Roadside foraging by kangaroo rats in a grazed short-grass prairie landscape

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The foraging behaviors of small mammals reflect trade-offs between locating and harvesting food efficiently, and minimizing exposure to predators (Lima and Dill 1990). Understanding the strategies employed by individual foragers may ultimately reveal mechanisms underlying patterns of local abundance (Bowers and Dooley 1993, Sutherland 1996, Smith and Litvaitis 2000). In northern short-grass prairie, Ord’s kangaroo rats (Dipodomys ordii) are typically restricted to areas with sandy soils, large shrubs, and high plant diversity (Stapp et al. in press). They also are seen in high numbers, however, along roads and on moonlit nights, suggesting that kangaroo rats recognize potential risks associated with roadsides on bright nights. The high numbers of kangaroo rats along roads suggest that benefits associated with these habitats (ease of digging, dust bathing, higher soil seed banks) exceed the costs associated with higher risk of mortality from predators and vehicles. Our results demonstrate how foraging decisions differ depending on the spatial and temporal contexts in which behaviors are measured, and underscore the potential value of less common habitats such as road margins for increasing landscape-scale diversity and wildlife habitat in grazed grasslands.

The foraging behaviors of small mammals reflect trade-offs between locating and harvesting food efficiently, and minimizing exposure to predators (Lima and Dill 1990). Understanding the strategies employed by individual foragers may ultimately reveal mechanisms underlying patterns of local abundance (Bowers and Dooley 1993, Sutherland 1996, Smith and Litvaitis 2000). In northern short-grass prairie, Ord’s kangaroo rats (Dipodomys ordii) are typically restricted to areas with sandy soils, large shrubs, and high plant diversity (Stapp et al. in press). They also are seen in high numbers, however, along unimproved gravel roads (Flake 1971, Abramsky 1978, Kaufman and Kaufman 1982), where vegetation and levels of disturbance differ markedly from the surrounding landscape, which is grazed by cattle. These granivorous rodents may be attracted to seeds produced by weedy plants growing in the ungrazed road margins (Betz 2001), or to the loose soils that facilitate dust bathing, burrow construction, and foraging (Brock and Kelt 2004). However, roads are also used extensively by carnivores such as coyotes (Canis latrans) and foxes (Vulpes velox; Roell 1999, Engeman et al. 2002, Kamler et al. 2003), and utility poles and fence posts that are associated with roads provide owls and raptors with an abundance of hunting perches (Marti 1974). Diets of Great Horned Owls (Bubo virginianus) in short-grass prairie suggest that they hunt along roadsides (Zimmerman et al. 1996); Ord’s kangaroo rats comprised 16%–25% of the small rodents consumed by these owls. Foraging in the exposed areas of roads thus may be profitable, but also dangerous, especially on moonlit nights, when aboveground activity of rodents is more conspicuous (Kotler 1984, Longland and Price 1991).

If predation risk influences the surface activity of kangaroo rats, and ultimately, population density, individuals should concentrate foraging activities beneath large shrubs and other tall vegetation, and avoid exposed areas between shrubs. The difference in perceived risk between shrub and open microhabitats may contribute to this species’ relative absence in upland areas of short-grass prairie (Stapp et al. in press), where the dominant vegetation is
very short (<15 cm). *Dipodomys ordii* is the only kangaroo rat found in grasslands of the Great Plains (Garrison and Best 1990), but it has never been captured in 10 years of live-trapping (nearly 30,000 trap-nights) in three 3.14-ha grassland sites in northern Colorado. It is among the most common rodents, however, in areas with shrub cover and sandier soils (Stapp et al. in press).

We used a combination of live-trapping and foraging trials in artificial seed trays to (1) investigate the use of roadside habitats by Ord’s kangaroo rats in short-grass prairie and (2) evaluate the potential importance of predation risk as an explanation for the microhabitat affinities and local abundance patterns of kangaroo rats. We compared giving-up densities (GUDs; Brown 1988) in paired seed trays placed beneath the cover of shrubs and in the open on nights with different moonlight levels to determine the importance of predation risk as a determinant of microhabitat use. We compared foraging rates in seed trays along road margins with those from trays located far (120 m) from the road into the adjacent pasture to determine if perceived predation risk differed by landscape position (roadside vs. pasture). If kangaroo rats behave so as to minimize predation risk, seed consumption should be higher (and GUDs lower) in shrub versus open microhabitats, in pasture versus roadside locations, and on dark versus moonlit nights.

**Methods**

The study area was the Central Plains Experimental Range (CPER), located in northern Colorado, (40°49′N, 104°46′W). The climate at the CPER is semiarid, with cold, dry winters and most of the 322 mm of annual precipitation falling as rain between April and September (Lauenroth and Milchunas 1991). Vegetation was classified as short-grass prairie, and was dominated by 2 perennial warm-season grasses (*Bouteloua gracilis* and *Buchloe dactyloides*). Low-lying areas with loamy sand and sandy soils comprised a relatively small fraction (<15%) of the study area, but often supported taller vegetation and a higher diversity of grasses, forbs, and woody shrubs such as four-wing saltbush (*Atriplex canescens*). The roads on the CPER were a mixture of dirt and gravel, and fine soil accumulated on the road margins as a result of regular grading. Vehicle traffic was very light, especially at night, when kangaroo rats were active. Because most pastures on the CPER were fenced, road margins were not grazed, except briefly when cattle were moved between pastures in late spring and autumn. Therefore vegetation along road-sides was much taller and cover was much greater than in the adjacent pasture (Betz 2001), and included native grasses (*Pascopyron smithii*, *Sporobolus cryptandrus*, and *Stipa comata*) and forbs (*Helianthus annuus*, *Euphorbia glyptosperma*, and *Psoralidum teretiflorum*), as well as numerous exotic weeds (*Eragrostis cilianensis*, *Kochia scoparia*, *Salsola iberica*, *Melilotus officinalis*, and *Verbena bracteata*). Roadside habitats were largely maintained by cattle grazing and regular road grading; on open range in the nearby Pawnee National Grasslands, where there were no barbed-wire fences, vegetation along roads resembled that in the adjacent prairie.

Rodents were live-trapped on 3 occasions (30 September–3 October 1997, 20–23 April 1998, 17–24 August 1998) to estimate patterns of relative abundance along roadsides and in adjacent pastures. Pasture locations included upland grasslands, with short vegetation typical of short-grass prairie, and in mixed grassland-shrub areas dominated by saltbush (canopy cover = 3%–7%; Stapp unpublished data). Most grasslands were grazed at a moderate stocking rate during summer, whereas saltbush areas were moderately grazed in winter. Trapping transects consisted of 12–15 large Sherman live-traps, set approximately 10 m apart. Transects were set out in pairs, with 1 along the road and the other approximately 120 m into the pasture. Transects were separated by >100 m. For each trapping session, traps were set for 4 consecutive nights and were checked each morning at dawn. Captured individuals were marked with aluminum ear tags and released. We used the number of unique individuals captured per 100 trap-nights as an index of relative abundance for each trapping session. Values were log(x + 1)-transformed for analysis. All procedures were approved by the Animal Care and Use Committee at Colorado State University and met guidelines recommended by the American Society of Mammalogists (1998).

Because kangaroo rats were absent from grassland pasture transects, foraging trials were
conducted along roads and in pastures in salt-bush areas only. Four 110-m transects were established along gravel roads, and 4 parallel transects were established approximately 120 m into the pastures. Foraging stations were approximately 20 m apart along each transect (3–4 stations per transect). At each station, cardboard seed trays (40 × 26.5 × 5 cm) were placed in pairs, with trays separated by 3 m. One tray was placed beneath the canopy of a large saltbush and the other in open, short vegetation or bare soil, ≥3 m from any tall (>0.25 m) vegetation. Trays were placed near where kangaroo rats had been frequently live-trapped.

On each night of foraging trials, trays were supplied with 2 L of presifted, fine sand mixed with 4.00 g of millet. Millet was stored indoors in plastic film canisters, but was not oven-dried before weighing. Foraging trials lasted 2–3 nights and were conducted on nights of a full, new, and 1st-quarter (hereafter, half) moon between 8 October and 4 November 1997. Seeds remaining in trays in the morning were sifted, cleaned of sand and debris, and weighed to estimate rates of seed consumption. The remaining seeds were dried at 55°C for 24 hours and reweighed to estimate and correct for moisture gained by seeds overnight. We also counted the number and kinds (morphotypes) of seeds that fell or blew into the trays from the surrounding vegetation as a rough index of seed rain during the study period. Relative humidity was measured at a standard weather station on the CPER on the morning following each trial. For 1 night for which weather data were not available, we estimated relative humidity from the average wet and dry bulb temperatures for the previous and subsequent night, using an equation provided by the U.S. Water Conservation Laboratory (Barnes 2005). Calculations using this equation tended to underestimate the measured values of relative humidity by 2.5%.

Seed consumption values were used to calculate the giving-up density (GUD), the amount (g) of seeds left behind in each tray. Differences in GUDs between shrub and open trays reflect the animal’s perception of the relative costs (e.g., predation risk, energetic costs, or missed-opportunity costs) and benefits (high harvest rates) of foraging in these microhabitats (Brown 1988). This technique has been widely used to study foraging behavior and microhabitat use of granivorous rodents, including several Dipodomys species (e.g., Brown 1988, Brown et al. 1988, Brown et al. 1994, Bouskila 1995, Herman and Valone 2000, Price and Correll 2001, Sullivan et al. 2001). In our study, only pairs of trays in which there was evidence of kangaroo-rat foraging (e.g., tail drags, large tracks) in at least 1 tray were included in GUD analyses (at 66 g, D. ordii is about twice the size of other nocturnal rodents present). Pairs in which either tray had tracks or feces of other species (28 trays; primarily northern grasshopper mouse, Onychomys leucogaster) were omitted. Because trays were always set out in pairs (1 open tray and 1 shrub tray), we analyzed the mean difference between GUDs in open and shrub trays at a given station. Foraging by other species resulted in large differences in sample sizes among moonlight levels and landscape locations; therefore, we used paired t tests to determine whether mean differences in GUDs between shrub and open trays were significant in different locations during trials with different moonlight levels. SAS 9.0 was used for all statistical analyses. Means are presented with 1 standard error (sx).

**RESULTS**

Ord’s kangaroo rat was the most commonly captured species (46% of 124 individuals) and comprised 63% of the individuals captured in roadside transects and 19% of those in pasture transects. No kangaroo rats were captured in grassland pasture locations. The northern grasshopper mouse was the 2nd-most abundant species (30% of individuals) and was captured primarily in pasture transects (88% of captures). Deer mice (Peromyscus maniculatus; 21% of individuals) were much more common in road margins (81% of captures) than in pastures. Two western harvest mice (Reithrodontomys megalotis) and 1 prairie vole (Microtus ochrogaster) were captured only in road margins, and a single hispid pocket mouse (Chaetodipus hispidus) was captured in a saltbush pasture site.

There was no significant difference in relative abundance of kangaroo rats between trapping sessions ($F_{2, 32} = 0.61, P = 0.550$), nor any significant interactions between trapping session and vegetation type (grassland, salt-bush) or landscape location (roadside, pasture;
A 2-way ANOVA showed that kangaroo rats were more abundant along roadways than in pastures in both vegetation types \((F = 17.52, df = 1, P = 0.0002)\), that they were more abundant in saltbush than in grasslands \((F = 38.19, df = 1, P < 0.0001)\), and that they were 4–6 times more numerous along roads in saltbush areas than on other transects \((interaction: F = 4.26, df = 1, P = 0.047; overall: F_{3, 32} = 19.99, P < 0.0001; \text{Fig. 1})\).

A total of 186 artificial seed trays were set out in 93 shrub–open pairs on 7 nights during the 3 lunar phases. Seeds in the trays gained an additional 0.8%–25% of their initial weight overnight, apparently by absorbing moisture from the air and sand \((\text{Fig. 2})\). The amount of water absorbed was significantly related to the amount of seeds remaining in the trays \((\text{Spearman's } r = 0.94, P < 0.001)\); seeds in trays that were not visited increased more in weight \((14.0\% \pm 0.9\% \text{ of initial weight}, n = 42)\), whereas seeds and chaff remaining in heavily foraged trays absorbed relatively little water \((6.2\% \pm 0.4\% \text{ increase}, n = 144)\).

The amount of weight gain was significantly related to relative humidity \((\text{Fig. 2})\). There was no significant difference in amount of weight gained between shrub and open trays \((t = 1.17, df = 184, P = 0.244)\), regardless of whether trays were visited or not \((P > 0.406)\).

There was no significant difference in the proportion of shrub and open trays foraged or not foraged in a given location during any session \((\text{Fig. 3A}; \text{Proc FREQ}, \text{separate chi-square tests for roadside and pasture locations in each session}, P > 0.14)\), except on roadsides during half-moon trials, when visitation was greater at shrub trays \((\chi^2 = 5.87, P = 0.015)\). Pooling across all locations and microhabitats in a given session showed that kangaroo rats foraged in significantly more trays on new moon nights \((93.75\%)\) than during either full-moon \((75.93\%)\) or half-moon trials \((53.57\%; P < 0.014)\).

We present results based on differences in dry-weight GUDs, although we obtained similar results when we repeated analyses using the wet-weight values and estimates of the amount of seed consumed, which were calculated by subtracting the wet-weight GUDs of trays that were visited from the mean GUD value of all unforaged trays on a given night. GUDs were significantly higher in open microhabitats than in shrub microhabitats, but only near roadsides on full \((t = 2.75, P = 0.018)\) and half-moon \((t = 1.86, P = 0.083)\) nights \((\text{Fig. 3B})\). There was no difference in GUDs between open and shrub trays near roads on dark nights \((t = 1.06, P = 0.308)\). Farther into the pastures, GUDs were similar between open and shrub trays during full-moon \((t = 1.01, P = 0.341)\) and new-moon \((t = 0.61, P = 0.559)\) trials \((\text{Fig. 3B})\). On average, kangaroo rats foraged more in the open in pastures on half- and full-moon nights \((P = 0.021 \text{ and } 0.023, \text{respectively})\).

Deposition of wild seeds from the surrounding vegetation during foraging trials was higher in pasture trays and, especially, beneath shrubs \((\text{Fig. 4})\). Rates of seed fall were highest beneath shrubs in pasture trays, and lowest in open trays along the edge of the road \((\text{2-way ANOVA}: F_{3, 8} = 41.34, P < 0.0001; \text{microhabitat, location, and interaction significant}, P < 0.008)\). Although seeds were only identified to morphospecies, the diversity of seed types was...
similar between microhabitats and locations ($F_{3, 8} = 0.33, P = 0.802$; overall mean $= 1.56 \pm 0.20$ seed types $\cdot m^{-2} \cdot$ night$^{-1}$, $n = 12$).

**DISCUSSION**

From a methodological standpoint, we were surprised by the amount of water that seeds absorbed in foraging trays overnight, which was strongly related to relative humidity. In trays that were not visited or that were foraged only lightly, seeds gained much more weight than those in trays where few intact seeds remained, suggesting that the seeds, not the hulls, absorbed most of the moisture. Despite the potentially important implications for the use of GUDs in studies of granivore foraging behavior, to our knowledge this effect has not been quantified before. Kelt et al. (2004) adjusted seed removal rates for amount of moisture gained in an enclosed reference tray, but did not provide details on water gain. Absorption of water could lead to underestimates of seed consumption and overestimates of GUDs, particularly if relatively little seed is consumed, if the differences between GUDs in different microhabitats are small, or if data are pooled over multiple nights in areas where relative humidity varies greatly over time. Possible remedies include drying the remaining seeds before weighing (e.g., Morris and Davidson 2000), or using wet weights of seeds in trays that are not visited by rodents to correct GUDs for changes in water absorption. Others (Davidson and Morris 2001, Price and Correll 2001) have used changes in the number of seeds, as well as weight, to estimate GUDs, but this would increase processing time substantially. We obtained the same results whether we used dry or wet weights or whether we adjusted measures of seed consumption; however, we recommend drying seeds before and after foraging trials, especially if precise estimates of the amount of seed consumed are needed.

Ecological studies of roads have emphasized their negative effects, which include traffic noise, heavy metals pollution, vehicle mortality, and creation of barriers that fragment and isolate populations (Forman and Alexander 1998, Trombulak and Frissell 2000). In some systems, however, linear landscape features such as roads may also provide benefits such as corridors for dispersal.
as road margins may be important ecological features for small mammal populations. Road margins may be used as movement or dispersal corridors (Brock and Kelt 2004) and may provide opportunities for range expansion (Getz et al. 1978). Roadsides can also increase local diversity by providing habitats and resources that differ dramatically from the surrounding landscape (Adams and Geis 1983, Schwartz et al. 1994, Clark et al. 1996, Betz 2001). Finally, roadsides can serve as refugia when adjacent habitats are unsuitable (Bolger et al. 2001, Stapp et al. in press). The degree to which road margins function as corridors or suitable habitat, or act as sinks or ecological traps (Schlaepfer et al. 2002), varies among species and ecosystems.

In a structurally simple environment like short-grass prairie, vegetation associated with fenced roads contrasts markedly with the surrounding grazed grasslands that composed most of our study area. This contrast is less obvious in areas where the surrounding vegetation is taller and more productive, as was the case in the saltbush areas we studied. Ord’s kangaroo rat was absent from grassland pastures, but a few individuals were able to live in the adjacent road margin, where grasses and forbs were taller (Betz 2001). More kangaroo rats were captured in saltbush pastures, where they were consistently the 2nd- or 3rd-most-common species (Stapp et al. in press), but the highest densities were found along roads in saltbush areas. Other species of Dipodomys are known to use roadsides and to reach high numbers there (Roberts and Packard 1973, Abramsky 1978, Price et al. 1994). Brock and Kelt (2004) captured similar numbers of D. stephensi along dirt roads and in sparse grasslands, but found fewer along gravel roads and in dense grasslands. They attributed the high rates of population turnover along roads to dispersal or possibly to high mortality associated with vehicles or predators.

Although we did not measure predation directly, our indirect measure of risk (differences in foraging activity between shrub and open microhabitats) suggested that kangaroo rats are sensitive to perceived predation risk, but that their response depends on both landscape...
context (pasture vs. roadside) and the expected degree of risk (moonlight intensity). In pasture locations and on dark nights, foraging rates of kangaroo rats were similar in shrub and open trays, which is consistent with the prevailing view that bipedal rodents such as kangaroo rats are able to use both open and shrub microhabitats (Reichman and Price 1991). The relatively low visitation rates during half-moon trials, which were conducted first, may indicate that individuals were slow to locate trays, especially in pasture transects where there were fewer kangaroo rats. Several studies have shown that kangaroo rats reduce their activity (Lockard and Owings 1974) and increase their use of bush microhabitats in response to increased illumination, presumably to reduce exposure to aerial predators (e.g., Schwab 1986, Kotler 1984, Price et al. 1984, Brown et al. 1988). Bouskila (1995) found that moonlight depressed foraging activity and increased use of shrubs by *D. merriami* and *D. deserti*, but only during autumn, when rattlesnakes were not active. Using spotlight counts, Kaufman and Kaufman (1982) showed that moonlight reduced overall activity and the use of roadside cover by kangaroo rats along gravel and dirt roads in tallgrass prairie in Kansas.

In contrast, we observed no increase in the use of shrubs on full-moon or partially illuminated nights in pasture locations; kangaroo rats instead significantly reduced their foraging activity in the open in roadside habitats in response to increased moonlight. Bowers and Dooley (1993) reported similar landscape-scale effects on foraging behavior in response to moonlight and cover. Increases in use of cover by *Peromyscus leucopus* and *Microtus pennsylvanicus* on moonlit nights were only detectable at grassland patch edges, which they attributed to higher activity and hunting success of predators in ecotones. Although we had no information on predator abundance or activity during our study, we speculate that *D. ordii* perceives the open roadside to be riskier on bright nights than on dark nights, at least during the autumn period when we conducted our study. Conspicuous activity in the open can attract predators and increase mortality. Attacks by Great Horned Owls on *D. merriami* were ~40 times higher in open microhabitats than in shrub microhabitats under full-moon nights, compared to ~7 times higher on new-moon nights (Longland and Price 1991). Daly et al. (1990) reported that predator mortality for radio-tagged *D. merriami* was highest for the most mobile animals, such as those that might be expected in roadside habitats if these habitats act as dispersal corridors.

If roadsides are riskier than adjacent pastures, why are kangaroo rats so abundant there? We cannot rule out the possibility that roadsides are population sinks or traps, such that density and habitat quality are decoupled (Van Horne 1983), although we captured primarily adults on roadsides (85% of total individuals captured vs. 37% in pastures), many of which were in breeding condition. Alternatively, rodents may simply be able to perform all necessary activities within the cover of vegetation, limiting their use of the open during brighter periods. Although seed rain was higher in shrub trays than in the open, it was much higher in the saltbush pasture than in the road margin. Our seed data are limited because they were collected only at night and over a very short time compared to the entire period of seed production, and because we did not distinguish between seeds that might be preferred or avoided by kangaroo rats. We also did not sample rodent caches or the soil seed bank. Price and Joyner (1997) found little correlation between seed rain and soil seed bank in the Mojave Desert, and suggested that granivores consume most of a year’s seed crop before it enters the soil. Estimates of seed rain in their study (262 seeds m$^{-2}$d$^{-1}$) and during peak production in Arizona (215 seeds m$^{-2}$d$^{-1}$; Pulliam and Brand 1975) were more than 1 order of magnitude higher than the highest values in our study (19.8 seeds m$^{-2}$night$^{-1}$), but these authors measured seed rain continuously over periods of months.

Working at our study site, Betz (2001) found that soil seed densities, especially those of exotic plants, were significantly higher along roads than 25 m into pastures, which could explain why kangaroo rats forage intensively and reach high densities along roads. However, her study was conducted exclusively in and adjacent to grassland vegetation, where shrubs are rare, and where the plant communities are less productive and less diverse than in the saltbush areas where we worked. If our estimates of seed rain better reflect patterns of seed availability at microhabitat scales and in shrub-dominated habitats, they might explain the tendency of kangaroo rats to forage beneath
shrubs, but they cannot explain kangaroo rat abundance along roads. Our results are similar to those of Miller et al. (2003), who reported high densities of kangaroo rats in river bottoms, even though these areas were flooded seasonally and soil seed densities were much higher in upland habitats nearby. They suggested that the ease of digging in the sandy soils, combined with reduced competition from harvester ants, might explain high kangaroo rat densities in floodplains, despite the low seed densities and potential mortality from flooding. In our study, predation risk and, therefore, mortality may be greater along roadsides, but the benefits associated with these habitats apparently outweigh the costs.

There is increasing demand to manage western rangelands for values other than livestock production, including wildlife (Vavra 2005). Although disturbances associated with roadsides may facilitate invasion by exotic plant species (e.g., Forcella and Harvey 1983, Greenberg et al. 1997), the strong contrast between vegetation in road margins and the adjacent grazed pastures in short-grass prairie contributes to biodiversity at a landscape scale. Our results suggest that vegetation along rural roads with low vehicle traffic provides habitat for small mammals that are less common, such as Ord’s kangaroo rats, in locations where the natural prairie vegetation, under levels of grazing considered typical for the region, would not normally allow them to persist. In our study, the relative value of road margins as habitat for small mammals seemed to depend on vegetation in the surrounding pastures; that is, roadside vegetation in more productive, saltbush-dominated areas was, qualitatively, more similar to adjacent pastures, and relative numbers of kangaroo rats were, as a result, similar as well. As suggested by the differences in perceived risk by kangaroo rats between open and shrub microhabitats on moonlit nights, roadsides may also be important foraging habitats for vertebrate carnivores, including species of regional conservation concern such as the swift fox and Burrowing Owl (e.g., Haug and Oliphant 1990, Kintigh and Andersen 2005). Efforts to maintain and increase landscape-scale heterogeneity in grazed grasslands, as well as in areas with other agricultural lands (e.g., Saarinen et al. 2005), should include less-common landscape elements such as roadsides and fence rows to enhance local biodiversity.

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