POPULATION VIABILITY ANALYSIS FOR RED-COCKADED WOODPECKERS USING AN INDIVIDUAL-BASED MODEL

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Abstract. Red-cockaded Woodpeckers (Picoides borealis) are distributed in the southeastern United States among closed populations whose maximum size is limited. Previous population viability analyses for this species have been confined to examination of threats posed by catastrophes and loss of genetic variability, because of the lack of demographic models that incorporate the extreme spatial constraints on dispersal that characterize this species. We used a spatially explicit, individual-based simulation model to assess the vulnerability of Red-cockaded Woodpecker populations to demographic and environmental stochasticity. Vulnerability to these threats was relatively low, because the presence of a substantial nonbreeding class (i.e., helpers) ameliorated the impact of stochastic variation in mortality and reproduction on the size of the breeding population. Because dispersal of helpers is spatially restricted, this effect was most pronounced when territories were aggregated or at high densities. Populations of 250 and 500 territories were stable regardless of the level of territory aggregation at the densities examined, whereas populations of 25, 49, and 100 territories ranged from rapidly declining to stable, depending on territory density and level of aggregation. Techniques that enable managers to maintain existing territories and create new ones are well established for this species. Thus managers may reasonably expect to maintain even small populations of Red-cockaded Woodpeckers by increasing the density, level of aggregation, and number of territories.

Key words: cooperative breeding; demographic stochasticity; environmental stochasticity; individual-based model; management; Picoides borealis; population viability analysis; Red-cockaded Woodpecker; spatially explicit model.

INTRODUCTION

The relative viability of animal populations is a critical conservation issue. The USDA Forest Service has a legal mandate to preserve viable populations of vertebrate species, and impacts on viability are commonly an issue in decisions made by the U.S. Fish and Wildlife Service about the degree to which proposed actions jeopardize populations of species protected under the Endangered Species Act. Managers are concerned with the relative ability of alternative management actions to promote viability of populations on the lands for which they are responsible. Conservationists are concerned with the threats to viability posed by global habitat loss and fragmentation. Lacking sufficient relevant empirical data, viability usually is evaluated through mathematical modeling, a process known as population viability analysis, or PVA (Boyce 1992, Beissinger and Westphal 1998).

The Red-cockaded Woodpecker (Picoides borealis) is an endangered species endemic to pine habitats of the southeastern United States. It is well studied, being the subject of three symposia (Thompson 1971, Wood 1983, Kulhavy et al. 1995), several review articles (e.g., Walters 1990, Jackson 1994), and numerous other publications. It is an unusual species, in that it is a cooperative breeder and excavates cavities for roosting and nesting in living pine trees. Its biology generally is well known, and management needs are well understood in comparison to those of other endangered species (Walters 1991, Jackson 1994, Conner et al. 2001). In contrast, assessment of the viability of Red-cockaded Woodpecker populations is deficient relative to work on other endangered species.

The viability assessment paradigm is particularly appropriate for the Red-cockaded Woodpecker because of its space requirements and distribution. Family groups occupy permanent territories 50–100 ha in size (Walters 1990, Jackson 1994). The species is highly habitat specific, and habitat loss has produced a highly discontinuous distribution of the species across the southeastern United States (USFWS 1985, James 1995). Thus the species exists largely in closed populations whose maximum size is limited, which is the population structure to which viability analysis is most relevant.

Populations face four threats to their persistence: demographic stochasticity, environmental stochasticity, natural catastrophes, and genetic uncertainty (Shaffer 1981, 1987). The current recovery plan for the Red-
cockaded Woodpecker (USFWS 1985) attempts to address the latter two threats, and subsequent work has improved on those attempts. Hurricanes are the primary catastrophic threat to Red-cockaded Woodpecker populations, but their effects can be minimized through management, specifically construction of artificial cavities (Copeyan 1990, Allen 1991, Copeyan et al. 1991) to replace cavities lost to storm damage (Watson et al. 1995). Methods for calculating the rate of loss of genetic variation have been developed (Reed et al. 1993) to replace the primitive methods used in the recovery plan (USFWS 1985). However, translating rates of loss of genetic variation into reductions in viability remains problematic (Lande 1988, 1995, Reed et al. 1993).

Conspicuously lacking are attempts to assess vulnerability to demographic and environmental stochasticity, the cornerstone of most viability analyses (e.g., Lacy and Clark 1990, Ryan et al. 1993, Beissinger 1995). Models of Red-cockaded Woodpecker demography exist (Heppell et al. 1994, Maguire et al. 1995), but the utility of these models in simulating population dynamics is limited because they do not incorporate the extreme spatial constraints on dispersal that characterize this species. Many males and some females, instead of dispersing in their first year to acquire breeding positions, remain on their natal territory as nonbreeding helpers (Ligon 1970, Lennartz et al. 1987, Walters et al. 1988, Walters 1990). Helpers eventually become breeders by inheriting a breeding position on the natal territory or dispersing to fill a breeding vacancy, but they only compete for breeding vacancies in the immediate vicinity of their home territory (Walters et al. 1988, Daniels 1997). Helpers represent a pool of replacement breeders and thus can act as a buffer between breeder mortality and population productivity, but only if they are within dispersal range of the territories on which breeders perish.

Vucetich and Creel (1999) advocate the use of complex models when there are large effects of social structure on population dynamics (see also Boyce 1992). Red-cockaded Woodpeckers clearly meet this criterion. Previous population models for Red-cockaded Woodpeckers are deterministic (Heppell et al. 1994) or stochastic (Maguire et al. 1995) stage-based matrix models. They represent sophisticated versions of the types of single-sex, single-population model most commonly used in population viability analysis (Beissinger and Westphal 1998). These Red-cockaded Woodpecker models have yielded important insights, but their authors, too, have advocated the use of more complex models to simulate population dynamics, specifically spatially explicit, two-sex models (Heppell et al. 1994, Maguire et al. 1995).

Spatially explicit, individual-based models, because they track the performance and fate of individual organisms (Judson 1994) and their locations (Dunning et al. 1995), can incorporate constraints on movement. Previously, we described such a model for Red-cockaded Woodpeckers and used it to show that the buffering effect of helpers was sufficient to prevent demographic stochasticity from producing fluctuations in the size of the breeding population, provided territories were sufficiently aggregated (Letcher et al. 1998). This result is of limited significance to conservation, because demographic stochasticity is a relatively minor source of variability compared to environmental stochasticity. Here we assess the power and consequences of the buffering effect of helpers under more realistic conditions, by considering environmental stochasticity as well as demographic stochasticity in a formal viability analysis. Specifically, we examine effects of territory number and territory distribution on viability within the range of population densities currently characterizing Red-cockaded Woodpeckers.

**METHODS**

**Model structure**

A major limitation of complex simulation models is the amount of data required for parameter estimation (Murdoch et al. 1992, Conroy et al. 1995, Beissinger and Westphal 1998). Fortunately, extensive, detailed demographic data for Red-cockaded Woodpeckers were available from 15 yr of study of >200 groups of color-marked birds in the North Carolina Sandhills (Carter et al. 1983, Walters et al. 1988, Daniels 1997). These data drove the construction of our model and, to a large extent, dictated its structure.

The model is described in detail in Letcher et al. (1998). Briefly, the landscape in the model consists of breeding territories and nonbreeding space that the birds must cross to find the territories. In this species, territories may be occupied for decades (Doerr et al. 1989), and new territories are created primarily through the process of budding, in which one territory (and its cavities) is divided into two (Hooper 1983). In the model, accordingly, territory locations are fixed, but territories are lost if unoccupied for more than five consecutive years. New territories are created through budding and each existing territory has a probability of 1% of producing a bud each year. However, a territory can bud only if sufficient, unoccupied space is available in its vicinity. Thus the effective budding rate is <1% at high densities. Budding rates reported from natural populations vary from near zero to 2% (Doerr et al. 1989, Conner et al. 2001).

The model is configured for both males and females: female fledglings disperse in their first year, whereas male fledglings may either disperse or become helpers. In the model, dispersing fledglings move in a random direction and continue in that direction at a specified rate until they die, obtain a breeding position, or leave the population. At each time step, they compete for any breeding vacancy within 3 km of their current position. Males may compete for both breeding vacancies and empty territories, but females may only move into ter-
territories that contain a male. Helpers do not change status or location until they die or fill a breeding vacancy. The oldest helper inherits breeding status if the breeder on its territory dies and helpers compete for any breeding vacancies within 3 km of their home territory. The probability of mortality of each individual depends on its status. Mortality rates and helper dispersal behavior are well documented (Walters et al. 1988, 1992, Daniels 1997), as are dispersal probabilities and dispersal distances of fledglings (Walters et al. 1988, Daniels and Walters 2000), but the behavior of fledglings while dispersing is not.

The number of fledglings produced on each territory each breeding season is a function of breeder age and number of helpers present. In this species, young birds (i.e., <4 yr old) are less productive than older ones (Walters et al. 1988, 1992, Walters 1990), and helpers increase the productivity of their group (Lennartz et al. 1987, Walters 1990, Heppell et al. 1994).

Initial conditions

For each model run, the landscape contained a predetermined number and density of territories arranged according to a predetermined level of clumping. The territories were added as circles that ranged in radius from 0.3 to 0.5 km, with the final size being dependent on territory density (for additional details, see Letcher et al. 1998). The same initial conditions were applied randomly to each territory. Each territory had a 90% chance of beginning with a breeding pair, and territories without a breeding pair contained a solitary male. A number of helpers equal to one-half the number of territories was added randomly to the territories with pairs, so that about one-half of the territories had no helpers, and a few had more than one. The ages of the birds also were assigned randomly, but from a distribution designed to reproduce the age distribution of birds observed in the North Carolina Sandhills population in 1991, a typical year (J. R. Walters, J. H. Carter III, and P. D. Doerr, unpublished data). These initial conditions were independently applied to each run of each scenario. Thus variation among replicates was due to variation in the initial population, as well as variation in simulation dynamics.

Demographic stochasticity was incorporated into the simulations by applying to each individual annual survival probabilities, annual status transition probabilities of male fledglings, and probabilities of producing different numbers of fledglings. This was accomplished by drawing a deviate from a random uniform distribution and determining whether or not the deviate was less than the appropriate probability value. Environmental stochasticity was incorporated by varying survival probabilities and probabilities of producing different numbers of fledglings each year. This was accomplished by determining the variance among years in these parameters across 14 yr of data from the North Carolina Sandhills, and then drawing randomly each year from the resulting distribution to determine that year's probability value. This procedure was applied independently to each parameter, as we found no covariance in survival probabilities among status classes or between survival and productivity in the North Carolina Sandhills data.

Simulation scenarios

We ran 100 replicates, each of 100 yr duration, for each simulation scenario. We first simulated populations of five different sizes (25, 49, 100, 250, and 500 initial territories), each at two different levels of aggregation, on a landscape 32 × 24 km in size. The landscape size was chosen to match the range of densities observed among existing populations. Specifically, the 49 and 100 territory scenarios match current densities of two coastal North Carolina populations and the 500 territory scenario corresponds to the recommended density in recent management guidelines. The two levels of aggregation span the range of variability in clumping among real populations.

A random distribution of territories on x–y axes constituted one level of aggregation. We used the k parameter of the negative binomial distribution to generate the second level of aggregation. For a particular number of territories distributed across the landscape, we determined the k values when territories were minimally (i.e., randomly distributed, maximum k) aggregated and maximally (i.e., aggregated in one large patch, minimum k) aggregated. To create the second level of aggregation, we selected a k value that was 25% of the distance between log kmin and log kmax. We then generated spatial distributions of territories using those 25% k values, which represent a fairly high degree of clumping (Fig. 1).

In the first set of simulations, population size and territory density covary. To examine the effects of density independent of the effects of population size and aggregation, we conducted a second set of simulations. In the second set, we simulated population dynamics of the smaller populations (25, 49, and 100 territories) at each of the densities characterizing the larger populations (250 and 500 territories) in the first set of simulations. This was accomplished by changing the size of the landscape to produce the required overall density (Fig. 1). In the second set of simulations, territories were randomly distributed on the landscapes (i.e., the maximum k value was used).

For each simulation scenario, we measured the mean and variance among the 100 replicates in annual population growth rate and the percentage of the initial territories remaining after 100 yr. The latter was calculated as the number of surviving original territories plus the number of surviving territories formed by budding, divided by the number of original territories. We also calculated the number of replicates in which extinction, defined as loss of all territories, occurred, and the mean and distribution of time to extinction.
**RESULTS**

Population growth rate varied with both territory number and territory distribution (Fig. 2). The mean growth rate was close to 1.0 for populations of 250 and 500 territories, for both levels of aggregation. Although the mean growth rate of populations of 25, 49, and 100 territories was <1.0, growth rate among these populations was highly dependent on the level of aggregation. For example, the mean growth rate of populations of 100 randomly distributed territories was less than the mean growth rate of populations of 50 more aggregated territories (Fig. 2). Population density also had a pronounced effect on population growth rate. Mean growth rates of small populations were considerably higher under both high-density conditions. In fact, the mean growth rate of populations of 100 reached 1.0 at the highest density (Fig. 3).

The average number of surviving territories after 100 yr was as high or higher than the initial number for populations of 250 and 500 initial territories at all densities, for populations of 100 initial territories at the two highest densities, and for populations of 50 initial territories at the highest density (Figs. 4 and 5). For populations of 25, 49, and 100 initial territories, initial population size, aggregation, and density all had pronounced effects on the average number of surviving territories.

No populations of 250 or 500 groups went extinct at either level of aggregation. For smaller populations, the number of simulations in which extinction occurred varied with population size, aggregation, and density (Table 1). All populations of 25 and 49 groups went extinct quickly at the lowest level of aggregation, but populations of these sizes survived much longer, sometimes for 100 yr, when density or aggregation was higher (Table 1, Fig. 6). Most populations of 100 groups went extinct at the lowest level of aggregation, whereas all survived at the highest density (Table 1).

**DISCUSSION**

**Effects of population size and spatial distribution**

Our simulations resulted in two major findings. First, distribution and density of territories have as large an effect on population behavior as population size, at least within a certain range of population sizes (i.e., fewer than 250 territories). Second, effects of environmental stochasticity are relatively small compared to those evident in viability analyses of other species (Beissinger 1995, Wisdom and Mills 1997, Beissinger and Westphal 1998); as a result, relatively small populations could be relatively stable. Indeed, the behavior of populations in this study was quite similar to that of comparable populations in a previous analysis in which only effects of demographic stochasticity were considered (Letcher et al. 1998). These results can be related to the buffering effect of helpers on population
dynamics. The presence of a substantial nonbreeding class ameliorates the impact of stochastic variation in mortality and reproduction on the size of the breeding population. Such variation leads to fluctuations in the size of the helper class rather than to changes in the number of territories occupied. Evidently, the capacity of the helper class to absorb stochastic variation is sufficient to absorb not only demographic stochasticity, but also environmental stochasticity, at least down to a population size of 50 groups. However, the buffer effect depends on territories being within the dispersal range of helpers of one another. Hence its capacity is determined by population density and aggregation, as well as size. Vucetich et al. (1997) describe a similar

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**Fig. 2.** Annual population growth rate as a function of number of territories and distribution. The mean and standard deviation among 100 replicates are indicated for each condition; $k_{max}$ indicates random distribution of territories, and $k_{25}$ a more aggregated distribution (see Methods: Simulation scenarios for additional description).

**Fig. 3.** Annual population growth rate as a function of territory number and density. The mean and standard deviation among 100 replicates are indicated for each condition. The lower density ($d_{250}$) corresponds to 3.26 territories/km², the density of 250 randomly distributed territories in the first set of simulations ($k_{max}$ in Fig. 2). The higher density ($d_{500}$) corresponds to 6.51 territories/km², the density of 500 randomly distributed territories in the first set of simulations. Results for randomly distributed territories at the various densities examined in the first set of simulations are shown for comparative purposes ($k_{max}$).
Fig. 4. Percentage of initial territories surviving after 100 yr as a function of initial number of territories and distribution. The mean and standard deviation among 100 replicates are indicated for each condition; $k_{\text{max}}$ indicates random distribution of territories, and $k_{\text{agg}}$ indicates a more aggregated distribution.

Fig. 5. Percentage of initial territories surviving after 100 yr as a function of initial territory number and density. The mean and standard deviation among 100 replicates are indicated for each condition. The lower density ($d_{250}$) corresponds to 3.26 territories/km²; the density of 250 randomly distributed territories in the first set of simulations ($k_{\text{max}}$ in Fig. 4). The higher density ($d_{500}$) corresponds to 6.51 territories/km², the density of 500 randomly distributed territories in the first set of simulations. Results for randomly distributed territories at the various densities examined in the first set of simulations are shown for comparative purposes ($k_{\text{max}}$).
**Table 1.** Number of simulations out of 100 in which extinction of Red-cockaded Woodpecker populations occurred, and mean time to extinction as a function of territory number and distribution.

<table>
<thead>
<tr>
<th>No. territories</th>
<th>Distribution†</th>
<th>No. extinctions</th>
<th>Time to extinction (yr) (mean ± 1 SD)</th>
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<td>25</td>
<td>$k_{\text{max}}$</td>
<td>100</td>
<td>24 ± 6</td>
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<tr>
<td>25</td>
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<td>$d_{250}$</td>
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<td>25</td>
<td>$d_{584}$</td>
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<td>49</td>
<td>$k_{\text{max}}$</td>
<td>100</td>
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<td>$k_{250}$</td>
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<td>$d_{250}$</td>
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<td>$d_{584}$</td>
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<td>78 ± 20</td>
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<td>4</td>
<td>84 ± 12</td>
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<td>100</td>
<td>$d_{584}$</td>
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† Here, $k_{\text{max}}$ and $k_{250}$ indicate degree of aggregation of territories, $k_{\text{max}}$ representing a random distribution, and $k_{250}$ a more aggregated than random distribution. $d_{250}$ and $d_{584}$ indicate territory density, corresponding to densities of 250 and 500 territories per 762 km², respectively. See Methods: Simulation scenarios for details.

**Fig. 6.** Distribution of time to extinction for simulations in which extinction occurred as a function of number of territories and distribution, where $k_{\text{max}}$ indicates random distribution of territories within a 32 × 24 km landscape; $k_{250}$, a more aggregated distribution within this same landscape; $d_{250}$, a density of 3.26 territories/km² (equivalent to $k_{\text{max}}$ for 250 territories), and $d_{584}$, a density of 6.51 territories/km² (equivalent to $k_{\text{max}}$ for 500 territories).
appears to be relatively uncorrelated among status classes (Fig. 7). The range of variation observed in the Sandhills (Fig. 7) is likely to encompass all but the most severe catastrophic events. When the Camp Lejeune population in coastal North Carolina was struck by two hurricanes between the 1996 and 1997 breeding seasons, mortality of breeding males that year was 29%, compared to an average value over the previous 10 yr of 18% (J. R. Walters, unpublished data). During that same year, mortality of helper males was 39% compared to an average value of 23%, and that of breeding females was 31% compared to an average value of 24%. Mortality of fledglings was not affected. Hurricane Hugo, however, caused mortality well beyond this range when it struck the population on the Francis Marion National Forest in 1989 (Hooper et al. 1990).

How realistic are the densities that we examine? We have been involved in field studies of Red-cockaded Woodpeckers at four locations in recent years: Eglin Air Force Base in the Florida panhandle; Croatan National Forest and Camp Lejeune Marine Base in coastal North Carolina; and the Sandhills population in south-central North Carolina. The densities of the two coastal populations in North Carolina lie between those of our theoretical populations of 49 and 100, and these populations are ~250 territories in size. Territories in the real populations are more aggregated than random because of the distribution of suitable habitat.

Densities of most existing populations fall within the range represented by our theoretical populations of 49–250, and their level of aggregation probably is comparable to our k_{500}. Recommended densities in suitable habitat in current management documents correspond to populations of 475–950 territories on our 32 × 24 km landscape (USFWS 1985, USDA 1995, U.S. Army 1996). Our results suggest that even small populations (i.e., 25 territories) might be fairly persistent at these densities, but achieving these densities will be constrained by the proportion of suitable habitat in the total landscape. The higher density levels that we examined, however, should be achievable on most landscapes. These correspond to roughly one territory per 300 ha (our 250 territories) and one territory per 160 ha (our 500 territories).

Our population projections should not be taken literally until the model has been more thoroughly validated. The model is subject to a variety of inaccuracies that make it inappropriate to interpret our results in absolute terms. Some parameters may be inaccurate, notably dispersal speed and search range of fledglings. Considerable controversy exists over the sensitivity of spatially explicit models to errors in estimation of dispersal parameters (Wennergren et al. 1995, Ruckelshaus et al. 1997, 1999, Mooij and DeAngelis 1999, South 1999). Even those parameter values based on sufficient data come from a single population and thus may not be representative of all populations. Mortality rates are known to be lower in coastal North Carolina (J. R. Walters, unpublished data) and central Florida (DeLotelle et al. 1995) than in the North Carolina Sandhills, for example. Especially important in this respect is helper dispersal. Although helper search range is well documented for the North Carolina Sandhills (Walters et al. 1988, Daniels 1997), and helper life history is constant across populations, it is not known whether helper search range changes with population density. If it increases when population density is low, as seems likely, stability will decrease less, with decreasing density, than our simulations suggest. Fortunately, model performance is not highly sensitive to any of these parameters (Letcher et al. 1998). Therefore, we believe that the strong effects of density and aggregation evident in our results are real.

Implications for management

The obvious implication of these findings is that management actions that increase the aggregation and density of territories of this endangered species will be beneficial; those that increase the isolation of territories will be harmful. Although this may be viewed as a general principle, the benefits of improved spatial structure will only be manifest when the assumptions
of our analyses are met. Most importantly, we assume that territories will not be lost to habitat degradation, and that vital rates are not depressed by adverse anthropogenic factors. That is, we assume that the important conservation problems currently are those inherent to small, isolated populations (i.e., the small population paradigm; Caughley 1994) rather than the environmental problems that caused the species to become endangered (i.e., the declining population paradigm; Caughley 1994). This may seem a questionable approach for an endangered species, but in this case, there is ample justification. As recently as the late 1980s, most populations of Red-cockaded Woodpeckers were declining and the healthiest ones, with one exception (Hooper et al. 1991), were stable rather than increasing (Conner and Rudolph 1989, Costa and Esco 1989). Clearly, the declining population paradigm was the appropriate one in this era. However, breakthroughs in research and management provided an understanding of the factors driving populations to extinction and the means to combat them (Walters 1991, Conner et al. 2001). In the 1990s, well-managed populations, even small ones, have increased (Franzreb 1997, Conner et al. 2001). Our results apply to these well-managed populations. Where the critical environmental problems have yet to be addressed, populations will perform more poorly than suggested by our model projections because of violations of our assumptions about habitat quality and vital rates.

Especially relevant among the current management tools are prescribed growing-season fire to control the hardwood midstory, and construction of artificial cavities to replace cavities lost to wind damage, fire, lightning strikes, and other forces. In combination, these tools can eliminate the forms of habitat degradation that cause territories to be abandoned (Walters 1991). Also, construction of artificial cavities in unoccupied areas stimulates formation of new groups on new territories (Copeyon et al. 1991). Population growth rates of 5–10% can be achieved with artificial cavity construction, whereas growth rates through natural processes are only 1–2% (Conner et al. 2001). Thus populations may perform better than our model projections where artificial cavity construction is employed effectively. This technique is ideal for improving the aggregation and density of territories, and thereby also improving population behavior.

One possible concern in increasing the aggregation of territories is that vulnerability to catastrophe may increase. Hurricanes are the only significant catastrophic threat to Red-cockaded Woodpecker populations, but the threat is serious (Hooper et al. 1990, Hooper and McAdie 1995). In fact, Hooper and McAdie (1995) recommend against aggregating groups to reduce hurricane impacts. However, the scale of hurricane damage is much larger than the scale at which aggregation affects population behavior. Whether woodpecker territories are aggregated or scattered within a hurricane's path will have little effect on the damage sustained. More important are managing habitat to reduce vulnerability to wind damage (Hooper and McAdie 1995) and countering any losses of cavity trees by constructing artificial cavities. Through these means, impacts of hurricanes on Red-cockaded Woodpecker populations have been minimized in recent years (Conner et al. 2001). We conclude that the benefits of territory aggregation to population dynamics greatly outweigh any costs due to increased vulnerability to catastrophe.

The relative stability in our simulations of fairly small populations with favorable spatial distributions, coupled with the existence of management tools that can increase stability, offer hope that even small populations of Red-cockaded Woodpeckers can be maintained. There is empirical evidence that this hope might be realistic. Although many small populations have declined, some isolated populations of only 25 or even 10 territories have been remarkably persistent (James 1995). Our assessment of these data is that persistence is associated with high levels of territory aggregation.

Additional empirical evidence of the importance of spatial distribution of territories is emerging from well-managed populations. Where habitat is fragmented and, as a result, population density is low, populations tend to decline (Conner and Rudolph 1991). In large populations, territories tend to be lost at the edges of the population and gained (through budding) in the center. This has occurred, for example, at Eglin Air Force Base in Florida (J. R. Walters and K. E. Gault, unpublished data) and in the North Carolina Sandhills (J. H. Carter III, J. R. Walters, and P. D. Doerr, unpublished data) in recent years.

Population viability standards

Regardless of its advisability in theory, in practice, agencies routinely set population size objectives based on assessments of viability. Current management guidelines for the Red-cockaded Woodpecker stipulate that a population size of ~400 territories is required for viability (USDA 1995, U.S. Army 1996). This standard is based solely on projected rates of loss of genetic variability (USFWS 1985, Reed et al. 1993). The basis of such a criterion is sufficiently suspect (Lande 1988, 1995, Reed et al. 1993) that maintaining sufficiently large, closed populations should be abandoned in favor of other means to maintain genetic variability, such as translocation of individuals between populations. Translocation techniques are well developed for this species (DeFazio et al. 1987, Rudolph et al. 1992). Demographic stability is much more critical, but no demographic viability standard has been proposed previously, because of the limitations of previous models in projecting population dynamics. Our results do not suggest a strict population size standard. They do suggest that it may be possible to maintain populations much smaller than the previous standard, and that in-
creasing territory aggregation, density, and number are all important.

Only a few Red-cockaded Woodpecker populations are as large as the largest size classes (250–500 territories) that we analyzed, but many are of comparable size to the smaller size classes (25–250 territories) that we considered, and some are even smaller (James 1995). Our results suggest that the largest populations (i.e., 250–500 territories) can readily be maintained and that those of intermediate size (i.e., 49–250 territories) might be as well, if densities and levels of aggregation are favorable. In some cases, but perhaps not all, judicious management will be required to achieve stability. The critical point is that, over a wide range of population sizes and realistic levels of aggregation and density, projected population growth rates remained within the range in which losses can reasonably be countered by available management techniques. The size at which populations performed well over a range of reasonable densities, 100 territories, is considerably smaller than is typical of animal species. This provides reason to be optimistic about recovery efforts for the species, considering that large populations (i.e., 500 or even 250 territories) are not achievable on many management units.

Very small populations (i.e., ≤25 territories) present a more difficult problem. Many will probably decline faster than can be counteracted by management. Still, they may not be doomed if high levels of aggregation can be achieved. Indeed, when we simulated very small populations, we found that even populations of as few as 10 territories could be sufficiently stable to be maintained through management if territories were maximally aggregated (L. B. Crowder, J. A. Priddy, and J. R. Walters, unpublished report). Management of isolated groups seems a poor investment, but management of small aggregations of groups may not be.

In conclusion, our results suggest that populations of Red-cockaded Woodpeckers are less prone to the stochastic fluctuations in size that plague efforts to preserve isolated populations of many other species. This provides managers with additional incentive to invest in their populations, however small, as it increases the chances that their efforts will produce demonstrable benefits. Seldom does the fate of a species depend so much on how effectively appropriate management is implemented.

ACKNOWLEDGMENTS

We thank Selina Heppell and Karin Schiegg for helpful discussion, and Benjamin Letcher for his initial work on the simulation model. We also acknowledge Jay Carter, Phillip Doerr, and Kerry Brust, whose ongoing efforts have contributed greatly to the production of the North Carolina Sandhills data set on which we have relied in our modeling efforts.

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