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DIRECT AND INDIRECT EFFECTS OF HERBIVORES ON NITROGEN DYNAMICS: VOLES IN RIPARIAN AREAS

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Abstract. Herbivores can directly increase nitrogen mobility by increasing the quality of organic matter entering the decomposition cycle, but they also may decrease nitrogen mobility by decreasing the biomass of high-nitrogen species in the plant community. We assessed effects of voles (Microtus) on nitrogen dynamics using exclosures in two riparian meadows (Crystal Bench and Blacktail Deer Creek) in Yellowstone National Park (USA). At both sites, the quantity of plant litter was decreased by herbivory following a vole population peak in 1992. At Crystal Bench, removal of voles caused a decrease in the nitrogen concentration and an increase in the C:N ratio of plant litter over the four years of the study. The higher quality litter produced in the presence of voles at Crystal resulted in a larger pool of potentially mineralizable nitrogen in soil from control plots relative to soils from plots that had not been accessible to voles. At Crystal, vole removal did not cause a change in plant community composition. However, at Blacktail, after several years of vole exclusion, legumes became more common in exclosures than in control plots that were accessible to voles. Selective herbivory on high-nitrogen legumes kept the litter quality outside exclosures low, whereas higher legume biomass caused a decrease in C:N ratio of plant litter inside exclosures. The removal of voles at Blacktail caused a 15% increase in the fraction of the soil nitrogen that was rapidly mineralizable.

Our results show that voles increased nitrogen mobility, especially during and after population peaks. However, that increase was offset by decreases in nitrogen mineralization over longer periods when voles caused a decrease in high-quality plant litter produced by preferred forage plants, especially legumes. Thus, both the mechanisms by which voles affected nitrogen dynamics and the net effects of voles varied over time and space. The balance of direct and indirect effects may provide a general mechanistic explanation of whether herbivores increase or decrease the rate of nitrogen cycling.

Key words: herbivory; legumes; Microtus; mineralization; nitrogen cycling; nitrogen, potentially mineralizable; plant litter; riparian; soil organic matter; Trifolium; voles; Yellowstone National Park.

INTRODUCTION

Terrestrial herbivores often increase (McKendrick et al. 1980, Swank et al. 1981, Ruess and McNaughton 1987, Holland and Detling 1990, Shariff et al. 1994, Frank and Evans 1997, McNaughton et al. 1997), but sometimes decrease (Pastor et al. 1988, 1993, Ritchie et al. 1998), the rate of nitrogen cycling. The factors that determine whether herbivory has a net positive or negative effect on nitrogen cycling are not clear and are likely to vary in time and space. Studies that address the mechanisms by which herbivores affect nitrogen dynamics are needed to understand the net effects of herbivores on spatial and temporal patterns of nitrogen cycling.

Herbivores affect the rate of terrestrial nitrogen turnover largely by influencing the amount and quality of organic matter in the soil and on the soil surface (e.g., Ruess and McNaughton 1987, McNaughton et al. 1988, Holland and Detling 1990, Pastor et al. 1993). The size of the active pool of nitrogen in soil organic matter is an important control on nitrogen availability (Fig. 1). In general, the direct effects of herbivores on plants and on the physical environment, such as removing standing vegetation, disturbing soil, and depositing high-nitrogen waste products, accelerate nitrogen cycling by increasing the quality of the soil organic matter (Ruess and McNaughton 1987, Frank and Groffman 1998). Conversely, selective foraging by herbivores alters plant community composition, which indirectly decreases nitrogen cycling. Because herbivores selectively consume more palatable plants, increases in less palatable plant species change overall plant litter quality and ultimately reduce the quality of the soil organic matter; thus, net nitrogen mineralization is decreased (Pastor et al. 1988, 1993, McInnes et al. 1992). The direct effect, increased nitrogen cycling rate in the presence of herbivores, probably occurs in all ecosystems, but this effect may be overshadowed by the indirect decelerating effect, caused by changes in plant community composition, that often results from selective
herbivory (Fig. 1). The relative importance of direct and indirect effects should vary with spatial and temporal patterns of herbivore abundance.


Nevertheless, herbivores sometimes decelerate nitrogen cycling. This appears to occur when herbivores cause changes in plant community composition. Selective foraging on high-quality plants, e.g., plants with a relatively low C:N ratio or with low concentrations of recalcitrant chemicals, can decrease the abundance of these plants relative to plants of lower quality (McInnes et al. 1992, Ritchie et al. 1998). The relatively low-quality plant litter mineralizes nitrogen at a slower rate than would occur in the absence of herbivory (Pastor et al. 1988, 1993).

The objective of this study was to determine the effects of voles (Microtus, Rodentia; Microtinae) on nitrogen dynamics in riparian meadows in Yellowstone Park. Although much research has been done on the effects of ungulate herbivores, less is known about the effects of smaller animals on ecosystem processes. Voles are strict herbivores with high reproductive rates; their populations can fluctuate greatly both between and within years (Taitt and Krebs 1985). When populations are high, voles can consume much of the aboveground plant biomass (Summerhayes 1941, Batzli and Pitelka 1971, Moen et al. 1993, Virtanen et al. 1997). Thus, the effects of voles on nitrogen dynamics can be pulsed, or discontinuous, in time. Occasional high population densities may directly cause large pulses in nitrogen mobility. Conversely, chronic herbivory may indirectly lead to decreases in nitrogen mobility by causing changes in plant species composition.

We used fenced exclosures to assess the effects of voles on nitrogen dynamics. We hypothesized that nitrogen cycling would be slower in the absence of voles,
because studies in Yellowstone Park and elsewhere have suggested that herbivores do not affect plant species composition dramatically in heavily grazed ecosystems (Milchunas et al. 1989, Mack and Thompson 1982). Thus, we hypothesized that voles primarily affected nitrogen cycling by directly stimulating nitrogen flux pathways, for instance, by increasing quality of litter and depositing urine and feces. The incorporation of higher quality organic matter in the form of urine, feces, and low C:N plant litter should increase the quality of the soil organic matter in control plots, which are used by voles, relative to exclosures. Therefore, we expected that laboratory incubations would reveal that soil from control plots has a larger pool of rapidly mineralizable nitrogen than soil from vole exclosures.

**Methods**

**Study areas**

We studied the effects of voles on riparian meadows in northern Yellowstone Park (110°42’ W, 44°59’ N), Wyoming, United States. The two study sites were chosen to represent physiographically different types of riparian landscape (Table 1). Blacktail Deer Creek is a third-order montane stream with a narrow, well-defined riparian area bounded by sagebrush and open forest uplands. The soil is heterogeneous, consisting of layers of old sandbars, muddy deposits, and cobble beds. Crystal Bench is a wet meadow below a series of small seeps feeding an intermittent stream that is usually dry by August. The soil at Crystal Bench is deep, rich, and of a mostly uniform loamy texture. The elevation of both sites is between 1600 and 1800 m. Both sites were relatively free of woody vegetation and had evidence of voles. Pocket gophers (*Thomomys talpoides*) were present during the four years of the experiment, but few gopher mounds appeared on study plots during the experiment.

**Estimation of small-mammal population density**

Small-mammal populations were censused in the late summer of each year by live-trapping. At each enclosure site, we placed two live-trap mini-grids adjacent to the study plots. Each mini-grid consisted of three rows of three Sherman traps spaced 15 m apart and three rows of three coffee-can pitfall traps spaced 15 m apart, with the Sherman grid being offset 7.5 m diagonally to the pitfall grid. Sherman traps were baited with oats and peanut butter; pitfalls were not baited. Traps were checked every morning and evening for 4 d, and captured animals were marked and released. To estimate small-mammal density, the total number of animals captured over the 4-d period was entered into an equation that had previously been calibrated by regressing mini-grid captures on mark–recapture population estimates from large grids at several riparian sites in northern Yellowstone Park (R. Crabtree, unpublished data). The relative trap spacing in these large grids was the same as in the mini-grids, but the large grids consisted of 10 rows of 10 traps each. A separate regression equation was created for each common small-mammal species in the area. The resulting regression equation for *Microtus* had $R^2 = 0.85$. The mean of the regression estimates from the two mini-grids at each site was used as the density estimate of the site.

**Experimental design**

In the spring of 1992, we built eight vole exclosures at each of the two riparian meadow sites in the northern range of Yellowstone National Park. The exclosures were $5 \times 2.5$ m rectangles constructed of 0.6-cm mesh welded wire fence, extending from 15 cm below ground to a height of 45 cm. We added 15 cm of aluminum flashing to the top of the welded wire to prevent small mammals from climbing over the fence. Each exclosure was paired with an unfenced (control) plot of the same size. Small mammals initially present in the exclosures were trapped and removed within the first month of the study.

**Vegetation sampling**

Aboveground plant biomass was measured in each plot during the period of peak biomass in late July or early August. In each plot, two strips of vegetation (0.1 $\times$ 0.5 m) were clipped, sorted to species, dried, and weighed. Similarly, total standing dead biomass was measured in October of each year (except 1993). The mean biomass of the two strips was used in statistical procedures. The carbon and nitrogen concentration of

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Blacktail</th>
<th>Crystal</th>
</tr>
</thead>
<tbody>
<tr>
<td>1992 peak aboveground biomass (g/m²)</td>
<td>530</td>
<td>477</td>
</tr>
<tr>
<td>Dominant graminoids (relative biomass, %)</td>
<td><em>Carex</em> spp. (35)</td>
<td><em>Phleum pratense</em> (38)</td>
</tr>
<tr>
<td></td>
<td><em>Calamagrostis canadensis</em> (21)</td>
<td><em>Solidago canadensis</em> (13)</td>
</tr>
<tr>
<td></td>
<td><em>Phleum pratense</em> (9)</td>
<td><em>Geranium richardsonii</em> (11)</td>
</tr>
<tr>
<td></td>
<td><em>Carex</em> spp. (6)</td>
<td><em>Epilobium angustifolium</em> (9)</td>
</tr>
<tr>
<td>Dominant forbs (relative biomass, %)</td>
<td><em>Aster sibiricus</em> (7)</td>
<td><em>Arnica chamissonis</em> (7)</td>
</tr>
<tr>
<td></td>
<td><em>Geranium richardsonii</em> (11)</td>
<td><em>Epilobium angustifolium</em> (9)</td>
</tr>
<tr>
<td>Soil nitrogen concentration (% by mass)</td>
<td>0.29</td>
<td>0.70</td>
</tr>
<tr>
<td>Soil carbon:nitrogen (mass:mass)</td>
<td>17.6</td>
<td>13.2</td>
</tr>
</tbody>
</table>

TABLE 1. Vegetation and soil fertility characteristics of the two study sites in northern Yellowstone Park, Wyoming, USA.
plant litter was measured with a Carlo Erba CN1500 combustion C-H-N analyzer (CE Instruments, Milan, Italy).

**Soil incubations in the laboratory**

Potentially mineralizable nitrogen is the fraction of the organic nitrogen that is mineralized rapidly, or over an ecologically relevant time scale. The size of this pool is a function of the type of organic molecules in the soil; thus, it reflects the quality of the soil organic matter. The amount of potentially mineralizable nitrogen \( N_0 \) in soil can be roughly measured as the cumulative amount mineralized over a year-long laboratory incubation under conditions favoring optimal microbial activity (Stanford and Smith 1972). However, this cumulative total includes small amounts of nitrogen mineralized from more recalcitrant (i.e., relatively inactive) organic pools, and thus is not an accurate measure of rapidly mineralizable nitrogen. To more accurately assess \( N_0 \) in each soil sample, we used three nonlinear regression models (Deans et al. 1986). The three models hypothesize (1) a single pool of organic nitrogen mineralizing at a nonlinear, decreasing rate, (2) a fast pool and a slow pool, both mineralizing at a nonlinear, decreasing rate, and (3) a fast pool mineralizing at a nonlinear, decreasing rate and a slow pool mineralizing at a constant rate:

\[
N_i = N_0 (1 - e^{-kt}) 
\]

\[
N_i = N_0 S (1 - e^{-h t}) + N_0 (1 - S)(1 - e^{-kt}) 
\]

\[
N_i = N_0 (1 - e^{-kt}) + C t 
\]

where \( N_i \) is the cumulative nitrogen mineralized at time \( t \), \( N_0 \) is the hypothetical pool size of potentially mineralizable nitrogen, and \( t \) is time in weeks. In model 2, \( S \) and \( 1 - S \) represent the size of the fast and slow mineralizing subpools, respectively, and \( h \) and \( k \) represent the rate constants for these pools. In model 3, \( C \) is the size of the slow, constant pool.

Soil from two control plots at Crystal did not fit any of the three nonlinear models. In both cases and for all three models, either the iterations failed to converge on a least squares equation or the estimated parameters that produced a least squares equation were unreasonable. The two soils had very low total carbon and nitrogen and only a negligible amount of potentially mineralizable nitrogen. Because the behavior of these two soils was so different from all others in the study, we excluded them from statistical analysis.

Potential nitrogen mineralization was measured in the laboratory by incubating soil samples in microlysimeters under controlled conditions (Stanford and Smith 1972, Nadelhoffer 1990). Samples from the top 12 cm of the soil profile were collected in October 1995 from each control and enclosure plot, dried, and sieved. We put 30-g (dry mass) subsamples of each soil into separate microlysimeters and incubated them in a dark, 30°C environmental chamber for 40 wk.

During the initial weeks of the incubation, we extracted inorganic nitrogen from soil samples weekly by leaching each lysimeter with 100 mL of nutrient extractant solution (Nadelhoffer 1990). After 8 wk, we leached the lysimeters every 2 wk with 80 mL of nutrient extractant solution. As the nitrogen yield slowed, we leached at less frequent intervals until the last leaching interval of 13 wk. To ensure equal soil water tension, each lysimeter was suctioned to a tension of \(-0.05 \text{ MPa}\) for several minutes after the leachate was added. Ammonium concentration in leachates was measured colorimetrically at 630 nm by the indophenol blue method. Nitrate was reduced to nitrite with cadmium and was measured colorimetrically at 540 nm using a diazotized sulfanilamide EDTA method (Greenburg et al. 1992). All colorimetric analyses were done with an Alpkem RFA 300 autoanalyzer (OI Analytical, College Station, Texas, USA).

**Hypothesis testing**

Variables that were measured each year were analyzed with a two-factor (treatment and time), blocked (treatments spatially paired) repeated-measures design. Because we were interested in vole removal effects in individual years as well as in time-by-vole effects, we used Tukey’s hsd test within the repeated-measure design to assess enclosure effects in individual years. Several control–enclosure pairs were not sampled at Crystal Bench in October 1992. Consequently, repeated-measures analysis on these data could only include six replicates, and the repeated-measure analysis lacked power. These data were analyzed with blocked, one-way ANOVA on each year’s data separately; the resulting probabilities were evaluated against a Bonferroni-adjusted alpha value of 0.05. Variables that were not measured repeatedly were analyzed with blocked, one-way ANOVA. All statistical calculations were done with SYSTAT v. 5.1 (SPSS, Chicago, Illinois, USA).

**Results**

**Small-mammal density**

Three small-mammal species represented 95% of all individuals captured. Deer mice (*Peromyscus maniculatus*) and jumping mice (*Zapus princeps*) were frequently found at both Blacktail and Crystal. Total mouse densities remained fairly stable (5–15 individuals/ha) at both sites throughout the study. Except for one long-tailed vole (*Microtus longicaudus*) and one water vole (*Microtus richardsoni*), all microtine rodents caught were montane voles (*Microtus montanus*). Vole densities were high (>50 individuals/ha) at both sites in 1992 (Fig. 2). Vole populations remained high at Crystal Bench in 1993, whereas the Blacktail population declined. Only the Blacktail site had a measurable vole population in 1994 and 1995.
Effects of vole herbivory on plant litter and community composition

Early in the study, when vole densities were high, standing dead litter biomass was greater inside exclosures than in plots accessible to voles at Crystal, and was marginally greater at Blacktail (Fig. 3). Across all years, litter biomass was greater in exclosures than in the presence of voles at Blacktail (Fig. 3, Table 2). Early in the study, plant community composition did not differ between vole exclosures and control plots. By 1995, nitrogen-fixing legumes (Trifolium spp.) were more abundant in plots from which voles were excluded than in control plots at Blacktail, where the combined biomass of Trifolium hybridum and T. longipes in vole exclosures increased by over an order of magnitude, from <1% of the total aboveground biomass in 1992 to 13% in 1995 (Fig. 4). Trifolium biomass was greater in vole exclosures than in control plots at Crystal in 1994, but was small compared to Blacktail.

Initially, the concentration of nitrogen in plant litter did not differ between exclosures and control plots at Crystal. By 1995, however, nitrogen concentration was marginally greater and C:N was significantly lower in the presence of voles than in plots without voles (Fig. 3). These results were consistent with our predictions that vole herbivory would decrease the C:N ratio of plant litter. However, at Blacktail, repeated-measures ANOVA indicated that the pattern of change in litter nitrogen concentration over time differed between control and exclosure plots (Fig. 3, Table 2). The removal of voles at Blacktail caused an increase in the nitrogen concentration of plant litter, but only after several years, as the relative abundance of legumes increased in the absence of vole herbivory.

Laboratory measurement of potentially mineralizable nitrogen

Approximately twice as much nitrogen was mineralized over the 40-wk incubation from Crystal Bench soil as was mineralized from Blacktail soil (Fig. 5). Despite the large variation between sites in the absolute amount of nitrogen mineralized, most lysimeters mineralized between 4% and 7% of the total nitrogen in the soil. The pattern of cumulative yield of mineral nitrogen over time was best fit with the mixed exponential–linear model (Eq. 3), explaining >99.6% of the variation in both absolute and proportional nitrogen mineralization. We used Eq. 3 to estimate \( N_0 \) for all lysimeters. Consistent with our hypotheses, removal of voles caused a decrease in the rapidly mineralizable pool at Crystal. For Crystal Bench soils, this pool \( (N_0 \) measured in mg N/kg soil) was 27% larger in plots accessible to voles (238 mg N/kg soil) than in exclosures (187.5 mg N/kg soil; Table 3). However, at Blacktail, removal of voles caused an increase in rapidly mineralizable nitrogen. Although there was no statistical difference in absolute \( N_0 \), 3.17% of the total nitrogen in control soils was in the rapidly mineralizable pool, which was 15% less, a statistically significant difference, than the 3.74% in soils from plots that had voles removed (Table 3).

Discussion

Soil organic matter is an important integrating pool linking herbivore activities and nitrogen availability (Ruess and McNaughton 1987, Pastor et al. 1988, Holland and Detling 1990, Frank et al. 1994, Frank and Evans 1997, de Mazancourt et al. 1998, Frank and Groffman 1998) (Fig. 1). The soil organic matter pool is expected to be sensitive to herbivore effects because it integrates processes such as waste deposition, changes in plant growth and nutrient allocation patterns, and the decomposition environment, which may be affected by herbivores. Laboratory incubation of soil is a powerful tool to assess the net effects of herbivores on organic matter quality, because it eliminates the environmental variability that increases sample variance and decreases the power to detect differences in field incubations.

Many studies have documented the stimulation of nitrogen cycling by herbivores (e.g., Ruess and McNaughton 1987, Frank and Groffman 1998), and some have reported a deceleration of nitrogen cycling as an indirect consequence of selective herbivory (Pastor et al. 1993, Ritchie et al. 1998), but no studies have examined both mechanisms simultaneously in a site. We found that the net effect of vole herbivory on soil organic matter quality differed between two study sites. At Crystal, voles were responsible for inputs of low C:N plant litter into the soil organic matter, keeping the pool of rapidly mineralizable nitrogen larger than it would be in the absence of voles. Fertilizer effects of vole urine and feces also probably contributed to this effect. These differences in litter quality and rapidly mineralizable nitrogen pools were evident in 1995, even though vole densities at Crystal were very low in both 1994 and 1995, possibly reflecting a time lag be-
TABLE 2. Results of two-way repeated-measures ANOVA. Plots (exclosure or control) were sampled yearly from 1992 to 1995 (excluding October litter, which was not sampled in 1993).

<table>
<thead>
<tr>
<th>Site</th>
<th>Dependent variable</th>
<th>Factor</th>
<th>$F$</th>
<th>df</th>
<th>$P$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blacktail</td>
<td>Oct. litter biomass (g/m$^2$)</td>
<td>Exclosure treatment</td>
<td>15.95</td>
<td>1, 14</td>
<td>0.001</td>
</tr>
<tr>
<td>Blacktail</td>
<td>Oct. litter N concentration (% by mass)</td>
<td>Year</td>
<td>3.35</td>
<td>2, 28</td>
<td>0.050</td>
</tr>
<tr>
<td>Blacktail</td>
<td>Oct. litter C:N (mass:mass)</td>
<td>Year</td>
<td>2.76</td>
<td>2, 28</td>
<td>0.080</td>
</tr>
<tr>
<td>Blacktail</td>
<td>Trifolium biomass (% of total)</td>
<td>Treatment</td>
<td>5.18</td>
<td>1, 14</td>
<td>0.039</td>
</tr>
</tbody>
</table>

Notes: Group means and standard errors are presented in Figs. 3 and 4. All variables and factors yielding $P$ values of 0.10 or less are included in the table; $n = 8$ per treatment per year.
tween initial increases in litter quality and the appearance of this plant litter material in the soil organic matter pool. At Blacktail, however, the direct stimulating effects of vole herbivory on nitrogen cycling were overshadowed by herbivore-mediated changes in plant community composition. Removal of voles caused an increase in high-quality (high-nitrogen) plant litter entering the soil organic matter, which increased the pool of rapidly mineralizable nitrogen in the ab-

between initial increases in litter quality and the appearance of this plant litter material in the soil organic matter pool. At Blacktail, however, the direct stimulating effects of vole herbivory on nitrogen cycling were overshadowed by herbivore-mediated changes in plant community composition. Removal of voles caused an increase in high-quality (high-nitrogen) plant litter entering the soil organic matter, which increased the pool of rapidly mineralizable nitrogen in the ab-

Fig. 4. Relative biomass (percentage of peak above-ground biomass) of *Trifolium* spp. at Blacktail and Crystal. *P* values are probabilities associated with Tukey’s *hsd* test (*Q* calculated with the error term from repeated-measures ANOVA) for treatment differences within each year’s data. Only *P* values <0.10 are reported. Error bars represent ±1 se; *n* = 8 for all years and treatments.

Fig. 5. Cumulative laboratory mineralization (NO$_3^-$ + NH$_4^+$) of soils collected in 1995 from Blacktail (BT) and Crystal (CB) controls and exclosures: top, absolute mineralization; bottom, mineralization as a percentage of total nitrogen in the soil. The *P* value is associated with the treatment effect of blocked ANOVA on cumulative nitrogen yield (Crystal) at the end of the experiment (*F* = 8.87; df = 1, 5). Means for Crystal control plots exclude two outliers; otherwise, *n* = 8; error bars represent ±1 se.

Table 3. Mean parameter estimates (with 1 se in parentheses), both in absolute terms (mg N/kg soil) and proportional terms (% of the total soil N that is in the rapidly mineralizing pool), of the exponential/linear model (Eq. 3: $N_t = N_0[1 - e^{-kt}] + C$) from nonlinear curve-fitting.

<table>
<thead>
<tr>
<th>Site</th>
<th>Dependent</th>
<th>Treatment</th>
<th>$N_0$ (1 se)</th>
<th><em>k</em> (1 se)</th>
<th><em>C</em> (1 se)</th>
<th>$R^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>BT</td>
<td>mg N/kg soil control</td>
<td>89.4 (14.6)</td>
<td>0.181 (0.009)</td>
<td>1.54 (0.23)</td>
<td>0.996</td>
<td></td>
</tr>
<tr>
<td>BT</td>
<td>mg N/kg soil exclosure</td>
<td>105.4 (8.5)</td>
<td>0.182 (0.014)</td>
<td>1.18 (0.10)</td>
<td>0.997</td>
<td></td>
</tr>
<tr>
<td>BT</td>
<td>% of total N control</td>
<td>3.17 (0.19)</td>
<td>0.181 (0.009)</td>
<td>0.060 (0.009)</td>
<td>0.997</td>
<td></td>
</tr>
<tr>
<td>BT</td>
<td>% of total N exclosure</td>
<td>3.74 (0.27)</td>
<td>0.182 (0.014)</td>
<td>0.043 (0.006)</td>
<td>0.997</td>
<td></td>
</tr>
<tr>
<td>CB</td>
<td>mg N/kg soil control</td>
<td>237.6 (29.2)</td>
<td>0.200 (0.021)</td>
<td>2.50 (0.59)</td>
<td>0.996</td>
<td></td>
</tr>
<tr>
<td>CB</td>
<td>mg N/kg soil exclosure</td>
<td>187.5 (19.1)</td>
<td>0.162 (0.028)</td>
<td>1.99 (0.34)</td>
<td>0.997</td>
<td></td>
</tr>
<tr>
<td>CB</td>
<td>% of total N control</td>
<td>3.17 (0.32)</td>
<td>0.200 (0.021)</td>
<td>0.031 (0.005)</td>
<td>0.996</td>
<td></td>
</tr>
<tr>
<td>CB</td>
<td>% of total N exclosure</td>
<td>2.80 (0.29)</td>
<td>0.162 (0.028)</td>
<td>0.029 (0.004)</td>
<td>0.997</td>
<td></td>
</tr>
</tbody>
</table>

Notes: Site abbreviations are BT = Blacktail and CB = Crystal Bench. Units of $N_0$ and *C* are the same as the dependent variable ($N_t$), with *C* being per week; *k* is a unitless nonlinear rate constant, and *t* is the incubation time (weeks). Standard errors for $R^2$ were less than 0.001 for all data sets. The control means for Crystal soils exclude two outliers; *n* = 8 for all means except Crystal controls, where *n* = 6.

† Treatment effect significant (blocked ANOVA, *F* = 8.67, df = 1, 5, *P* = 0.032).

‡ Treatment effect significant (blocked ANOVA, *F* = 5.93, df = 1, 7, *P* = 0.045).
ence of voles relative to that of soil from vole-accessible plots.

Small mammals commonly change plant community composition (Summerhayes 1941, Huntly 1987, 1991, Noy-Meir 1988, Moen et al. 1993, Virtanen et al. 1997). Selective consumption of plants with high nitrogen concentration, low C:N ratio, or low concentrations of secondary chemicals is one mechanism by which plant community composition can be changed. Many herbivores select plants with high nitrogen concentrations, especially legumes, as preferred forage (Thompson 1965, Mattson 1980, Ritchie et al. 1998), giving less preferred plants competitive advantages over species that are consumed by herbivores. The possibility that changes in plant community composition lead to changes in nutrient dynamics in these systems deserves more consideration.

Why do statistically significant differences exist only in the absolute pool size of rapidly mineralizable nitrogen at Crystal, and in the proportional pool size at Blacktail, when the directional response of absolute and proportional pools to exclosure treatment was similar within each site? At Crystal, the rapidly mineralizable pool decreased 21% and the proportion of soil nitrogen that was rapidly mineralizable decreased 12% in the absence of voles, yet only the absolute pool size differed significantly in response to vole exclosure (Table 1). At Blacktail, both means increased by ~18% when voles were removed, but only the change in the proportion of total N that was in the rapidly mineralizable pool increased significantly. At Blacktail, the variation in the proportional $N_0$ was small compared to the variation in absolute $N_0$, so there was more power to detect this difference relative to the difference in absolute $N_0$. At Crystal, the variances of these two variables were similar, but the decrease in absolute pool size in response to vole removal was nearly twice as large as the decrease in the proportion rapidly mineralizable nitrogen. This suggests that the changes in response to vole removal at Crystal were primarily the result of decreased flow of nitrogen into the soil organic matter pool, rather than a large change in organic matter quality. For example, the increased biomass of standing dead litter in exclosures, compared to unexclosed plots, represents a pool of nitrogen that is not available for decomposition and has not entered the soil organic matter pool.

The rate at which nitrogen is mineralized from the soil organic matter pool is an important rate-limiting step in the cycling of nitrogen in terrestrial systems (Whitmore and Handayanto 1997). The net effect of herbivores on nitrogen flux depends upon the relative strength of direct and indirect mechanisms that affect mineralization in soil organic matter, and the balance of these no doubt varies through time and space. The appearance of important indirect effects at Blacktail, but not at Crystal, attests that the dominant mechanisms by which herbivores control nitrogen dynamics vary spatially. In addition, we hypothesize that, in the vole-riparian system, the relative importance of direct and indirect mechanisms may vary over time, correlated with changes in vole population density. Measurements of net nitrogen mineralization in the field in 1993, immediately following the vole population peak, revealed increased nitrogen mobility in the presence of high vole populations (Fig. 6). These differences were detected despite the high variance in nitrogen mineralization caused by uncontrollable factors in the field, such as local variation in the soil organic N pool, soil moisture,
and temperature. A study at the same sites in 1995, three years after the population peak, revealed no effect of vole removal on nitrogen mineralization in the field. These data suggest that herbivores with cycling or fluctuating populations may provide spikes of increased nutrient availability during population peaks, independent of the long-term positive or negative effects of herbivores on the rate of nitrogen cycling.

We found differences in the size of the potentially mineralizable nitrogen pool despite finding no significant differences due to exclosure treatment on total soil nitrogen concentration. This emphasizes that most nitrogen mineralization is from a very small pool of active organic nitrogen. Under conditions of moisture and temperature favorable to microbial mineralization, this laboratory incubation yielded a maximum mineralization of 7% of the total organic nitrogen (Fig. 5). Under field conditions at both Blacktail and Crystall, <1% of total nitrogen was mineralized from May to October 1995. Because the proportion of the soil nitrogen that is in a rapidly cycling pool is so small, the 15% larger fast pool in soil from Blacktail exclosures, relative to control soils, is very likely an important factor in differences in mineral nitrogen availability in the presence and absence of voles.

Conclusions

We found that voles affected nitrogen dynamics both directly and indirectly. Direct mechanisms included alterations in litter quantity and quality that increased the pool of soil organic nitrogen that was readily mineralizable. However, after several years and contrary to our initial hypotheses, indirect effects of voles became important to nitrogen dynamics. The removal of chronic grazing by voles increased the biomass of preferred food species, which were the most readily decomposable, inside exclosures. When these high-quality (low C:N) plants increased inside Blacktail exclosures, the quality of soil organic matter also increased. This change resulted in increased potential for nitrogen mineralization in the absence of herbivory. There was little change in plant community composition at Crystall. Litter quality at this site continued to be more heavily influenced by direct effects of herbivory.

The balance of direct and indirect effects of herbivores on nitrogen cycling appears to be a key to understanding whether herbivores accelerate or decelerate nitrogen cycling. This balance may fluctuate through time. For instance, direct effects of voles on nitrogen cycling occurred as pulses through time, being strong when populations were high, as they were in 1992 and 1993. Indirect effects, resulting from longer term changes in plant community composition, may overshadow direct effects, as we found at the Blacktail site when vole populations were low.

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Literature Cited


