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FOOD WEBS AND INTRAGUILD PREDATION: COMMUNITY INTERACTIONS OF A NATIVE MESOCARNIVORE

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Abstract. Trophic level interactions between predators create complex relationships such as intraguild predation. Theoretical research has predicted two possible paths to stability in intraguild systems: intermediate predators either outcompete higher-order predators for shared resources or select habitat based on security. The effects of intraguild predation on intermediate mammalian predators such as swift foxes (*Vulpes velox*) are not well understood. We examined the relationships between swift foxes and both their predators and prey, as well as the effect of vegetation structure on swift fox–coyote (*Canis latrans*) interactions, between August 2001 and August 2004. In a natural experiment created by the Pinon Canyon Maneuver Site in southeastern Colorado, USA, we documented swift fox survival and density in a variety of landscapes and compared these parameters in relation to prey availability, coyote abundance, and vegetation structure. Swift fox density varied significantly between study sites, while survival did not. Coyote abundance was positively related to the basal prey species and vegetation structure, while swift fox density was negatively related to coyote abundance, basal prey species, and vegetation structure. Our results support the prediction that, under intraguild predation in terrestrial systems, top predator distribution matches resource availability (resource match), while intermediate predator distribution inversely matches predation risk (safety match). While predation by coyotes may be the specific cause of swift fox mortality in this system, the more general mechanism appears to be exposure to predation moderated by shrub density.

Key words: asymmetrical competition; Colorado, USA; food web; intraguild predation; safety match; Sobel test; swift fox; *Vulpes velox*.

INTRODUCTION

Over the past 65 years, the study of predator–prey relationships has steadily progressed from relatively simple interactions to complex, multi-trophic level relationships (Solomon 1949, Hairston et al. 1960, Paine 1966, Gratton and Denno 2003, Ives et al. 2005). Recently there has been “an explosion of interest in the complexities that arise from interactions within diverse predator–prey assemblages” (Ives et al. 2005). This has included the influence of habitat complexity (Finke and Denno 2004, Gruner 2004, Langelotto and Denno 2004), temporal or seasonal shifts in regulatory processes (Gratton and Denno 2003, Hampton 2004), direct vs. indirect effects (Fox and Olsen 2000, Navarrette et al. 2000), and guild-level effects (Sih et al. 1998, Finke and Denno 2004). A general synthesis has emerged that top-down and bottom-up forces act simultaneously (Hunter and Price 1992, Denno et al. 2002) and in concert with both biotic and abiotic forces (Power 1992, Meserve et al. 2003).

Concurrent with this explosion, interest in intraguild predation (IGP), predators eating predators, has increased as well. Most natural food webs are not linear; predators must forage under the risk of predation by similar or higher-order predators (Polis et al. 1989, Rosenheim 2004). Trophic interactions between predators can lead to complex system dynamics (Polis et al. 1989, Navarrette et al. 2000, Heithaus 2001), and have forced ecologists to acknowledge less discrete trophic levels (Polis and Strong 1996, Sih et al. 1998). The vast majority of empirical evidence regarding the effect of intraguild predation on predator–prey dynamics comes from biological control studies and arthropod communities (Rosenheim et al. 1995, Snyder and Ives 2001). Such experiments have indicated that intraguild predation pressure is influenced by characteristics of the top predator (Wise 1993, Denno et al. 2002), behavior of the intermediate predator (Lucas et al. 1998), and habitat complexity (Finke and Denno 2002, Langelotto and Denno 2004). Local extinctions of intermediate predators due to intraguild predation pressure are often predicted (Holt and Polis 1997) and observed (Rosenheim 2001).

While there is ample evidence that intraguild predation dynamics exist in terrestrial mammalian systems (Ralls and White 1995, Creel and Creel 1996, Durant

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2000, Fedriani et al. 2000, Switalski 2003, Macdonald and Sillero-Zubiri 2004), there is almost no evidence regarding its effects on community dynamics or the influence of environmental heterogeneity. In particular, the effects on the intermediate predator, or intraguild prey, are largely unknown. While many of the predictions from arthropod studies are applicable to mammalian carnivore interactions, there are crucial differences. For example, in arthropod systems an increase in basal prey abundance often reduces the intensity of intraguild predation pressure (Rosenheim 2001, Denno et al. 2004). However, in mammalian systems intraguild predation is more characteristic of an extreme version of competition; intermediate predators are often killed but not consumed (Polis et al. 1989, Macdonald and Sillero-Zubiri 2004). Therefore the availability of basal prey may be expected to exert little influence. Mammalian intraguild predation is also often characterized by large differences in body size, creating strongly asymmetrical dynamics (Macdonald and Sillero-Zubiri 2004). These differences indicate that while arthropod research is useful in understanding mammalian predator interactions, these predictions must be viewed cautiously.

The potential for landscape or environmental factors influencing the relative strength of predator-prey interactions has also been widely discussed. The terms “mediating” or “moderating” are casually used and frequently interchanged in ecological research to indicate the effect of a third variable on a relationship. However, in biomedical research these terms not only have precise definitions, but rigorous statistical tests have been designed to evaluate the influence of a third variable on the relationship between dependent and independent variables. Here we introduce the use of the Sobel test to quantify the influence of landscape or environmental variables on species interactions (K. J. Preacher and G. J. Leonardelli, *unpublished software*). The Sobel test is relatively simple, using the coefficients and standard errors of linear regression to generate a Wald test statistic and an associated P value. The P value represents whether or not the inclusion of a second explanatory variable significantly altered the relationship between the dependent variable and the first explanatory variable (Fig. 1). For additional information on this or other tests associated with mediating variables, or examples from biomedical literature, see Sobel (1982) or Kenny et al. (1998). For precise definitions of moderating and mediating variables, see Baron and Kenny (1986).

In order to better understand the effect of intraguild predation on swift foxes (*Vulpes velox*), an intermediate mammalian predator of concern, illuminate the relative influence of top-down and bottom-up forces in terrestrial population dynamics, and to determine how these forces interact to determine swift fox distribution, we studied the ecology and demography of swift foxes in southeastern Colorado between August 2001 and August 2004. We monitored the influence of higher

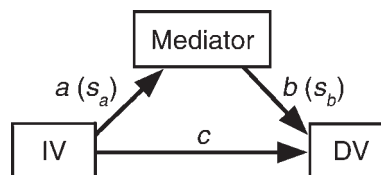


FIG. 1. Outline of the Sobel test from Preacher and Leonardelli (*unpublished software*) showing the influence of a third variable (mediator) on the relationship between an independent variable (IV) and dependent variable (DV). The regression coefficients are a , b , and c ; s_a and s_b indicate model standard error.

and lower trophic levels as well as vegetation structure on swift fox population dynamics. We used several multivariate analytical techniques, including the Sobel test, AIC (Akaike’s information criterion), and hierarchical partitioning, to evaluate the relative strength of food web linkages. Our research centered on the Pinon Canyon Maneuver Site (PCMS), a U.S. Army owned mechanized infantry training facility in southeastern Colorado. Relatively recent changes in the local disturbance regimes, initiated following purchase of the PCMS by the U.S. Army in 1982, have created a natural experiment on the interactions of wildlife communities under different landscapes. While there was no true experimental control of treatments in our study, due to the temporal and spatial scale of terrestrial vertebrate research, observational studies following landscape-level changes are often the only available option. We therefore use the term “natural experiment” cautiously; our research was observational yet capitalized on a well-defined change in land-use practices and the resulting changes in landscape structure and food web dynamics.

METHODS

Study area and system

We conducted research on and around the 1040-km² Pinon Canyon Maneuver Site (PCMS) located in northern Las Animas County, ~50 km northeast of Trinidad, Colorado (Fig. 2). North of the PCMS, in Otero County, the study area extended into the United States Forest Service (USFS) Comanche National Grassland. The study area also extended southward onto private ranchlands. The study area was bordered by the Purgatoire River canyon complex to the east and U.S. Highway 350 to the west.

The region was classified as semiarid grassland steppe, with ~60% of the PCMS categorized as shortgrass prairie dominated by blue grama (*Bouteloua gracilis*), western wheatgrass (*Agropyron smithii*), and galleta (*Hilaria jamesii*) (Shaw et al. 1989). Shrublands interspersed throughout the study area included four-winged saltbrush and greasewood (*Sarcobatus vermiculatus*), as well as prickly pear cactus (*Opuntia phaeacantha*), tree cholla (*Cylindropuntia imbricata*), and yucca (*Yucca glauca*). The remaining landscape was dominated by

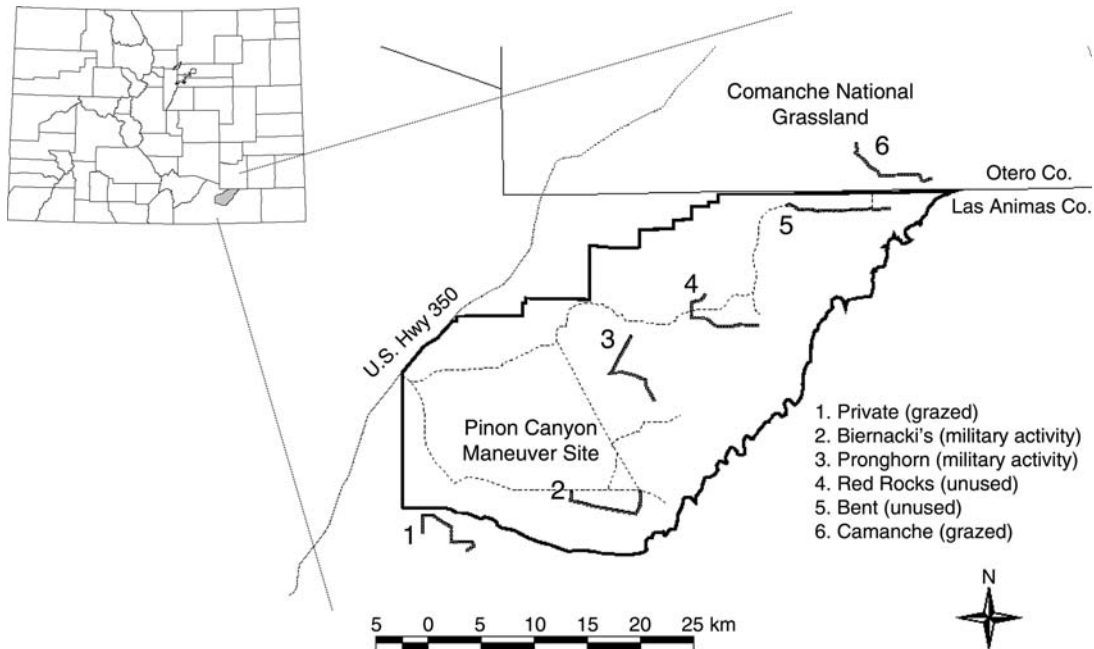


FIG. 2. Six study sites on and around the Pinon Canyon Maneuver Site in southeastern Colorado, USA. Locations of six transects are indicated, as well as the associated dominant land use for that site.

pinon–juniper woodland (*Pinus edulis*, *Juniperus monosperma*). Elevation varied from 1310 to 1740 m, average temperatures range from 1°C in January to 23°C in July, and precipitation averages 30 cm but fluctuates widely (Shaw and Diersing 1990). Monthly precipitation is highest in July with an average of 4.3 cm of rain, though the 35% of the annual precipitation that falls during the cool season (March–May) has a proportionally greater impact on productivity (Milchunas et al. 1999).

Prior to 1982, the region was used extensively for domestic livestock production and fires were infrequent due to the lack of available fuel. In 1982 the PCMS was purchased by the U.S. Army; grazing ceased and fire suppression efforts were initiated. Fire suppression was considered necessary because the release from grazing resulted in rapid biomass accumulation and an increase in fire frequency. In 1985 the U.S. Army commenced large-scale mechanized infantry training on the base. The shift from grazing to mechanized military training has had profound effects on the plant community, including reduced shrub density, increased bare ground, and decreased litter (Shaw and Diersing 1990, Milchunas et al. 1999). At the same time, portions of the PCMS unsuitable for training were left undisturbed and accumulated extensive basal coverage and litter. Examples of unsuitable areas included those containing archeological sites, sensitive plant species, or sections of the base separated from the main training area by enough topographic variation to preclude tank travel. Livestock production continues on the private and public land surrounding the PCMS. The abrupt shift in management practices and the well-documented

effects on vegetation structure, in concert with the discrete boundaries of the base and the overall limited spatial scale, formed the basis for our natural experiment approach.

Southwestern grasslands are characterized by a nonlinear predator guild subject to intensive asymmetrical intraguild predation. Coyotes (*Canis latrans*) are the numerically dominant predator, though they compete with bobcats (*Lynx rufus*) and badgers (*Taxidea taxus*). Swift foxes are an intermediate predator, subject to predation by all three higher-order predators as well as golden eagles (*Aquila chrysaetos*). Despite the diversity of predators, coyote predation is the main cause of swift fox mortality (Sovada et al. 1998, Matlack et al. 2000, Schauster et al. 2002a, Thompson 2006). The basal prey for these predators is characterized by a diverse small-mammal community (Ribble and Samson 1987); however the three most abundant species, Northern grasshopper mice (*Onychomys leucogaster*), Ord's kangaroo rat (*Dipodomys ordii*), and deer mice (*Peromyscus maniculatus*), dominate the community (Thompson 2006). Black-tailed jackrabbits (*Lepus californicus*) and desert cottontails (*Sylvilagus audubonii*) also form a large portion of the higher-order predators' diet, and are infrequently killed by swift foxes (Kitchen et al. 1999).

Study design

In order to deal with the range of spatial scales used by predators and prey, we developed a hierarchical study design. We identified six study sites in areas subjected to three land-use regimes: livestock grazing,

mechanized military training, and unused (Fig. 2). Two study sites were located in each land-use regime. At each site, we established a 10-km trapping transect. Within each site, we randomly placed 50×70 m sampling grids within 1 km of the transect and at the rate of four grids per site per season over three years. We used a random number generator to create a distance along the transect, a direction (right or left), and a distance from the transect. This point became the northwest corner of the grid. These grids served as sampling units for both small-mammal trapping and vegetation structure surveys. We defined seasons as: winter, 15 December–14 April; summer, 15 April–14 August; and fall, 15 August–14 December, corresponding to swift fox breeding, pup-rearing, and dispersal behavior. New random locations were selected each season, resulting in 12 grids sampled per site per year.

While we assumed that differences in vegetation structure resulted primarily from differences in land use, each study site was considered an experimental unit due to the intrinsic small-scale variation between them. We attempted to minimize the effect of within-site heterogeneity through replication and the distribution of sites; however additional uncontrollable and confounding factors such as disturbance intensity, cattle stocking rates, and the degree of fire suppression precluded the use of a treatment–control design. Throughout the study, dispersing foxes regularly moved between sites indicating that all sites were potentially available habitat.

Field methods

We captured foxes using double-door box traps (Tomahawk Live Trap Company, Tomahawk, Wisconsin, USA) baited with raw chicken (Karki 2003). Traps were placed 500 m apart along each 10-km trapping transect bisecting each study site, resulting in 21 trap locations per study site. Each trap was oriented and covered with brush to provide protection from exposure. Traps were set in the late afternoon, checked early the following morning, and left closed throughout the day. Each site was trapped for four consecutive nights three times per year. For recollaring or targeting animals, a trap-enclosure system was used at den sites (Kozłowski et al. 2003). Captured foxes were handled without anesthesia, weighed, sexed, and aged through tooth wear (adult, juvenile). Foxes were considered juvenile until the pup-rearing season following their birth (15 April). Foxes were ear-tagged and collared with 30–50 g radio transmitters (Advanced Telemetry Systems, Isanti, Minnesota, USA).

We located foxes a minimum of three times per week, twice during nighttime hours when animals were actively hunting and once during daylight hours to locate den sites. Mortality sensors within transmitters indicated when a collar had been stationary for 4–6 hours. When a mortality signal was detected, the transmitter was recovered immediately and the location was recorded.

Efforts to determine the cause of death included searching the area for tracks and other sign as well as through necropsy of any remains (Disney and Spiegel 1992). We used Program MARK to calculate density and survival rates based on the known fate, logistic regression model (White and Burnham 1999). Known fate models based on telemetry data effectively accommodate staggered entry designs and use locations as recapture events to maximize estimate accuracy.

Coyote abundances were estimated using a scent station relative abundance index (Knowlton 1984). This has been shown to be a reliable, noninvasive method for estimating abundances of shy or nocturnal predators (Harrison et al. 2002). Scent stations consisted of a 1 m circle of sifted soil baited with a plaster disk soaked with fish oil. Stations were placed at 500-m intervals along each trapping transect, resulting in 21 stations per study site. Stations were baited and monitored for three consecutive nights, once each season. Tracks were identified and erased each morning; unknown tracks were digitally photographed for further identification. We estimated lagomorph abundances based on vehicle-based spotlight surveys conducted along each transect (Barnes and Tapper 1985, Ralls and Eberhardt 1997). Surveys were done over three consecutive nights once each season, and nightly totals were averaged for a seasonal count (Schauster et al. 2002b).

On each sampling grid, we evaluated vegetation structure using eight 50-m, north–south line transects following the short axis of the grid and spaced 10 m apart. Along each transect, we measured vegetation type and height at 1-m intervals (Dale 1999:41). Parameters estimated for each grid included live basal percent cover, percentage bare ground, percentage litter, mean grass height (centimeters), mean shrub height (centimeters), and shrub density (shrubs/100 m²). We identified shrubs to species but not grasses or forbs.

Immediately following vegetation sampling, we placed 35 Sherman live traps with 10-m spacing throughout the grid. Traps were baited with equine sweet feed, a mix of corn, oats, and molasses. Trapping grids were run for four consecutive nights, checked and closed each morning, and reset each afternoon. This method reduced our likelihood of capturing diurnal rodents such as ground squirrels, which due to high summer temperatures was required. Captured rodents were marked with Sharpie pens on the tail and abdomen allowing for identification of recaptures over the four-day trapping period. Relative abundance for each species was estimated based on the number of individuals captured. Available prey biomass for each grid was calculated as either total (number of individuals \times average mass) or per capita (total biomass/total number of captures).

Data analysis

We used one-way ANOVA (SASv8, SAS Institute, Redlands, California, USA) to test whether transects differed in vegetation structure and predator communi-

ty, followed by two-tailed *t* tests for unequal variances to determine statistically similar groupings. Ratios of predator abundance were regressed against vegetative structure and prey base variables to assess the relative influence of lower trophic levels on the predator guild. In all analyses, we used annual estimates of survival and density for each site as dependent variables. We used the Sobel test (K. J. Preacher and G. J. Leonardelli, *unpublished software*) to evaluate the influence of vegetation structure on the relationship between coyote abundance and either swift fox density or survival.

We evaluated the relative strength of food web linkages using two methods: hierarchical partitioning (Chevan and Sutherland 1991) and information theoretic model averaging (Burnham and Anderson 2002). We used two methods due to the inherent weakness of AIC model averaging when dealing with a large number of candidate models, the tendency of different multivariate techniques to give slightly different answers, and our overall interest in interpreting food web patterns as opposed to the statistical significance of a particular relationship. We used hierarchical partitioning code for R written by Chris Walsh and available online at the Comprehensive R Archive Network (*available online*).⁴ This analysis calculates the goodness of fit for a single dependent variable to all possible combinations of independent variables in a multivariate data set, and partitions the explained variance. This analysis was conducted for each of the seven species (deer mice, northern grasshopper mice, Ord's kangaroo rat, desert cottontail, black-tailed jackrabbit, swift fox, coyote) independently using lower trophic level variables as independent, explanatory variables.

For each of the seven species, we then constructed a suite of candidate models consisting of all possible one, two, or three explanatory variables from lower trophic levels. We calculated AIC weights for each candidate model and summed the weights for all models in which a specific independent variable occurred as described by Burnham and Anderson (2002:167) in order to quantify the importance of individual model parameters. AIC weights depend upon the suite of models used; therefore values represented the relative importance of individual connections to a particular species but are not directly comparable between species. While using a large suite of candidate models is dangerous in AIC model selection, our intention was to explore the role of community web components using model averaging and not to create explanatory models. Under such an objective, an "all possible subsets" approach helps to explain individual variable contributions and avoid problems with multicollinearity (Graham 2003).

Due to the difficulties in combining data across spatial and temporal scales and the influence of spatial scale on food web topology (Navarrette et al. 2000, Thompson

and Townsend 2005), we used several data sets to illustrate food web dynamics. Interactions involving coyotes or swift foxes were evaluated at the spatial scale of study sites and temporal scale of the three-year study period due to their capacity for movement and relatively long generation time. We evaluated the relationship of lagomorphs to vegetative structure at the spatial scale of study sites but the temporal scale of individual years due to their rapid population dynamics. Finally, we evaluated the relationship between small mammals and vegetative structure at the scale of individual sampling grids. This hierarchy of scales allowed us to most appropriately represent interactions between different trophic levels. Interactions between species operating at different scales were scaled up to the coarser of the two scales.

RESULTS

Site differences

Between 20 November 2001 and 27 November 2004, 116 swift foxes were captured 238 times; 109 foxes were fitted with radio collars. Captures were not distributed equally among sites ($\chi^2 = 26.6$, $df = 5$, $P < 0.001$), with 86% of all captures occurring on the grazed or military sites and only 14% occurring on unused sites. Fifty-five confirmed deaths occurred (38 adult, 17 juvenile). Of these deaths, 22 (40%) were coyote predation, three (5%) badger predation, three (5%) vehicle collision, two (4%) golden eagle predation, one (2%) bobcat predation, and 24 (44%) unknown causes. Survival estimates did not differ significantly between seasons ($F = 0.01$, $df = 2$, 27 , $P = 0.99$), by year ($F = 0.98$, $df = 2$, 27 , $P = 0.386$), by age ($F = 0.02$, $df = 2$, 27 , $P = 0.891$), or by site ($F = 0.57$, $df = 5$, 24 , $P = 0.721$) (Table 1). Population density estimates of swift foxes differed by season and site (Table 1). Site was the most important factor for density estimates ($F = 5.78$, $df = 5$, 24 , $P = 0.004$, $R^2 = 0.385$). Season was a marginally significant variable influencing fox density ($F = 3.07$, $df = 2$, 27 , $P = 0.057$); however its inclusion raised the R^2 from 0.385 to 0.467.

We evaluated vegetation structure on 185 sampling grids across the six study sites between December 2001 and August 2004. Basal percent cover did not vary significantly between sites, while percentage litter, mean grass height, and shrub density were significantly different between sites (Table 2). With respect to mean grass height, the Bent and Private sites were significantly different from each other as well as the other four sites. The remaining four sites did not differ. With respect to shrub density, three significantly different groups were indicated: Bent, Biernacki's, Private, and Red Rocks; Pronghorn and Private; and Comanche. Percentage litter also indicated three significantly distinct groups: Bent, Biernacki's, and Private; Biernacki's, Comanche, and Red Rocks; and Comanche, Pronghorn, and Red Rocks. These groupings did not correspond to land-use treatments, indicating heterogeneity between local landscapes and disturbance regimes.

⁴ (<http://www.r-project.org>)

TABLE 1. Estimates (mean, with SE in parentheses) of population density and survival rates for adult swift foxes on six sites in southeastern Colorado, USA, 2001–2004.

Site	Density (no./km ²)	Annual survival	Seasonal survival rates		
			Breeding	Pup rearing	Dispersal
PRV	0.18 (0.10)	0.54	0.81 (0.09)	0.83 (0.08)	0.81 (0.09)
COM	0.04 (0.05)	0.92	0.92 (0.08)	1.0 (0.0)	1.0 (0.0)
BTS	0.11 (0.08)	0.54	0.84 (0.08)	0.78 (0.10)	0.82 (0.08)
PRN	0.09 (0.06)	0.54	0.73 (0.11)	0.94 (0.06)	0.79 (0.09)
RRK	0.05 (0.03)	0.50	0.80 (0.18)	0.83 (0.15)	0.75 (0.22)
BNT†	0.03 (0.05)				

† Throughout the study, only one animal remained and/or survived on the Bent site throughout a full season. As a result we were unable to estimate survival rates for that site.

Predator distribution

Swift fox densities on PCMS are negatively correlated with grass height, while coyote abundance is positively correlated with shrub density (Thompson 2006). Our simple linear regression results indicated that the ratio of swift fox density to coyote abundance was negatively related to all four vegetation structure variables measured (Fig. 3). However, three of the four relationships were statistically insignificant. Only shrub density was significantly negatively related to the predator ratio ($R^2 = 0.35, P = 0.02$).

Prey distribution

Lagomorph abundance was significantly positively correlated with shrub density ($R^2 = 0.33, P = 0.006$; Fig. 4). All other lagomorph/vegetation structure relationships were positive relationships; however, they were either statistically insignificant ($P > 0.05$) and/or had R^2 values ≤ 0.03 . These relationships did not vary seasonally, with the exception of black-tailed jackrabbits being observed more often in areas with taller grass during the winter season ($R^2 = 0.37, P = 0.04$).

Small-mammal communities were sampled on 185 grids throughout the study. Northern grasshopper mice, Ord’s kangaroo rat, silky pocket mice (*Perognathus flavus*), western harvest mice (*Reithrodontomys megalotis*), white-footed mice (*Peromyscus leucopus*), southern plains woodrat (*Neotoma micropus*), 13-lined ground squirrels (*Spermophilus tridecemlineatus*), deer mice, and spotted ground squirrels (*Spermophilus spilosoma*) accounted for >99% of all captures. Three species,

Northern grasshopper mice, deer mice, and Ord’s kangaroo rat, accounted for 76% of all captures. Total small-mammal biomass was highest at moderate grass height (Fig. 5), decreasing in either direction. Small-mammal biomass peaked at either low or high shrub densities, primarily due to ground squirrel captures in areas of low shrub density and southern woodrat captures in areas of high shrub density. Total captures, another indicator of prey availability, was also highest at moderate grass height and peaked in areas of high shrub density (Fig. 5).

Hierarchical partitioning results indicated a variety of relationships between grassland prey species and vegetation structural characteristics (Fig. 6). Vegetation structure accounted for 14% of the variance in desert cottontail abundance, and >90% of this resulted from a negative association with grass height and shrub density. Vegetation structure also accounted for 14% of the variance in black-tailed jackrabbit abundance, with 65% of this stemming from a positive association with shrub density. Relationships of the three most common small-mammal species with vegetation structure varied; northern grasshopper mice were associated with areas of increased grass height and basal percent cover while Ord’s kangaroo rats were negatively associated with basal percent cover and shrub density. Deer mice favored areas of increased basal cover and shrub density but avoided areas with high percentage litter cover.

Food web interactions

Hierarchical partitioning results indicated a variety of relationships between predators and both lower trophic

TABLE 2. ANOVA results indicating differences in vegetation structural variables between study sites in southeastern Colorado, 2001–2004.

Site	Basal percent cover		Litter (%)		Grass height (cm)		Shrub density (no./100 m ²)	
	Mean	Variance	Mean	Variance	Mean	Variance	Mean	Variance
Bent	0.44	0.05	0.17	0.01	12.79	23.54	1.15	0.90
Biernacki’s	0.40	0.03	0.18	0.02	9.53	25.99	0.71	0.98
Comanche	0.45	0.04	0.25	0.02	10.60	9.80	2.71	4.07
Pronghorn	0.35	0.02	0.26	0.02	8.92	18.24	0.26	0.08
Private	0.38	0.04	0.18	0.01	6.69	15.58	0.92	4.98
Red Rocks	0.44	0.03	0.23	0.02	9.36	8.91	0.68	0.72

Note: For basal cover, $F_{5,179} = 1.38, P = 0.23$; for litter, $F_{5,179} = 3.30, P = 0.01$; for grass height, $F_{5,179} = 6.84, P = <0.01$; and for shrub density, $F_{5,179} = 12.01, P = <0.01$.

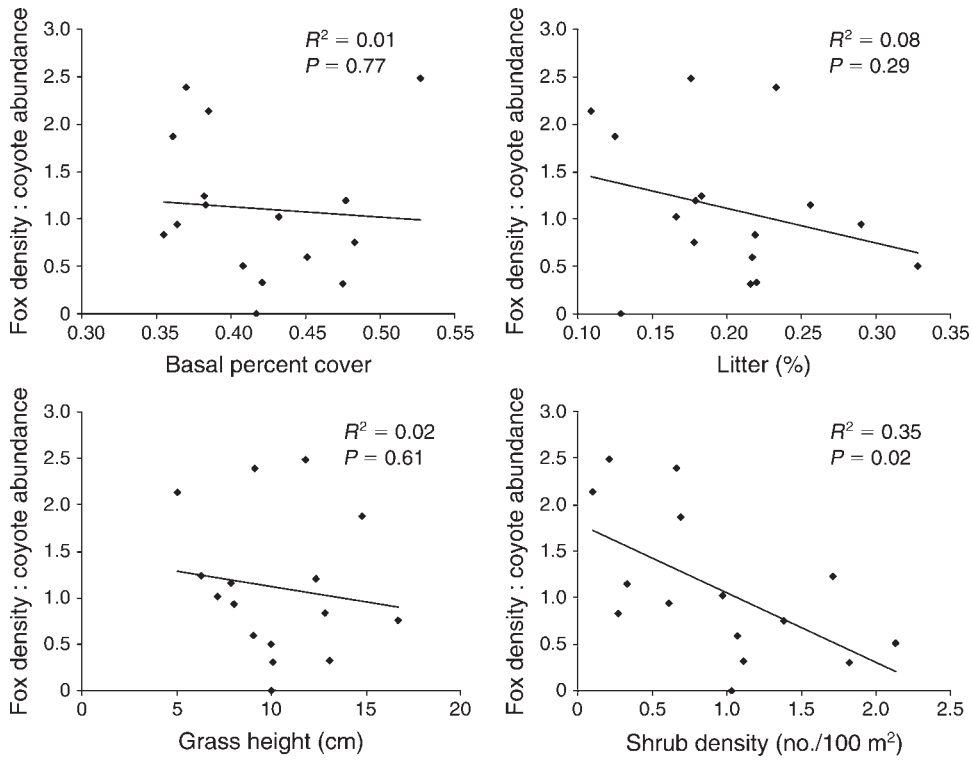


FIG. 3. Influence of vegetation structure variables on the ratio of swift fox density to coyote abundance in six study sites in southeastern Colorado, USA. Values represent seasonal averages for each study site between 2001 and 2004.

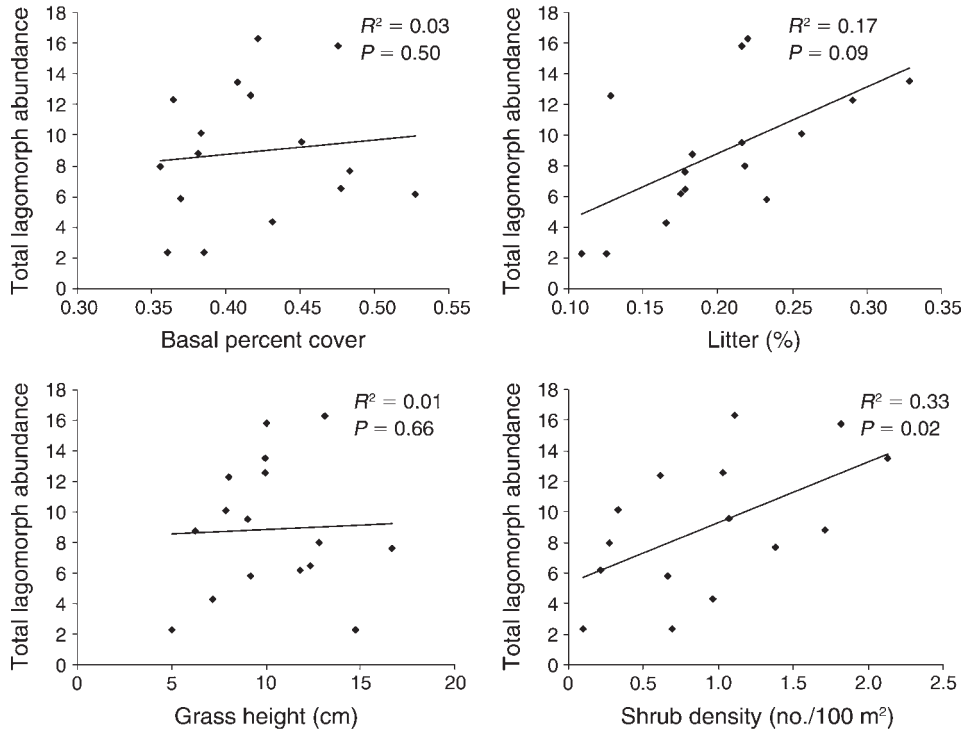


FIG. 4. Relationship between total lagomorph abundance (black-tailed jackrabbit and desert cottontail) and vegetation structure on the Pinon Canyon Maneuver Site, Colorado, USA. Values represent seasonal averages for each study site between 2001 and 2004.

levels and vegetation structure (Fig. 7). Of the total variance in swift fox density, 44% was explained by the independent and joint contribution of prey species distribution, and another 34% was explained by vegetation structure variables. Of these, the dominant relationship was a negative association with black-tailed jackrabbit abundance, which accounted for 25% of the total explained variance. Forty-five percent of the variation in coyote abundance was explained by the same food web variables; of this, 68% was contributed by positive relationships with all four vegetation structure variables and another 17% was contributed by a positive relationship with black-tailed jackrabbit abundance. More importantly, coyotes and swift foxes showed a nearly perfect (eight of nine variables) pattern of opposite interactions; when coyotes were positively associated with a variable, swift foxes were negatively associated with the same variable and vice-versa (Fig. 7). Only percentage litter was positively associated with both predators.

This pattern of positive and negative interactions in food web dynamics was also indicated in the AIC model averaging results (Fig. 8). A negative relationship between one predator and a prey species was always balanced by a positive relationship between the other predator and the same prey species. Coyote abundance was positively related to three of five prey species and to all structural measures, though only the coyote:jackrabbit and coyote:shrub density relationships were statistically significant ($R^2 = 0.21$ and 0.96 , $P = 0.005$ and 0.0005 , respectively). In contrast, swift fox density was positively related to the remaining two prey species and negatively related to three of four structural measures, though again only one of each was statistically significant (swift fox:jackrabbit, $R^2 = 0.18$, $P = 0.01$; swift fox:grass height, $R^2 = 0.76$, $P = 0.02$). Only black-tailed jackrabbit abundance was significantly related to both predator populations, positively to coyotes and negatively to swift foxes.

The majority of interactions remained consistent, with respect to sign, significance, and magnitude, across seasons with two exceptions. First, the magnitude of the coyote:jackrabbit:swift fox relationship increased during winter. The R^2 value of the coyote/jackrabbit relationship increased from 0.21 to 0.54, and from 0.18 to 0.61 for the jackrabbit/swift fox relationship. Second, predator-prey relationships weakened during fall months with R^2 values dropping an average of 0.08. This dilution most likely represents a combination of enhanced movement during the dispersal season as well as the increased availability of arthropod prey during fall months (Kitchen et al. 1999).

Sobel test results indicated that only shrub density significantly altered the relationship between swift fox survival and coyote abundance. Fox survival was not significantly related to either coyote abundance or shrub density independently. However including shrub density in the fox survival:coyote abundance model raised the

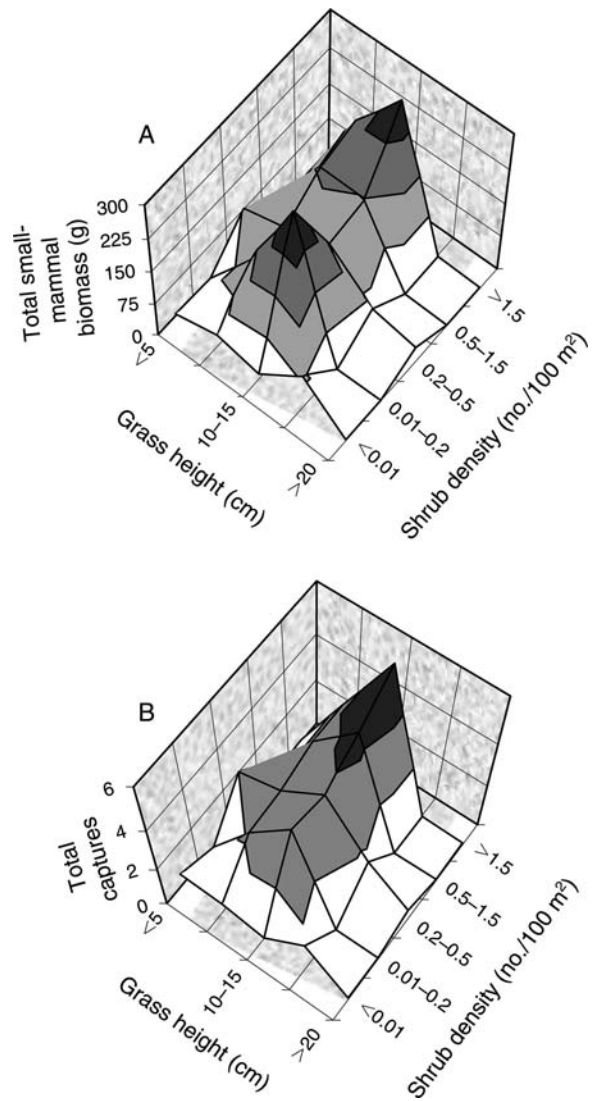


FIG. 5. Distribution of small-mammal (A) biomass and (B) captures in relation to vegetation structure in southeastern Colorado, 2001–2004.

R^2 value from 0.180 to 0.512 and lowered the P value from 0.080 to 0.007, a difference significant at the $P = 0.015$ level (Table 3). Biologically, this indicates that in shrub-dominated habitat, swift fox survival is negatively related to coyote abundance, while in more open habitat it is not. No other vegetation structure variable significantly altered either the fox survival:coyote abundance or fox density:coyote abundance models.

DISCUSSION

Traditional predator-prey models often predict what Sih (1998) termed the “leapfrog” effect, in which predators are most abundant in high quality prey habitat. This elevated predation risk moderates the advantage of habitat quality, and prey are more evenly distributed. This pattern has been observed in many

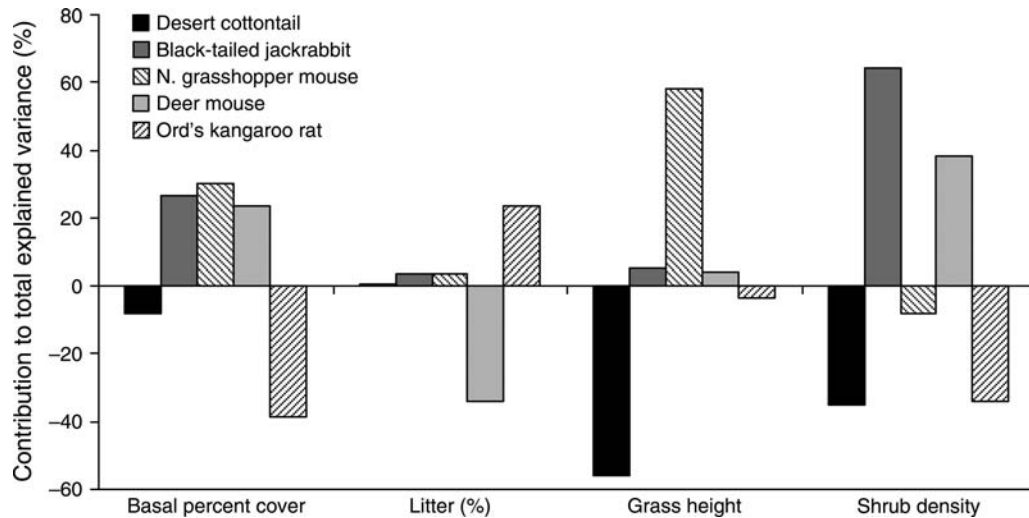


FIG. 6. Hierarchical partitioning results for prey species. Values indicate the percentage of the total explained variance for each species contributed by each explanatory variable. Negative values indicate a negative relationship.

linear trophic relationships (Kitchell et al. 1994). However, nonlinear relationships appear to be more common in natural systems (Rosenheim et al. 1995, Polis and Strong 1996), and behavior such as intraguild predation has the capacity to complicate predator–prey dynamics. In addition, while vegetation structure has been suggested as influential on arthropod predator–predator interactions (Roda et al. 2000, Finke and Denno 2002), evidence from vertebrate systems is almost nonexistent.

Recent theoretical work has outlined two paths to system stability under intraguild predation. One is that intermediate predators must specialize and outcompete top predators for the shared basal prey in order to persist (Holt and Polis 1997). The second is that intermediate predators must disassociate from the more

linear top predator–basal prey relationship and do what has been termed “safety-matching,” selecting habitat based on security from predation rather than resource availability (Heithaus 2001, Rosenheim 2004).

In our study area, predator–prey dynamics appear to strongly conform to the second prediction, that top predators “resource match,” intermediate predators “safety match,” and shared basal prey are more evenly distributed. Both coyotes and swift foxes are opportunists, sharing prey species and capable of exploiting a range of habitats. While the majority of the predator–prey relationships were statistically insignificant, the repeated pattern of positive–negative relationships across multiple analytical techniques is strong indication of biological significance.

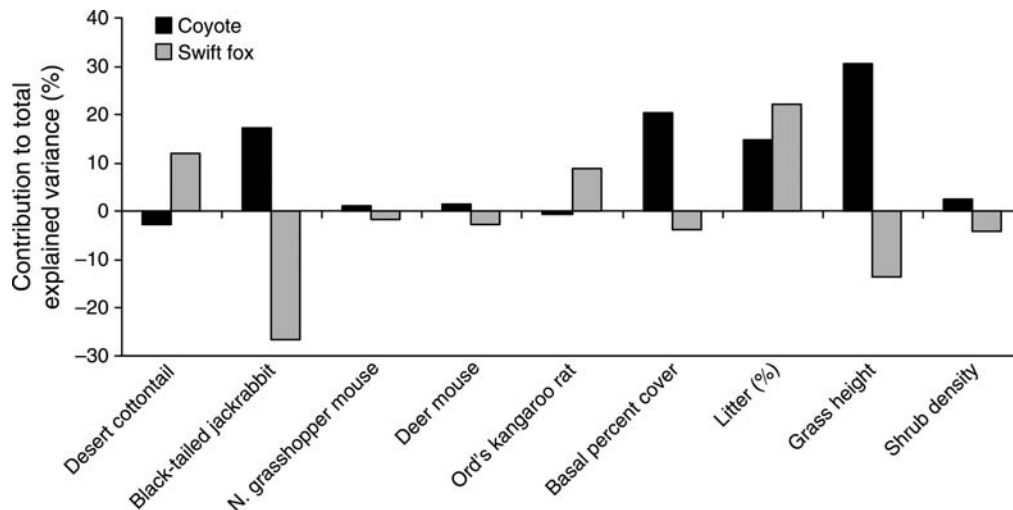


FIG. 7. Hierarchical partitioning results for two predator species. For each predator species, values indicate the percentage of the total explained variance in either abundance (coyote) or density (swift fox) contributed by each explanatory variable. Negative values indicate a negative relationship.

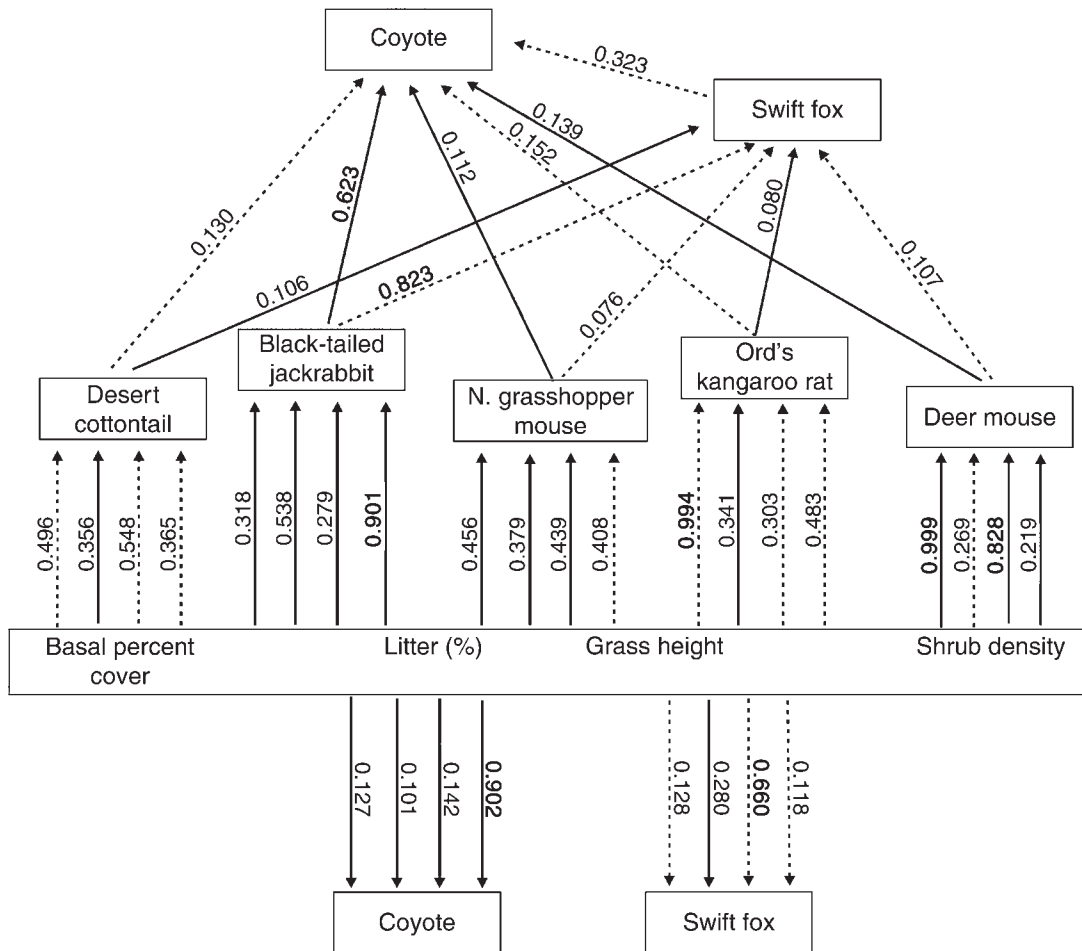


FIG. 8. Simplified food web diagram for southern Colorado showing the relationships between (A) all trophic levels and (B) vegetation structure and higher-order predators. Numbers indicate the summed AIC weights for all models in which that relationship occurred, and reflect the importance of an independent variable in the distribution of a particular species. Values >0.60 are indicated in boldface type. Solid arrows indicate positive relationships; dashed arrows indicate negative relationships. Arrows leaving the vegetation level indicate the four vegetation structure measurements in order. All values associated with arrows entering a particular species are directly comparable and reflect the relative importance of individual parameters. Values associated with arrows entering different species are not directly comparable due to the use of different candidate model suites for each species.

Both coyotes and swift foxes showed strong responses to vegetation structure. As predicted based on the leapfrog effect, coyotes were most abundant in areas of enhanced structural diversity where prey are typically more abundant. In contrast, swift fox density was negatively related to three of four structural measurements, significantly so with grass height. While they are capable of exploiting a wide variety of landscapes (Matlack et al. 2000), swift foxes are a fossorial species and depend on visually detecting predators and quickly escaping to a nearby den (FaunaWest, unpublished report). Landscapes where increased vegetation structure restricts line-of-sight, in the presence of higher-order predators, are risky habitats that are avoided. This behavior is validated based on the results of the Sobel test: shrub density significantly influenced the relationship between swift fox survival and coyote abundance. The idea that plant resources or habitat complexity

moderates the strength of predator–predator and predator–prey interactions under intraguild predation has been well established in arthropod (Gratton and Denno 2003, Langelotto and Denno 2004) and aquatic (Hampton 2004) systems, but not in terrestrial vertebrate systems.

Our work suggests that changes in the disturbance regime have the capacity to exclude swift foxes from potential habitat due to subsequent changes in the vegetation structure. On the PCMS, the change in land use resulted in two interacting landscape trajectories; an increase in basal cover and grass height following the release from grazing and a reduction in basal cover, shrub height, and shrub density associated with military training (Shaw and Diersing 1990, Milchunas et al. 1999). The trajectory of landscape change appears to regulate the exposure of swift foxes to coyote predation more strongly than coyote abundance. Therefore while

TABLE 3. Linear regression and Sobel test results outlining the influence of coyote abundance and/or vegetation structure on swift fox density and survival in southeastern Colorado, 2001–2004.

Model	R^2	Regression P	Sobel test P †
Fox density = coyote abundance	0.020	0.575	
Fox density = shrub density	0.030	0.508	
Fox density = grass height	0.354	0.009	
Fox density = basal percent cover	0.179	0.080	
Coyote abundance = shrub density	0.581	0.0004	
Coyote abundance = grass height	0.0004	0.934	
Coyote abundance = basal percent cover	0.643	0.310	
Fox density = coyote abundance + shrub density	0.038	0.803	0.745
Fox density = coyote abundance + grass height	0.371	0.031	0.933
Fox density = coyote abundance + basal percent cover	0.180	0.225	0.371
Fox survival = coyote abundance	0.180	0.080	
Fox survival = shrub density	0.012	0.679	
Fox survival = grass height	0.037	0.440	
Fox survival = basal percent cover	0.001	0.905	
Fox survival = coyote abundance + shrub density	0.512	0.007	0.015
Fox survival = coyote abundance + grass height	0.226	0.146	0.934
Fox survival = coyote abundance + basal percent cover	0.199	0.188	0.597

Note: $N = 18$: fall, winter, and summer averages across six study sites over three years.

† Sobel tests are used to evaluate the relative change in regression coefficients and standard errors due to the inclusion of a third variable in the model. A value of $P < 0.05$ indicates that the inclusion of the third variable significantly altered the relationship between the dependent and independent variables.

the most visible source of swift fox mortality is coyote predation, the underlying cause is exposure to predation risk due to changes in landscape structure. This differentiation between direct sources of mortality and underlying causes may help explain the variation in published studies regarding the effect of coyote removal on swift fox population density (Kamler et al. 2003, Karki 2003).

Mammalian carnivores are relatively rare, wide ranging, and secretive, making empirical research on community interactions difficult (Fedriani et al. 2000, Switalski 2003). In particular, intraguild predation in mammalian carnivores, where the temporal and spatial scale of two or more such species must be considered, poses a significant challenge to wildlife ecologists (Creel and Creel 1996, Fedriani et al. 2000). As a result, while there is rapidly accumulating evidence that intraguild predation exists and plays an important role in shaping terrestrial predator guilds (see Macdonald and Sillero-Zubiri 2004 for a review) most evidence stops at documenting the interaction. There is currently very little information on the behavioral implications for either top or intermediate predators or the role of moderating factors such as landscape structure or habitat complexity.

Intraguild predation is a powerful evolutionary force capable of influencing species abundance, distribution, and behavior (Polis et al. 1989, Navarrette et al. 2000). In systems where the intraguild predation pressure is strongly asymmetrical, selection can be expected to favor behavioral responses by intermediate predators that reduce the probability of an interaction with a higher-order predator (Polis and McCormick 1987, Gerber and Echternacht 2000). Such behavioral changes have been documented for scorpions (Polis and Mc-

Cormick 1987), spiders (Wilder and Rypstra 2004), and lizards (Losos and Spiller 1999). Similar responses have been suggested in red fox (*Vulpes vulpes*)/coyote (Voigt and Earle 1983), coyote/wolf (*Canis lupus*; Switalski 2003), cheetah (*Acinonyx jubatus*)/lion (*Panthera leo*; Durant 2000) interactions; however the behavioral response of the intermediate predator and the mechanism of avoidance were not directly addressed. To our knowledge, only one study concerning intraguild predation between mammalian carnivores has specifically addressed the behavioral response of the intraguild prey. The density of African wild dogs (*Lyacon pictus*) is lowest where their primary prey, impala (*Aepyceros melampus*) and kudu (*Tragelaphus strepsiceros*), is most abundant (Mills and Gorman 1997). The authors concluded that wild dogs avoided prey-rich areas in order to avoid lions, the primary source of wild dog mortality, a behavior identical to that displayed by swift foxes in our study. In a more recent experimental study, Morris (2005) found that red-backed voles (*Clethrionomys gapperi*) altered their habitat selection to avoid food supplements, a paradox explained only when black bear (*Ursus americanus*) predation is taken into account.

While this was an observational study, the spatial and temporal scale of terrestrial carnivore behavior precludes most experimental approaches. The abrupt change in land management practices and the resulting shift in vegetation structure over the last 20 years, as well as the discrete boundaries maintained by the U.S. Army at the PCMS, created a heterogeneous system at a spatial scale appropriate for terrestrial carnivore research. While the intraguild predation of coyotes on swift foxes has been well documented, the subsequent effect on fox populations and behavior has not been addressed (Moehrenschrager et al. 2004). In this system,

coyotes and swift foxes appeared to follow the predictions of resource and safety-matching outlined by Heithaus (2001) and Rosenheim (2004). Mesocarnivores, subject to a variety of selective forces, are often species of concern, and the mechanism by which they avoid higher-order predation is of direct management concern. Our results support Hunter and Price's (1992) contention that top-down (predation) and bottom-up (vegetation structure) act simultaneously and interact to control population densities (Leibold 1989), and suggest that vegetation structure plays a crucial role in moderating intraguild predation pressure on intermediate predators in terrestrial systems.

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