Successful island reintroductions of New Zealand robins and saddlebacks with small numbers of founders

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Abstract
Populations established with a small number of founders are thought to have a high risk of extinction due to Allee effects, demographic stochasticity, inbreeding and reduced genetic variation. We tested whether the initial number of birds released was related to persistence in reintroductions of saddlebacks (Philesturnus carunculatus) and robins (Petroica australis) to New Zealand offshore islands. Data were analysed for 31 populations that had been observed for at least 3 years since reintroductions. The numbers released ranged from 5–188. Most of the populations (26) survived and grew, including five from less than 15 founders, and four out of the five extinctions were attributable to introduced mammalian predators. The number of individuals released did not significantly affect extinction probability. The ability of these small releases to establish populations can be attributed to the closed nature of the islands (allowing birds to find mates), low mortality rates following release and high growth rates at low density. Stochastic simulation models based on data from two reintroduced populations suggested that populations with four founders (two male, two female) would have a negligible chance of extinction through demographic stochasticity and would be able to grow even if there were high rates of egg failure through inbreeding.

INTRODUCTION
Comparative analyses of bird and mammal reintroductions has shown that the probability of success is correlated with the number of animals released (Griffith et al., 1989; Wolf et al., 1996; Wolf, Garland & Griffith, 1998). Griffith et al.’s (1989) model suggested that about 80 animals needed to be released to approach the maximum probability of success. While some minimum number is needed to ensure a high probability of success in any reintroduction, it may not be a general rule that as many as 80 animals is necessary.

Griffith et al.’s (1989) asymptote may overestimate release size because managers are unlikely to translocate large numbers of animals unless they judge the reintroduction to have a high probability of success. More importantly, the number of animals needed will depend on circumstances. Small populations will have a high risk of extinction through demographic stochasticity if they suffer a phase of high post-release mortality, meaning the number of founders is much less than the number released, and if their expected rate of increase is low. Small populations can also be driven to extinction through Allee effects such as post-release dispersal that prevents individuals from finding mates, or increased susceptibility to predation at low densities (Lande, 1999). Small populations may also be prone to extinction due to inbreeding or loss of genetic diversity (Lande, 1999). The impact of these factors will depend on the characteristics of the species and release sites, hence the number of animals that need to be released may vary greatly.

Bird reintroductions in New Zealand have had a high success rate in comparison to the rest of the world (Armstrong & McLean, 1995). Most reintroductions have been to offshore islands following forest regeneration or eradication of introduced mammalian predators, two factors that have produced favourable habitat. Favoured habitat coupled with high pairing success due to the closed character of island systems and the sedentary nature of these endemic birds (i.e. flightless/reduced flight and non-migratory) are expected to produce high initial growth rates.

These factors all suggest that the numbers of animals needed for island reintroductions of New Zealand birds may be somewhat less than the numbers needed for many other species and systems. The small number of birds used to successfully re-introduce populations has also been cited as evidence that New Zealand birds are less vulnerable to inbreeding depression (Craig, 1991, 1994; Craig et al., 2000). We assess whether a small number of birds can establish populations in New Zealand as well as the impact that inbreeding depression may have on population growth rates. We used data for
two passerines, the saddleback (Maori name = tieke) *Philesturnus carunculatus* and the New Zealand robin (Maori name = toutouwai) *Petroica australis*, where good data are available from reintroductions to multiple islands. We first analysed success/failure of reintroductions versus number released. We then used stochastic models developed for individual robin and saddleback populations to predict the relationship between the probability of successful reintroduction, numbers released and egg failure rates.

**METHODS**

Saddlebacks (80 g) and robins (30 g) are small to medium-sized forest passerines, each with a modal clutch size of two eggs and up to four clutches per season (Heather & Robertson, 1996; Armstrong et al., 2000; Hooson & Jamieson, 2003a). They are territorial and sedentary, become sexually mature in their first year and form monogamous pairs. Saddlebacks and robins are relatively weak fliers and migration between current island populations is unlikely. After their release, neither species receives any substantial hands-on management. Saddlebacks are classified as secondary cavity nesters but nest in a variety of protected sites other than tree cavities (Heather & Robertson, 1996; Armstrong et al., 2002; Hooson & Jamieson, 2003a). Three islands with saddlebacks (Kapiti, Mokoia and Tiritiri Matangi) had nest boxes erected for use but were not used to the exclusion of natural nesting sites (Lovegrove, 1996a; Armstrong et al., 2002; Stamp, Brunton & Walter, 2002). Both species show evidence of low genetic variation within and between island populations (Livingston, 1994; Arderne & Lambert, 1997; Lambert et al., 2005; Tepolt, 2005), suggesting the current populations are inbred.

We obtained data for 36 island reintroductions of robins and saddlebacks where success or failure could be assessed. We excluded one robin and four saddleback reintroductions to islands where stoats (*Mustela erminea*), feral cats (*Felis domesticus*), Norway rats (*Rattus norvegicus*) or ship rats (*R. rattus*) were present. Research has shown that saddlebacks cannot coexist with these predators (Lovegrove, 1996a,b), hence these failed reintroductions provide no information on the importance of numbers of birds released. Other potential predators such as Polynesian rats or kiore (*Rattus exulans*), possums (*Trichosurus vulpecula*), mice (*Mus musculus*) and weka (*Gallirallus australis*), a native flightless rail, appear to have much less devastating effects (Lovegrove 1996a,b; Hooson & Jamieson, 2003b). We therefore included the possible effect of the presence/absence of the lesser predators on the various islands. These predators were ranked according to the risk they posed: 2 = possum, kiore; 1 = mice, weka; 0 = no predators.

Both saddlebacks and robins are relatively easy to survey because they are noisy, respond readily to playback and are attracted to human observers. If subsequent surveys of islands after release events failed to record any birds, we categorised these populations as extinct.

We also considered a population to be quasi-extinct if the number of individuals decreased by more than 50% 3 or more years after the initial release. We did not consider any populations that had been established for less than 3 years, but did include the number of years after release as an explanatory variable in our analysis. In some cases, the same species was translocated to the same island more than once. If these translocations were within 3 consecutive years, we used the total number of released birds as the initial population. If the translocations were separated by more than 3 years, we used the first translocation only (i.e. each island population only occurred once in the analysis).

The effect that the number of birds released had on extinction probability (success/failure) was tested for saddlebacks using logistic regression with predation risk, years since release and island area included as other explanatory variables. None of the six robin populations went extinct and the lack of variation for this variable meant: (1) we were unable to combine the robin and saddleback data in an overall model with species as an explanatory variable, or (2) run a separate model for the robin data (see Results, below). Data were obtained from published papers, personal communication with New Zealand’s Department of Conservation staff and the website of the Oceania Section of the Reintroduction Specialist Group (http://www.massey.ac.nz/~Darmstro/nz_projects.htm). Means and standard deviations are given throughout and statistical analyses were performed using SPSS 11.5 with significance set at $\alpha = 0.05$.

We also assessed extinction probability using stochastic matrix models based on Armstrong & Ewen’s (2002) and Armstrong et al.’s (2002) analysis of the reintroduced robins on Tiritiri Matangi Island and Armstrong et al.’s (2005) analysis of the reintroduced saddleback population on Mokoia Island. Both models included demographic stochasticity in births, deaths and sex ratio of offspring, with births sampled from the Poisson distribution and deaths and sex sampled from the binomial distribution. The models tracked both males and females and assumed strict monogamy (i.e. the number of breeding pairs is the number of males or the number of females, whichever is less). The expected finite rates of increase ($\lambda$) were initially 1.80 and 2.35 for the robin and saddleback populations, respectively, but declined with density. The log odds of juvenile survival was inversely correlated with the number of pairs in both models and for the saddlebacks, the number of fledglings per female declined both with density and with the quality of territories available. We ran each model 500 times and obtained the 2.5% and 97.5% percentile for the number of females present at the start of each year.

We used the simulation models both to predict probability of extinction as a function of initial population size and to model the effects that inbreeding depression might have on reintroduced robin or saddleback populations. We examined the effect of egg failure rates (caused by infertility and embryo deaths) because it is a well-established indicator of inbreeding depression (Spottiswoode & Moller, 2004). No data were available on
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egg failure rates for populations of different initial sizes, so we simply examined the effect of hypothetical increases in egg failure rate. We assumed that a reduction in the proportion of eggs hatching would cause a proportional reduction in the number of young fledged.

RESULTS

Effects of initial release numbers on extinction probability

The average number of robins and saddlebacks released during each translocation was 31.3 ± 26.5 (range = 5–66, n = 6) and 34.0 ± 34.3 (range = 6–188, n = 25) respectively. Little Barrier Island was an outlier with respect to the number of birds released (188) and island area (3083 ha). However, its inclusion in the analysis did not affect the results.

Only five out of 24 island populations of saddlebacks went extinct or quasi-extinct, while none of the six robin populations failed (Table 1). Furthermore, four out of the five failed saddleback populations were on islands with a high risk of predation (Table 1). Two robin releases involved five birds each (in one case, only one pair was reported to have bred successfully: Flack, 1977) and produced populations of 60 birds on 15 ha Allports Island and over 400 on 59 ha Motuara Island (Byrne, 1999).

In the full logistic regression model for saddlebacks, years since release and predation risk were correlated (correlation matrix r = 0.825), preventing model convergence. When these variables were considered singly, predation risk was significant (Wald = 4.920, d.f. = 1, P = 0.027) but years since release was not (Wald = 2.971, d.f. = 1, P = 0.085) so we removed years since release from the model. In the final model, none of the variables were significant, i.e. island area (Wald = 0.016, d.f. = 1, P = 0.39, odds ratio 95% confidence intervals (CI) = 0.955–1.018; Fig. 1(a)), predation risk (Wald = 2.610, d.f. = 1, P = 0.066, odds ratio CI = 0–1.363; Table 1), or the number of birds released (Wald = 0.074, d.f. = 1, P = 0.3032, odds ratio CI = 0.801–1.071; Fig. 1(b)). The

<table>
<thead>
<tr>
<th>Species</th>
<th>Outcome</th>
<th>Release no. (mean ± SD)</th>
<th>Predation risk</th>
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<tbody>
<tr>
<td>Saddleback (n = 25):</td>
<td>Failed</td>
<td>21.0 ± 15.1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>Succeeded</td>
<td>37.2 ± 37.2</td>
<td>17</td>
</tr>
<tr>
<td>Robin (n = 6):</td>
<td>Failed</td>
<td>–</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Succeeded</td>
<td>31.3 ± 26.5</td>
<td>6</td>
</tr>
</tbody>
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†Predators were ranked according to the risk they posed (0 = no predators, 1 = mice, weka, 2 = possum, kiore), with categories 0 and 1 combined because of small sample sizes.

Fig. 1. Number of failed and successful saddleback (●) and robin (□) populations in relation to: (a) island area and (b) the number of birds released. The successful release and establishment of 188 saddlebacks on Little Barrier Island (3083 ha) is not shown.
estimated coefficient for the number of birds released was –0.077, which corresponds to an increase in the probability of extinction from 0.0008 to 0.003 when the number of birds released decreases by 50% from 34 (the mean number of saddlebacks released) to 17, assuming a mean island area of 294 ha and low predation risk.

The stochastic simulation models also supported the conclusion that few founders were needed to ensure a high probability of reintroduction success for saddlebacks and robins. Under the parameters estimated for the Mokoia saddleback population and the Tiritiri Matangi robin population, a founder size of only two males and two females was predicted to have 0% and 2% chance of extinction in 15 years for observed egg failure rates of 22% and 15% respectively (Figs 2(a) and 3(a)).

Modelling the effects of inbreeding depression on extinction probability of saddlebacks and robins

The simulation models also suggest that such populations could experience high levels of egg failure and still grow. Doubling or tripling the egg failure rate of the Mokoia saddleback population slowed growth rates of simulated populations, but produced negligible extinction risk (Figs 2(b) & (c)). Doubling and tripling the egg failure rate of the Tiritiri Matangi robin population had a more pronounced effect than in the saddlebacks, reflecting the lower finite rate of increase for the robin population, but the simulated populations still had a 67% chance of persisting with the egg failure rate tripled (Figs 3(b) & (c)).

DISCUSSION

The above results are necessarily limited to the period of time over which the island populations have been monitored. Transfer failures were known to have occurred within 3 years of birds being released and so 3 years was set as the lower limit for our analysis. Almost half of the populations have persisted for more than 20 years, or at least an estimated 6–7 generations, which we believe should be long enough to have detected declines in populations if they were occurring. Given our conservative definition of a failed population – the initial release population declined by 50% or more – there appeared to be relatively few failures. Introduced predators such as stoats, cats and rats are still the primary drivers of extinction in saddlebacks and robins, as they are for most of New Zealand’s endemic bird fauna (Clout, 2001; Duncan & Blackburn, 2004). However, when the adverse effects of these major predators are removed, far more translocated populations established successfully than failed and were successful across a wide range of initial populations sizes. This suggests that surviving New Zealand bird populations on small islands are not exceptions to many unobserved or unrecorded extinct populations, as suggested by some authors (Frankham, Ballou & Briscoe, 2002). Furthermore, relatively small numbers of released birds do not appear to prevent population growth recovery to carrying capacity as long as major introduced mammalian predators are absent or are controlled.

Although small island populations of New Zealand native birds appear to persist, they may still suffer from the detrimental consequences of inbreeding. There is increasing evidence that New Zealand’s native birds show a reduction in fitness associated with close inbreeding (Jamieson, Roy & Lettink, 2003) or severe bottlenecks (Jamieson & Ryan, 2000; Briskie & Macintosh, 2004). In addition, Briskie & Macintosh (2004) have found that hatching failure rates of native birds that have gone
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Fitness traits such as egg fertility and hatchability can still grow even when inbreeding depression in the population is occurring. Saddleback and robin populations can grow in spite of potential inbreeding problems such as high rates of egg failure. Nevertheless, these populations could have lost significant genetic variation through founder events, drift or inbreeding, or a combination of these factors, placing the new populations at greater risk to invading pathogens or environmental perturbations over the long term. The issue of long-term viability of translocated populations established with a small number of founders clearly requires further research.

**CONCLUSIONS**

The results of this study appear to support the perception that translocations of New Zealand native birds to predator-free islands are frequently successful even when relatively few individuals are transferred and released. We have shown that reintroduced saddleback and robin populations can grow in spite of potential inbreeding problems such as high rates of egg failure. Nevertheless, these populations could have lost significant genetic variation through founder events, drift or inbreeding, or a combination of these factors, placing the new populations at greater risk to invading pathogens or environmental perturbations over the long term. The issue of long-term viability of translocated populations established with a small number of founders clearly requires further research.

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**REFERENCES**


