HOME RANGES OF RUSTY BLACKBIRDS BREEDING IN WETLANDS: HOW MUCH WOULD BUFFERS FROM TIMBER HARVEST PROTECT HABITAT?

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Abstract. We calculated the home ranges and core areas of 13 adult Rusty Blackbirds (Euphagus carolinus) in Maine to determine (1) the area requirements of breeding adults, (2) whether area requirements of the sexes and of colonial and noncolonial individuals differ, and (3) the proportion of the home range and core area that would be protected by a buffer of no logging of 50–100 m around occupied wetlands. Mean home ranges (37.5 ± 12.6 ha) and core areas (11.1 ± 2.8 ha) were large in comparison to those of other breeding icterids, and adults often foraged in multiple unconnected wetlands. Rusty Blackbirds that were part of a loose colony had home ranges and core areas three times larger than those of pairs that nested solitarily, which we speculate may be due to adults following one other to feed on unpredictable emergences of aquatic insects. Home ranges and core areas included a surprisingly small amount of wetland habitat, only 12% and 19% respectively, but adults often foraged in small wet patches (<16 m²) in otherwise upland habitat. The 75-m buffers around wetlands that we recommended in a concurrent study may help protect the Rusty Blackbird’s nesting habitat, but such buffers contained less than half the average home range, suggesting that they may be of only limited benefit as a conservation strategy for protecting foraging habitat.

Key words: boreal wetlands, Euphagus carolinus, home range, radio telemetry, Rusty Blackbird, wetland buffers.

INTRODUCTION

Understanding how animals use landscapes is critical in formulating strategies to conserve their habitats (Storch 1995, Caro 1999, Powell and Bjork 2004). Home range is defined as the area an individual uses during a specific period of its life (Burt 1943) and can be useful in comprehending a species’ social structure, habitat, and area requirements (Bingham and Noon 1997). Understanding home ranges also can be useful for determining the size of buffers necessary to protect important habitats from human disturbance (Milam and Melvin 2001).
Our understanding of the breeding ecology of the Rusty Blackbird (*Euphagus carolinus*) has improved markedly since Greenberg and Droge (1999) first publicized the species’ decline (Matsuoka et al. 2010a, Powell et al. 2010a), yet we still know little about breeding birds’ social organization, area requirements, or how the quality and distribution of foraging habitat affects each of these. Rusty Blackbirds breed exclusively in boreal wetlands, where they occur at low densities, nest predominately in conifers (Matsuoka et al. 2010a, Powell et al. 2010a), and forage on aquatic insect larvae and crustaceans by probing in vegetation in or near shallow water (Forbush 1927, Ellison 1990, Avery 1995). Although breeding Rusty Blackbirds have been commonly observed taking flights of several hundred meters (Machtans et al. 2007, Matsuoka et al. 2010b), there is no prior information on their home ranges. Orians (1985) described the species as loosely colonial, yet there is only one such report from New England (Powell et al. 2010b). Consequently, it is unclear if area requirements vary with social organization.

Powell et al. (2010a) found that when Rusty Blackbirds nested in wetlands with recent timber harvests, they suffered 2.5x more nest predation than when they nested in wetlands without recent timber harvests. Retaining unlogged buffers in upland habitats around wetlands may help mitigate these negative effects on reproduction (Powell et al. 2010a). Spatial patterns in breeding blackbirds’ use of wetlands may reveal the potential for upland habitat buffers around wetlands to minimize disturbance of wetlands in which the birds nest or forage. In this study, we radiotracked adult Rusty Blackbirds during the breeding season in northern Maine to (1) estimate sizes of home ranges and core areas, (2) compare home ranges and core areas by sex and of colonial and noncolonial individuals, and (3) determine the proportion of the average home range and core area that would be protected by buffers of 50, 75, and 100 m of unharvested upland habitat around occupied wetlands.

**METHODS**

**STUDY AREA**

We captured, marked, and radiotracked Rusty Blackbirds at four sites located between 15 and 28 km of Moosehead Lake and within industrially managed forests in Somerset and Piscataquis Counties, Maine. Site A was centered on a 70-ha, 13-year-old stand of regenerating spruce (*Picea* sp.) and balsam fir (*Abies balsamea*) that included a loose colony of Rusty Blackbirds described in detail by Powell (2010b; Fig. 1). This stand contained two 0.6-ha scrub-shrub wetlands (Cowardin et al. 1979), a well-maintained gravel road, and many wet patches of *Sphagnum* sp. A hill just east of the regenerating stand was covered by a mosaic of older (>40 years) conifers, many recent partial cuts dominated by raspberry brambles (*Rubus* sp.), and a 1.3-ha forested wetland dominated by northern white cedar snags (*Thuja occidentalis*). Site B was centered on three small and shallow vernal pools (0.7 ha total, <20 cm deep) with gravel bottoms that originated from borrow pits created during road construction (Fig. 2a). Rusty Blackbirds often foraged in these vernal pools, which contained many tadpoles (L. L. Powell, pers. obs.). Wetland vegetation was dominated by speckled alder (*Alnus rugosa*); the uplands included red spruce (*P. rubens*), balsam fir, paper birch (*Betula papyrifera*), and many recent partial cuts.

Site C (Fig. 3) was centered along a 1-m-wide stream that connected a complex of terraced wetlands impounded by beaver (*Castor canadensis*). An abandoned logging road, overgrown with vegetation, bisected this site. Wetland vegetation included black (*P. mariana*) and red spruce, speckled alder, and *Sphagnum* sp. The surrounding forest was composed primarily of 15-year-old regenerating spruce and fir 3–4 m tall.

Site D was centered along a well-maintained and active logging road that bisected the two wetlands regularly used by Rusty Blackbirds (Fig. 2b). South of the road, 16- to 18-year-old spruce and northern white cedar dominated a >50-ha regenerating fen with *Sphagnum* sp. and puddles of open water. North of the road, a 20-m-wide strip of mature spruce, balsam fir, and paper birch buffered a 1-ha emergent and forested wetland. The surrounding forest had been partially harvested within the last three years.

**FIGURE 1.** Home ranges of four male Rusty Blackbirds at a loose colony, site A, in Maine, 2007. Triangles represent active nests, white space represents forested upland, and polygons of different shades represent individual males’ home ranges. Females are not shown, nor are two isolated relocations of the male with the black home range. The two lighter gray polygons did not reach an asymptote.
From late May to mid-June 2007, we captured adult Rusty Blackbirds in mist nets placed near nests or wetlands used for foraging (Powell 2008). We banded eight males and seven females each with a U.S. Geological Survey aluminum band and a unique combination of colored leg bands. We fitted each bird with a radio transmitter (1.6 g for females, 1.9 g for males; <3% of body weight) from Holohil Systems Ltd. (Ontario; model BD-2) by means of a cotton-thread harness (Rappole and Tipton 1991). Following each capture, we waited at least 24 hr before recording locations to allow each bird adequate time to adjust to the transmitter and bands. We relocated the birds from 7 June to 11 July 2007.

We radiotracked each individual for a minimum of 3 non-consecutive days in a 5-day period. The one exception was a female that we radiotracked intensively for one day (05:20–15:00). We included data from this bird in our statistical analyses because her home range reached an asymptote. We categorized the daylight hours into five equal time blocks, randomized which time block to target for a given radio-marked bird, and then relocated the target bird at 10-min intervals. We chose the latter to provide biological independence of locations by ensuring “a sampling interval long enough to allow the animal to move from any point in its home range to any other point” (Lair 1987:1099). Rusty Blackbirds can fly approximately 5.7 km in 10 min (Wood 1933), more than twice as far as the diameter of the largest home range we identified (Fig. 1). Thus our sampling design controlled for diurnal patterns in bird activity over the study period and minimized autocorrelation between consecutive relocations (Otis and White 1999).

During each 10-min interval, two or three technicians each used a hand-held Yagi antenna and simultaneously recorded (1) the compass direction of the strongest radio signal from the bird that they were tracking and (2) the location

FIGURE 2. Home ranges of three Rusty Blackbirds at (a) site C and (b) site D, in Maine, 2007. Triangles represent active nests, white space represents forested upland, and bold black and gray polygons represent home ranges of one male and two females, respectively. In (a), the dashed-line polygon represents the female’s core area (76% KDE), and the open water resulted from borrow pits created during road construction. In (b), the male’s home range did not reach an asymptote.

FIGURE 3. Home range of a breeding pair of Rusty Blackbirds at site B, in Maine, 2007. The triangle represents the pair’s nest, white space represents forested upland, and gray contour lines represent buffers of 50, 75, and 100 m around wetlands. Bold black and gray polygons represent the home ranges of the male and female, respectively.
where they received it with a hand-held GPS receiver. Technicians arranged themselves so that bearings were as close to 90° from each other as feasible and as close to target birds as possible without noticeably affecting behavior. We triangulated bearings with LOCATE III (Nams 2006). We tested for observer error in measuring compass bearings to radio-marked birds by randomly placing a test transmitter in and around wetlands used by radio-marked Rusty Blackbirds. Naïve observers then recorded bearings to the test transmitter by the same techniques described above.

HOME RANGE AND CORE AREA
We visually inspected each individual's locations in chronological order and then removed locations representing obvious range shifts to meet the assumption that individuals show fidelity to a given area over the course of the sampling period. We estimated 95% fixed-kernel densities (KDE; referred to hereafter as “home range”) with Home Range Tools for ArcGIS (Rodgers et al. 2007). We estimated fixed rather than adaptive kernels on the basis of Seaman and Powell's (1996) conclusion that fixed kernels outperform adaptive kernels. We used the Gaussian (bivariate normal) kernel form and a least-squares cross-validation for automated bandwidth selection. Kernels had a grid-cell resolution of 10 m, and contouring was performed by volume. We used a scaling factor of 1,000,000 and rescaled home ranges to unit variance. We calculated core areas for each individual with Powell's (2000) objective and area-independent method (Laver and Kelly 2008) in ABODE for ArcGIS (ESRI 2006, Laver 2006). Rather than arbitrarily using 50% KDE, we used ABODE to calculate the probability of use for each cell of the KDE and defined the core range as the area in which the probability density was significantly greater than expected by a random distribution.

We used ABODE for ArcGIS (ESRI 2006, Laver 2006) and Laver's (2005) recommendations to determine the number of relocations at which home-range size reached an asymptote. We recalculated each individual's home range after the addition of each randomly added location and repeated this procedure 10 times for each home range. We considered the home range to have reached an asymptote when the confidence interval fell within 20% of the final home-range size for five consecutive points.

BUFFERS AROUND WETLANDS
We used a digital coverage of the National Wetlands Inventory (NWI; Cowardin et al. 1979) in ArcGIS 9.2 (ESRI 2006) to examine whether buffers around wetlands might be useful for protecting Rusty Blackbird breeding habitat. We first verified the accuracy of the NWI on the ground and then used ArcGIS to add previously unmapped beaver-impounded wetlands to site C (Fig. 3). We then calculated the percentage of each Rusty Blackbird’s home range and core area that was encompassed by wetlands alone and by wetlands with buffers of 50, 75, and 100 m around their perimeters. We selected these three widths (Fig. 3) on the basis of the following justifications: (1) 50 m, because rates of predation on bird nests are highest within 50 m of habitat edges (Paton 1994); (2) 75 m, because Vander Haegen and Degraaf (1996) reported the effects of predation on forest birds extend 75 m into riparian buffers in Maine, and because 75 m was the minimum buffer suggested by Powell et al. (2010a) for breeding Rusty Blackbirds; and (3) 100 m, because 43 Rusty Blackbird nests in northern New England found from 2006 to 2008 were 0–95 m away from wetlands (Powell et al. 2010a).

STATISTICAL ANALYSES
We log-transformed home range and core area as needed to meet the assumptions of normality (Shapiro–Wilk tests) or equal variances (Levene's tests) and then ran t-tests (α = 0.10) for equal variances in Program R (R Development Core Team 2009) to test for differences in mean home-range and core-area size by sex and coloniality. Similarly, we ran paired t-tests for equal variance to test for differences between the sexes in the size of home ranges and core areas of breeding pairs. We present all estimates ± SE unless otherwise noted.

RESULTS
We captured and radiotagged 15 adult Rusty Blackbirds for which we recorded 493 relocations at four sites. We calculated home ranges for the 13 individuals (6 males and 7 females) for which we had ≥25 relocations (̄x = 38 ± 3 relocations). Individuals pecked at their leg bands for the first few hours after capture, but we did not observe any evidence that leg bands or transmitters impeded the ability of adults to fly, forage, or feed young. We collected 54 bird-days of data: three during incubation, 11 during the nestling stage, 25 during the fledging stage (defined as ≤6 days after fledging), and 15 on one male and one female at site A whose breeding status we could not determine. We excluded all data from 8 to 11 July because family groups began to wander during this time (Palmer 1949) and violate the assumption of site fidelity. Thus, we used only relocations from 7 to 26 June in quantifying home ranges. The mean angular error of triangulated locations on the test transmitter was 13.2° (SD = 9.0, n = 38 bearings).

The mean size of the home ranges of all 13 individuals was 37.5 ± 12.6 ha (range 3.8–172.8 ha), and the mean core area was 11.1 ± 2.8 ha (range 1.5–34.8 ha). Powell's (2000) method for estimation of the core area produced a mean of 63 ± 3% KDE, which was considerably larger than the arbitrary 50% KDE used in most core-area analyses (Laver and Kelly 2008). Radio-marked birds included an average of 2.8 ± 0.4 wetlands within their home ranges and 1.4 ± 0.3 wetlands within their core areas. The size of the home range of the four individuals whose home-range size did not reach an asymptote (̄x = 17.7 ± 3.1 ha) was not statistically different from that of the nine individuals whose home-range size did reach an asymptote.
The average percentage (± 1 SE) of the 95% fixed-kernel home ranges and core areas of 13 Rusty Blackbirds that included wetlands and wetlands surrounded by buffers of 50–100 m. Rusty Blackbirds were tracked by radio telemetry on the breeding grounds in Maine, 2007.

\( \bar{x} = 46.3 \pm 17.6 \text{ ha}; t_{11} = -0.64, P = 0.53 \). Furthermore, number of relocations to asymptote (\( \bar{x} = 31.6 \pm 2.4 \) points) was similar to total number of relocations we recorded for the birds for which an asymptote was not reached (\( \bar{x} = 29.3 \pm 1.4 \) points). Because of the similarity in those groups and our small sample size, we pooled data for home ranges that did and did not reach an asymptote.

The sizes of males’ home ranges were variable (\( \bar{x} = 48.0 \text{ ha} \pm 26.2 \), range 3.8–172.8 ha, \( n = 6 \)) and not different from that of females (\( \bar{x} = 28.4 \pm 8.1 \) ha, range 4.4–71.6 ha, \( n = 7 \); \( t_{11} = -0.11, P = 0.91 \)). Among breeding pairs (\( n = 5 \)), the mean size of the home range of the male (\( \bar{x} = 23.1 \pm 8.9 \) ha) again was not different from that of the female (\( \bar{x} = 22.0 \pm 4.6 \) ha; \( t = 0.12, P = 0.91 \)). Size of the core area did not differ by sex for all birds pooled (\( \bar{x}_{\text{male}} = 12.0 \pm 5.5 \) ha; \( \bar{x}_{\text{female}} = 10.3 \pm 2.6 \) ha; \( t = -0.30, P = 0.77 \)) or within pairs (\( \bar{x}_{\text{male}} = 7.4 \pm 3.3 \) ha; \( \bar{x}_{\text{female}} = 8.2 \pm 2.0 \) ha; \( t = -0.17, P = 0.87 \)). Home ranges of colonial-nesting birds (\( \bar{x} = 51.8 \pm 18.8 \) ha, \( n = 8 \)), however, were on average 3.5× larger than those of noncolonial birds (\( \bar{x} = 14.6 \pm 5.8 \) ha, \( n = 5 \); \( t_{11} = -2.5, P = 0.03 \)). The mean size of the core area of colonial birds (\( \bar{x} = 14.9 \pm 3.8 \) ha) also was 3× larger than that of noncolonial birds (\( \bar{x} = 4.9 \pm 1.5 \) ha; \( t_{11} = -2.0, P = 0.07 \)).

The wetlands mapped by NWI encompassed on average only 12 ± 3% of home ranges and 19 ± 6% of core areas (Fig. 4). However, adding buffers around wetlands increased this area substantially; for example, wetlands with a 75-m buffer encompassed 44 ± 7% of home ranges and 51 ± 9% of core areas (Fig. 4).

**DISCUSSION**

**HOME RANGE AND THE EFFECTS OF SEX AND COLONIALITY**

Rusty Blackbirds had mean home ranges and core areas of 37.5 and 11.1 ha, respectively, but individuals varied considerably in the amount of space they used (home range 4–179 ha; core area 2–35 ha). Breeding Rusty Blackbirds appear to require more space than breeding Red-winged Blackbirds (*Agelaius phoeniceus*) or Yellow-headed Blackbirds (*Xanthocephalus xanthocephalus*), which generally have territories of <0.6 ha (Gori 1988, Searcy and Yasukawa 1995; but see Anich et al. 2009). Rusty Blackbirds likely had large home ranges because adults often used multiple wetlands for foraging (range 1–5; Fig. 1–3). We found no difference between the size of males’ and females’ home ranges or core areas, possibly because of the small sample size or wide variation among individuals.

Despite the variation in space use, home ranges and core areas of Rusty Blackbirds in the loose colony were approximately 3× larger than those of noncolonial individuals. Horn (1968) suggested that Brewer’s Blackbirds (*Euphagus cyanocephalus*), which also have a variable social organization, nest colonially to take advantage of each other’s success in finding spatially and temporally unpredictable emergences of damselflies. Brewer’s Blackbirds have relatively small territories (0.05–0.20 ha; Stepney 1971), but like Rusty Blackbirds, may fly up to several kilometers to forage away from their nesting colonies (Fig. 1; Martin 2002, Machtans et al. 2007). Rusty Blackbirds, both colonial and noncolonial, may benefit from flights away from nesting wetlands to feed on ephemeral but rich aquatic insect emergences. Noncolonial birds may have fewer opportunities to learn of such sites on their own. Social interactions may improve foraging efficiency or group defense of nest sites (Powell et al. 2010b) and may help explain the larger home ranges we observed in colonial individuals. More colonies must be located and studied before we can test hypotheses related to colony formation or space use by colonial Rusty Blackbirds.

**SAMPLING RECOMMENDATIONS**

The nine of 13 Rusty Blackbird home ranges that reached an asymptote required an average of 32 relocations to do so. This result accords with the recommendation of Seaman et al. (1999) to collect 30–50 points per individual. Although we did not follow Laver and Kelly’s (2008) recommendation to include in statistical analyses only home ranges that reached asymptotes, the sizes of the four home ranges that did not reach an asymptote were not statistically different from those that did. Rusty Blackbird studies requiring that all home ranges reach asymptotes should set an a priori goal of at least 50 points per individual or consider calculating asymptotes during the study to confirm when home range sizes stabilize and data collection can cease.

Eleven of 13 Rusty Blackbirds had home ranges larger than 10 ha, and adults often visited wetlands and wet patches not connected to nesting wetlands (Fig. 1, 2b). We doubt that individuals can be effectively tracked over such large areas without telemetry and believe that territory mapping without telemetry would both underestimate the area the species uses (Anich et al. 2009) and underrepresent the potential value of habitats beyond the wetland in which the birds are nesting. Thus, telemetry studies minimize bias in estimates of territory.
size (Anich et al. 2009) and are more useful in identifying habitats appropriate for conservation of this species.

We suspect that home-range size varies with a bird's breeding status, but our small sample limited our ability to test this hypothesis directly. Pairs with fledglings or older nestlings likely have home ranges smaller than those of unpaired birds, as they must meet the constant demand for food by their young. The nine adults we tracked with nestlings or fledglings had relatively small home ranges (17.0 ± 3.4 ha). Incubating females also probably had relatively small home ranges, but with only three bird-days of data during this stage, we were unable to confirm this. Conversely, unpaired or non-breeding individuals have more freedom to wander and, thus, should have larger home ranges. The two adults we studied for which breeding status was unknown had the largest home ranges (72 and 173 ha); we suspect that these individuals were not breeding when we tracked them. We recommend that future telemetry studies of breeding Rusty Blackbirds begin to capture and radiotrack adults in early May when nest building begins (Avery 1995). An early start will provide researchers with the opportunity to better quantify how space use varies across the entire breeding season.

EVALUATION OF BUFFERS FOR HABITAT PROTECTION

The Rusty Blackbird is considered an obligate wetland breeder (Avery 1995); in New England its site occupancy is best predicted by shallow, independent pools of water (Powell 2008). We were therefore surprised that upland habitats constituted 88% of home ranges and 81% of core areas in this study. These high percentages could be due to errors in triangulation or an artifact of fixed KDE smoothing when the home ranges were calculated (Silverman 1986). However, we often observed Rusty Blackbirds foraging in small wetlands such as wet seeps with patches of Sphagnum sp. and depressions resulting from logging and road construction such as sites of removed stumps, skidder ruts, roadside ditches with alders, and borrow pits. These were often too small (<16 m²) to be delineated as wetlands by NWI, but they appeared to be important foraging habitats for Rusty Blackbirds. Thus, we likely underestimated the importance of wetlands in home ranges because the resolution of NWI was too coarse for us to accurately map many of the smaller wetlands in which the birds foraged.

The effects of timber harvests on breeding Rusty Blackbirds are complex, as logging appears to attract the species by increasing both the number of small water bodies available for foraging and the density of young conifers selected for nesting (Powell et al. 2010a). However, when Rusty Blackbirds nested in regenerating timber harvests that extended into or to the edges of wetlands, they suffered twice the rate of nest predation of pairs nesting in wetlands without recent logging. In this respect, harvested stands within or adjacent to wetlands may act as ecological traps (Powell et al. 2010a). Of 43 nests found by Powell et al. (2010a), 29 were within wetlands, and the 14 in uplands averaged only 19 m from wetlands. A 75-m buffer of no logging around wetlands would clearly encompass most nests and thus likely minimize nest predation (Powell et al. 2010a). Nevertheless, 75-m buffers and their wetlands encompassed an average of only 51% of core areas and completely encompassed the core area of only three of 13 individuals. The Rusty Blackbird’s mobility, coupled with its use of small and perhaps ephemeral wetlands, likely explains this. Therefore, the 75-m-wide buffers recommended by Powell et al. (2010a) to protect nesting habitat may be of only limited benefit as a conservation strategy for protecting foraging habitat. Fortunately, Rusty Blackbirds appear flexible in their use of areas with surface water for foraging, so buffers may be best suited for protecting nesting habitat; however, more research is needed to determine if disturbed and undisturbed foraging habitats are equally suitable.

Landscapes that maximize the Rusty Blackbird’s fitness likely contain combinations of wetlands with patches of shallow water that are rich in aquatic prey and undisturbed wetlands with small conifers for safe nesting (Batáry and Báldi 2004, Powell et al. 2010a). The challenge ahead is to implement meaningful conservation measures that accommodate the species’ requirement for a large area, need for wetlands that are appropriately buffered from the effects of nest predators, and protection from other potential limiting factors such as high exposures to methylmercury (Edmonds et al. 2010). Time for implementing such measures is short, particularly in New England and southern Canada, where Rusty Blackbirds are rare and the breeding range is retracting quickly (Powell 2008, Greenberg et al., in press; Maritime Breeding Bird Atlas, unpubl. data).

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