Attributes of rodent burrows include measures of tunnels (e.g., diameter, angle of incline, depth), density of surface openings, configurations of surface openings (e.g., presence or absence of soil mounds), connectivity of openings via tunnels, and blockage of connecting tunnels with soil plugs. Burrow plugging is observed in many rodent species that occupy burrow systems. For instance, in Columbian ground squirrels (*Otospermophilus columbianus*) and Merriam’s kangaroo rats (*Dipodomys merriami*), plugging behavior is commonly observed during the breeding season, where it may shield preweaned litters in natal dens from predators.

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infanticide (Reynolds 1960, McLean 1978). Werner et al. (2005) also noted plugging behavior in pocket gophers (Thomomys bottae navus) in response to the presence of cold temperatures and light, while Thomas (1974) reported that eastern chipmunks (Tamias striatus) plug food caches underground and appear to use earthen plugs as protection against predators. Burrow plugging seems to be a natural manifestation of “defensive burying” and other substrate movement responses that have been measured in psychometric experiments on captive rodents over many years (De Boer and Koolhaas 2003)

We investigated several attributes of the burrows of black-tailed prairie dogs, including density and configuration of surface openings (discussed below), but connectivity of openings as influenced by plugging was a primary focus. Black-tailed prairie dogs (Cynomys ludovicianus, henceforth “prairie dogs”) surface-plug openings to their burrow systems (Smith 1958, Henderson et al. 1969, Sheets et al. 1971) and sometimes construct subsurface plugs within tunnels (Sheets et al. 1971). Various explanations have been proposed for burrow plugging by prairie dogs. Female prairie dogs plug “auxiliary” openings to nesting chambers containing recently born pups, presumably in defense against infanticide (Hoogland 1995: 29). Shooting and poisoning induce plugging behavior by prairie dogs (Campbell and Clark 1981), suggesting prairie dogs plug burrow openings to entomb dead conspecifics below-ground (Smith 1958). The presence of black-footed ferrets (Mustela nigripes, henceforth “ferrets”), which are semifossorial obligate predators of prairie dogs (Sheets et al. 1972, Campbell et al. 1987), and snakes also elicit plugging by prairie dogs (Hillman 1968, Henderson et al. 1969, Forthenbery 1972, Halpin 1983, Eads and Biggins 2012). Subsurface plugs that result from changes in tunnel design by the prairie dogs (Wilcomb 1954) might reduce prairie dogs’ risk of predation by semifossorial predators, such as ferrets.

Prairie dogs must expend energy to dig and plug burrows. Burrowing can be 360–3400 times more energetically expensive than walking an equal distance (Veleck 1979, but see Bozinovic et al. 2005), and costs can vary by a factor of 9 depending on whether the soil is sand or clay (Veleck 1981). Costs of soil manipulation can be split into those associated with shearing packed soil and those associated with transport of loose soil. Shearing costs are likely low when prairie dogs use loose soil and litter around openings to create surface plugs, but these costs could be high when subsurface plugs are created using soil excavated from within a tunnel. Transport costs can vary due to steepness and length of the incline (Luna and Antinuchi 2007). Complete replacement of plugged burrows with new burrow construction, as observed by Halpin (1983), would involve high energy expenditures.

The indirect impacts of ferret presence likely extend beyond the energetic expenditures of digging and plugging. Prairie dogs may be trading time spent plugging for time that they might use for foraging, resting, or other activities. Also, the presumed benefits of multi-opening burrows, such as air circulation (Vogel et al. 1973, King 1984) and predator avoidance (Hoogland 1995), are likely compromised by surface and subsurface plugging. For example, American badgers (Taxidea taxus) commonly dig into burrows to capture prey (Michener 2004) and may be able to detect single-opening systems, where prey presumably would be most vulnerable (Eads and Biggins 2008 and other citations therein). Surface plugs would also seem to reduce the number of burrow openings that are available to prairie dogs seeking refuge from raptors, coyotes (Canis latrans), or other predators that attack aboveground. If prairie dog plugging behavior increases risk of predation for prairie dogs, it might also do so for ferrets, and this behavior might be an example of a prey species (prairie dog) trading increased vulnerability to generalist predators for reduced vulnerability to a presumably more efficient specialist predator (Hassell and May 1986, Hanksi et al. 1991).

We add anecdotally that our own excavations by shovel and steel bar to recover radio transmitters and the remains of ferrets killed by badgers (Biggins 2000, Biggins et al. 2006b), along with our experience with sampling prairie soils using hydraulic soil machines (Eads personal observations), have left a lasting impression regarding the difficulties and energy required to penetrate ≥2 m vertically through prairie sod and compacted subsoils. These experiences have generated great respect for the animals that routinely create burrow systems in this environment. Because of the difficulties
in excavating prairie dog burrows (King 1955, Sheets et al. 1971), subsurface attributes of prairie dog burrow systems have rarely been studied. A few studies have documented complex burrow systems with multiple connections (Wilcomb 1954, Sheets et al. 1971, Verdolin 2008), as well as simple tunnels with a single opening at the surface (Stromberg 1978).

In this study we aimed to further examine influences of ferrets and recreational shooting on rates and locations of burrow plugging by prairie dogs. To investigate influences of ferrets, we examined surface plugging and subsurface burrow connectivity in prairie dog colonies occupied by ferrets and in colonies with no known ferret presence. We also examined plugging and connectivity in prairie dog colonies where shooting was allowed and in colonies protected from shooting.

We anticipated that prairie dog colonies with resident ferrets would exhibit higher rates of surface and subsurface plugging than colonies with no known ferrets (because prairie dogs might use both forms of plugging in defense against ferrets). On colonies where shooting was permitted, we anticipated higher rates of surface plugging compared to colonies protected from shooting (because prairie dogs might bury the remains of their shot counterparts). In addition, presence of shooting or ferrets might reduce rates of burrow connectivity because these factors tend to reduce numbers of prairie dogs (i.e., excavators that maintain burrow systems), perhaps leaving burrow systems in disrepair.

Presence of ferrets or shooting might also affect density and configuration of prairie dog burrow openings. Configurations of openings have been classified according to presence or absence of soil mounds (Hoogland 1995). During construction of a new burrow system, prairie dogs leave dome-shaped deposits of soil around the opening at which the excavation began (Smith 1958, Hoogland 1996). Systems with multiple openings are thought to be created when prairie dogs dig upward at a steep angle and break through the surface at new locations (King 1955, Sheets 1970). The additional openings created do not have mounds until the prairie dogs construct them by scraping surface soil into crater-shaped mounds surrounding the new openings. Some openings with dome-shaped mounds remain as single-opening burrows. We thus hypothesized that openings with dome-shaped mounds would have less frequent subsurface connections than would other types of openings. That relationship might be modified, however, if prairie dogs dig new burrow systems soon after they surface-plug existing burrows as reported by Halpin (1983). Newly added burrows would be expected to alter ratios of mound types present and rates of connectivity associated with various types of openings. With increased prairie dog mortality (due to shooting or predation by ferrets) or increased frequency of burrows occupied by ferrets, we thus predicted an increase in the ratio of burrow openings with dome-shaped mounds compared to other types of mounds due to increased construction of new burrows. On colonies occupied by ferrets or on colonies where shooting was allowed, we might expect that openings with crater-shaped mounds or no mounds would have lower rates of connectivity to other openings than on colonies without ferrets or shooting.

METHODS

We sampled 15 prairie dog colonies on Badlands National Park and adjacent portions of the Buffalo Gap National Grasslands, South Dakota (Fig. 1). On many sites, vegetation is dominated by western wheatgrass (Pascopyrum smithii), blue grama (Bouteloua gracilis), and buffalograss (Bouteloua dactyloides). Where cattle graze with prairie dogs, various species of forbs dominate the landscape. We examined effects of ferrets and shooting on surface plugging rates and subsurface connectivity of burrow openings by contrasting colonies with ferrets but no shooting, colonies with shooting but no ferrets, and colonies with neither shooting nor ferrets (i.e., “control” colonies, Table 1). Shooting was prohibited on colonies believed to be occupied by ferrets, so effects of shooting and ferret occupancy could be separately evaluated. However, interactions between effects of ferrets and shooting could not be adequately assessed.

We defined colony boundaries using outermost burrow openings and the density criteria of Biggins et al. (1993). We classified the colonies to be sampled as occupied by ferrets if female ferrets with litters were detected on them via spotlight surveys (Biggins et al. 2006a) during summer 2009, but we did not attempt to estimate ferret densities. Unoccupied colonies
Fig. 1. Black-tailed prairie dog colonies in Badlands National Park and adjacent areas of Buffalo Gap National Grassland, South Dakota, on which connectivity and surface plugging of burrow openings were sampled during 2009. Colony numbers correspond to those in Table 1.

Table 1. Number of burrow openings sampled at black-tailed prairie dog colonies in Badlands National Park and adjacent areas of Buffalo Gap National Grassland, South Dakota. Treatment groups are colonies known to be occupied by black-footed ferrets (ferret), colonies where prairie dog shooting was allowed (shooting), and colonies with neither ferrets nor shooting (control). Numbers correspond to the locations on Figure 1.

<table>
<thead>
<tr>
<th>Colony</th>
<th>Number</th>
<th>Treatment</th>
<th>0.073-ha plots</th>
<th>Line surveys</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Focal burrows</td>
<td>Openings</td>
<td></td>
</tr>
<tr>
<td>Antelope</td>
<td>1</td>
<td>Control</td>
<td>12</td>
<td>223</td>
</tr>
<tr>
<td>East Quinn</td>
<td>2</td>
<td>Control</td>
<td>12</td>
<td>255</td>
</tr>
<tr>
<td>Pinnacles</td>
<td>3</td>
<td>Control</td>
<td>18</td>
<td>208</td>
</tr>
<tr>
<td>Prairie Winds</td>
<td>4</td>
<td>Control</td>
<td>12</td>
<td>239</td>
</tr>
<tr>
<td>West Basinger</td>
<td>5</td>
<td>Control</td>
<td>17</td>
<td>299</td>
</tr>
<tr>
<td>Agate North</td>
<td>6</td>
<td>Ferret</td>
<td>12</td>
<td>95</td>
</tr>
<tr>
<td>Conata West</td>
<td>7</td>
<td>Ferret</td>
<td>12</td>
<td>87</td>
</tr>
<tr>
<td>North Exclosure</td>
<td>8</td>
<td>Ferret</td>
<td>12</td>
<td>151</td>
</tr>
<tr>
<td>Roberts A</td>
<td>9</td>
<td>Ferret</td>
<td>12</td>
<td>176</td>
</tr>
<tr>
<td>Roberts B</td>
<td>9</td>
<td>Ferret</td>
<td>12</td>
<td>142</td>
</tr>
<tr>
<td>South Exclosure</td>
<td>10</td>
<td>Ferret</td>
<td>12</td>
<td>283</td>
</tr>
<tr>
<td>West Quinn</td>
<td>11</td>
<td>Ferret</td>
<td>12</td>
<td>201</td>
</tr>
<tr>
<td>Bigfoot EP 6</td>
<td>12</td>
<td>Shooting</td>
<td>12</td>
<td>99</td>
</tr>
<tr>
<td>Bigfoot EP 10</td>
<td>13</td>
<td>Shooting</td>
<td>12</td>
<td>106</td>
</tr>
<tr>
<td>White River</td>
<td>14</td>
<td>Shooting</td>
<td>13</td>
<td>293</td>
</tr>
<tr>
<td>TOTAL</td>
<td>192</td>
<td></td>
<td>2895</td>
<td>651</td>
</tr>
</tbody>
</table>
were those that lacked detection of ferrets by spotlight surveys or snow tracking (Biggins et al. 2006a) during the previous 12 months. We did not attempt to quantify the amount of shooting on colonies where shooting was allowed.

Surface and Subsurface Plugging

Burrow sampling involved examination of burrow openings for surface plugging and subsurface connectivity. We tallied surface plugs and total numbers of burrows on plots and line surveys. To determine burrow connectivity, we tested burrow systems with forced air from a gasoline-powered blower, a noninvasive means of determining connectivity (Eads and Biggins 2008, Biggins 2012).

From July through October 2009, we sampled 192 randomly selected focal burrow openings (i.e., unplugged openings) in the 15 study colonies (Table 1). We counted the total number of burrow openings and the number of surface-plugged openings within 15.24 m of the focal opening, creating a burrow-centered circular plot of 0.073 ha. Surface-plugged burrow openings were those for which the opening was completely filled with soil and litter (Henderson et al. 1969). As weathering degrades the fresh soil deposited into a burrow opening by prairie dogs, plugged openings presumably lose their definition, becoming nonvegetated mounds and depressions (without evidence of openings) that later support vegetation. Because these transitions are continuous, defining categories is difficult. We classified as plugged all nonvegetated mounds and small depressions lacking openings, without regard to evidence of recent soil disturbance.

At a burrow-centered circular plot, we used the blower on the focal burrow opening to test for an open connection to other burrow openings within the plot. We inserted the delivery tube of the blower into the focal burrow opening, sealed the space between the tube and tunnel walls with plastic bags and soil, and started the blower to create positive air pressure within the burrow system. Feathers were sequentially placed into all other burrow openings that were not surface-plugged; airflow within a tunnel caused feathers to be expelled from the opening. We sequentially plugged (with plastic bags and soil) burrow openings within the plot that were found to be connected to the focal opening. This action increased airflow to any remaining openings that were connected. After each opening was plugged, all remaining openings were reexamined with feathers. Because we had no noninvasive method to measure burrow structure in 3 dimensions, we measured the 2-dimensional Euclidian distance between the openings for focal burrows and burrows that exhibited airflow. Those distances underestimate actual distances through the connecting tunnels because tunnels must descend and ascend and often take circuitous routes laterally. During sampling, burrow openings were categorized with regard to the type of mound surrounding the opening (dome, crater, or nonmounded; sensu Sheets 1970).

We used Mann–Whitney tests to compare total burrow densities on colonies with ferrets or shooting to densities on colonies with neither source of mortality. We used logistic regression to compare plots on ferret-occupied colonies to plots on control colonies with regard to connectivity of burrow openings and with regard to surface plugging. The logistic regression procedure was repeated to compare plots on colonies where shooting was allowed to plots on control colonies. Binomial response variables were presence or absence of openings connected to the focal burrow and presence or absence of surface-plugged burrow openings. Because densities of burrow openings would be expected to influence the probability that a focal burrow would be connected to other burrow openings, we retained the total number of burrow openings that were not surface-plugged within a 0.073-ha plot as a covariate in the statistical models of connectivity and surface plugging rates. Predictor variables of primary interest were black-footed ferret occupancy of a colony and whether or not prairie dog shooting was allowed. We initially evaluated multivariate models using an information theoretic approach, with models ranked via Akaike’s information criterion (AIC). We used likelihood ratio (LR) tests to further compare certain nested submodels to more general models. For data from the 0.073-ha plots, we created point estimates and confidence intervals using simple models that did not include the relationship involving total burrow openings.

Numbers and Types of Burrow Openings

Because openings having crater-shaped mounds or no mounds are thought to be
terminus openings for systems initiated from an opening with a dome-shaped mound, their connectivity is expected to be more responsive to factors that alter the frequency of subsurface plugging (such as shooting and ferret-induced mortality). We thus separately compared these categories of colonies with respect to connectivity rates for openings with dome-shaped mounds and openings with combined crater-shaped mounds or no mounds. Separation of the sample into categories of burrow mounds produced inadequate sample sizes in some groups, so we compared pooled data from colonies with shooting and colonies with ferrets to data from the control colonies with neither of these sources of mortality. We used 2 simple logistic regression models: one for openings with dome-shaped mounds and one for openings with craters or no mounds.

Our 0.073-ha plots did not allow unbiased estimates of rates of surface plugging or ratios of types of burrow openings because the plots were centered on a burrow opening that was not surface-plugged. Thus, we also walked along lines of 1-m width and variable length that completely crossed the colonies, starting at a random point at the edge of the colony, following a predetermined azimuth, and ending at the opposite edge of the colony. Along the lines, we classified burrow openings as crater, dome, or nonmounded and noted surface plugs. In total, we classified 651 burrow openings on the 15 colonies where the 0.073-ha plots were sampled (Table 1).

We used contingency table analyses to compare the 3 categories of colonies with respect to ratios of types of burrow openings. And we used logistic regression as described above to assess rates of surface plugging at burrow openings with dome-shaped mounds and nonmounded openings. We censored crater-shaped mounds from this logistic regression analysis because the rarity of plugging there caused convergence problems during iterative maximum likelihood estimation. For all analyses, we assessed the effect of ferret occupancy using colonies with no shooting and the effect of shooting using colonies with no ferrets.

**RESULTS**

Density of burrow openings was 239.0 ha⁻¹ on colonies with neither ferrets nor shooting, compared to 159.9 ha⁻¹ on colonies with shooting (Mann–Whitney $\chi^2_{1} = 23.904, P < 0.001$) and 167.2 ha⁻¹ on colonies with ferrets (Mann–Whitney $\chi^2_{1} = 31.138, P < 0.001$). These estimates are likely higher than true densities of burrow openings because the 0.073-ha plots were always centered on an opening. Focal burrow openings were connected to a maximum of 9 additional openings in the plots, and there were up to 8 surface-plugged burrow openings in these plots. Because a few connected burrow openings were near the perimeters of the plots, it is likely that a few additional subsurface connections existed beyond the 15.24-m radius of sampling.

In the analysis of ferret effects on connectivity of burrow openings within 0.073-ha plots, competitive models (ΔAIC within 3 units of model with lowest AIC) retained the ferret variable (Table 2). The ferret variable was also highly influential when the general model with effects of ferrets and total burrows was compared to the reduced model with total burrows only (LR $\chi^2_{1} = 7.814, P = 0.005$). Presence of ferrets was associated with relatively few plots having connected burrow openings (Fig. 2). In a separate analysis, effect of shooting was not influential (Fig. 2) as judged by the most supported model with constants only (Table 2), and explanatory power of shooting was weak when assessed by model contrasts as above (LR $\chi^2_{1} = 0.664, P = 0.415$).

In the analysis of ferret effects on surface plugging within plots, competitive models again retained the ferret variable (Table 2), and a contrast of models as above strongly supported the contention that ferret presence influenced the frequency of surface plugging (LR $\chi^2_{1} = 16.928, P < 0.001$). Ferret presence was associated with an abundance of plots with burrow openings that were surface plugged (Fig. 2). Similar to results for subsurface connections, there was little evidence for effect of shooting on frequencies of surface plugging in plots. The most supported model had only the constant (Table 2), and explanatory power of shooting was weak when assessed by model contrasts as above (LR $\chi^2_{1} = 0.275, P = 0.600$).

We found support for our expectation that differences in rates of connectivity due to shooting and ferret occupancy would be more detectable for burrow openings with crater-shaped mounds and those without mounds.
than for openings with dome-shaped mounds. On colonies with assumed prairie dog mortality from shooting or ferrets, the rate of connectivity for burrow openings with crater-shaped mounds or no mounds was 0.051 (95% CI, 0.019–0.127), compared to 0.179 (95% CI, 0.092–0.317) for such openings on control colonies (LR $\chi^2_{1} = 5.069$, $P = 0.024$). In contrast, there was little difference (LR $\chi^2_{1} = 0.025$, $P = 0.874$) between these categories of colonies regarding rates of connectivity for openings with dome-shaped mounds (0.214, 95% CI 0.115–0.363; 0.231, 95% CI 0.108–0.428).

Logistic regression analyses of line-survey data were consistent with plot data with respect to surface plugging of burrow openings (Fig. 3). In the assessment of ferret effects, the most supported model contained only the ferret variable (Table 3), and all competitive models also had the ferret variable. Ferret effect was highly influential, as suggested by comparison of the model with ferrets and mound types to the model with only mound types (LR $\chi^2_{1} = 37.034$, $P < 0.001$). Thus, the lower-ranking competitive models relate to questions about additional effects of mound types (main effect and interaction with

### Table 2. Evaluation of frequencies of subsurface connections and surface plugging rates at burrow openings on colonies with and without black-footed ferrets (BFF) or shooting (SHOOT). Total numbers of nonplugged burrow openings (TB) was used as a control variable. Data were derived from 0.073-ha plots.

<table>
<thead>
<tr>
<th>Model</th>
<th>Log-likelihood</th>
<th>AIC</th>
<th>$\Delta$AIC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subsurface connection, ferret</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant + BFF + TB</td>
<td>-51.323</td>
<td>108.646</td>
<td>0.000</td>
</tr>
<tr>
<td>Constant + BFF + TB + (BFF × TB)</td>
<td>-50.532</td>
<td>109.064</td>
<td>0.418</td>
</tr>
<tr>
<td>Constant + TB</td>
<td>-55.230</td>
<td>114.459</td>
<td>5.814</td>
</tr>
<tr>
<td>Subsurface connection, shooting</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant + TB</td>
<td>-55.751</td>
<td>115.503</td>
<td>0.000</td>
</tr>
<tr>
<td>Constant + SHOOT + TB</td>
<td>-55.419</td>
<td>116.839</td>
<td>1.336</td>
</tr>
<tr>
<td>Constant + SHOOT + TB + (SHOOT × TB)</td>
<td>-55.419</td>
<td>118.839</td>
<td>3.336</td>
</tr>
<tr>
<td>Surface plug, ferret</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant + BFF + TB</td>
<td>-98.917</td>
<td>203.835</td>
<td>0.000</td>
</tr>
<tr>
<td>Constant + BFF + TB + (BFF × TB)</td>
<td>-98.899</td>
<td>205.799</td>
<td>1.964</td>
</tr>
<tr>
<td>Constant + TB</td>
<td>-107.381</td>
<td>218.763</td>
<td>14.928</td>
</tr>
<tr>
<td>Surface plug, shooting</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant + TB</td>
<td>-67.058</td>
<td>138.115</td>
<td>0.000</td>
</tr>
<tr>
<td>Constant + SHOOT + TB + (SHOOT × TB)</td>
<td>-65.840</td>
<td>139.680</td>
<td>1.565</td>
</tr>
<tr>
<td>Constant + SHOOT + TB</td>
<td>-66.920</td>
<td>139.840</td>
<td>1.725</td>
</tr>
</tbody>
</table>
Because the model with the interaction was somewhat competitive ($\Delta AIC = 1.076$), we generated estimates (Fig. 3) from that model. The interaction was likely produced by the disproportionately larger difference in plugging between colonies with and without ferrets for openings with dome-shaped mounds as compared to nonmounded openings (Fig. 3). Line-survey data showed somewhat equivocal effects of shooting on surface plugging. The most supported model retained mound effect and shooting effect, and a contrast of nested models as above supported an influence of shooting on the rate of surface plugging ($LR \chi^2_{1} = 3.912, P = 0.048$). A model that was closely competitive ($\Delta AIC = 0.074$), however, had only the constant (Table 3).

Ratios of various types of burrow openings varied among categories of colonies. On colonies without ferrets or shooting, nonmounded openings were most common (Fig. 4). Proportions of each mound type were much different from control colonies on colonies with ferrets but no shooting (Fig. 4; $\chi^2_{1} = 23.863, P < 0.001$) and on colonies with shooting but no ferrets (Fig. 4; $\chi^2_{1} = 77.583, P < 0.001$). Dome-shaped mounds were prevalent on colonies with ferrets or shooting. Because crater mounds were relatively uncommon, differences in their ratios contributed less to overall variation than differences in ratios of other types of openings (e.g., craters produced 3.4% of the $\chi^2$ value in the evaluation of ferret effects and 15.9% in the evaluation of shooting effects).

Table 3. Evaluation of surface plugging rates at burrow openings with dome-shaped mounds and nonmounded openings (MOUND) on colonies with and without shooting (SHOOT) or black-footed ferrets (BFF). Data were from line surveys.

<table>
<thead>
<tr>
<th>Model</th>
<th>Log-likelihood</th>
<th>AIC</th>
<th>$\Delta$AIC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Black-footed ferret</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant + BFF</td>
<td>-200.519</td>
<td>405.038</td>
<td>0.000</td>
</tr>
<tr>
<td>Constant + BFF + MOUND</td>
<td>-200.044</td>
<td>406.088</td>
<td>1.051</td>
</tr>
<tr>
<td>Constant + BFF + MOUND + (BFF $\times$ MOUND)</td>
<td>-199.057</td>
<td>406.113</td>
<td>1.076</td>
</tr>
<tr>
<td>Constant + MOUND</td>
<td>-218.561</td>
<td>441.122</td>
<td>36.084</td>
</tr>
<tr>
<td>Constant only</td>
<td>-220.759</td>
<td>443.519</td>
<td>38.481</td>
</tr>
<tr>
<td>Shooting</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constant + SHOOT + MOUND</td>
<td>-52.592</td>
<td>111.183</td>
<td>0.000</td>
</tr>
<tr>
<td>Constant only</td>
<td>-54.629</td>
<td>111.258</td>
<td>0.075</td>
</tr>
<tr>
<td>Constant + SHOOT</td>
<td>-53.558</td>
<td>111.258</td>
<td>0.075</td>
</tr>
<tr>
<td>Constant + MOUND</td>
<td>-54.548</td>
<td>113.095</td>
<td>1.912</td>
</tr>
<tr>
<td>Constant + SHOOT + MOUND + (SHOOT $\times$ MOUND)</td>
<td>-52.384</td>
<td>113.167</td>
<td>1.984</td>
</tr>
</tbody>
</table>
Of the 48 openings found to be connected to the focal opening (the one with the blower), only 35.4% of connections were the dome-to-crater and dome-to-nonmounded openings that would be expected if prairie dogs construct burrow systems in the manner described. Dome-to-dome connections were unexpected but comprised 50.0% of the connections. The crater-to-crater and crater-to-nonmounded openings (14.6%) might have been associated with complex multi-opening systems with several terminus openings (craters and non-mounded) that are therefore connected to each other and also to the origin at a dome.

**DISCUSSION**

Our study design was constrained by a tendency for clustering in geographic distribution of the treatments. For example, ferrets inhabited relatively large colonies, and many of these colonies were in the southwest portion of the study area (Fig. 1). In addition, shooting was allowed only in areas away from Badlands National Park and where ferrets had not been released. Other factors might have influenced the results, including densities of prairie dogs, soil types, and vegetation composition. However, our replication within treatments should have helped to reduce effects of these variables because we expected variation in these factors within treatments. Also, in an attempt to account for prairie dog density, we used total burrow density as a control covariate in models that evaluated data from plots. Thus, although we could not control or measure some variables, the likely increase in statistical variation due to those factors did not seem to prevent detection of at least some relationships, and the potential for confounding interpretations did not seem high.

Within our study area, prairie dog colonies occupied by ferrets had burrow attributes that differed from colonies where ferrets were not recently detected. Overall, densities of burrow openings on ferret-occupied colonies were lower than those on control colonies. Although ferrets likely affect prairie dog population characteristics, the degree of impact has been debated (Henderson et al. 1969, Hillman and Linder 1973). Under the conditions of our study, our data suggest a 30% reduction in density of prairie dogs due to ferret occupancy if density of burrow openings is linearly correlated with density of prairie dogs (Biggins et al. 1993, 2006c).

Some effects of ferret presence are less direct than actual predation. Our data augment observations that occupancy of prairie dog colonies by black-footed ferrets causes increased surface plugging of burrow systems by prairie dogs (Smith 1958, Henderson et al. 1969, Sheets et al. 1971). Excavations of burrow systems (Sheets 1970) and rates of connectivity of
Burrow openings suggest prairie dogs create plugs belowground also. The effects of ferrets on plugging rates are not trivial. Surface plugging rates increased by a factor of 15.4 for dome openings and by a factor of 4.2 for crater and nonmounded openings, as suggested by our comparison of colonies with and without ferrets (Fig. 3). Considering remaining burrow openings that were not surface-plugged, subsurface connections had relative frequencies 4.1 times higher on colonies without ferrets than on colonies with ferrets (Fig. 2).

Prairie dogs tend to graze near burrow openings, perhaps to reduce risk of predation from animals that attack them aboveground (e.g., birds of prey). Spatially restricted foraging (Van Horne 2007) depletes vegetation surrounding burrow openings (Koford 1958). A reduction in the density of burrow openings due to surface plugging by prairie dogs would appear to influence the trade-off between quick access to safety and access to high quality and quantity of forage. With fewer available openings for escape, prairie dogs must choose whether to range farther from safety or consume less vegetation (or vegetation of lower quality). Thus, where surface-plugging rates are high due to presence of ferrets, prairie dogs might be presented with more difficult choices regarding nutrition and safety from other predators. Nutrition has been linked to reproductive performance (King et al. 1991, Hoogland 1995), to successful energy conservation via torpor (Lehmer et al. 2006), and to infanticide (Hoogland 1995, Ebensperger and Blumstein 2007). If nutrition is sacrificed to reduce risk of predation, additional indirect consequences of ferret presence on a colony might include reduced reproduction by prairie dogs and increased mortality of prairie dogs from infanticide and during torpor. The possible combinations of reduced nutritional input, increased energy expenditures, and increased rates of predation could substantially alter population growth rates of prairie dogs when ferrets are present. If so, the estimate of a 30% reduction in density of prairie dogs due to ferret occupancy might reflect not only the direct cost of predation by ferrets on prairie dogs, but also behaviorally mediated, indirect interactions (review in Creel and Christianson 2005) that elevate the costs of ferret presence.

Burrow plugging by prairie dogs might create other intriguing scenarios. By plugging burrows and reducing escape routes, prairie dogs might be trading decreased risk to ferrets for increased risk to more generalized predators, but they also might be increasing the risk of their specialized predator, the ferret, to badgers that prey on both prairie dogs and ferrets. Ferrets tend to use multi-opening burrow systems, perhaps as a defense against badgers (Biggins 2012). Although ferrets seem quite capable of quickly removing shallow surface plugs (Henderson et al. 1969), subsurface plugs can be long (Sheets 1970) and seem to reduce the availability of multiopening systems that ferrets prefer. Ferrets likely remove long plugs at times as suggested by removal of >20 kg of soil during a single excavation event (Biggins et al. 2012) and removal of 45 kg of soil during a single night (Clark 1989). Nevertheless, ferrets do not seem to construct their own burrow systems, so we might assume that they are less efficient excavators than prairie dogs are. If so, plugs created by prairie dogs might be more quickly removed by prairie dogs than by ferrets, creating the possibility of differential risk for each species to excavation by badgers. If a prairie dog can reopen an alternative opening by plug removal at a rate faster than a pursuing badger can enlarge a burrow through the undisturbed soils surrounding the tunnel, an escape might be available for the prairie dog. The same might be true for a prairie dog being pursued by a digging ferret. The plausibility of this digging speed hypothesis as a defense is supported by observations on use of a similar tactic by kangaroo rats (Dipodomys spp.). These rodents appear to avoid threats within burrows by digging quickly through the soil remaining at the ends of tunnels that they have constructed with a terminus near the surface (White 2009).

We cannot be certain that the comparatively low rates of opening connectivity that we detected on prairie dog colonies occupied by ferrets are due to subsurface plugging by prairie dogs. That explanation is likely, however, because others have reported surface plugging in response to ferrets (Hillman 1968, Martin et al. 1984, Eads and Biggins 2012). We have observed captive and free-ranging prairie dogs in the act of plugging burrow openings when the associated burrow systems were known to be occupied by ferrets, and excavated burrow systems formerly occupied by ferrets often contained long subsurface plugs (Sheets 1970). Subsurface plugging by prairie dogs would seem to be a logical extension of
surface plugging. It is unlikely that ferrets would do the plugging, because ferrets seem to prefer burrow systems with multiple openings (i.e., multiple routes of escape; Biggins 2012). Stromberg (1978) suggested that ferrets might prefer complex burrow systems, based on the excavations by Sheets et al. (1971) of ferret-occupied systems that had multiple openings in 15 of 18 cases. In that example, however, 10 of the 15 multi-opening systems were plugged with soil that likely would have prevented air circulation, and 2 of the remaining 3 were incompletely excavated (Sheets 1970). Thus, 28% of the burrow systems (i.e., 5 of 18) examined by Sheets (1970) might have had connected openings not detectable with our method or that of Stromberg (1978). Our testing strategy resulted in detection of 22% of burrow openings with at least one connection on colonies without ferrets (Fig. 1), much higher than the 8% rate that led Stromberg (1978:175) to conclude that “complex burrow systems were rare.”

Perhaps our estimates of changes induced by presence of ferrets are underestimated. First, relatively recent occupancy by ferrets in the Roberts Colony might have allowed sufficient time for only small effects on the prairie dog population, and changes in burrow configurations likely have even greater lag times. Ferrets were released in 2004 on the Roberts Colony in Badlands National Park, and that colony likely did not attain ferret densities similar to other ferret-occupied colonies until 2007. Nevertheless, there were relatively stable ferret densities maintained for >8 years on other colonies we sampled. Second, we classified colonies (Table 1) with respect to their status in 2009, but ferrets had been present at least once previously on 2 of the small colonies where they were not known to be present in 2009. Burrow attributes on those colonies might to some degree reflect transitory occupancy by ferrets. Third, lack of detection does not assure that there were no ferrets present on colonies.

Effects of ferret presence on plugging by prairie dogs could have implications for species other than prairie dogs, ferrets, or badgers. For instance, air circulation becomes more restricted in plugged burrows. How might this affect burrow climate and how might those changes affect other associates of prairie dog burrows? Perhaps plugging causes average temperatures within burrows to be warmer in winter and cooler in summer, and it may cause humidity to be higher. Humidity and temperature are known to affect flea ecology (Krasnov 2008) and the flea-borne bacterium (Yersinia pestis) that causes plague (Gage and Kosoy 2005), a disease to which prairie dogs and ferrets are highly susceptible (Biggins et al. 2010, Matchett et al. 2010). Also, perhaps there are effects of these microclimatic influences on Y. pestis in soil, one possible maintenance environment for the bacteria (Baltazard 1964). The tendency for prairie dogs to plug burrows containing dead conspecifics also might lead to a positive feedback cycle that increases rates of plague transmission during an epizootic, wherein the process of plugging exposes prairie dogs to infected fleas that have left the underground carcasses of their counterparts and congregated near burrow openings.

Effects of shooting were less easily interpreted than effects of ferret presence. Overall burrow densities were lower on colonies where shooting was allowed than on colonies without shooting, and the ratios of the several types of mounds were clearly different. Although the line-survey data gave some support for more surface plugging on colonies with shooting than on those without shooting, the plot data did not provide additional evidence. Nevertheless, these data collectively are consistent with the hypotheses that (1) shooting reduces average densities of prairie dogs (Reeve and Vosburgh 2006), leading to reduced densities of burrow openings overall; (2) some prairie dogs that are shot die in burrows, rather than aboveground; (3) survivors tend to surface-plug the burrows containing carcasses; and (4) increased numbers of plugged burrows lead to new burrow construction, creating a higher ratio of dome-shaped mounds. Effects of shooting on burrow plugging might be less dramatic than effects of ferrets simply because shooting instantly kills many prairie dogs aboveground and entombment by survivors is not possible, whereas ferrets tend to consume prairie dogs belowground and presumably leave portions of carcasses in burrows for varying periods.

Openings with crater-shaped mounds or no mounds are thought to be terminus openings usually associated, at least initially, with dome-shaped mounds where the burrow construction originated. Lack of connectivity at these
 terminus openings should be a better indica-
tor of subsurface plugging than lack of connect-
itivity at dome-shaped mounds because the lat-
ter could also be indicative of single-opening
burrows that are being constructed or that
were discontinued before any connections
were created. The different rates of connectiv-
ity for openings without dome-shaped mounds,
compared to lack of variation in such rates for
openings with domes, thus provide support for
the hypothesis that at least some of the varia-
tion in connectivity of burrow openings
on colonies with and without ferrets and
shooting is due to subsurface plugging rather
than construction of new systems that do not
yet have multiple openings.

The preponderance of dome-to-dome con-
nections we detected was unexpected given
the hypothesized sequence of burrow con-
struction (Sheets 1970). We presume that
prairie dogs would find it impossible to exca-
vate soil vertically through the nonmounded
terminus burrows exemplified in the maps of
Sheets (1970). There are several possible expla-
nations for dome-to-dome connections. Per-
haps prairie dogs dig some moderately sloped
(rather than vertical) exit tunnels, which are
later modified into domes by the delivery of
subsoil from additional tunnel extensions to
the surface at those formerly nonmounded
openings. Perhaps erosion degrades dome-
shaped mounds that are later modified into
labeled-shaped mounds, or crater-shaped mounds
might erode into domes if not continuously
maintained as craters. We have also watched
badgers destroy crater-shaped mounds. Per-
haps prairie dogs eventually convert craters or
nonmounded openings to domes because high
domes serve as vantage points in close prox-
imity to escape cover. Finally, prairie dogs
might intersect existing tunnels as they dig
new burrows, either by chance or by some
method of detecting the existing tunnels. For
instance, if prairie dogs follow certain soil
strata when digging (King 1984), the chance of
intersecting another tunnel might increase.
Also, prairie dogs might detect existing tun-
els via sound or odor. Indeed, such detec-
tions seem possible in prairie dog habitats, as
exemplified in a study involving Siberian pole-
cats (Mustela eversmanii) used as research sur-
rrogates for ferrets (Biggins et al. 2011); an
American badger seemed to locate a prairie
dog burrow with a polecat in it and began dig-
ging directly toward a radio-tagged polecat
that was below it rather than excavating at the
opening of the burrow, which was several meters
distant (DEB unpublished observation). Some
burrowing mammals may be able to perceive
seismic cues (Reichman and Smith 1990).

Burrow plugging and excavation in the
prairie dog ecosystem is of theoretical interest,
providing opportunities to investigate several
modes of direct and indirect interactions
among multiple mesopredators and their prey.
Within the geographic area of this study, sev-
eral findings seem noteworthy, collectively
suggesting ferrets have impacts on the Bad-
lands–Conata Basin complexes of prairie dog
colonies that go far beyond the direct killing of
prairie dogs as prey. Nevertheless, general
conclusions in the context of burrow relation-
ships among prairie dogs, ferrets, and other
predators await replication of these kinds of
studies and others at sites over the broad
range of the black-tailed prairie dog and at
sites with other prairie dog species.

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