Alpine Grassland CO2 Exchange and Nitrogen Cycling: Grazing History Effects, Medicine Bow Range, Wyoming, U.S.A

Authors: Welker, Jeffrey M., Fahnestock, Jace T., Povirk, Kyra L., Bilbrough, Carol J., and Piper, Robert E.

Source: Arctic, Antarctic, and Alpine Research, 36(1) : 11-20

Published By: Institute of Arctic and Alpine Research (INSTAAR), University of Colorado

Alpine Grassland CO₂ Exchange and Nitrogen Cycling: Grazing History Effects, Medicine Bow Range, Wyoming, U.S.A.

Jeffrey M. Welker,* Jace T. Fahnestock,† Kyra L. Povirk,‡ Carol J. Bilbrough,§ and Robert E. Piper§

*Natural Resource Ecology Laboratory, Colorado State University, Fort Collins, CO 80523, U.S.A.
†Northwind Environmental Consulting, Twin Falls, ID 83405, U.S.A.
‡Division of Environmental Quality, State of Wyoming, Cheyenne, WY 82002, U.S.A.
§Rock Creek, Wyoming, U.S.A.

Abstract

Our study examined carbon dioxide exchange and nitrogen cycling over two consecutive years (winter and summer) in a grazed alpine grassland and in an embedded long-term grazing exclosure to ascertain whether grazing history had resulted in divergent soil carbon attributes, CO₂ exchange rates, and different vegetation C and N and soil N processes. Soil C and N concentrations and masses were significantly higher in the grazed than in the ungrazed area, though grass leaf N was higher in the ungrazed area, as was vegetation biomass. Detectable amounts of CO₂ were lost from the grazed and ungrazed areas of this grassland during the winters of 1998, 1999, and 2000, and at 6 of 15 winter flux sample dates, CO₂ efflux was greater in the grazed area than in the ungrazed area. The ungrazed area consistently gained more C during the summer months than the grazed area, with net CO₂ exchange peaking in mid-July 1998 at nearly 5 μmol m⁻² s⁻¹ in the ungrazed area compared to 0.2 μmol m⁻² s⁻¹ in the grazed area. During the 2-yr study period, the grazed area was a carbon source of 170 g C m⁻², while the ungrazed area was a carbon sink of 83 g C m⁻². Lower N mineralization rates early and late in the summer (1999) in the grazed area at Libby Flats corresponded to reductions in net CO₂ exchange and lower plant N content compared to the ungrazed exclosure. Based on these results, we suggest that: (1) long-term grazing in high-altitude rangelands can alter annual CO₂ exchange and N dynamics; (2) temporal synchrony in C and N processes occur during the summer; that is, increased C exchange rates accompany increased N mineralization rates; and (3) integrative (total soil C and N) and instantaneous (CO₂ exchange and vegetation N) measures of C and N dynamics may not necessarily lead to the same interpretation regarding C sequestration and N cycling in alpine grasslands.

Introduction

Alpine ecosystems are an important forage resource for livestock and wildlife, especially in the Rocky Mountain region of North America and in the Alps of Western Europe, where they are used for summer pasture (Billings, 1979; Thienhuis, 1979; Bliss, 1985; Diemer, 1992; Schappi and Körner, 1997; Povirk et al., 2000). Globally, grazing systems in the alpine life zone represent 3% of the global land area (5.7 × 10⁶ km²) and at least 1% of the global terrestrial C pool (Körner, 1995). Although a multitude of studies have examined biogeochemical cycling in low-altitude rangeland ecosystems (Milchunas and Lauenroth, 1993; Kelly et al., 2002; Smith et al., 2002), very few studies have compared carbon and nitrogen processes between high-altitude alpine grasslands that have been grazed as opposed to adjacent areas that have been either ungrazed or released from grazing. Understanding the role of grazing in alpine grasslands C and N biogeochemistry is important today as land managers are considering means by which they may alter carbon sequestration characteristics with grazing management practices (Follett et al., 2000; Povirk et al., 2000, Conant and Paustian, 2002). Recently it has been suggested that mountain regions in the central United States are important carbon sinks in global carbon budgets (Schimel et al., 2002). While coniferous forests may be accounting for a large portion of these carbon budgets, alpine grasslands and tundra are spatially an important part of high-elevation systems in the Rocky Mountain region, and their carbon budgets may be vulnerable to changing climates (Walker et al., 1994, 1999; Welker et al., 2001).

Grazing by large herbivores can alter ecosystem processes directly and indirectly through biomass and nutrient redistribution (Frank et al., 2002). Biomass removal by grazers reduces the amount of standing litter affecting soil temperatures. Grazing may alter the magnitude of carbon cycling by shifting plant biomass allocation from roots to shoots or vice versa (Holland and Detling, 1990; Holland et al., 1992; Frank et al., 2002) and by delaying senescence. Urine and fecal deposition by herbivores can also alter soil nitrogen processes, thus affecting net soil N mineralization, plant N content, and photosynthetic processes (Holland and Detling, 1990; Frank and Groffman, 1998; Lal et al., 1998; Frank et al., 2000; LeCain et al., 2000). When herbivory results in greater N mineralization and higher leaf N contents (Holland and Detling, 1990; Seagle et al., 1992), then the short-term carbon exchange rates and carbon sequestering ability of grazed areas may become greater than ungrazed or lightly grazed areas due to stimulation of photosynthesis (Painter and Detling, 1981). However, generalizations such as these are not always certain or possible because of different herbivory intensities, intra- and interseasonal variations in temperature and precipitation, differing experimental designs, and statistical interpretations (Wilson and Jeffries, 1996; Tracy and Frank, 1998), along with intrinsic differences in rangeland ecosystems.

In this study, we quantified the CO₂ exchange (net ecosystem flux, ecosystem respiration, gross photosynthesis) and the soil N mineralization patterns as well as total biomass and C and N of soils and plants in an alpine grassland where grazing has been maintained and where it has been eliminated 45 yr ago with an exclosure (i.e., ungrazed area). Our measurements were conducted over two consecutive years, including winter, because non-growing-season C and N dynamics were shown to be important in alpine and winter-dominated ecosystems and C and N
biogeochemistry (Sommerfeld et al., 1993, 1996; Brooks et al., 1996; Brooks et al., 1997; Fahnestock et al., 1998, 1999; Bilbrough et al., 2000).

**Materials and Methods**

**SITE DESCRIPTION AND HISTORY**

Our study was conducted at Libby Flats (41°20'N, 106°17'W), located on the east side of the Medicine Bow Range in southeastern Wyoming (U.S.A.) approximately 64 km west of Laramie. The research site is an open alpine grassland located at 3230 m with a 3° slope to the north. The area became subject to intense unregulated grazing (e.g., herds numbering 200,000 in the Medicine Bow National Forest, with 60,000 in the Libby Flats—Medicine Bow Peak vicinity) with the introduction of sheep and cattle in the late 1890s, supporting the local mining and timber industries. Sheep grazing since the mid-1900s has been much more moderate, with 1 to 3 herded bands of ~2000 sheep rotating within the area each grazing season (July to September). The sheep allotment was not leased after 1995; thus, no livestock were present during the study period. A grazing exclosure (60 m × 25 m) was constructed in the center of the grassland by the U.S. Forest Service 45 yr prior to our study to serve as a visual reference site for the surrounding land that continued to be grazed. We used the exclosure and the grazed area surrounding it for this study. The exclosure is located in the center of a large grassland area (>3 ha), and no apparent micrometeorologic variance is evident between the exclosure and the grazed area within a 50-m circumference of the exclosure. Because this study was a comparison of 1 ungrazed, or released from grazing, area compared to 1 grazed area (pseudoreplication), we attempted to overcome this deficiency with repeated sampling of replicate plots within each area over a 2-yr period and conducted soil measurements at a second, subsidiary high-elevation grassland study site: Hay Creek. The Hay Creek study site (3000 m a.s.l.) is located 1.6 km north of the Big Horn National Forest Experimental Pasture, near Sheridan, WY. This exclosure was officially established in 1924 but had been in place and used as an observational plot since 1913 (May, 1954). The area adjacent to the Hay Creek exclosure was, and still is, grazed from June to September by cattle. This grassland is dominated by Festuca idahoensis, Bromus pumpellianus, Geum triflorum, Myosotis alpestris, Cerastium arvense, Lapinus sericeus, Phlox multiflora, Agoseris glauca, Poa canbii, Achillea millefolium, and Carex spp. (May, 1954). Our findings are primarily limited in scope to the study site at Libby Flats, but because the flora, parent material, and land-use history are representative of other alpine grasslands in the Rocky Mountain region (Billings, 1979; Walker et al., 1994), our findings may have implications for surrounding alpine grasslands.

Forb and graminoid species dominate the vegetation inside and outside the exclosure at Libby Flats and include Deschampsia cespitosa, Poa alpina, Polygonum bistortoides, Trifolium spp., Artemesia scopulorum, Geum rossii, Achillea millefolium, Cerastium arvense, Phleum alpinum, Solidago multidivarsata, and several Carex species. Soils at Libby Flats are fine, loamy, mixed Entic Humic Dystrochrepts (USDA, 1999).

Precipitation during the summer in southeast Wyoming is generally in the form of intermittent thunderstorm activity or, more rarely, snowfall. Annual precipitation, recorded at the nearby Glacier Lakes Ecosystem Experiments Site (GLEES), is approximately 120 cm, with summer precipitation accounting for less than 20% (NADP, 1999). Precipitation during our study period is summarized in Table 1, with winter snow being almost 50% greater in 1999 than in 1998; summer precipitation was 25% lower in 1999 than in 1998. If we compare years on a “water-year” basis (October–September), precipitation was greater in 1999 than in 1998. Over our 2-yr study period, this area received 302 cm of precipitation, 33% more than the average for the 6 previous 2-yr periods (NADP, 1999).

**CARBON DIOXIDE EXCHANGE MEASUREMENTS**

Carbon dioxide exchange patterns at Libby Flats were quantified by taking measurements during the growing (snow-free) and non-growing (i.e., snow-covered) seasons from January 1998 to January 2000. During the nongrowing season when snow was present, we used a CO₂ concentration gradient approach with a simple diffusion model to estimate wintertime losses of C from grazed and ungrazed areas through snowpack. This method has been documented elsewhere (Sommerfeld et al., 1993, 1996; Brooks et al., 1996; Fahnestock et al., 1998, 1999), but briefly, it entails measuring CO₂ concentrations at the snow-atmosphere and snow-soil interfaces with an infrared gas analyzer (PP Systems EGM-1) and calculating diffusional CO₂ loss from the soil to the atmosphere. Values are subsequently corrected for temperature, pressure, and nonlinear diffusional flow (i.e., tortuosity) resulting from snow density differences throughout the snowpack. Measurements were taken during the winters of 1998 (1 January–12 May), 1999 (12 December–3 June), and early 2000 (6 January). Weather permitting, 50 randomly located measurements were taken both inside and outside the exclosure during each sample date. Snow depths were recorded at each measurement location. The snow depth increased throughout the winter of 1998, was generally consistent in the winter of 1999, and was not different between the grazed and ungrazed areas (Fig. 1a). Profiles of snow density and temperature were made from snow pits dug at each site during each sample date, where soil temperatures were recorded (Fig. 1b). In addition to our CO₂ flux estimates, these measurements of snow depth and soil temperature allowed us to evaluate their relationship (Fig. 1c). Soil temperature may be one of the controlling factors of soil respiration and CO₂ efflux in winter (Brooks et al., 1996; Schimel and Clein, 1996; Fahnestock et al., 1998).

Growing-season CO₂ exchange measurements were taken with an infrared gas analyzer (LiCor, LI-6200) connected to a transparent chamber (Vourlitis et al., 1993) that was placed over preselected plots at the time of each measurement. Experimental plots (0.75 m × 0.75 m) were selected in July 1997 within the Libby Flats exclosure (8 plots) and in the adjacent grazed area (7 plots). Square polyethylene basins (75 cm × 75 cm × 20 cm high with a 5-cm horizontal lip) were set 15 cm into the soil to provide a seal for CO₂ flux measurements (Vourlitis et al., 1993). All grazed plots were located between 10 m and 30 m from the exclosure so that the differences between the grazed and ungrazed areas could not be attributed to differences in slope or aspect.

Carbon dioxide exchange was measured during the late spring, summer, and fall of 1998 (12 June–12 September) and in 1999 (20 June–12 November). Diurnal CO₂ measurements were taken ca. every 4 hr on each sampling date, and sampling occurred approximately
every 1 to 4 wk during the snow-free season. A 75 cm × 75 cm × 40 cm transparent chamber was sealed to each chamber base at the time of each measurement. Four small fans continuously mixed air in the chamber during measurements. Once CO₂ concentrations within the chamber stabilized (typically 30–60 s), net CO₂ exchange determinations were made for each plot. After these net CO₂ exchange measurement (net ecosystem exchange [NEE]), the chamber was covered with an opaque cloth to prevent photosynthesis, and measurements of ecosystem respiration rates were determined. Whole ecosystem C assimilation (i.e., gross photosynthesis) was subsequently calculated by subtracting the respiration measurements from the net CO₂ exchange measurements, similar to techniques we have used for carbon exchange characteristics of other alpine (Welker et al., 1999) and arctic (Jones et al., 1998; Welker et al., 2000) systems.

SOIL AND VEGETATION MEASUREMENTS

In July 1997, 10 soil samples were collected in the ungrazed exclosure and grazed area at Libby Flats, and in August 1998 at the Hay Creek site (May, 1954), from 3 depths (0–7.5 cm, 7.5–15 cm, 15–30 cm) for analyses of total C and N content using a 2-m King Tube. Soils were air dried, processed through 2 sieves (0.168 cm and 0.084 cm), and analyzed for total C and N concentration on a Carlo-Erba CHN analyzer. Soil bulk density was determined to a depth of 5 cm, as herbivore effects are often confined to the upper soil profiles (Elliott et al., 1999; Schuman et al., 1999). At the Hay Creek site only soil C and N measurements were conducted.

Net N mineralization rates were measured using an in situ soil core incubation method (DiStefano and Gholz, 1986). Eight sets of mineralization measurements were made from 24 June through November 1999 in the grazed and ungrazed areas. For each measurement period, paired soil cores were installed in close proximity to each other. One soil core was harvested immediately for an initial measurement of inorganic N and gravimetric water content. An ion-exchange resin was placed at the bottom of the second core as a leachate trap, and the core was returned to the soil for an incubation period of 2 to 4 wk. Entire soil cores were returned to the lab and processed within 4 hr of sample collection. Coarse roots and gravel were removed, and gravimetric soil water content was determined on a 10-g subsample. Nitrogen was extracted from a second 10-g subsample with 2M KCl for 1 hr. Filter extracts of both soils and resins were analyzed for NH₄⁺ and NO₃⁻ using a continuous-flow autoanalyzer. Leachate N values from resin bags were added to the N values from the incubated soil cores. The difference in N concentration between the initial (T₀) and final (T₁) measurement of each period was divided by the number of incubation days to yield daily net N mineralization rates. At each sample date, 3 soil temperature measurements were recorded adjacent to the soil cores using a thermocouple placed 5 cm below the surface.

Vegetation samples for C and N concentration determination and total biomass were collected in mid-July 1999 from 10 randomly placed quadrats (0.25 m²) in the grazed and ungrazed areas. All vegetation in each quadrat was harvested to the soil surface and separated in grass and forb categories. Green leaves were separated from dead leaves and stems, all vegetation was oven-dried at 60°C for 48 hr, and total biomass was measured to the nearest 0.1 g. Green leaf biomass was ground to a fine powder using a Wiley mill and a mortar and pestle, and subsamples were analyzed for total C and N concentration on a Carlo-Erba CHN analyzer.

STATISTICAL ANALYSIS

A two-way, repeated-measures analysis of variance (SAS Institute, 1994) was used to detect significant differences (P < 0.05) in summer CO₂ exchange between sample dates and between the ungrazed and grazed treatments, with plots treated as replicates. Separate analyses were performed on the net CO₂ exchange rate and the rates of photosynthesis and ecosystem respiration. General Linear Model procedures (SAS Institute, 1994) were used to detect differences (P < 0.05) in rates of wintertime CO₂ efflux and total soil and plant C and N concentration between grazed and ungrazed treatments, with the soil data separated by soil depth. When significant interactions occurred between sample date and grazing treatment, individual t-tests were used to test for significant (P < 0.05) grazing treatment effects for CO₂ exchange characteristics at individual sample dates. Soil inorganic N pools and net N mineralization rates were each analyzed with a two-
factor ANOVA using General Linear Models with grazing treatment and sampling date as the main effects. Box plots and normal probability plots of residuals were used to assess normality and outliers. Where appropriate (i.e., $P < 0.05$ in the main analysis), means were compared using the Tukey-Kramer test.

**Results**

**WINTER CO$_2$ EXCHANGE**

Detectable levels of CO$_2$ were lost from these grasslands to the atmosphere in all three winters on Libby Flats. Carbon dioxide efflux was measurable even when air and soil temperatures were fairly cold (e.g., on 1 January 1998, air and soil temperatures were $-7.0$ and $-2.9^\circ$C, respectively) (Fig. 2) and snow depth was relatively shallow (ca. 20 cm; Fig. 2). Snow depth and temperature were found to be linearly related (Fig. 1c) ($r^2 = 0.4$). Overall, there was a significant ($P < 0.01$) interaction between sample date and grazing treatment effect on winter CO$_2$ efflux. At several sample dates during the winter measurement periods (1 January, 22 April, and 12 May 1998; 3 March and 3 June 1999; and 6 January 2000), the grazed area lost significantly more carbon (i.e., was a greater C source) than the ungrazed area (Fig. 2). The range of carbon lost during winter varied between $-0.2$ and $-1.5$ mol CO$_2$ m$^{-2}$ s$^{-1}$ (note that negative numbers signify loss of C to the atmosphere, while positive values indicate net C gain by the ecosystem).

**GROWING SEASON CO$_2$ EXCHANGE**

The grazed area at Libby Flats generally had lower rates of net ecosystem CO$_2$ exchange (i.e., lower C-sink activity) during the snow-free period (≈ early June to mid-October) than the ungrazed area, especially during the first summer of study (Fig. 2). A significant ($P < 0.01$) interaction between sample date and grazing treatment was found for net CO$_2$ exchange in both the 1998 and 1999 growing seasons. In 1998, net CO$_2$ exchange was significantly higher in the ungrazed than in the grazed area on half of all sample dates, while in 1999 this difference was evident at 25% of the sample dates. Lower growing-season carbon gain in the grazed relative to the ungrazed area was due to a combination of lower photosynthetic rates and higher respiration rates (Fig. 3). These higher respiration rates correspond to higher soil temperatures in the grazed as opposed to the ungrazed area (Table 1).

Diurnal patterns of NEE depict the short-term temporal nature of carbon exchange at this site in addition to the effects of grazing history (Fig. 4). Early in the summer, both of these systems are net C sources to the atmosphere, but by mid-July, during all hours of the day, both grazed and ungrazed areas net C sources, being greater in the ungrazed. At sunset, while the grazed area is becoming a C source, the ungrazed area remains a C sink. Maximum rates of daily NEE occur in late July in both the grazed and ungrazed systems, being almost two-fold higher in the grazed.

We estimated the carbon budget of these systems by taking the mean daily CO$_2$ flux for each sample date (average of the diurnal or average of efflux measured during the day in winter) and multiplying it by the number of days assigned to represent a period between sampling dates (Welker et al., 1999, 2000; Bubier et al., 2002). We then converted the mass of CO$_2$ to the mass of CO$_2$-C to estimate the CO$_2$-C exchange in the grazed and ungrazed areas over this 2-yr period (Welker et al., 1999, 2000). The grazed area was a carbon source of 170 g C m$^{-2}$, while the ungrazed area was a carbon sink of 83 g C m$^{-2}$ during the 2-yr period.
SOIL AND VEGETATION PROPERTIES

Total soil C and N concentrations in the upper soil layer (0–7.5 cm) were 67% and 60% greater in the grazed than in the ungrazed area, respectively (Table 2). However, there were no significant differences in soil C and N between grazed and ungrazed areas in the lower soil layers, and no differences between bulk density or soil C:N ratios at any depth (Table 2). This pattern of higher C and N in the upper profile was also observed for Hay Creek, though the values were slightly higher than those from Libby Flats (Table 2). Total N of live green leaves of aboveground vegetation, on the other hand, was significantly (P < 0.05) higher in the ungrazed than grazed area. Total biomass was significantly (P < 0.04) higher in the ungrazed area (225 g m⁻²) compared to total biomass in the grazed (177 g m⁻²) (Table 2).

Rates of nitrification varied significantly over the 1999 growing season and were highest in spring and lowest in late fall. Grazing history had no detectable effect on nitrification rates (Fig. 5a). However, there was a significant (P < 0.05) effect of grazing history on net N mineralization rates at certain dates in 1999 (Fig. 5b). Rates of N mineralization were highest early in the growing season for both grazing treatments, with significantly higher rates in the ungrazed than grazed area in late June to early July (Fig. 5b). Rates declined rapidly with the onset of summer, and no significant differences between grazing treatments were observed during this period. Net N mineralization rates increased in early fall and were once again higher in the ungrazed than grazed area between August 30 and September 23 (Fig. 5b). This pattern switched, however, between late September and early October, with the ungrazed area having lower net N mineralization rates than the grazed area, though the absolute rates were exceedingly low.

There was a significant (P = 0.01) date by grazing treatment interaction for extractable soil NH₄⁺, but extractable soil NO₃⁻ levels were only affected by sample date (P = 0.02) (Fig. 6). Soil NH₄⁺ levels were significantly (P < 0.05) higher in the ungrazed grassland in early and mid-July when concentrations were at their greatest. Soil nitrate levels did not differ between grazing treatments and were also at their greatest concentrations in late July. There was a significant date by grazing treatment interaction (P = 0.01) on gravimetric soil moisture levels (Fig. 7a), with higher soil moisture content in the ungrazed than the grazed area in early summer. Soil temperatures, measured at 5 cm depth, were similar in grazed and ungrazed areas (Fig. 7b).

Discussion

The C and N characteristics of this grazed and ungrazed area were different in the patterns and magnitudes of carbon exchange, carbon sequestration budgets, and in the carbon and nitrogen concentrations of soils and vegetation. Based on our initial discovery that the soil carbon and nitrogen contents were significantly higher in the grazed area (1997), our studies began with the expectation that the grazed area would have higher rates of net CO₂ exchange that would be accompanied by higher vegetation N contents and higher soil N

TABLE 2
Soil and vegetation properties of ungrazed and grazed alpine grassland at Libby Flats and at Hay Creek, Wyoming

<table>
<thead>
<tr>
<th>Soil property</th>
<th>Ungrazed</th>
<th>Grazed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk density (g cm⁻¹)</td>
<td>0.59 ± 0.10</td>
<td>0.58 ± 0.10</td>
</tr>
<tr>
<td>% Carbon</td>
<td></td>
<td></td>
</tr>
<tr>
<td>−7.5 cm</td>
<td>6.6 ± 0.3*</td>
<td>11.0 ± 1.0*</td>
</tr>
<tr>
<td>7.5–30 cm</td>
<td>3.6 ± 0.7</td>
<td>3.7 ± 0.3</td>
</tr>
<tr>
<td>30–60 cm</td>
<td>1.0 ± 0.1</td>
<td>1.7 ± 0.3</td>
</tr>
<tr>
<td>% Nitrogen</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–7.5 cm</td>
<td>0.6 ± 0.1*</td>
<td>0.9 ± 0.1*</td>
</tr>
<tr>
<td>7.5–30 cm</td>
<td>0.3 ± 0.1</td>
<td>0.3 ± 0.1</td>
</tr>
<tr>
<td>30–60 cm</td>
<td>0.1 ± 0.1</td>
<td>0.1 ± 0.1</td>
</tr>
<tr>
<td>C:N ratio</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–7.5 cm</td>
<td>11.5 ± 1.5</td>
<td>12.1 ± 1.1</td>
</tr>
<tr>
<td>7.5–30 cm</td>
<td>12.8 ± 2.1</td>
<td>11.9 ± 2.0</td>
</tr>
<tr>
<td>30–60 cm</td>
<td>15.8 ± 1.5</td>
<td>11.8 ± 1.0</td>
</tr>
<tr>
<td>Vegetation N (%)</td>
<td>1.53 ± 0.03*</td>
<td>1.41 ± 0.03*</td>
</tr>
<tr>
<td>Mean Summer Soil Temperatures (°C)</td>
<td>7.7 ± 0.9</td>
<td>8.9 ± 0.7</td>
</tr>
<tr>
<td>Plant Biomass</td>
<td>225 ± 12.7*</td>
<td>177 ± 15.9</td>
</tr>
<tr>
<td>Hay Creek</td>
<td></td>
<td></td>
</tr>
<tr>
<td>% C (0–7.5 cm)</td>
<td>6.1 ± 0.5</td>
<td>7.2 ± 0.3*</td>
</tr>
<tr>
<td>% N (0–7.5 cm)</td>
<td>0.5 ± 0.1</td>
<td>0.7 ± 0.1*</td>
</tr>
<tr>
<td>C:N ratio (0–7.5 cm)</td>
<td>12.2 ± 1.4</td>
<td>10.0 ± 0.9</td>
</tr>
</tbody>
</table>

* indicates significant (P < 0.05) differences between grazing regimes.
availability (N mineralization). These postulates were, however, not manifested in our winter and summer carbon exchange measurements, our estimates of CO₂-C budgets, or our measurements of soil and plant N contents. For instance, the grazed area was a C source over this sampling period while the ungrazed area was a C sink, with the grazed area having lower biomass. These differences correspond to the summer temperature measurements whereby soils were warmer in the grazed area, which would account for the higher rates of ecosystem respiration, lower C-sink strength, and overall CO₂-C source activity of the grazed area during our study period.

Our measurements of carbon exchange depicted three important findings: (1) Carbon loss in winter was detectable at all measurement periods (winter soil temperatures were always above −5°C, a theoretical threshold of microbial activity (Schimel and Clein, 1996; Brooks et al., 1996) and there were measurement periods during the snow-covered period, when the grazed area had significantly greater rates of carbon loss (carbon source) are likely the result of the higher soil C contents in the grazed area (Table 2), as opposed to being the result of warmer winter soil temperatures (Schimel et al., 2004). (2) There were also periods during the snow-free period (summer) when the rates of net CO₂ flux were lower in the grazed area compared to the ungrazed system, which is the result of generally lower rates of ecosystem photosynthesis in the grazed (primarily 1998) as opposed to the ungrazed area, in conjunction with higher rates of ecosystem respiration in the grazed area (1998 and 1999). The lower rates of gross photosynthesis correspond to the lower plant N content, which typically equates to lower rates of leaf photosynthesis (Field and Mooney, 1986). Collectively, these fluxes equate to a biannual carbon budget in the grazed area of 170 g C m⁻² (CO₂-C source) (an average of 85 g C m⁻²); while the ungrazed area was a CO₂-C sink of 83 g C m⁻² over the biannual period (an average of 41 g C m⁻²). These estimates of annual CO₂-C flux are very similar in magnitude to the mass of C acquired by alpine systems in GLEES (40.9 g C m⁻² in 1991, 47.9 g C m⁻² in 1992) (Sommerfeld et al., 1993), though higher than the values reported for a suite of alpine ecosystems on Niwot Ridge, Colorado, in 1993 and 1994 (0.3–16.9 g C m⁻² yr⁻¹) (Brooks et al., 1996). The higher elevation of Niwot Ridge (>3000 m a.s.l.) and its associated shorter growing season could account for the lower magnitudes of annual carbon exchange. In addition, the magnitudes of

FIGURE 5. Growing-season (1999) patterns of (a) net nitrification and (b) net N mineralization in an ungrazed and a grazed area of Libby Flats, Wyoming. Shown are means ± 1 SE. Asterisks indicate significant (P < 0.05) differences between treatments at a sampling date.

FIGURE 6. Seasonal (1999) patterns of extractable inorganic soil N in grazed and ungrazed areas of Libby Flats, Wyoming. Shown are means ± 1 SE of (a) NH₄⁺, (b) NO₃⁻, and (c) total N. Asterisks indicate significant (P < 0.05) differences between treatments at a sampling date.

FIGURE 7. Seasonal pattern of summer (a) gravimetric soil water content, measured from 0–10-cm depth, and (b) soil temperature, measured at 5-cm depth in 1999. Shown are means ± 1 SE. Asterisks indicate significant (P < 0.05) differences between treatments at a sampling date.
the Rocky Mountains (e.g., —80 g C m\(^{-2}\) yr\(^{-1}\), Schimel et al., 2002). (3) Our carbon exchange and sequestration findings indicate that integrative measures of ecosystem carbon budgets (soil C contents) may not necessarily correspond to shorter-term, instantaneous measures of CO\(_2\) exchange. Clearly the accumulation of soil carbon is indicative of prolonged periods where carbon gains exceed carbon losses, especially in grasslands where secondary stem growth is absent (Follett et al., 2000). However, the time periods by which soil carbon properties are manifested often exceed the measurement periods of short-term studies, such as this and others (Frank et al., 1995; LeCain et al., 2000). We propose that the nature of ecosystem carbon sequestration is dynamic, and while the trajectory for this grazed grassland is positive (accumulating soil carbon), periods of net soil carbon activity are interspersed with periods when these grazed grasslands are net carbon sinks (Fig. 8). Oechel et al. (1993, 1995) report that the North Slope of Alaska (a cold Carex-dominated grassland) has oscillated between being a C sink for extended periods, interspersed with periods of being a net carbon source of CO\(_2\) to the atmosphere and being in carbon balance (Oechel et al., 2000).

The interpretation of the carbon dynamics of our ungrazed area (grazing removal exclosure) requires qualification. We have relied on the assumption that the exclosure was placed in a representative area that was not distinct from the surrounding grassland. This assumes, then, that at the time of construction, the soil carbon and nitrogen contents were uniform across the entire study area. If this assumption is true (likely), then we are left with deciphering how the elimination of grazing by livestock and wildlife (no observational or feral evidence of wildlife grazing) in the exclosure could lead to lower soil carbon and nitrogen pools in the upper soil profile. Soil carbon losses could be the result of greater rates of decomposition and associated greater rates of soil and ecosystem carbon loss; CO\(_2\), or methane (Hobbie et al., 2000). If soil temperatures were warmer in the exclosure, then it is possible that greater carbon losses could account for these differences. However, our measurements of soil temperatures indicate that, on the contrary, soils are warmer in the grazed area where standing dead leaf litter is eliminated by herbivores (Fig. 2).

Another possible explanation for the higher soil C and N in the grazed as opposed to the ungrazed area may be the interactions that occur between herbivory (grazing) intensity and carbon allocation to above- and belowground tissue by grasses (Welker et al., 1985; Holland et al., 1992; Welker and Briske, 1992; Holland et al., 1996; Frank et al., 2002). In some instances, grazed grasslands allocate greater proportions of carbon to roots, as compared to ungrazed grasslands, and have higher root turnover, which in turn contributes to greater amounts of soil carbon and nitrogen (Sims and Singh, 1978; Frank et al., 2002). It is likely that reducing the grazing intensity on this alpine grassland by erecting the exclosure may have lead to a reduction in carbon allocation belowground, which would lead to lower rates of root turnover and lower soil carbon accumulation and sequestration over time (Fig. 8).

However, there appears to be a disconnect between our short-term measures of ecosystem carbon cycling (i.e., CO\(_2\) flux measurements) and the longer-term soil carbon dynamics. This disconnect between short- and longer-term processes is not uncommon, and other studies in low-altitude grasslands with divergent grazing histories also indicate that instantaneous rates of CO\(_2\) exchange and soil C contents may not be consistent, i.e., grasslands with the highest soil C contents do not necessarily exhibit the highest rates of instantaneous CO\(_2\) exchange (LeCain et al. 2000). The processes controlling the linkages between instantaneous rates of net CO\(_2\) exchange and soil carbon contents and soil carbon sequestration require further investigation and represent one of the greatest uncertainties in grassland carbon cycling, grazing management, and its interaction with climate.

Our observation that curtailing grazing in a mountain alpine grassland may change in the short term the sign of CO\(_2\) exchange from a CO\(_2\)-C source to a CO\(_2\)-C sink has implications for the study of carbon sequestration in the western U.S. mountains (Conant and Paustian, 2002; Schimel et al., 2002,?). In this presentation, Schimel et al. (2002) indicates that changes in land use practices, the recovery of forests from intensive harvesting, and active fire suppression are contributing to the strong C sinks found in the higher elevations of the Rocky Mountain region, approaching 80 g C m\(^{-2}\) yr\(^{-1}\). Our findings suggest that changing grazing practices may also contribute to these “hot spots” of carbon sequestration at high altitudes, as there has been a progressive decline of sheep grazing in high-altitude grasslands, due primarily to reduced consumer demand for red meat, and especially lamb, over the past 40 yr (Yav ra et al., 1994; Laycock et al., 1996). Considering the rates of C sequestration of these high-altitude grasslands may approach 40 g C m\(^{-2}\) yr\(^{-1}\), the sign of these systems within the Rocky Mountains may have a large influence on regional carbon processes. While these alpine grassland systems may continue to be used as summer forage by Cervus (elk), the reduction of heavy grazing by sheep may contribute to the strong C sinks observed in the western U.S. mountains. As indicated by others, land use history and carbon cycling are closely linked (Falkowski et al., 2000) and may or may not have just as large an effect on carbon sequestration as changes in climate at high altitudes (Schuman et al., 1999; Welker et al., 1999, Conant and Paustian, 2002).

Livestock grazing history at our site appears to have had a long-term positive effect on soil N and C content, especially in the upper soil horizons, similar to that reported by Schuman et al. (1999). This is in contrast to the findings of Frank and Groffman (1998), who found that long-term grazing by native ungulates in Yellowstone National Park, another high-altitude Wyoming grassland, had no effect on any grassland soil variable except gravimetric soil moisture. However, a later study reports that root biomass is higher in grazed than in ungrazed grasslands (Frank et al., 2002), which should in turn equate to higher soil C and N. The higher soil N and C contents we observed at Libby Flats were not, however, mirrored in plant N content, nor were mineralization and nitrification rates consistently higher in grazed sites, as might have been expected. In fact, several times during the growing season, plants in grazed sites had reduced N availability relative to plants in ungrazed sites. This also is in contrast to the findings of Frank and Groffman (1998), who found that net N mineralization in grazed areas was double that of ungrazed areas.
The difference between our observations (grazing having a positive effect on soil C and N contents) and those of Frank and Groffman (1998) may be attributed to the dispersed nature of bison in Yellowstone and thus the patchy nature of microsites where urine and defecation may have had a positive effect on soil N, net primary productivity, and soil C. In contrast, Libby Flats had historical grazing intensities that exceeded thousands of animals in a single open grassland surrounded by subalpine forest, so more uniform grazing, defecation, and urination occurred (Duthie, 1916; Povirk et al., 2001). Bands of sheep are typically herded as very large units to minimize predatory attack; thus, uniform impacts on plants and soils are more likely.

While many studies suggest that large herbivores generally have a positive effect on N cycling (e.g., McNaughton, 1985; Frank and Evans, 1997; Frank et al., 2000), our study, and those of others, provides contrasting examples. In both our study and those in northwest Wyoming (Frank and Groffman, 1998), there were periods during the growing season when rates of N mineralization were indeed lower in grazed as opposed to ungrazed areas. In our study, leaf N was also lower in the grazed area, supporting our observations of reduced N mineralization rates. Tracy and Frank (1998) found that net N mineralization was not significantly \( (P < 0.05) \) higher in grazed than in ungrazed areas in northwest Wyoming, nor was shoot N significantly greater in grazed areas. Therefore, it appears that grazing effects on higher-altitude grassland nitrogen mineralization rates are not spatially uniform and merit further investigation before thematic generalizations can be adopted.

Lower N mineralization rates early and late in the summer (1999) in the grazed site at Libby Flats corresponded to lower rates of net CO\(_2\) exchange and lower plant N content compared to the ungrazed enclosure. However, this apparent relationship between C and N cycling was not observed in the middle of the growing season; while CO\(_2\) gain was much lower in grazed than ungrazed sites at this time, there were no differences in N mineralization or nitrification rates. Because mineralization and nitrification rates are so close to zero, linkages between C and N processes are difficult to detect. It does, however, merit reinforcing that there are indeed phases of the growing season (early summer) when lower rates of nitrification and mineralization (N availability indices) are in line with lower magnitudes of net CO\(_2\) flux, suggesting that coupling can occur between C and N processes in this alpine grassland.

Taken together, important insights regarding soil and plant N dynamics and the impact of grazing can be drawn from this and related studies (Dormaar et al., 1977; Dormaar and Willms, 1990; Bauer et al., 1987; Wilson and Jefferies, 1996; Bardgett et al., 1997, 1998; Frank and Groffman, 1998; Schuman et al., 1999; Stohlgren et al., 1999; LeCain et al., 2000):

1. Grazing history can affect the magnitudes, patterns, and amounts of C exchanged with the atmosphere in both predictable and unpredictable ways.
2. Reducing, and in this case eliminating, livestock grazing may result in time periods when the annual or biannual estimated carbon budgets are different from those where grazing is maintained and may cause the carbon budget sign to switch from being a carbon source to being a carbon sink (Conant and Paustian, 2002). However, one needs to keep in mind that these short-term estimates of carbon budgets are not necessarily a reflection of the longer-term carbon sequestration trajectories and may not completely account for changes in belowground carbon processes.
3. Positive effects of grazing on N mineralization are not necessarily manifested consistently throughout the growing season but may be episodic in nature. There are times when the effect of grazing is positive, periods in which grazing may have a negative effect, and times when there is no grazing effect on N mineralization. These are likely controlled, in part, by the allocation of carbon belowground by plants, the use of root exudates by soil microbes, and their effect on soil N mineralization and precipitation and soil water conditions (Holland and Detling, 1990; Holland et al., 1992, 1996).

4. The effects of grazing on N concentrations and fluxes may not be the same for all graminoid-dominated systems; thus, converting ungrazed systems to grazed systems will not necessarily have a beneficial effect on soil N mineralization or plant-available N and may not lead to increases in carbon sequestration or forage production in all systems (McNaughton, 1985; Coughenour, 1991; Milchunas and Lauenroth, 1993), nor is it certain that converting heavily (over)grazed systems to ungrazed or lightly grazed systems will alter short- and longer-term carbon processes in predictable ways.
5. There appear to be periods during the summer when the magnitudes of net carbon flux correspond to the indices of plant nitrogen availability and there are periods when there is very low correspondence between C and N processes, indicative of the complexity in biogeochemical cycling in terrestrial ecosystems.

### Acknowledgments

Funding for this research was provided in part by the USDA National Research Initiative Soil and Soil Biology Program (2001789) and the NSF Joint Program for Terrestrial Ecology and Global Change (IBN-97227796).

### References Cited


Conant, R. T., and Paustian, K., 2002: Potential soil carbon se-


Ms submitted June 2002
Revised ms submitted June 2003