CHAPTER FOUR

Home Range Size and Movements of Greater Prairie-Chickens

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**Abstract.** Size of a home range is key to a species’ conservation and management. Estimates of home range size vary with movement patterns, which in turn vary with sex, age class, season, time of day, and habitat configuration, particularly extent of fragmentation. We describe variation in home range and movements in a grouse endemic to North American prairie, the Greater Prairie-Chicken (*Tympanuchus cupido pinnatus*). Our study area included a large, contiguous block of tallgrass prairie. We found that daylight movements varied with time of day: typically, birds were least active in the heat of midday and most active in the relative cool of morning and evening, a pattern consistent with sunrise and sunset, particularly in autumn, winter, and spring. The species’ lek and nesting biology predicted observed lulls in male movement in spring and female movement in summer; sexes are equally mobile at other seasons. Females had larger home ranges than males, moved more frequently between activity centers, and moved greater maximum distances; therefore, females may be more susceptible to the negative effects of habitat fragmentation. Yearlings of both sexes tended to move more than adults. A synthesis of home range estimates from our work and past studies suggests there may be an inverse relationship between habitat continuity and home range sizes. Our results underscore the need to consider various environmental and other factors when estimating home range size. We also present preliminary evidence that habitat fragmentation may force prairie grouse to expand their home range, potentially decreasing survivorship through increased mortality from predation risk or energy expenditure.

*Key Words:* circadian rhythm, fragmentation, home range, movement, seasonality, *Tympanuchus cupido.*

The home range—the amount of physical space individuals need, on average, to survive, grow, and reproduce—is a fundamental aspect of a species' ecology and is crucial to an understanding of a species' place in the ecosystem (i.e., its ecological niche). Moreover, it is difficult to develop meaningful management and conservation strategies for rare species if we lack a basic knowledge of their spatial needs (Belovsky 1987). Both movements and home ranges depend on a variety of endogenous and exogenous factors, including demographic status and local habitat and conditions (Southwood 1977). From the organism's view, habitat can be continuous, patchy, or isolated, and large-bodied organisms generally require larger home ranges (Kelt and Van Vuren 1999, Peery 2000).

Few habitats in North America are more fragmented and depleted than tallgrass prairie: Only ∼4% of this biome remains (Samson and Knopf 1996), with most remnants being small, widely scattered, and altered by human activity. As a result of this extensive alteration, the Greater Prairie-Chicken (Tympanuchus cupido pinnatus)—a species emblematic of tallgrass prairie—now survives on native grassland embedded in a matrix of pastures, cultivated fields, roads, fences, homesteads, and woodlands. Most prairie chicken populations are of conservation concern, underscoring the need for a clear understanding of the species’ home range requirements and movement patterns (Niemuth, this volume, chapter 1). In this species, movement frequency and distance varies temporally with season or time of day (Robel et al. 1970, Hamerstrom and Hamerstrom 1973, Drobney and Sparrowe 1977); endogenously with demographic factors such as sex, age, or breeding status (see Toepfer 1988); and exogenously with habitat extent and fragmentation.

Early studies of movements and home ranges of birds relied on band recoveries, fortuitous sighting of marked individuals, or following individual flocks through a day (Hamerstrom and Hamerstrom 1949). These methods produced important data but only allowed study of short-term, short-distance movement. The advent of lightweight radio transmitters in the 1960s greatly benefited the study of both the frequency and the distance of animal movement. Still, there have been relatively few telemetry studies on prairie grouse, most of them involving prairie chickens in fragmented habitat or small blocks of prairie (Hamerstrom and Hamerstrom 1949, Burger 1988, Toepfer 1988, Schroeder and Braun 1992a).

Our objective was to estimate home range size for Greater Prairie-Chickens on a large block of unfragmented tallgrass prairie. We further sought to examine movement patterns at several time scales, ranging from within a day to among seasons to over the life span of an individual. This last effort allowed us to identify whether a prairie chicken's center of activity—by which we mean the extent of the principal area used—was stable or changed over a bird's life. We compared our findings to results from previous studies elsewhere in the species’ range, allowing us to postulate how home range size might be affected by habitat fragmentation. Given that our study area was a contiguous block of tallgrass prairie, we predicted our home range estimates would be smaller than in previous studies because in fragmented areas energetic needs for maintenance, growth, and reproduction can be met equally in either a smaller contiguous block of suitable habitat or in a larger mosaic of suitable and unsuitable habitats (Reiss 1988). We also examined how movements vary with sex, age, season, and time of day, thus generating a better understanding of the spatial ecology of the Greater Prairie-Chicken.

**METHODS**

**Study Area**

Our study area encompassed ∼450 km² of tallgrass prairie in the Flint Hills of north-central Osage County, Oklahoma, its north edge abutting Kansas (36°46'–37°00' N, 96°22'–96°40' W). The Flint Hills ecoregion consists largely of unplowed tallgrass prairie, although much of this region is grazed heavily and burned annually (Zimmerman 1997, With et al. 2008). Habitat in our study area was relatively homogenous prairie, with no cultivation (<1% of the area has ever been cultivated), no significant development, and few fences. The few roads were primarily graded dirt or gravel without bordering ditches or embankments. Deciduous woodland (<5% of the area) occupied a small portion of the southeast corner of the area, chiefly occurring in narrow corridors along two creeks.
Prescribed fires burned 60–80% of the area annually (Patten et al. 2007), generally in early spring (March–April). Cattle grazing usually followed burning, the predominant system being early intensive stocking: Steers are brought to the ranches for ∼100 days from April to July, allowing the range vegetation to recover in late summer and autumn (Smith and Owensby 1978). Cow-calf operations occupied ∼10% of the study area; such operations avoid annual burns and graze at a lower stocking rate throughout the year. A low density of American bison (Bison bison) grazed ∼5% of the study area year-round, all on the Nature Conservancy’s Tallgrass Prairie Preserve (in the southeastern quadrant), and blocks (<100 ha) of this preserve burned sporadically. A small fraction (1–5%) of the study area was hayed each year, chiefly in August. Rainfall during the study (1997–2000) exceeded (z = 0.44–1.42) the long-term (1949–2003) average, but annual temperature centered on the average (z = −0.45–1.40).

Tracking

We radio-tagged and tracked Greater Prairie-Chickens year-round for a 3-year period, from April 1997 to July 2000. Birds were trapped at leks using walk-in funnel traps (Schroeder and Braun 1991) connected by 8-m zigzags of plastic drift fence. At first capture, birds were fitted with a bib-mounted radio transmitter and a loop antenna (AVM and Telemetry Solutions Inc.) weighing 18 g, which was ∼2% of body mass (800–1000 g). We used feather wear and replacement to identify age classes (Wright and Hiatt 1943, Ammann 1944). Yearlings were birds <1 year old (we treated 1 July as the “birthday” for all birds), adults were birds >1 year old, some of which may have been 3–4 years old. Tracking equipment consisted of five-element, handheld Yagi antennas and ATS® model R-2000 or R-4000 receivers. We began tracking a bird the day it was captured; it was tracked thereafter as often as possible, averaging once every 3 days, at varying times of day. Over 99% of bird locations were from direct homing (<1% triangulation) and from <50 m distance. Roughly 17% of birds flushed, the majority during night trapping or being females at nests or with broods; no birds were pushed into new areas, and we have no reason to suspect that a flushed bird left its home range. Two person-days per week were devoted to finding “lost” birds, defined as individuals not detected for two weeks, and we conducted study area–wide aerial transects for lost birds 5–6 times per year, extending 3–8 km beyond any tracked bird and often >25 km from previous locations.

There are several sources of bias when estimating home range and maximum distances moved. Unless a marked bird is found dead, a bird’s life span is unknown, particularly if it is lost from the study area; range size and distance moved for lost birds may be larger than estimated (Sharp 2009). We determined if estimates of home ranges for birds eventually found dead were systematically smaller than estimates for lost or dispersed birds, in each case using movement data up to our last confirmed location. Birds were also tracked for different durations. We evaluated sources of potential bias by regressing range size against two tracking measures: (1) the number of tracking records and (2) days elapsed between when a bird was first and last tracked. We log10-transformed home range estimates before performing each regression. We evaluated tracking records versus home range by fitting a loess smooth (f = 0.5) to the data (see Cleveland and Devlin 1988).

Movements

We analyzed daily movement data from a subset of individuals tracked at 30-min intervals from within 1 hour of sunrise to within 1 hour of sunset. Because sunrise and sunset vary through the year, we obtained a variable number of locations—generally between 20 and 30—for each day of tracking. Of 185 tagged birds, we used only those 32 individuals with extensive data: 22 males (17 adult, 5 yearling) and 10 females (8 adult, 2 yearling). Linear distances were calculated between consecutive tracking locations. Maximum distance refers to the greatest distance between any two consecutive locations that a given individual moved.

Because movement distances are not comparable when tracking intervals vary widely, we used only locations from nonoverlapping 3-day intervals. Our approach eliminates bias by restricting data to equal intervals over which distances can be calculated between consecutive locations.

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For these analyses, we selected only individuals with >25 records, used only individuals for which there were at least four 3-day tracking intervals in a season (n = 617 tracking events), and used only the first location of a day if a bird was tracked more than once that day. We avoided bias in first locations by tracking a given set of birds across different periods in a 5-day rotation: day 1, 1400–2300 H; day 2, 1200–2100 H; day 3, 1000–1900 H; day 4, 0800–1700 H; and day 5, 0600–1500 H.

In some cases, we divided analyses into four 3-month seasons, with prairie-chicken biology defining the seasons: Spring corresponded with lekking activity (15 February–14 May), summer with nesting and brood rearing (15 May–14 August), autumn with late-season lekking (15 August–14 November), and winter accounting for the rest (15 November–14 February). We averaged movements for individual birds within a season and smoothed hourly and seasonal data using loess regression with $f = 0.5$ (Cleveland and Devlin 1988). Differences in movements were assessed with the Mann–Whitney U-test.

### Annual Home Range

We estimated annual home range using both a 100% minimum convex polygon (MCP) and a kernel density estimator (see Powell 2000). We used ArcGIS 9.2 (ESRI, Redlands, CA) and program Abode (P. N. Laver 2005, http://filebox.vt.edu/users/plaver/abode/contact.html) to estimate MCP of all points, centered on their median, to compare our estimates of home range size to MCP values reported in prior studies of Greater Prairie-Chickens (Robel et al. 1970, Newell 1987, Toepfer 1988, Burger 1988, Schroeder and Braun 1992a). For other comparisons, home ranges for each bird were estimated using kernel methods at 95% and 50% isopleths, smoothed with least-squares cross-validation. We considered the 50% isopleth to be a center of primary activity that may be spread over more than one area. We minimized temporal autocorrelation by including only one tracking location per day, selected at random.

We restricted initial analyses to individuals tracked at 20 unique locations (all had >50 tracking records across >70 days), yielding a set of 100 individuals, 29 females (14 adult, 6 yearling, 9 unknown) and 71 males (27 adults, 36 yearlings, 8 unknown). The number of locations at which an individual was recorded ranged from 20 to 159, for a sum of 4,925 locations across all birds. The mean (±SE) duration between which bird was first and last tracked was 354.6 ± 21.6 days. Differences in home range size were assessed with the Wilcoxon two-sample test.

### Habitat Continuity

In order to place our estimate of annual home range size in context, we compiled other MCP-based estimates for the Greater Prairie-Chicken.
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(T. c. pinnatus subspecies only) from Newell (1987), Toepfer (1988), and Burger (1988); the last two studies present estimates from two sites each. Neither Robel et al. (1970) nor Schroeder and Braun (1992) reported annual home range sizes, but we used their data on seasonal home range size. For each study site, we used the largest block of unbroken prairie-chicken habitat as an index for habitat continuity. We assumed that the relationship between this continuity index and home range size would be an inverse polynomial curve of the form

\[ f(x) = y_0 + \frac{a}{x} \]

where \( y_0 \) marks the inflection point and \( a \) the slope parameter.

RESULTS

Potential Sources of Bias

Estimated home range size, regardless of method, was not related to the number of tracking locations (95% kernel: \( r^2 = 0.01, F_{1,98} = 0.85, P > 0.35 \); MCP: \( r^2 = 0.001, F_{1,98} = 0.10, P > 0.70 \)), although the slope flattened only after ~30 locations had accumulated (Fig. 4.1). Accordingly, our home range estimates below were based on individuals for which >30 tracking locations were available (\( n = 71 \)). The estimate of home range size increased with the number of days a bird was tracked (95% kernel: \( r^2 = 0.08, F_{1,98} = 8.02, P < 0.01 \); MCP: \( r^2 = 0.04, F_{1,98} = 4.36, P < 0.05 \)), up to one year, after which the relationship flattened to a slope of zero. Range size estimates thus increase over the short term, but our long-term data should mitigate this problem. Estimates of home range size did not differ for birds lost from the study (1,302 ± 172 ha) versus those eventually found dead (1,496 ± 395 ha; Wilcoxon \( C = 1506, P > 0.40 \)).

Hourly Movements

At the latitude of our study site there are ~5 hours more daylight at summer solstice than at winter solstice, so movements per hour of individuals in a single day should be computed only within season. In autumn, winter, and spring, prairie chickens moved most often in early morning and late evening, and least often at midday and early afternoon (Fig. 4.2). The same pattern

Figure 4.2. Mean hourly movement of the Greater Prairie-Chicken across seasons (see text for definitions). Curves were smoothed using a loess procedure (\( f = 0.5 \)). Note that the pattern of summer movements differs fundamentally from patterns in other seasons, and that male and female behavior differs markedly in spring and summer.

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held when we used sunrise and sunset as common reference points, suggesting movements may track ambient temperature, at least during non-breeding seasons (Fig. 4.3). Male movements in spring were low in the morning, an expected consequence of lek attendance. Although the pattern within days was similar, birds moved farther in winter than in autumn and spring (see below); activity in summer was different, with a male peak near midday and a female peak in early afternoon (Fig. 4.2).

**Movements by Season**

Individual movements across the year averaged 429 m (range 0–1,706 m) per 3-day period. Distances varied substantially among four seasons (Fig. 4.4). Average autumn (±SE = 839 ± 142 m) and winter (845 ± 121 m) movements were 2–3 times greater than those from spring (289 ± 40 m) and summer (391 ± 43 m). Males tended to move less (266 ± 41 m) than females (461 ± 132 m) in spring (Fig. 4.5; \( U_{18.5} = 138, 0.05 < P < 0.10 \)) but more (469 ± 57 m) than females (272 ± 55 m) in summer (Fig. 4.5; \( U_{23.5} = 265, P < 0.01 \)). Because our sample size of females was small, we restricted our analysis of movements...
by age classes to males. Average movements of adults (375 m) and yearlings (426 m) did not differ, but adult movement tended to vary less (0–1,138 m, SE = 43 m) than yearling movement (0–1,405 m, SE = 102 m; variance test: $F_{17,45} = 2.05$, $P < 0.06$). None of the pairwise comparisons between seasons was significant.

**Annual Home Range Size**

Greater Prairie-Chickens moved, on average, 5.0 km from one end of their MCP home range to the other (Fig. 4.6); the minimum distance was only 1.1 km, whereas the maximum was >15 km. Females moved twice as far as males on average (Fig. 4.6; 8 vs. 4 km; $U_{13,12} = 313$, $P < 0.01$); minimum and maximum distances moved by females were also twice those of males, half of the females having a maximum distance greater than that for all but a few males.

Annual 95% kernel home range size averaged 1,203 ± 219 ha ($n = 71$; median = 554 ha; range 36–10,433 ha). Many of the largest home ranges were split into multiple activity centers, although the largest with a single activity center was 4898 ha. The average annual home range for females (2,593 ha) was >3 times larger than that for males (731 ha; Fig. 4.6; $C = 722$, $P < 0.001$). By contrast, the proportion of females and males with split ranges—in which a bird moved between two or more clearly defined activity centers—did not differ for 95% kernel home ranges (males: 0.793; females: 0.704; $X^2 = 4.40$, $P > 0.20$); however, on the basis of 50% kernels, significantly more males had a single core area (males: 0.103, females: 0.366; $X^2 = 9.81$, $P < 0.05$), presumably their lek. We detected neither a seasonal nor a daily pattern to movements among activity centers, and home range size was not associated with the number of activity centers ($r^2 = 0.02$, $F_{3,96} = 0.49$, $P > 0.60$). In both sexes, annual home range did not vary with body mass (males: $r^2 < 0.01$, $F_{3,17} = 0.11$, $P > 0.70$; females: $r^2 < 0.01$, $F_{5,51} = 0.05$, $P > 0.80$) and did not differ between adults and yearlings (males: $C = 39$, $P > 0.10$; females: $C = 423$, $P > 0.30$), although yearlings tended to have larger home ranges (males: 4792 ha vs. 1614 ha; females: 801 ha vs. 637 ha).

**Habitat Continuity**

The annual 100% MCP home range for the Greater Prairie-Chickens in our contiguous block of prairie averaged 1,371 ha, a value almost exactly at the mean for six other studies (=1,370 ha).
across a range of fragmented prairie (Fig. 4.7). An inverse polynomial ($y_0 = 1.025.4, a = 986,465.0$) indicated that extent of fragmentation explained much of the variance in home range size ($r^2 = 0.83$). The curve’s transition point implies that home range size is stable at $>4,000$ ha of contiguous habitat but increases sharply below $\sim 3,000$ ha.

**DISCUSSION**

On average, the Greater Prairie-Chicken’s pattern of movements across a day is like that of most diurnal endotherms: extensive morning and evening activity bracketing a midday lull (Rensing and Ruoff 2002). Hamerstrom and Hamerstrom (1949) noted a similar pattern, with movement in morning from the feeding area to a midday loafing area, then back to the feeding area in evening shortly before going to roost. We found diurnal patterns to be fairly consistent even when day length varied among seasons. Accordingly, it seems reasonable to conclude that, at least outside the breeding season, the movements are driven partly by thermoregulation and, if so, should track ambient temperature (Fig. 4.3), which climbs from dawn until early afternoon, then drops to a low in the middle of the night. Yet it is possible that the birds respond directly to photoperiod, meaning the association between temperature and movement is spurious—both measures simply vary with time of day. Still, we noted more movement in winter, perhaps because food is dispersed more widely and cold temperatures reduce the risk of heat stress.

**Seasonal Movements**

Variation in seasonal movements cannot be attributed solely to climate—it depends also on the Greater Prairie-Chicken’s biology and their environment. For example, movement by both sexes is reduced during the late summer molt of flight feathers (Schroeder and Braun 1992a). Burger (1988) noted that hourly movement peaked in autumn and winter, a pattern we also documented (Fig. 4.4). Between spring and summer, males and females reverse their propensity to move (Fig. 4.5). A male prairie grouse spends most of his time at a lek in spring (Hamerstrom and Hamerstrom 1949, Giesen 1997), limiting his movement. Lekking male sage grouse lose a great deal of fat during the energetically demanding courtship period (Vehrencamp et al. 1989), so in summer they primarily forage. Females visit leks only briefly in spring, but by summer a female is tied to her nest or brood, limiting her movement at that season. Greater Prairie-Chickens have precocial young, but chicks cannot fly until two weeks of age, so even after completing incubation a female’s movements are restricted by the distance her chicks can walk safely. Burger (1988) reported a similar reduction in female movement in summer, about half as much as in autumn and winter. He further reported a significant difference in summer movement between females with and without
broods; Schroeder and Braun (1992a) also found that females with broods tended to have smaller home ranges than females without broods.

Robel et al. (1970) reported a somewhat different pattern of seasonal movement, with maximum movements in February, corresponding to the early part of our spring. Some studies of the Lesser Prairie-Chicken (\textit{T. pallidicinctus}), which occurs in arid shortgrass prairie, have also reported the most movement in early spring (March; Jamison 2000). Our data suggest that prairie chickens moved the least in spring, although this result may be partly the result of different definitions of the seasons, as our results correspond well with reports of reduced movement in summer and increased movement in autumn and winter for both the Greater (Hamerstrom and Hamerstrom 1949, Robel et al. 1970) and Lesser Prairie-Chickens (Taylor and Guthery 1980, Jamison 2000). Likewise, our data were consistent with previous findings that female prairie chickens move less than males in summer and early autumn (Robel et al. 1970, Riley et al. 1994).

Finally, we did not detect a difference between movements of adult and yearling males of the Greater Prairie-Chicken, a finding consistent with other studies of prairie chickens (Robel et al. 1970, Taylor and Guthery 1980). As with our study, Robel et al. (1970) noted that movement of yearling males was more variable than movement of adult males, perhaps the result of yearling males visiting multiple leks before settling to establish a territory (Rippen and Boag 1974, Schroeder and Braun 1992b).

Migration

Schroeder and Braun (1993) reported that Greater Prairie-Chickens in northeastern Colorado were partly migratory. We found no evidence that the population in northern Oklahoma migrated in the strict sense (\textit{sensu} Gauthreaux 1985), that is, moved between separate geographic locations or habitats. Birds in our study were sedentary, remaining in their home range until they died or were “lost,” either because they moved out of the study area or their transmitter ceased to work (as evidenced by recaptures). A few females moved among different activity centers, but we feel that such movement would be better termed \textit{reciprocal dispersal}, a tendency to move among local sites regularly but without distinct seasonality. We had no birds disappear completely only to reappear later. Milder winters in northern Oklahoma than in northern Colorado may account for lack of migratory behavior. Still, even in northern populations that ought to be prone to migration, most Greater Prairie-Chickens appear to spend

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**Figure 4.7.** Relationship between approximate extent of habitat fragmentation, estimated as the largest block of contiguous prairie in the study and home range size (100\% MCP) for Greater Prairie-Chickens using our data and results reported by Newell (1987), Toepfer (1988), and Burger (1988). The open circle is for the present study.
their entire lives in a small area (Hamerstrom and Hamerstrom 1949).

Annual Home Range

In interspecific comparisons of lekking grouse, male territory size is inversely related to female home range size (Bradbury et al. 1986). If home range size is proportional to territory size, then this relationship may imply that a male’s home range is smaller than a female’s. In our study, female Greater Prairie-Chickens had larger home ranges than males. In Wisconsin and northeastern Colorado, female Greater Prairie-Chickens likewise tended to have larger home ranges than males (Toepfer 1988, Schroeder and Braun 1992a); however, Giesen (1997) found no difference in home range size between sexes of the Sharp-tailed Grouse (T. phasianellus).

Relative to males, female Greater Prairie-Chickens moved significantly farther and more often between activity centers, a finding in agreement with past research (Hamerstrom and Hamerstrom 1949, 1973; Toepfer 1988; Schroeder and Braun 1993). Increased movements may have both costs and benefits. Annual survival and reproductive success of female prairie chickens may be related negatively to their amount of movement (Burger 1988), although a female may place her nest farther from the lek where she was inseminated than from the nearest lek (Wakkinen et al. 1992, Aldridge and Brigham 2001). Neither we nor Schroeder and Braun (1992a, 1993) found significant differences in home range size among age classes, although in each case yearlings tended to have larger home ranges.

Habitat Continuity

Fragmentation depends on scale and on the organism under study. We follow Franklin et al. (2002) in defining habitat fragmentation as “the discontinuity, resulting from a given set of mechanisms, in the spatial distribution of resources and conditions present in an area at a given scale that affects occupancy, reproduction, or survival in a particular species.” Under this view, anthropogenic habitat alteration that results in discontinuity can have a profound effect on an animal’s home range requirements (Haskell et al. 2002). Habitat alteration and fragmentation may unduly influence an organism’s movements, either by restricting them or by forcing the organism to move farther to find food, shelter, or a mate (McNab 1963, Herfindal et al. 2005). This restriction includes features that affect an organism’s perception of contiguity, such as erection of high-tension power lines across prairie (Pruett et al. 2009a, 2009b).

We could only approximate extent of fragmentation in other studies, but our survey suggests a negative relationship between extent of contiguous habitat and size of the home range (Fig. 4.7). Excepting when blocks become too small to sustain a population (Winter and Faaborg 1999), we posit that a prairie chicken requires a larger home range in fragmented habitat because it must move farther (see Ryan et al. 1998) and more often to locate food, suitable cover, and safe nesting and roosting sites. This pattern has been implied elsewhere for the Greater Prairie-Chicken: Toepfer (1988) noted that home range was smaller with a “closer year-round proximity of food and cover”; Schroeder and Braun (1992a) reported a tendency for increased home range with greater distance between cover and nests or leks; Svedarsky and Van Amburg (1996) attributed larger egg-laying ranges to wider spacing of cover, and Ryan et al. (1998) observed that “greater dispersion of native prairie in the mosaic area was associated with . . . greater movement of broods.” Beyond energetic costs, extensive movement may increase mortality risk, whether from increased exposure to predators or increased probability of collision with fence lines or other anthropogenic structures (e.g., Patten et al. 2005a, Wolfe et al. 2007, McNew et al., this volume, chapter 19).

CONCLUSIONS

An apparent negative relationship between habitat continuity and home range size suggests a possible mechanism by which populations of the Greater Prairie-Chicken decline as habitat fragmentation increases: It is an indirect result of increased mortality imposed on the birds because they must expand their home ranges. Females, the more peripatetic sex (Fig. 4.6), may be particularly susceptible. On the basis of seasonal and circadian movements (Figs. 4.2–4.5; see also Robel et al. 1970), we predict that the dangers of increased movement in fragmented landscapes may be magnified in winter, in the morning and evening, and for females. A substantial increase in female mortality could doom a population to extinction (Patten et al. 2005a). Indeed, a preliminary analysis of our data implies that survival probability decreases as home range increases (Cox regression: Wald
$\chi^2 = 5.98, \ P < 0.02)$. Increasingly fragmented habitat could also impair a yearling male’s ability to locate leks with available territories, jeopardizing long-term lek persistence, and may limit the extent to which a brooding female can locate enough food and shelter for growing chicks.

Management strategies should focus on minimizing further fragmentation of the tallgrass prairie, including fragmentation from the birds’ perspective (e.g., Pruett et al. 2009b). Limiting direct loss of habitat is an obvious step, but continuity may be reduced by other means, such as extensive spring burning or an increase in fences (Patten et al. 2005a, Reinking 2005). Our data also suggest a potential impact of reducing cover: Thermoregulation may play a key role in habitat selection by prairie grouse (figure 3 of Patten et al. 2005b). Habitat restoration may address these potential problems by ensuring that there is adequate vegetative cover, particularly at key seasons, for long-term survival and reproduction.

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