-resight survival model was stagestructured and time invariant (Table S2). There was also statistical support for the next eight models ($\Delta \text{AIC}_c < 2$), yet the majority of model structural deviance was explained by the most parsimonious model (88% compared to 99%). The annual survival estimates so derived were 0.612 ± 0.0388 SE for juveniles and 0.913 ± 0.0123 SE for adults. All of the topranked 10 survival models incorporated stage structure with two age classes. There was little evidence for differences in survival between the sexes over the study period from the markresight data. Models including environmental covariates were suboptimal regardless of stage structure. All covariate models with no stage structure had wAIC_c < 0.01.

We used a mean annual fecundity estimate of 0.232 ± 0.0053 SE female nestlings produced per female of breeding age, and 0.296 ± 0.0068 SE male nestlings produced per female of breeding age, from 1996 to 2008, such that the finite rate of increase in the resultant matrix model was 1.0345, indicating a population increasing deterministically by 3.5% per year (Table 1; Appendix S2). The elasticities suggest that the rate of increase is most sensitive to adult survival.

SPATIAL RESULTS

There was considerable overlap between *Allocasuarina verticillata* patches and GBC presences. Approximately, 32% of GBC presences (feeding, nesting and band observations) were inside an *A. verticillata* patch, and 79% of presences were within 1 km of an *A. verticillata* patch (only 19% of the island is within 1 km of a patch).

The bioclimatic envelope modelling predicts that most of A. verticillata's range (and consequently the GBC's habitat) will remain intact under the reduced emissions (LEV1) scenario, whilst the range is likely to contract substantially under the high-CO₂-concentration scenario (WRE750) (Fig. 2). The majority of suitable habitat that is predicted to remain at the end of the century under the WRE750 emissions scenario is on the island's higher-elevation western plateau (Figs 1, 2). By 2100, total habitat suitability declined substantially (decreasing by 12%) in the WRE750 scenario, whereas suitability decreased by just 1% under LEV1 (Fig. 3). Range area was inversely related to average cockatoo density per cell (Fig. 3). This was especially evident for WRE750, where range area contracted by 77% and predicted density increased by 57% by 2100. Range area declined by only 6% in the LEV1 scenario (Fig. 3).

POPULATION VIABILITY

Habitat changes caused by unmitigated climate change had a strong effect on population viability, with simulated final population size and expected minimum abundance always < 160 birds, which is roughly equivalent to a return to the population

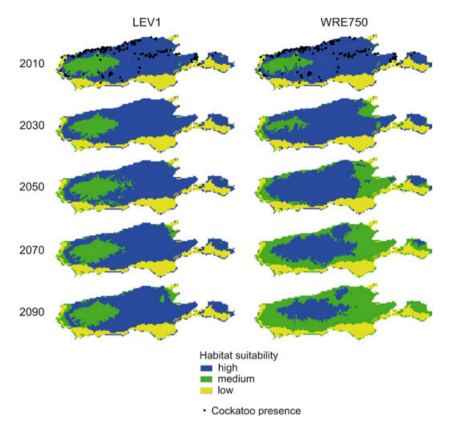


Fig. 2. Climate-change-driven maps of habitat suitability for *Calyptorhynchus lathami halmaturinus* according to a greenhouse gas mitigation policy scenario (LEV1) and a high- CO_2 -concentration stabilisation reference scenario (WRE 750). Recent cockatoo presences are shown on the 2010 maps. Habitat suitability is classified from a continuous variable into three categories to aid visual interpretation: high (above the habitat-suitability threshold), medium (below threshold) and low (unsuitable substrate for *A. verticillata*) suitability.

bottleneck of the 1980s (Fig. 4, Fig. S2). In contrast, all simulations in the no climate change (control) case had final population sizes >635, and EMA >350, unless brush-tail possum management ceased. The strong mitigation (LEV1) simulations had slightly lower final population sizes than the no climate change case, but still had all final population sizes > 595 unless there was no possum management. The simulations predicted that stopping possum management would have a serious effect on the population with all EMAs below 90 birds. Scenarios that ceased possum management were the only cases when the population did not stay close to carrying capacity. Unlike all other scenarios, possum scenarios had considerable probabilities of quasi-extinction (falling below 50 individuals): 10% for no climate change, 11% for LEV1 and 36% for WRE750. Stopping all management actions caused severe declines, with EMAs <26 birds for each scenario. The other catastrophes and changes in management had much more minor effects compared to possum management, although they did impact the population in the hypothesised directions (e.g. increased fire management caused slightly higher population sizes in LEV1 and no climate change). In this group of scenarios, beak-andfeather-disease outbreak had the strongest effects, but still only resulted in final population size reductions of 13, 12 and 1, compared to the baseline for no climate change, LEV1 and WRE750, respectively.

SENSITIVITY ANALYSIS

The Latin Hypercube sensitivity analysis indicated that model results were most heavily influenced by parameterisation of adult survival (top-ranked in each climate scenario) and carrying capacity (ranked second in each scenario; Fig. 5; Table S4). The standardised regression coefficients show that adult survival (low + high values from the segmented model) accounted for 35% (WRE750) to 52% (no climate change) of total sensitivity, whilst carrying capacity accounted for 21-32% of total sensitivity, respectively (Table S4). Decreased adult survival resulted in severe declines in GBC final population size, whilst increased adult survival had only slight or moderate effects because the modelled population, with the current survival estimate of 0.913, tracks carrying capacity with a positive population growth rate. Accordingly, varying carrying capacity also had substantial effects on final population size, especially for the WRE750 scenario where range area declines sharply. The other input parameters had small effects with sub-adult survival, fecundity and dispersal listed in order of decreasing importance. The additional disease outbreak sensitivity analysis indicated that increasing disease frequency or impact did not have substantially different effects on the population unless they were combined in the same scenario (Table S5).

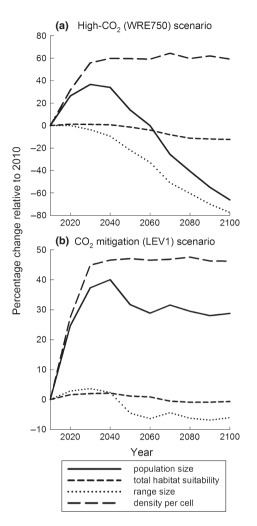


Fig. 3. Percentage changes in total habitat suitability (output from RAMAS GIS), range size (area of suitable habitat), cockatoo density per cell and population size according to two climate change scenarios: (a) high-CO₂-concentration stabilisation reference scenario (WRE750), (b) greenhouse gas mitigation policy scenario (LEV1).

Discussion

The population viability analysis for the South Australian glossy black-cockatoo illustrates the type of applied management questions that can be addressed using coupled demographic-bioclimatic approaches, as well as a method for incorporating dynamic vegetation-driven habitat change into animal population forecasts. The modelling indicates that the outlook for this small population depends strongly on continuous funding for management and global efforts to mitigate CO₂ emissions. The simulations suggest that GBC population size will increase under a low emissions future even if disease outbreaks were to occur, most management actions were reduced, and fire frequency was to increase. The gradual increase in the population over the last 15 years, combined with the large stands of underused Allocasuarina verticillata on the island, shows the potential for continued population growth. In contrast, a failure to mitigate CO₂ emissions could severely reduce GBC range area, critically threatening long-term population viability. Regardless of emissions scenario, our predictions indicate that the GBC's insular geographic constraints and low population size, which is wellbelow estimates of minimum viable estimates for most species (Traill *et al.* 2010), may leave the species vulnerable to decline.

Climate change under high CO₂ emissions (WRE750) caused a large reduction in range area, and contraction to the cooler and wetter western plateau, whilst habitat changes under low emissions (LEV1) were minimal, with range area decreasing modestly and habitat suitability remaining almost constant. Under high emissions, population size did not decrease as rapidly as range area because habitat suitability and cockatoo density initially increased in the remaining habitat (Fig. 3). These results indicate that range area is unlikely to be linearly related to GBC abundance. Habitat differences translated into much lower EMA for all high emissions scenarios compared to low emissions and no climate change. A population of 150 animals is inherently at risk of extinction from stochastic small-population processes (Traill et al. 2010). We did not run simulations beyond 2100 because of uncertainty in climate projections, but such small population sizes at the end of the century do not bode well for the GBC's persistence under a high-CO₂-concentration scenario.

Simulating reduced brush-tail possum management had a profound impact on GBC EMA, whilst reduction in little corella management was almost negligible because of the resilient GBC population. The absence of a strong response to corella management indicates that culling could be experimentally stopped in some areas in an adaptive management framework to save resources. Simulated psittacine beak-andfeather-disease outbreaks also had only slight effects on the GBC population. If mortality rates become higher and outbreak frequency is increased, disease could become a potent threat (Table S5). We suggest that continued vigilance and communication with organisations involved with disease management in other threatened parrots (e.g. orange-bellied parrot *Neophema chrysogaster* Latham) is needed.

Our results indicate that revegetation is only having small effects on the population at present, but altered spatial patterns of A. verticillata abundance from climate change and the carrying capacity of 653 individuals will probably necessitate revegetation in the future. Our model assumed full dispersal and establishment of habitat trees (with implicit instantaneous seed production), which may overestimate A. verticillata's ability to colonise new areas. Given the strong likelihood that emissions will exceed LEV1 levels (IPCC 2007) and that A. verticillata recruitment is limited by herbivores such as tammar wallaby Macropus eugenii Desmarest, managers will probably need to revegetate to maintain A. verticillata and GBC populations. Although revegation effort could be reduced over the short term, key model assumptions (full dispersal and unlimited recruitment of A. verticillata) and model sensitivity to variation in carrying capacity (driven by climate related changes in A. verticillata) mean that managers should be ready for intensive revegetation in the future.

Management and monitoring should focus on maintaining adult survival and fecundity at their current levels. The acute

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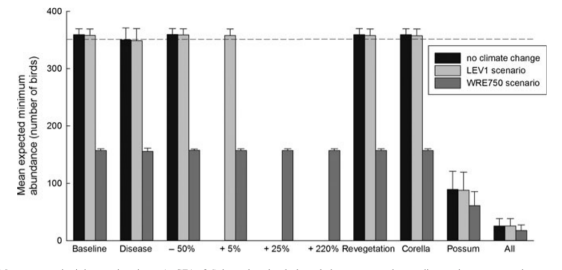


Fig. 4. Mean expected minimum abundance (\pm SD) of *Calyptorhynchus lathami halmaturinus* under no climate change, a greenhouse gas mitigation policy scenario (LEV1) and a high-CO₂-concentration stabilisation reference scenario (WRE750). The initial population size was 350 individuals (dashed line). Baseline = baseline scenario that includes observed fire frequency and ongoing use of current population management methods; disease = beak-and-feather-disease outbreak; -50% indicates 50% reduction in fire frequency from increased management; +5%, +25% and +220% (i.e. 2·2-fold increase) indicate increasing fire frequency from climate change. It was not realistic to model some fire increases for the no climate change or LEV1 scenarios. The last four groups of bars show the effects of ceasing management. 'Revegetation', 'corella' and 'possum' indicate stopping revegetation, little corella *Cacatua sanguinea* and brush-tail possum *Trichosurus vulpecula* management, respectively. 'All' indicates stopping all management actions.

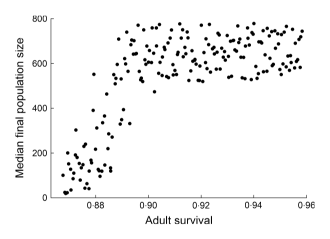


Fig. 5. Relationship between uncertainty in adult survival and median final population size in a Latin Hypercube sensitivity analysis for the no climate change scenario. The breakpoint for the segmented generalised linear model was 0.89, and the slopes were 78-9 and 0.76 for the low and high parameters, respectively. The mean estimate for adult survival from the mark-resight analysis is 0.913 (95% confidence interval from 0.88 to 0.93).

sensitivity of the model to lower (but still plausible) values of adult survival in the range of 85–90% emphasises the importance of monitoring adult survival over time. Predation from raptors such as wedge-tailed eagles *Aquila audax* Latham, climate variation, fire frequency and food availability may be important drivers of adult survival (Mooney & Pedler 2005), but there was no evidence of changing survival during the study period, and these relationships are incompletely known. Threats to the GBC may change over time, and the effects of climate variation on survival can be difficult to detect without monitoring data sets that span decades (Grosbois *et al.* 2008). Therefore, we suggest that mark-resight and reproductive data should continue to be collected to build this unique data set and allow ongoing analysis of the drivers of adult survival.

In addition to collecting data on the GBC, studies of *A. verticillata* are needed to improve forecasts of the GBC's extinction risk. In particular, studies on the effects of drought, warmer temperatures and fire on *A. verticillata* survival, recruitment and seed production are needed, especially given that climate change is likely to cause more extreme environmental events that would affect the life cycle of this food plant. New data could then be integrated with analyses that combine demographic models of both *A. verticillata* and the GBC.

Our approach minimised uncertainty by combining a comprehensive demographic data set with rigorous methods, including mark-resight estimation of survival and ensemble bioclimatic and global climate modelling, yet the model's assumptions should be considered when interpreting our results. The projected range contraction of *Allocasuarina verticillata* under the high emissions scenario assumes that the species' distribution–climate relationship remains the same as today and that climate is the main driver of range changes (species interactions are not considered for this plant). In addition, our model assumes that the relationship between *A. verticillata* and the GBC will remain strong in the future.

In conclusion, the results of our coupled demographic-BEM simulations suggest that the GBC is likely to continue its population increase over time until carrying capacity is reached, provided the climate remains similar to today and intensive possum control continues. However, should unmitigated climate change or reduced adult survival occur, severe declines are probable. We recommend continued intensive life-history monitoring on the GBC, possum management and research on A. verticillata, to promote the persistence of the GBC. The methods illustrated here demonstrate how species interactions can be included in coupled demographic-bioclimatic modelling approaches to add realism to forecasts of population viability under climate change for well-studied species of conservation concern. Furthermore, our analysis shows how coupled models can provide practical management advice in the face of broader issues and uncertainties such as global emissions mitigation.

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Supporting Information

Additional supporting information may be found in the online version of this article.

Fig. S1. Annual dispersal-distance curve for the *Calyptorhynchus lath-ami halmaturinus* population on Kangaroo Island.

Fig. S2. Mean final population size of persisting runs (\pm SD) of *Calyptorhynchus lathami halmaturinus* under no climate change, a greenhouse gas mitigation policy scenario (LEV1) and a high-CO₂-concentration stabilisation reference scenario (WRE750).

Table S1. Covariates and their data sources for the mark-recapture survival analysis of *Calyptorhynchus lathami halmaturinus* on Kangaroo Island.

 Table S2. Comparison of survival model results from Cormack-Jolly-Seber models in program MARK.

Table S3. Results of binomial GLMs relating spatial variables to *Calyptorhynchus lathami halmaturinus* presences on Kangaroo Island.

Table S4. Latin Hypercube sensitivity analysis results.

Table S5. Sensitivity of results to parameterisation of disease outbreaks.

Appendix S1. Previous modelling studies on the Kangaroo Island GBC.

Appendix S2. Detailed population-modelling methods.

Appendix S3. Climate change forecasts and bioclimatic envelope modelling methods.

Appendix S4. Integrating population and distribution models (methods).

Appendix S5. RAMAS scenarios and sensitivity analysis methods.

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1 Supporting Information

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3 Appendix 1. Previous modelling studies on the Kangaroo Island GBC

Two previous studies used population models to estimate the viability of the GBC 4 5 population, although neither considered climate change. Pepper (1996) used survival estimates from Carnaby's black-cockatoo (C. latirostris) and fecundity data from the little 6 7 reproductive research that had been done on Kangaroo Island by that time. Using VORTEX 8 software (Lacy 1993), Pepper (1996) calculated a mean time to extinction of 5.8 years. Pepper (1996) doubted the results and suggested that the assumptions of the model were 9 10 incorrect. Southgate (2002) used mark-recapture data from 1996-2001 to estimate survival, without explicitly modelling recapture probability. He calculated survival to be 0.296 for egg 11 to age 1, 0.77 for age 1 to 2, 0.83 for age 2 to 3, and c.0.85 for age 3+. Southgate (2002) used 12 13 data on sex ratio, clutch size, and percent of females breeding to estimate fecundity to be equal to 0.4 for female nestlings. Southgate (2002) used the software ALEX (Possingham & 14 Davies 1995) to estimate that the GBC population was declining by 10% a year. This finding 15 conflicted with census data which showed the population was increasing by c. 4% annually. 16 Southgate (2002) attributed the discrepancy to inaccurate survival data. 17

18

19

Appendix 2. Detailed population modelling methods

20 *Demographic structure*

We used life history data and expert knowledge from the GBC recovery program to parameterise the model (Crowder *et al.* 1994; Table 1). Breeding age for females is three years and for males is five years (LPP, pers. obs.; Mooney & Pedler 2005), and the species

forms permanent or semi-permanent monogamous pairs (Garnett et al. 2000). Black-

cockatoos probably show minimal reproductive senescence (Heinsohn *et al.* 2009). Thus, we
developed a stage- and sex-structured model with composite age classes for breeding female
(3+) and male age (5+) classes. Changes in mortality related to senescence are unknown in *Calyptorhynchus lathami* but we simulated the possible effects of senescence by adding a
senescent stage (age 20+), whereby mortality in this oldest stage was doubled. We found that
the growth rate (lambda) was reduced from 1.035 to 1.011.

31 Survival estimates

We estimated survival from 950 observations of 317 individuals marked between 1996–2008, using the Cormack-Jolly-Seber model for live recaptures in Program MARK (Cooch & White 2008). We used a two-stage modelling approach for mark-recapture data, whereby recaptures were initially modelled in combination with the most parameterised survival model, so as to retain as much power as possible for testing likely drivers of survival parameters (see Pardon *et al.* 2003 for justification). After the optimal recapture model was selected, a parsimonious survival model was sought.

39 Initially, we were interested in testing the effects of 13 covariates on annual cockatoo survival. We tested for correlations among covariates with a Spearman correlation matrix and 40 excluded five correlated variables (all remaining variables had all Spearman coefficients 41 <0.65; most were <0.3). The final analysis tested the effects of eight covariates on survival 42 (Table S1). The covariates for extreme events (drought, river flow, and repeated fire) were 43 best represented by thresholds in order to model GBC tolerance to low levels of these 44 45 variables. Therefore we converted these covariates into a binary format-ones or zeros if the values were above or below the median, respectively. Models were tested from an *a priori* 46 47 candidate set of 27 ecologically plausible models, which were developed based on our

48 experience with the species in the field. We used a hierarchical approach when testing for the optimal survival model (using likelihood) (Cooch & White 2008). We first tested for a cohort 49 effect but found no evidence for this. Then we tested different stage structures (two, three, or 50 51 four age classes) and found two stages was optimal. As the final step we compared models with no stage structure to those with two stages. Both classes of models included constant, 52 time-variant, and environmental covariate models. The only difference was that models with 53 54 no stage structure compared eight covariates (Table S1), while stage-structured models compared the three covariates (available protected hollows, number of hollows treated for 55 56 bees, and number of little corellas Cacatua sanguinea culled) that were likely to have a stronger effect on sub-adults than adults (Mooney & Pedler 2005). Models with wAIC < 0.01 57 are not included in Table S2. 58

We used parametric bootstrapping to estimate goodness-of-fit in the mark-recapture data (White 2002). We calculated $\hat{c} = 1.08$ by dividing the observed deviance for the most parameterised model by the mean deviance from 1,000 bootstrap simulations. This low value suggests little overdispersion and requires no adjustment (White, Burnham & Anderson 2001).

For model comparisons, we report $-2*\log(\text{likelihood})$ as the measure of deviance. We calculated an R² statistic from an analysis of deviance based on the following formula from Le Bohec *et al.* (2008): R² = (DEV(constant model) - DEV(covariate model)) /

(DEV(constant model) - DEV(time-dependent model)), where DEV is deviance. The
advantage of this method is that it assesses the relative effects of covariates on survival and
recapture rates. We used MARK to calculate weighted averages of the parameter estimates
from the Akaike weights (Burnham & Andersen 2002). Mark-resight data area continually
collected by the recovery program. Researchers wishing to use GBC survival estimates
should contact the recovery program for the latest figures.

73 Table S1. Covariates and their data sources for the mark-recapture survival analysis of 74 *Calyptorhynchus lathami halmaturinus* on Kangaroo Island. **availprot** = available protected hollows (artificial + natural); **bee** = number of hollows with honeybee Apis mellifera 75 76 deterrent inserted; **corella** = number of little corellas *Cacatua sanguinea* culled; **drought** = drought index (total rainfall in previous five years); **heat** = number of summer days \ge 35 °C; 77 **flow** = flow in Rocky River; **revegetation** = area revegetated with *A. verticillata* (with a six 78 79 year delay because A. verticillata cones require a minimum of six years to mature; PAM pers. obs.); **fire** = repeated fire index (area burned in previous 5 years) 80

Covariate Source		Possible effect on cockatoos		
availprot	GBCRP data*	Nest predation by possums		
bee	GBCRP data	Hollow competition		
11.				
corella	GBCRP data	Hollow competition/nest predation		
drought (threshold)	BOM, mean of 7 stations†	<i>A. verticillata</i> seed production and drinking water		
	BOM, mean of			
heat	3 stations	Heat stress on adults:		
		Proxy for available surface water for		
flow (threshold)	DWLBC¶	cockatoo drinking		
rovagatation	GBCRP data	A wartigillata soud production		
revegetation		A. verticillata seed production		
fire (threshold)	GBCRP/DENR data	Reduction of nesting and feeding habitat		
*Glossy black-cockatoo recov				

*Glossy black-cockatoo recovery program. See Mooney & Pedler (2005) for details.

[†]Bureau of Meteorology. We used data from weather stations with the most complete collection histories: stations 22800, 22801/23, 22803, 22817, 22835, 22836, & 22839 for rain; stations 22801/23, 22803, & 22841 for temperature. <u>http://www.bom.gov.au</u>

Summer is defined as December of the previous year and January and February of the current year. See Cameron (2008), Saunders, Mawson & Dawson (2011) for information on heat stress in *Calyptorhynchus*.
Department of Water, Land, and Biodiversity Conservation. Flow of Rocky River at gorge falls, site A5130501. http://e-nrims.dwlbc.sa.gov.au/swa/.

	82	Table S2.	Compariso	n of su	ırvival	model	results	from	Cormack	c-Jolly	-Seber mod	dels in
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Model	$\Delta \operatorname{AIC}_{c}$	Wi	k	LL	R^2
subad(.) ad(.)	0	0.20	15	2601.1	0.88
subad(corella) ad(.)	0.2	0.18	16	2599.2	0.90
subad(bee) ad(bee)	1.0	0.12	16	2600.0	0.89
subad(availprot) ad(.)	1.5	0.09	16	2600.5	0.88
subad(.) ad(.) + sex	1.6	0.09	16	2600.6	0.88
subad(availprot + corella) ad(.)	1.7	0.09	17	2598.6	0.90
subad(availprot) ad(availprot)	1.7	0.08	16	2600.7	0.88
subad(corella) ad(corella)	1.8	0.08	16	2600.8	0.88
subad(bee) ad(.)	2.0	0.07	16	2601.0	0.88
subad(t) ad(t) + sex	10.4	0	27	2586.6	1
constant	103.4	0	14	2706.5	0
t	104.4	0	25	2684.7	0.18
sex + t	105.5	0	26	2683.8	0.19

83 program MARK. The optimal recapture model was stage-structured and time-dependent.

84 *t* represents time. subad represents sub-adults, ad represents adults. Explanatory variables (Table S1) are 85 availprot = available protected hollows, bee = hollows treated for bees, corella = number of corellas culled, 86 repfire = repeated fires in the last five years. *k* indicates the number of parameters, AIC_c is Akaike's Information 87 Criterion corrected for small samples sizes, Δ AIC_c shows the difference between the model AIC and the 88 minimum AIC in the set of models, AIC weights (*w_i*) show the relative likelihood of model *i* and % DE is 89 percent deviance explained by the model.

90 *Fecundity*

We used the number of known fledglings in the population from 1996–2008 to
measure reproductive output in the population. This number is calculated each year by
summing the number of large nestlings seen at the nest up to a week before fledging, and
additional fledglings noted during the census. Sex ratio of fledglings and adults is 1.3 and 1.5
males to females, respectively (GBC recovery program data, 1996–2008). Fecundity was
calculated thus (Brook & Whitehead 2005):

97

x *

Number of nestlings of one sex produced per pair in year *i* =

Number of known fledglings in year i

 $_{98}$ population estimate in year *i* * proportion of females in breeding age class

The denominator represents the number of pairs alive in year *i* which is defined by the 99 100 number of breeding females in the population because females are limiting; the proportion of 101 females of breeding age (0.31) comes from the stable age distribution. x, the fledgling sex proportion, is equal to 0.4 and 0.6 to estimate the number of females and males produced per 102 breeding female, respectively (LPP pers. obs.). We then multiplied the number of fledglings 103 per female with adult survival to calculate fecundity based on a post-breeding census. This 104 resulted in a lambda < 1, whereas the observed population change indicated an annual rate of 105 106 increase (R) of 1.035. We thus adjusted the fecundities so that the eigenvalue of the stage matrix is 1.035. 107

108 Environmental stochasticity

109 RAMAS GIS simulates environmental stochasticity by sampling distributions as
110 specified by the mean and standard deviation of each stage matrix element (Akçakaya & Root
111 2005). To estimate standard deviation of fecundity we followed Akçakaya's (2002) approach

of subtracting the weighted average of demographic variance from the total variance. These
methods are commonly used to separate demographic and environmental variability for
population viability analyses (Lambert *et al.* 2006, Zeigler *et al.* 2010, Aiello-Lammens *et al.*2011). For the standard deviation of survival estimates, we used the square root of the process
error (sigma) reported by MARK (White, Burnham & Anderson 2001).

Appendix 3. Climate change forecasts and bioclimatic envelope modelling

119 Climate change forecasts

Spatial layers describing present day climate (0.01° x 0.01° latitude/longitude) were created
by interpolating between weather station records sourced from the Queensland Government
SILO patched point data base (Jeffrey *et al.* 2001), following the approach described in detail
by Fordham *et al.* (in press-a).

We used MAGICC/SCENGEN v5.3 (http://www.cgd.ucar.edu/cas/wigley/magicc), a 124 coupled gas cycle/aerosol/climate model used in the IPCC Fourth Assessment Report (IPCC 125 126 2007), to generate an annual time series of future climate anomalies for (2000-2100) for annual, austral winter and summer precipitation and temperature (0.5° x 0.5° 127 latitude/longitude; annual rainfall, January temperature, and July temperature in this study). 128 Projections were based on two emission scenarios: a high-CO₂-concentration stabilisation 129 reference scenario, WRE750, and a policy scenario that assumed substantive intervention in 130 131 CO₂ emissions LEV1 (Wigley, Richels & Edmonds 1996; Wigley et al. 2009). Models were chosen using an assessment of model convergence and skill in predicting seasonal 132 133 precipitation and temperature (see Fordham *et al.* in press-a for details). The nine skilful GCMs used to generate the multi- climate model ensemble average forecasts were GFDL-134

135 CM2.1, MIROC3.2(hires), ECHAM5/MPI-OM, CCSM3, ECHO-G, MRI-CGCM2.3.2,

UKMO-HadCM3, GFDL-CM2.1, MIROC3.2 (medres) (model terminology follows the
CMIP3 model database; http://www-pcmdi.llnl.gov/ipcc/about_ipcc.php). Although there is
no standard procedure for assessing the skill of GCMs (Fordham, Wigley & Brook 2012), by
using an ensemble model set of greater than five GCMs, the influence of model choice on
model prediction skill is lessened (Murphy *et al.* 2004; Pierce *et al.* 2009).

We downscaled the climate anomalies to an ecologically relevant spatial scale (0.01 x
0.01° longitude/latitude), using the "change factor" method, whereby the low-resolution
change from a GCM is added directly to a high-resolution baseline observed climatology
(Hulme, Raper & Wigley 1995). One advantage of this method is that, by using only GCM
change data, it avoids possible errors due to biases in the GCMs' baseline (present-day)
climate (Fordham, Wigley & Brook 2012; Fordham *et al.* 2012).

147 Bioclimatic envelope modelling

150

148 Allocasuarina verticillata presence data

149 We modelled the bioclimatic envelope of *Allocasuarina verticillata* (drooping she-

oak) because it provides the primary habitat and 98% of the diet of the GBC. A. verticillata

151 presences came from Department of the Environment and Natural Resources (DENR)

152 biological survey records across South Australia

153 (http://www.environment.sa.gov.au/Knowledge_Bank/Information_and_data/Biological_data

bases_of_South_Australia). The presences were carefully cleaned before inclusion; only

records with an accuracy of 1 km or better were retained, duplicate and erroneous records

were removed, and no opportunistic records were included, which left 572 presences for the

- analysis. Much of A. verticillata's range has been cleared, which may influence our ability to
- model the species's distribution. Using presences from across the species's South Australian

range and requesting validation from local plant ecologists helped address this issue. An
equal number of pseudoabsences were generated randomly within the study region; random
pseudoabsences were appropriate in this case because of the difficulty of intensively
sampling the study area (South Australia) (Wisz & Guisan 2009). Plant ecologists identified
three climate variables as having the greatest general influence on *A. verticillata* survival and
recruitment: mean annual rainfall, mean January temperature, and mean July temperature
(Stead 2008).

166 *Ensemble forecasting*

The potential distribution of A. verticillata was modelled with an ensemble 167 168 forecasting approach (Araújo & New 2007) based on seven BEM techniques: BIOCLIM (Busby 1991), Euclidian and Mahalanobis distances (Farber & Kadmon 2003), generalised 169 170 linear models (GLMs; McCullagh & Nelder 1989); Random Forest (Breiman 2001), Genetic 171 Algorithm for Rule Set Production (Stockwell & Noble 1992), and Maximum Entropy (Phillips & Dudík 2008) in BIOENSEMBLES software (Diniz-Filho et al. 2009). Internal 172 evaluation of the models was performed with a data split procedure, whereby 70% of the 173 occurrence data were randomly split and used for calibration of the models, and the 174 remaining 30% were used for cross-evaluation of the models. This procedure was repeated 10 175 176 times, thus generating a 10-fold cross-validation of model results. The observed prevalence of species was maintained in each partition, and for each partition we obtained alternative 177 models by projecting ranges after performing a full factorial combination of the 178 environmental variables used as predictors. The fitting and projection of alternative models 179 180 using data partition and multiple combinations of variables was used to account for uncertainties arising from the initial conditions and model parameterization (sensu Araújo & 181 182 New 2007). Model accuracy was measured using the average True Skill Statistic (Allouche, Tsoar & Kadmon 2006). This analysis was performed to check if a grossly implausible 183

projection was being made (i.e. TSS < 0.3). However, because measures of accuracy on nonindependent data do not provide a reliable benchmark for evaluation of projections of species distributional changes under climate change (Araújo *et al.* 2005), we instead used an unweighted consensus of the seven modelling techniques. The resulting map of the current distribution was validated by an expert botanist (P. Lang, DENR). We then ran the distribution models with the climate layers for 2011–2100 (described above) to create a combined time series of 91 climatic suitability maps for each year from 2010 to 2100.

The climate projected for 2100 on Kangaroo Island was within the range of variation in the training data for 2010. This was true for all three climate variables in both emissions scenarios. Therefore the bioclimatic model did not extrapolate to novel climates, which reduces uncertainty in projections (Pearson & Dawson 2003).

195 Appendix 4. Integrating population and distribution models

196 Calculating the habitat suitability function

The *A. verticillata* probability of occurrence maps for 2010–2100 (hereafter 'AVS')
were added to edaphic spatial layers (substrate, slope, and native vegetation) to mask out
unsuitable areas and delineate more suitable areas for *A. verticillata* and the GBC (Pearson,
Dawson & Liu 2004). Substrate and slope are specific to *A. verticillata*, while native
vegetation affects *A. verticillata* and the GBC.

Substrate, or geology, strongly influences soil type and is an important predictor of *A*. *verticillata* presence (Specht & Perry 1948; Green 1994). We collapsed category classes in
the Surface Geology of Australia dataset (1:1 million scale; Raymond & Retter 2010) into 17
classes in South Australia. Expert knowledge was used to define which substrate classes are
unsuitable for *A. verticillata* (mainly Holocene sands, and floodplain alluvium; P. Lang

unpubl. data). We treated areas with native vegetation (National Vegetation Information
System; http://www.environment.gov.au/erin/nvis/index.html) as having twice the suitability
of areas without native vegetation (Crowley, Garnett & Carruthers 1998). Because *A*. *verticillata* prefers to grow on steep, rocky slopes (Crowley, Garnett & Carruthers 1998;
Crowley *et al.* 1998), we created a slope layer from a digital elevation model (DEM-9S,
http://www.ga.gov.au/meta/ANZCW0703011541.html) in Arc GIS v9.3 (Arc GIS,

213 Environmental Systems Research Institute, Redlands, CA, USA).

We used binomial GLMs to relate the spatial layers to cockatoo presences and 214 generate the habitat suitability function. Presence data for the GBC (349 points) came from 215 216 active nest locations (n = 157; GBC recovery program data), band observations (n = 100; GBC recovery program data), known feeding sites (n = 18; GBC recovery program data), and 217 the South Australian Biological Survey (n = 74). No reliable absence points were available 218 219 for the GBC, so we were forced to generate psuedoabsences. Considering that the island has been well surveyed for GBCs, and that we wanted the model to focus on the factors 220 221 determining its distribution within the landscapes in which one might reasonably expect to 222 survey, we generated pseudoabsences using a positive distance weighting function that favours areas away from presences when creating pseudoabsences (Phillips et al. 2009; Wisz 223 & Guisan 2009). We tested models from an *a priori* candidate model set generated using our 224 knowledge of probable factors limiting the occurrence of GBCs. We primarily relied on 225 Akaike's Information Criterion corrected for small sample sizes (AIC_c) for model selection 226 (Burnham & Andersen 2002), but we also calculated the Bayesian Information Criterion 227 (BIC) because it is more conservative (tends to fit fewer tapering effects) and requires 228 substantially better fit before selecting a more complex model (Bolker 2008). 229

230 <u>Habitat suitability function</u>

231 Our selected covariates adequately predict GBC occurrence, explaining 38.5% of the

variance (Table S3). The best model (habitat suitability ~ substrate*slope +

vegetation*AVS; wAIC of 0.954) became the habitat suitability function for the RAMAS

234 model. Thus, habitat suitability is defined as:

235 habitat suitability = ((4.61*[substrate] + 1.49*(thr([slope],0.01366)) - 2.11*[vegetation] -

236 0.454*(thr([AVS],0.399)) - 0.8818*[substrate]*(thr([slope],0.01366)) +

237 **3.784*[vegetation]*(thr([AVS],0.399)))*[substrate])** / **5.34375**

The coefficients were estimated from the binomial model. The entire equation is multiplied by substrate in order to mask out areas with unsuitable substrate, and then divided by 5.34375 to scale habitat suitability from 0 to 1 in each grid cell. We applied thresholds (thr) to slope and AVS such that this part of the equation was equal to zero unless the grid cell's value was greater than the lower fifth percentile of the variable where GBCs occur. Thresholds used in this manner better capture species' responses to continuous spatial variables in metapopulation models (DAF unpubl. data).

We used a threshold to determine a lower habitat suitability limit below which we 245 would not expect an occurrence. Threshold selection affects range area predictions, and the 246 247 choice of a threshold depends on the goals of the modelling exercise (Liu et al. 2005). The GBC population on Kangaroo Island has been carefully censused so we had high confidence 248 that the distribution was well-represented by the point locality data. We aimed to characterise 249 250 the current extent of medium to high quality habitat and predict the potential distribution of suitable habitat patches in the future which we did by selecting cells with a HS value higher 251 than the value recorded for the lowest 5% of GBC presences. We used our knowledge of the 252 253 species in the field to validate the resulting habitat suitability maps.

254

Table S3. Results of binomial GLMs relating spatial variables to Calyptorhynchus lathami 255

halmaturinus presences on Kangaroo Island. AVS stands for climatic suitability of 256

Allocasuarina verticillata (the cockatoo's food plant). The global model had the strongest 257

AIC $_c$ and BIC support, explaining 38.5% of model structural deviance. Of the single term 258 models, slope had greatest support explaining 26.5% of model deviance. Models in bold had 259 wAIC >0.01.

260

Model	% DE	wAIC _c	ΔAIC_c	wBIC	Δ BIC	k
	70 DL	\mathcal{M}		,, DIC		N
substrate*slope + vegetation*AVS	38.5	0.954	0	0.497	0	7
substrate*slope + vegetation + AVS	35.9	0.022	7.5	0.065	4.1	6
substrate + vegetation +						-
slope + AVS	35.1	0.015	8.4	0.245	1.4	5
substrate*slope + AVS	34.9	0.010	9.2	0.161	2.3	5
substrate*slope	31.4	0	20.0	0.004	9.5	4
substrate + slope	30.8	0	20.0	0.024	6.1	3
substrate + vegetation + slope	31.0	0	21.7	0.002	11.3	4
AVS*slope	30.8	0	22.1	0.001	11.6	4
slope	26.5	0	34.1	0	16.6	2
vegetation*slope	27.3	0	35.4	0	25.0	4
substrate + vegetation*AVS	20.0	0	64.6	0	57.6	5
substrate	10.8	0	92.6	0	75.1	2
vegetation*AVS	6.7	0	111.9	0	101.5	4
AVS	3.3	0	120.2	0	102.7	2
null	0	0	130.5	0	109.5	1
vegetation	0.04	0	132.4	0	114.9	2

262 <u>Carrying capacity</u>

263	Estimates of carrying capacity were based on previous research on A. verticillata
264	productivity and extent on Kangaroo Island, and known density of GBCs in A. verticillata
265	stands. One hectare of moderate quality she-oak habitat (334,000 cones) supports
266	approximately 7.5 birds (Crowley, Garnett & Pedler 1997; Chapman & Paton 2002). The
267	current area of A. verticillata on Kangaroo Island is 4,900 ha (SA DENR data), so the
268	approximate carrying capacity of the island is 653 birds. This is a maximum estimate of
269	current carrying capacity given that GBCs only feed on c. 10% of available A. verticillata
270	(Chapman & Paton 2005). In RAMAS we used a scaling constant (0.233) to relate the known
271	carrying capacity to the number of suitable cells (noc). We applied a threshold to the equation
272	to eliminate very small unviable patches with carrying capacity <10 birds:

273 K = thr(0.233*noc,10)

274

275 <u>Initial abundance</u>

Initial abundance was calculated in a similar way. Annual censuses of the population
estimated the current population size at c. 350 individuals, so we used a lower scaling
constant to approximate this:

279 $N_{initial} = thr(0.125*noc,10)$

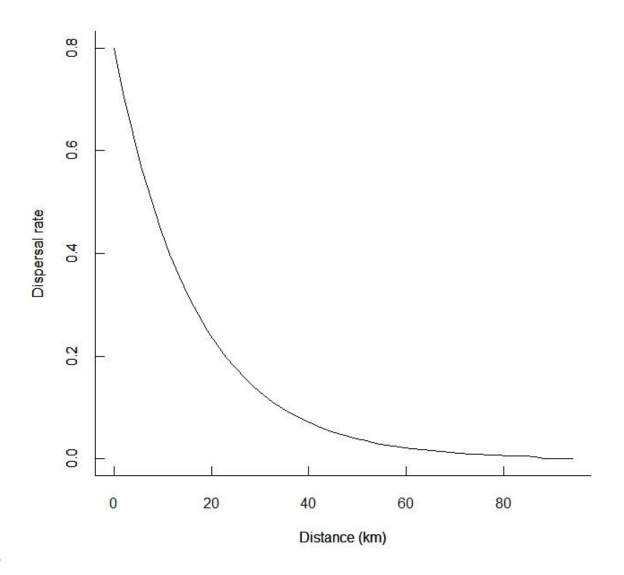
We ran trial scenarios with initial abundances of 100 and 200 birds and found that the population showed the same general responses as with 350 birds. These trials, combined with the carrying capacity of 653 under ceiling density dependence, suggest that the model was not very sensitive to initial population size.

284

285 <u>Dispersal</u>

Data on movements of marked birds were used to estimate annual dispersal. Available 286 information suggests that approximately 73% of birds leave the general natal area annually 287 and 23% of these leave the wider flock region, so c. 17% of birds disperse annually 288 289 (Southgate 2002; Mooney & Pedler 2005). Dispersers moved an average of 44 km and up to 78 km (Southgate 2002). This high rate of dispersal supports our use of mark-recapture-290 derived survival estimates even though only a portion of the island is covered by the mark-291 recapture surveys. Our dispersal function had 17% of birds dispersing \geq 28 km annually and 292 1% of the population (4 birds) dispersing 78 km annually (Fig. S1). We modelled dispersal as 293 294 a function of the distance between the centres of suitable habitat patches.

295 **dispersal** ~ **a** = **0.8**, **b** = **16.5**, **c** = **1**



296

Figure S1. Annual dispersal-distance curve for the *Calyptorhynchus lathami halmaturinus*population on Kangaroo Island.

299

300 <u>Correlation among grid cells</u>

Environmental variability was set to be correlated between populations depending on their spatial separation. Pairwise correlations were calculated using an exponential function, P =a.exp($D^{c/b}$), where D is the distance between centroids of habitat patches and a, b and c are constants. Following Keith et al. (2008), we used regional variation in year-to-year annual rainfall across South Australia to approximate environmental variability (a = 0.79, b = 1266, c = 1).

307

Appendix 5. RAMAS scenarios and sensitivity analysis

308 *Fire*

309 <u>Baseline fire frequency</u>

Four fires burned >10% of Kangaroo Island from 1950–2008, which yields an annual 310 probability of severe fire of 6.8% (GBC recovery program data). Our vital rates estimates 311 312 included the effects of past severe fires so we included observed fire frequency in the baseline scenarios. We modelled fire probability as being the lowest after a fire (0.1% 313 probability) and then increasing with mounting fuel loads until the maximum probability 314 (6.8%) is reached after seven years (Keith et al. 2008). To maintain structural simplicity of 315 the model, it was assumed that fires burnt entire patches (i.e. no fire heterogeneity within 316 317 patches)

318 Impacts of fire on the GBC

The best data on the effects of a severe fire on the GBC come from 2007 when fires burned 85,920 ha (19.5% of the island), destroying five known nest sites and 425 ha of *A*. *verticillata* feeding habitat (Sobey & Pedler 2008). Based on nesting data from 1997–2003, if five nests are lost, fecundity is reduced by 8–12%. Therefore we modelled the effects of a severe fire as having a 10% reduction in fecundity. Reduction in feeding habitat from severe fires is expected to have a minor, delayed impact on survival (DCP pers. obs.), so we modelled this effect by reducing sub-adult and adult survival by 3% after a severe fire.

326

327 <u>Climate change and increased fire management</u>

Climate change is predicted to cause a substantial increase in the number of days with 328 very high to extreme fire danger on the Fleurieu Peninsula (Lucas et al. 2007). These 329 predictions suggest that severe fire danger will increase by 5% or 25% by 2050 for low and 330 high emissions scenarios, respectively. We interpreted these changes as percent increases in 331 base probability of fire on Kangaroo Island and used the 2050 estimates as guidelines. 332 Making the conservative assumption that there is a linear correlation between fire frequency 333 and fire days, increases of 5% and 25% would yield annual fire probabilities of 7.1% and 334 8.5% on Kangaroo Island. We also considered a nonlinear example where a 2.2-fold increase 335 336 in fire frequency yielded a 15% annual fire probability on Kangaroo Island (approximately doubling the current probability). We also investigated the impact of increasing fire 337 management to reduce the annual probability of severe fire by half to 3.4%. 338

339

340 Disease

Psittacine beak-and-feather disease typically kills juvenile parrots only (DEH 2005). 341 Virulence of the disease varies; major epidemics with high mortality can occur in isolated 342 parrot populations with little immunity, while populations with previous exposure to the 343 344 disease are more resilient (DEH 2005; Khalesi 2007). There have been no recorded cases of beak-and-feather disease on Kangaroo Island (LPP pers. obs.), so we assumed low immunity 345 and high mortality. Little corellas regularly cross from the mainland to Kangaroo Island 346 (Mooney & Pedler 2005) and could serve as vectors of the disease (DEH 2005). We modelled 347 a possible outbreak by reducing survival of zero year olds and one year olds by 50%. We set 348

349	the annual probability of an outbreak at 5% and the probability of an infected dispersing bird
350	transmitting the disease at 75%. While the values of these parameters are poorly known in the
351	wild (Khalesi 2007) an expert on beak-and-feather disease confirmed that our
352	parameterisation was realistic (M. Holdsworth, pers. comm.).
252	

353

354 Active management

355 Brushtail possum management

The GBC recovery team manages nest-predating brush-tail possums Trichosurus 356 vulpecula by placing metal collars around the trunks of GBC nest trees and pruning 357 overlapping tree crowns to prevent access to nest trees (Mooney & Pedler 2005). Possum 358 359 management can increase fecundity by 78% (the probability of an egg producing a fledgling increases from 23% to 41%; Garnett, Pedler & Crowley 1999). If possum management were 360 361 stopped, fecundity would decrease by approximately 44%. We assumed a linear decrease in 362 fecundity after stopping management in 2010. By 2025 (15 years after stopping management) all benefits from protected hollows are modelled as being lost (no new hollows protected, tree 363 crowns overlap, and metal collars rust and fall off trees; LPP pers. obs.) and fecundity is 44% 364 365 lower.

366

367 <u>Corrella management</u>

The little corella *Cacatua sanguinea* population on Kangaroo Island has increased substantially over time, probably as a result of land clearance and grain cropping (Garnett *et al.* 2000). Corellas compete with GBCs for nests and kill GBC nestlings. As a result, corellas found near GBC nests have been culled since 1998. If corella management were stopped, it

has been estimated that approximately two GBC nestlings would be lost per year (Garnett,

Pedler & Crowley 1999; PAM pers. obs.), so we modelled stopping corella management as

causing a 7% drop in fecundity. We simulated stopping management in 2010 and assumed a

linear decrease in fecundity that took five years to reach the 7% reduction.

376

377 <u>Revegetation</u>

Volunteers and the GBC recovery team have planted *A. verticillata* on Kangaroo
Island since 1988 in an effort to augment GBC food sources. From 1996–2007, 39.3 ha were
revegetated which amounts to 3.5 ha per year on average. Most revegetation is now done near
<u>traditional nesting areas</u> where remnant *Allocasuarina verticillata* has been reduced
considerably by clearing. Consequently, the current revegetation rate can be approximated as
boosting fecundity by approximately 3% annually (PAM pers. obs.). We modelled stopping
revegetation as causing a linear decline in fecundity that lead to a 3% drop in five years.

We also simulated the effects of stopping all management actions (possum, corella, and revegetation in 2010). This lead to a 24.7% decrease in fecundity in five years and a 54% drop in 15 years.

388

389 Sensitivity analysis

For the Latin Hypercube sensitivity analysis we took samples from 200 equal-width strata (following the method described in Brook, Griffiths & Puckey 2002) along the following ranges of parameter values relative to the value used in the RAMAS models: adult survival (± 5 %), sub-adult survival (± 10 %), fecundity (± 10 %), carrying capacity (± 20 %), and annual dispersal (± 20 %) (Brook, Griffiths & Puckey 2002). The range for fecundity is equivalent to the standard error around the parameter estimate. The ranges for survival
needed to be larger than the standard errors to evaluate the model's sensitivity over a
plausible range. We used large ranges for carrying capacity and dispersal for the same reason.

Standardised regression coefficients, calculated by dividing the coefficient of each 398 parameter by its standard error, and then weighting the resulting coefficients to sum to 1 399 (Conroy & Brook 2003), were used to assess the sensitivity of the model to the input 400 parameters. The coefficients were estimated by fitting a quasiPoisson GLM (to correct for 401 overdispersion) with all of the sensitivity analysis parameters (adult survival, sub-adult 402 survival, fecundity, carrying capacity, and annual dispersal). The non-linear, near-threshold 403 404 relationship between adult survival and final population size was broken into two parts and was best dealt with by fitting a segmented model (Fig. 5; Muggeo 2012). Therefore, the GLM 405 406 included a segmented fit for adult survival which resulted in two parameters, one above and 407 one below the breakpoint. The breakpoints were estimated at 0.893 ± 0.00081 SE for no climate change (6 iterations to reach convergence), 0.895 ± 0.0011 SE for LEV1 (8 408 409 iterations), and 0.886 ± 0.0010 SE for WRE750 (4 iterations). Bootstrapping with 10,000 samples was used to estimate the 95% confidence intervals for the parameter estimates. 410

Table S4. Latin Hypercube sensitivity analysis results. Standardised regression coefficients
 were calculated from generalised linear models to rank six sensitivity parameters in order of

their importance on *Calyptorhynchus lathami halmaturinus* mean final population size. "adult

survival-low" is the parameter below the break point in the segmented model and "adult

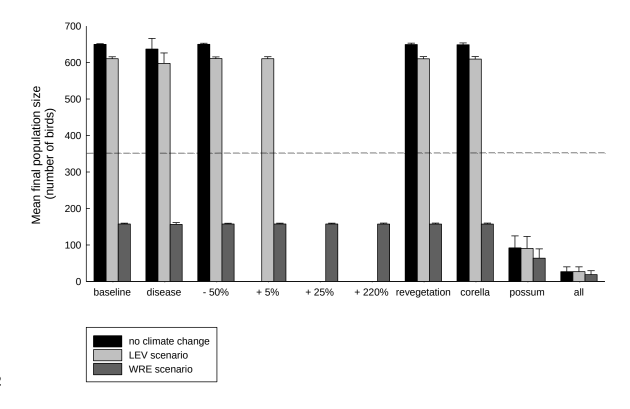
416 survival-high" is the above the break point.

-	. 1 1 1		1	
	standardised coefficient	coefficient	lower CI	upper CI
no climate change	coefficient	coefficient	CI	CI
adult survival-low	0.485	78.9	65.8	103.4
carrying capacity	0.211	0.0011	0.0009	0.0014
juvenile survival	0.110	1.26	0.76	1.86
fecundity, daughters	0.087	2.63	1.15	4.37
dispersal	0.041	-0.18	-0.39	0.01
fecundity, sons	0.033	0.79	-0.22	1.88
adult survival-high	0.033	0.76	0.27	1.78
LEV1				
adult survival-low	0.412	64.4	50.5	152.8
carrying capacity	0.246	0.96	0.75	1.14
sub-adult survival	0.154	1.98	1.24	2.77
fecundity, daughters	0.093	3.10	1.41	5.04
fecundity, sons	0.060	1.58	0.18	2.86
dispersal	0.022	0.11	-0.11	0.34
adult survival-high	0.013	0.35	-0.49	4.26
WRE750				
adult survival-low	0.327	67.7	45.2	131.4
carrying capacity	0.319	1.05	0.90	1.19
sub-adult survival	0.141	1.50	0.85	2.18
fecundity, sons	0.076	1.69	0.43	3.16
fecundity, daughters	0.071	1.99	0.51	3.59
dispersal	0.039	-0.16	-0.39	0.07
adult survival-high	0.026	0.49	-0.31	1.81

417

419 Table S5. Sensitivity of results to parameterisation of disease outbreaks.

Scenario	Mean final population size \pm SD		
baseline	649.66 ± 1.9		
disease outbreak, 5% annual probability, sub-adult survival reduced by 50% ¹	636.79 ± 29.1		
disease outbreak, 10% annual probability, sub-adult survival reduced by 50%	605.35 ± 65.3		
disease outbreak, 5% annual probability, sub-adult survival reduced by 75%	607.02 ± 69.6		
disease outbreak, 10% annual probability, sub-adult survival reduced by 75%	449.25 ± 164.6		
is the parameterisation used in the present study (see Appendix 5).			



422

Figure S2. Mean final population size of persisting runs (± SD) of *Calyptorhynchus lathami* 423 halmaturinus under no climate change, a greenhouse gas mitigation policy scenario (LEV1), 424 and a high-CO₂-concentration stabilisation reference scenario (WRE750). The initial 425 426 population size was 350 individuals (dashed line). Baseline = baseline scenario that includes observed fire frequency; disease = beak-and-feather disease outbreak; - 50% indicates 50% 427 reduction in fire frequency from increased management; +5%, +25%, and +220% (i.e., 2.2-428 fold increase) indicate increasing fire frequency from climate change. The last four groups of 429 bars show the effects of ceasing management. "Revegetation", "corella", and "possum" 430 indicate stopping revegetation, little corella Cacatua sanguinea, and brush-tail possum 431 Trichosurus vulpecula management, respectively. "All" indicates stopping all management 432 actions. 433

434

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