

Effects of increased soil water availability on grassland ecosystem carbon dioxide fluxes

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Abstract There is considerable interest in how ecosystems will respond to changes in precipitation. Alterations in rain and snowfall are expected to influence the spatio-temporal patterns of plant and soil processes that are controlled by soil moisture, and potentially, the amount of carbon (C) exchanged between the atmosphere and ecosystems. Because grasslands cover over one third of the terrestrial landscape, understanding controls on grassland C processes will be important to forecast how changes in precipitation regimes will influence the global C cycle. In this study we examined how irrigation affects carbon dioxide (CO₂) fluxes in five widely variable grasslands of Yellowstone National Park during a year of approximately average growing season precipitation. We irrigated plots every 2 weeks with 25% of the monthly 30-year average of precipitation resulting in plots receiving approximately 150% of the usual growing season water in the form of rain and supplemented irrigation. Ecosystem CO₂

fluxes were measured with a closed chamber-system once a month from May–September on irrigated and unirrigated plots in each grassland. Soil moisture was closely associated with CO₂ fluxes and shoot biomass, and was between 1.6% and 11.5% higher at the irrigated plots (values from wettest to driest grassland) during times of measurements. When examining the effect of irrigation throughout the growing season (May–September) across sites, we found that water additions increased ecosystem CO₂ fluxes at the two driest and the wettest sites, suggesting that these sites were water-limited during the climatically average precipitation conditions of the 2005 growing season. In contrast, no consistent responses to irrigation were detected at the two sites with intermediate soil moisture. Thus, the ecosystem CO₂ fluxes at those sites were not water-limited, when considering their responses to supplemental water throughout the whole season. In contrast, when we explored how the effect of irrigation varied temporally, we found that irrigation increased ecosystem CO₂ fluxes at all the sites late in the growing season (September). The spatial differences in the response of ecosystem CO₂ fluxes to irrigation likely can be explained by site specific differences in soil and vegetation properties. The temporal effects likely were due to delayed plant senescence that promoted plant and soil activity later into the year. Our results suggest that in Yellowstone National Park, above-normal amounts of soil moisture will only stimulate CO₂ fluxes across a portion of the ecosystem. Thus, depending on the topographic

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location, grassland CO₂ fluxes can be water-limited or not. Such information is important to accurately predict how changes in precipitation/soil moisture will affect CO₂ dynamics and how they may feed back to the global C cycle.

Keywords Net ecosystem CO₂ exchange · Plant assimilation · Ecosystem respiration · Soil respiration · Plant respiration · Closed chamber · Irrigation · Soil moisture · Climate change · Altered precipitation regime

Introduction

Global temperature and precipitation regimes are expected to change as a consequence of increasing levels of atmospheric carbon dioxide (CO₂) and other greenhouse gases (Oreskes 2004). Predicted changes in the amount and frequency of precipitation at both seasonal and annual scales will affect the variation in soil moisture and the plant and soil processes influenced by soil moisture (Lauenroth and Sala 1992; Knapp et al. 2002). Generally, under non-saturated soil conditions, water additions trigger short-term increases in microbial activity and plant assimilation/respiration, with plants reacting somewhat slower (hours to days) compared to microbes (hours; e.g., Sala and Lauenroth 1982; Huxman et al. 2004a, b). Longer-term (annual—decadal) responses of non-saturated systems to water additions are, in contrast, related to increases in soil and plant CO₂ fluxes associated with changes in plant community composition and/or structure and associated changes in the composition of the decomposer community (e.g., Brown et al. 1997; Knapp et al. 2002; Tylor et al. 2004). Once a system becomes saturated, non-water limited or even anoxic soil conditions may predominate and both soil and plant CO₂ fluxes decrease (e.g., Kozłowski 1984; Bouma and Bryla 2000). Thus, changes in precipitation regimes will affect the assimilation and loss of carbon (C) depending on how water-limited photosynthetic and metabolic processes are among members of the ecosystem.

Grasslands cover approximately one-third of the earth's terrestrial surface area (Lieth 1978), and store between 10% and 30% of the world's soil C (Anderson 1991; Eswaran et al. 1993). Consequently, small precipitation-associated changes in C pools and

fluxes within these systems could have a considerable impact on the global C cycle. Most studies that investigated the effect of variable precipitation on ecosystem CO₂ fluxes in grasslands limited their investigation to (i) differences found across large scales (regional, continental; e.g., see articles in Soussana et al. 2007), or (ii) to the variability at a single location only (e.g., Fay et al. 2000, 2003; Meyers 2001; Liu et al. 2002; Suyker et al. 2003; Hunt et al. 2002, 2004; Huxman et al. 2004a; Chimner and Welker 2005; Flanagan and Johnson 2005; Harper et al. 2005; Jaksic et al. 2006).

Yet, grasslands are spatio-temporally heterogeneous landscapes, where topography creates small-scale spatial differences in soil texture and moisture (e.g., Schimel et al. 1985; Frank et al. 1994), C and nitrogen content (e.g., Frank et al. 1994; Turner et al. 1997), and plant composition and production (e.g., Abrams et al. 1986; Frank and McNaughton 1992; Lauenroth and Sala 1992; Knapp et al. 1993; Briggs and Knapp 1995). Additionally, temporal (i.e., monthly and annual) changes in air temperature and precipitation alter plant and soil processes (Lauenroth and Sala 1992; Briggs and Knapp 1995; Epstein et al. 2002; Knapp et al. 2002). Consequently, the responses of grassland ecosystem CO₂ fluxes to changes in precipitation are likely to be spatio-temporally heterogeneous. To our knowledge, there is no information available on how the spatio-temporal variation that is characteristic of most grassland ecosystems may interact with changes in precipitation to control grassland CO₂ fluxes.

The objective of this study was to measure the effect of irrigation on ecosystem CO₂ fluxes among five widely variable grasslands in Yellowstone National Park during a year of approximately average growing season precipitation. Therefore we assessed whether grassland CO₂ fluxes were water-limited during conditions of average soil moisture at different locations within the park, and, if not, where in the landscape ecosystem CO₂ fluxes were not water-limited.

Methods

Study area and sampling design

The study was conducted in the northern part of Yellowstone National Park, located in the

northwestern corner of Wyoming, USA (44°55' N to 45°10' N and 110°10' W to 110°50' W). Yellowstone National Park encompasses nearly 9,000 km² and is home for large migratory herds of elk (*Cervus elaphus* L.), bison (*Bison bison* L.), and pronghorn (*Antilocarpa americana* (Ord.)). Elevations range from roughly 1,500–3,500 m. The climate features long cold winters and short warm summers with an average annual temperature of approximately 1°C. The average annual precipitation ranges from 260 mm per year at the north boundary to 2,050 mm per year in the south west corner of the Park. We selected five sites located on an elevational gradient (Table 1, Fig. 1). They ranged from dry to mesic grassland with different soils and species composition (Table 1).

Within each site, we selected six 2 × 1 m plots in homogenous vegetation at the beginning of the 2005 growing season. Three of these plots were randomly chosen for irrigation. The irrigated plots were evenly watered with a watering can every other week from mid-April through late-September 2005 with 25% of the average monthly 30-year average of precipitation for a given month (Fig. 2). Consequently, irrigated plots received 50% of average precipitation as supplemental water. Precipitation data used to calculate the monthly 30-year averages were taken from Mammoth Hot Springs for site 4, Tower Falls for sites 1 and 2, and Lake for sites 3 and 5 (Fig. 1). This assured that differences in mean precipitation among the sites was accounted for in our experiment. Both the control and watered plots were subject to “regular” precipitation events (Fig. 2). Average precipitation totals during the winter preceding the study (November 2004–March 2005) and during the 2005 growing season (April–October) were between 63–89% and 106–111% of the 30-year average, respectively.

Ecosystem CO₂ fluxes

At each irrigated and control plot we measured NEE under different light conditions on a 50 × 50 cm area in the center of one half of the 2 × 1 m plot. Soil respiration was measured in the center of the other half of the 2 × 1 m plot. The measurements were conducted once a month from May to September on clear, sunny days near the solar maximum (between 9.30 h and 15.30 h). CO₂ fluxes were measured

Table 1 Location, soil moisture (growing season average), elevation (Elev.), fine fraction soil bulk density, soil pH, soil textural class, soil depth, soil N and C concentrations, average shoot biomass during the 2005 growing season (May–September), and dominant grass species of the five study sites. Sites are in order from the driest (site 1) to the wettest (site 5)

Site	Location	Soil moisture (%)	Elev. (m)	Fine fraction (<2 mm) soil bulk density (g cm ⁻³)	Soil pH	Soil textural class	Soil depth (cm)	Soil N (%)	Soil C (%)	Average shoot biomass (g m ⁻²)	Dominant grass species
1	Crystal Bench	14	1,890	1.18	5.2	Loamy sand	10–30	0.5	6.2	64	<i>Poa sandbergii</i> Vasey, <i>Stipa comata</i> Trin. & Rupr., <i>Carex stenophylla</i> Wahlenb.
2	Specimen Ridge	16	1,650	0.91	6.0	Sandy loam	>90	0.1	1.2	196	<i>Festuca idahoensis</i> Elmer
3	Canyon	33	2,430	0.78	3.9	Loam	>90	0.5	8.9	185	<i>Phleum pratense</i> L., <i>Danthonia intermedia</i> Vasey, <i>Deschampsia caespitosa</i> Beauv.
4	Mammoth Hot Springs	38	1,920	0.89	7.4	Clay loam	>90	0.3	3.5	240	<i>Poa pratensis</i> L., <i>Phleum pratense</i> L.
5	Norris	47	2,260	1.06	7.1	Sandy loam	>90	0.5	6.2	241	<i>Carex praegracilis</i> W. Boott, <i>C. pellita</i> Muhl. ex. Willd., <i>C. nebrascensis</i> Dewey, <i>Juncus balticus</i> Willd., <i>Muhlenbergia richardsonis</i> (Trin.) Rydb.

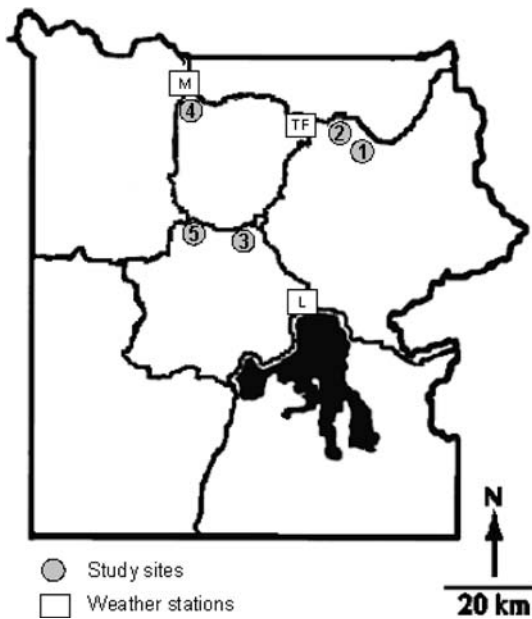


Fig. 1 The locations of the five study sites (grey dots) and three weather stations (white rectangles) in relation to roads in Yellowstone National Park (NOAA daily weather data). 1 = Crystal Bench, 2 = Specimen Ridge, 3 = Canyon, 4 = Mammoth Hot Springs, 5 = Norris; TF = Tower Falls, M = Mammoth Hot Springs; L = Lake

between 3 and 7 days after an irrigation event (Fig. 2). We are aware that by choosing such a measuring design we did not include short-term responses (1–2 days) of the CO_2 fluxes to irrigation, which normally includes increased soil respiration as a consequence of water infiltrating soil pores and purging the soil of CO_2 (e.g., Huxman and others 2004b). However, our goal was to specifically examine the ecosystem CO_2 exchange response to irrigation after the potential CO_2 flush from the soil to determine, biological responses of the plant and soil communities to increases in moisture availability. We could not sample CO_2 fluxes at site 3, located at the highest elevation, before July because of snow covering the site till mid June.

We sampled NEE using a temperature controlled 50 cm \times 50 cm \times 50 cm closed chamber system (chamber construction followed Wilsey et al. 2002, for specific chamber design used see Risch and Frank 2006). Fluxes were measured for 180 s after placing the chambers onto foam pads (to seal base) mounted to aluminum frames. Frames were driven three centimeters deep into the soil at least one hour before

the first measurement. The chamber was vented for approximately 120 s after each measurement. NEE ($\mu\text{mol CO}_2 \text{ m}^{-2} \text{ s}^{-1}$) was calculated using

$$\text{NEE} = d\text{CO}_2/dt \cdot P / [R \cdot (273.15 + T)] \cdot V / A \quad (1)$$

where $d\text{CO}_2/dt$ ($\mu\text{mol mol}^{-1} \text{ s}^{-1}$) = CO_2 accumulation rate inside the chamber during t seconds, based on linear regression, P = atmospheric pressure (kPa), R = gas constant, $8.314 \text{ (kPa m}^3 \text{ K}^{-1} \text{ mol}^{-1})$, T = chamber temperature ($^{\circ}\text{C}$), V = chamber volume (m^3), and A = chamber bottom area (m^2). CO_2 accumulation rates were corrected for water vapor (Hooper et al. 2002); air pressure correction was conducted automatically by the LICOR 6262. Pressure changes within the chamber were assumed to be minimal due to the large volume of the chamber, and the short measurement intervals (Healy et al. 1996).

We determined light-response curves for each grassland plot and sampling date by making two NEE measurements each under full light, 70% light, 50% light, 30% light (measured as photosynthetic photon flux density = PPFD) using different degree shade screens over the chamber, and under 0% light using a black cloth over the chamber. NEE and PPFD measurements were then used to determine relationships between CO_2 flux and light. We fitted linear

$$\text{NEE} = \alpha \cdot \text{PPFD} - R_{\text{eco}} \quad (2)$$

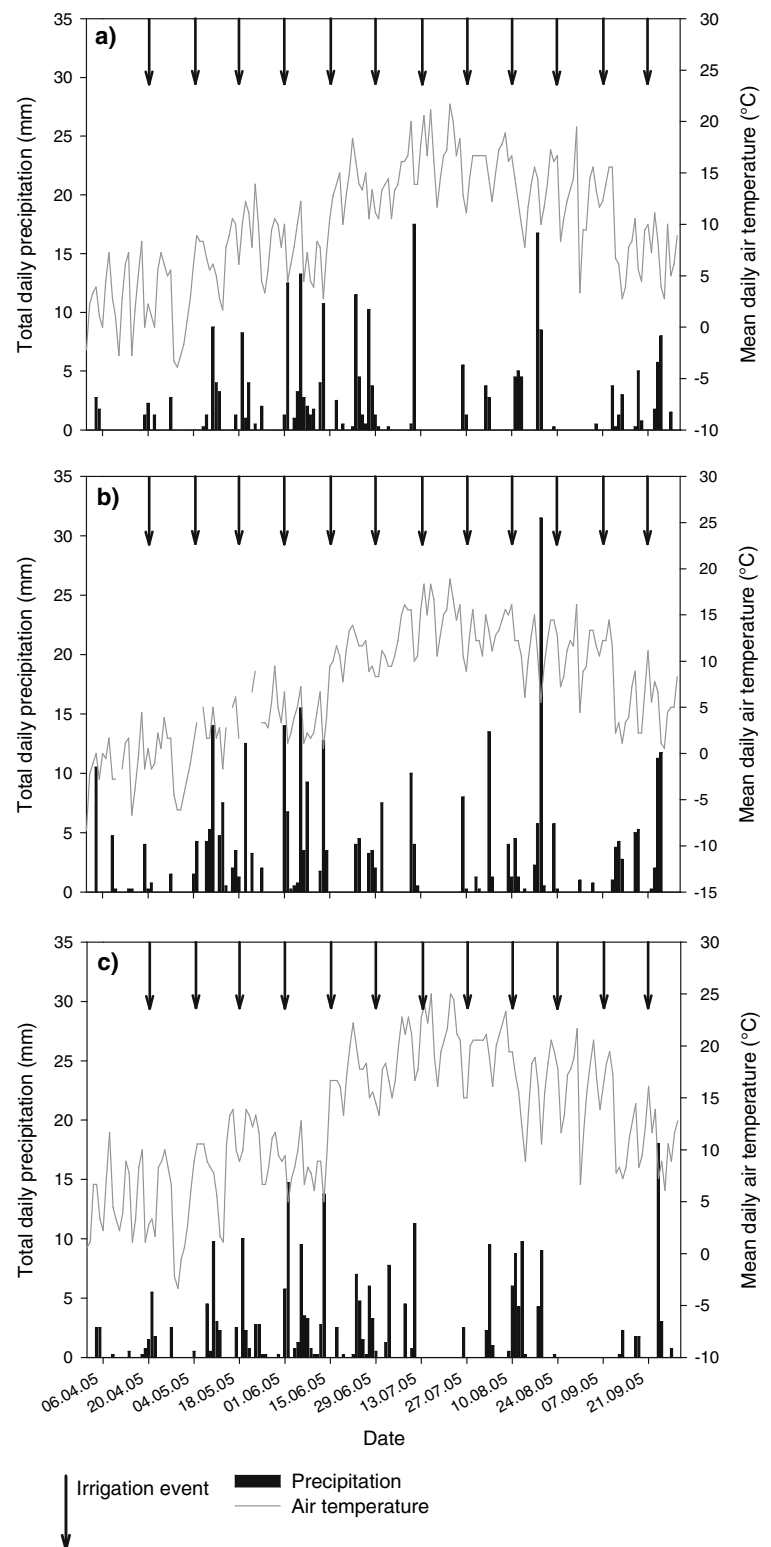
and hyperbolic

$$\text{NEE} = (\alpha \cdot \text{NEE}_{\text{max}} \cdot \text{PPFD} / (\text{NEE}_{\text{max}} + \alpha \cdot \text{PPFD})) - R_{\text{eco}} \quad (3)$$

equations following Ruimy et al. (1995), where α = quantum yield (initial slope in the hyperbolic equation), NEE_{max} = NEE maximum (asymptote of the hyperbola), R_{eco} = dark respiration (NEE at zero PPFD). Departure from linearity (D) was calculated as r^2 hyperbolic – r^2 linear (Ruimy et al. 1995). If D was <0.01 , the linear equation was used (Ruimy et al. 1995). We then calculated NEE_{1800} , which was NEE at full light ($1,800 \mu\text{mol s}^{-1} \text{ PPFD}$).

Soil respiration was measured with a 20 cm (high) by 10 cm (diameter) closed chamber system (for more detail see Risch and Frank 2006). The chamber was slid into PVC collars (10 cm diameter) driven five centimeters deep into the mineral soil

Fig. 2 Daily precipitation and air temperature values between April 1, 2005 and September 30, 2005 recorded at the Park's weather stations in closest proximity to our study sites: **(a)** Tower Falls, located at 1,910 m (sites 1 and 2), **(b)** Lake, located at 2,388 m (sites 3 and 5), **(c)** Mammoth Hot Springs, located at 1,899 m (site 4). Vertical arrows indicate time of irrigation events (bi-weekly)



after clipping the vegetation ≥ 40 min before the first measurement. New locations were randomly assigned for the rings every month. R_{soil} was calculated as the mean of two 3 min measurements using Eq. 1. The chamber was calibrated against the LICOR 6400-09 soil respiration system across a wide variety of soils in Yellowstone National Park to explore potential effects of pressure disequilibrium inside the chamber on respiration measurements (e.g., Davidson et al. 2002). We found a significant linear relationship between the two methods (eq. 2: $y = 1.061 \cdot x$, $r^2 = 0.72$, $P < 0.001$, $n = 30$, where y is the LICOR 6400-09 soil respiration system and x our own chamber). More pertinent to the aims of this calibration was that the X intercept was zero and there was no significant difference between the slope of the regression line and 1.0, indicating that the rates measured with our system were unbiased measures of soil respiration. In order to adjust for potential measurement differences between the different sized soil and NEE chamber, we calibrated these chambers against each other across a wide variety of soils to accurately calculate canopy respiration (R_{canopy}) by subtracting R_{soil} from R_{eco} (Eq. 3: $y = 0.4833x + 2.5128$, $r^2 = 0.74$, $P < 0.001$, $n = 33$, where y is the NEE chamber and x our soil chamber). R_{soil} measurements were then corrected using Eq. 3. Gross CO_2 assimilation (Assimilation) was calculated by summing NEE_{1800} and total ecosystem respiration (R_{eco}). Negative CO_2 flux values were CO_2 sources (CO_2 leaving the ecosystem), positive values were CO_2 sinks (CO_2 absorbed by the vegetation).

Biomass and environmental variables

We estimated aboveground biomass (biomass) non-destructively on three irrigated and three unirrigated 70×70 cm plots at each site with a canopy intercept method (Frank and McNaughton 1990). Biomass was estimated within 1 day of the CO_2 flux measurements.

Air temperature was recorded every five seconds during the CO_2 measurements with an air temperature sensor (HOBO weather station 8-bit temperature/RH smart sensor, Onset Computer, Bourne MA, USA) attached to the top of the NEE chamber and connected to a data logger (Fifteen channel H21

HOBO weather station data logger, Onset Computer, Bourne MA, USA). PPFD also was determined during the CO_2 measurements using a photosynthetically active radiation (PAR) sensor (HOBO weather station photosynthetic light (PAR) smart sensor, Onset Computer, Bourne MA, USA) attached to the inside surface of the top of the chamber and was recorded every five seconds on the data logger. Soil temperature (HOBO weather station 8-bit temperature smart sensor, Onset Computer, Bourne MA, USA) also was recorded on the data logger during the CO_2 measurements at both plots every five seconds at 20 cm mineral soil depth, while soil moisture was measured with a time domain reflectometer (TDR 100 soil moisture probe, Spectrum Technologies, Plainfield IL, USA) at six randomly selected locations to a depth of 12 cm immediately before the CO_2 measurements. Average soil moisture for each plot was then derived by averaging the six values. Both soil temperature and moisture were measured within the 2×1 m plots, 15 cm from the edge, but outside the area used for CO_2 measurements.

Statistical analyses

We averaged the monthly data for the ecosystem CO_2 exchange variables (R_{eco} , R_{soil} , Assimilation, R_{canopy} , NEE_{1800}), aboveground biomass, and soil moisture and used linear regression analyses to evaluate the relationship between the independent variable soil moisture, and the dependent ecosystem CO_2 exchange parameters (R_{eco} , R_{soil} , Assimilation, R_{canopy} , NEE_{1800}) for both the treatments separately. Linear regression analysis also was used to explore the relationships between the variables soil moisture and biomass, biomass and ecosystem CO_2 fluxes, ecosystem CO_2 fluxes and soil C, and between ecosystem CO_2 fluxes and air temperature. Irrigation effects on ecosystem CO_2 fluxes at each site were determined with two tailed paired t -tests ($\alpha = 0.1$) using monthly measurements of both irrigated and unirrigated plots at each site. To address the temporal (among months) variation in each of the ecosystem CO_2 exchange variables, we used repeated measure ANOVAs. Two tailed paired t -tests ($\alpha = 0.1$) were used to assess differences in ecosystem CO_2 fluxes between the irrigated and unirrigated plots for each month separately.

Results

Ecosystem CO₂ fluxes, biomass and soil moisture varied markedly among the five sites. Lowest fluxes generally were found at the driest site (site 1) and highest fluxes occurred at the site where average growing season soil moisture was highest (site 5; Table 1, Fig. 3). Soil moisture was positively and linearly correlated with assimilation, R_{eco}, R_{soil} and R_{canopy} measured in both irrigated and unirrigated grasslands across the Yellowstone landscape (Table 2). No correlation was found between NEE₁₈₀₀ and soil moisture for either of the treatments (Table 2). Soil moisture also was positively related to shoot biomass ($r^2 = 0.41$, $P = 0.047$), and shoot biomass was positively related to all the ecosystem CO₂ fluxes ($P < 0.05$ for all linear regressions). The fluxes were also closely correlated to air temperature (recorded throughout the measurements; $P < 0.05$ for all linear regressions). In contrast, none of the flux measurements was correlated to 0–20 cm mineral soil C concentrations.

Irrigating semi-monthly with a total of 50% of the 30-year average precipitation resulted in higher May–September soil moisture by an average of 11.5% at site 1, 5.2% at site 2, 3.2% at site 3, 2.7% at site 4 and 1.6% at site 5, respectively (at times of measurements). Thus, soil moisture content (see Table 1) of unirrigated plots was negatively correlated with increases in soil moisture content due to irrigation. Even though the snow melted much later at site 3 compared to the other sites, average soil moisture during the three months of measurement (July, August, September) was lower than at site 4 and 5 during the same period. At the two driest sites (1 & 2) R_{eco} was 7–39% (site 1) and 1.5–29% (site 2) higher at the irrigated compared to the

unirrigated plots during the growing season (P -values from paired t -tests: site 1: $P = 0.02$, site 2: $P = 0.007$; Fig. 4). At site 1, where soils were shallow (Table 1), the stimulation of R_{eco} by irrigation was due to a stimulation in R_{soil} ($P = 0.009$), which was between 4% and 36% higher at the irrigated compared to the unirrigated plots during May to September. At site 2, with very deep soils (Table 1), differences in R_{eco} were associated with significant increases in R_{canopy} ($P = 0.05$), which were between 26% and 52% higher at the irrigated compared to the unirrigated plots (Fig. 4). Irrigation also increased Assimilation by 8–64% ($P = 0.08$; Fig. 4) and NEE₁₈₀₀ by 20–2200% ($P = 0.08$; Fig. 4). NEE₁₈₀₀ also increased at the wettest site (site 5), which accumulated 1–41% more net CO₂ (NEE₁₈₀₀) when watered ($P = 0.06$; Fig. 4). In contrast, no irrigation effects on ecosystem CO₂ exchanges were detected at the two sites with intermediate soil moisture (sites 3 and 4; Fig. 4).

Assimilation (irr: $P = 0.008$, unirr: $P = 0.03$), R_{eco} (irr: $P = 0.01$, unirr: $P = 0.02$), and R_{soil} (irr: $P = 0.005$, unirr: $P = 0.007$) significantly differed over the course of the growing season, with highest fluxes at peak biomass (sites 1,2,4: July; sites 3,5: August) and lowest fluxes in May and September. NEE₁₈₀₀ and R_{canopy} values trended to be lower in May and September compared to June, July and August; however, no statistical differences were detected (NEE₁₈₀₀ irr: $P = 0.11$, unirr: $P = 0.43$; R_{canopy} irr: $P = 0.28$, unirr: $P = 0.15$), because of large measurement variations. All the ecosystem CO₂ fluxes measured during a specific month of measurement were closely related to shoot biomass ($P < 0.05$ for all linear regressions). When examining the effect of irrigation on CO₂ fluxes for each month separately, we found no significant difference in any ecosystem CO₂ flux variable measured until late in the season: in September, irrigation increased R_{eco} by 5–40% ($P = 0.05$), Assimilation by 13% and 61% ($P = 0.006$), R_{soil} by 4% and 36% ($P = 0.07$), and NEE₁₈₀₀ by 120–2200% ($P = 0.008$) among sites (Fig. 5). No significant difference was detected between irrigated and unirrigated plots for R_{canopy} in September (Fig. 5).

Table 2 Correlation coefficients and p -values from linear regressions between soil moisture and ecosystem CO₂ fluxes

	r^2		P -value	
	Unirr	Irr	Unirr	Irr
NEE ₁₈₀₀	0.47	0.55	0.2	0.15
Assimilation	0.79	0.80	0.04	0.04
R _{eco}	0.89	0.87	0.02	0.02
R _{soil}	0.85	0.94	0.02	0.006
R _{canopy}	0.78	0.71	0.05	0.07

$n = 5$, $\alpha = 0.1$, unirr = unirrigated, irr = irrigated

Discussion

Similar to this study, we previously found (Risch and Frank 2006) a close association between soil moisture

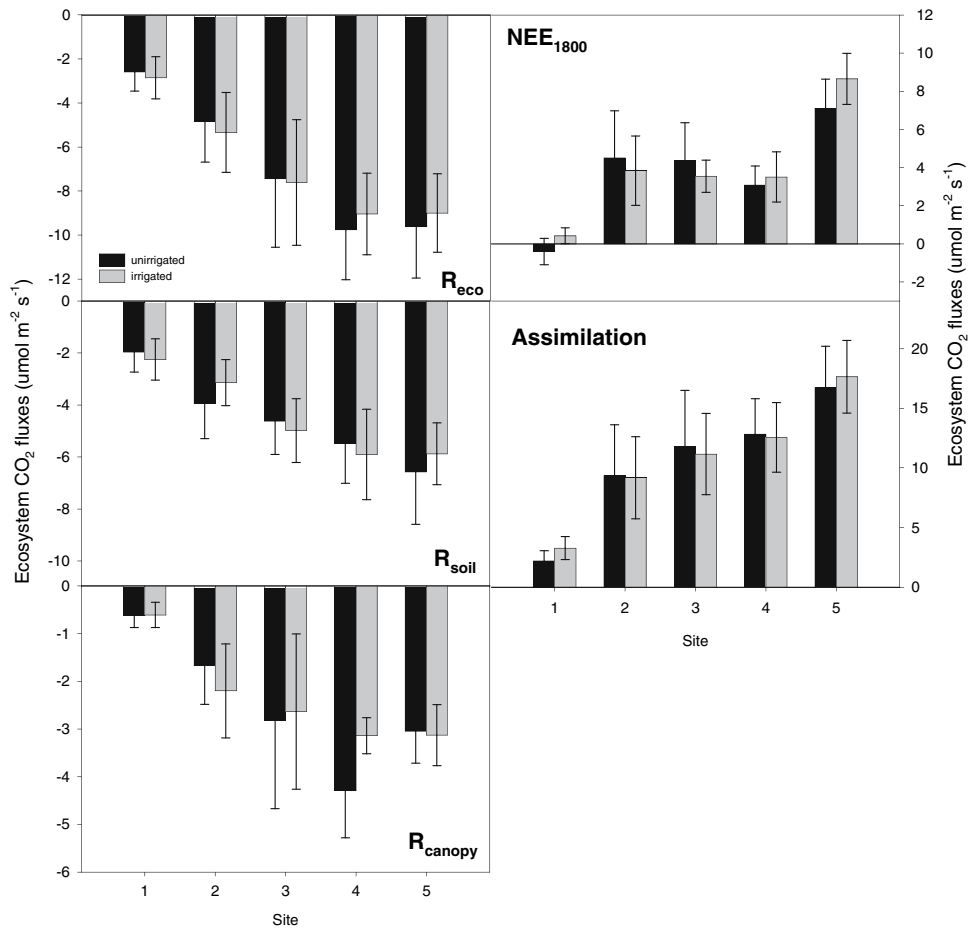


Fig. 3 Average net ecosystem CO₂ exchange (NEE₁₈₀₀), gross assimilation (Assimilation), and ecosystem (R_{eco}), soil (R_{soil}) and canopy (R_{canopy}) respiration for irrigated and unirrigated plots at each of the five study sites. Negative values are defined as CO₂ leaving the ecosystem, while positive values indicate

CO₂ taken up by the vegetation. Error bars indicate the standard error calculated from the measurements conducted over the course of the season at each site. Site 1 = Crystal Bench, Site 2 = Specimen Ridge, Site 3 = Canyon, Site 4 = Mammoth Hot Springs, Site 5 = Norris

and CO₂ fluxes when measuring ecosystem CO₂ exchange in grazed and ungrazed grassland across the landscape in Yellowstone National Park (USA) during the 2004 growing season. Strong correlations between these parameters were also detected in studies conducted in other grassland ecosystems (e.g., Meyers 2001; Flanagan et al. 2002; Xu and Baldocchi 2004; Kato et al. 2004; Flanagan and Johnson 2005; Li et al. 2005). Thus, changes in precipitation regimes, as a result of climate change that alter plant and soil processes (Laurenroth and Sala 1992; Knapp and Smith 2001), may have significant consequences on the amount of C stored and released from grassland ecosystems. However, given the characteristically large amount of spatial

variation in plant and soil properties in grassland habitat, ecosystem responses to changes in precipitation regimes may be heterogeneous. We found that increasing precipitation by 50% of the monthly 30-year average only had an effect on increasing CO₂ respiration (R_{eco}, R_{soil}, R_{canopy}) at the two driest sites (sites 1 and 2), and net CO₂ storage (NEE₁₈₀₀) at the driest (site 1) and the wettest site (site 5). The two sites with intermediate soil moisture, 3 and 4, were unaffected by supplemental water. Thus, CO₂ flows were water-limited at sites 1, 2 and 5, and non-limited at sites 3 and 4 during the average moisture conditions of the 2005 growing season.

Potential explanations for the differences in patterns of water-limitation across the Yellowstone landscape

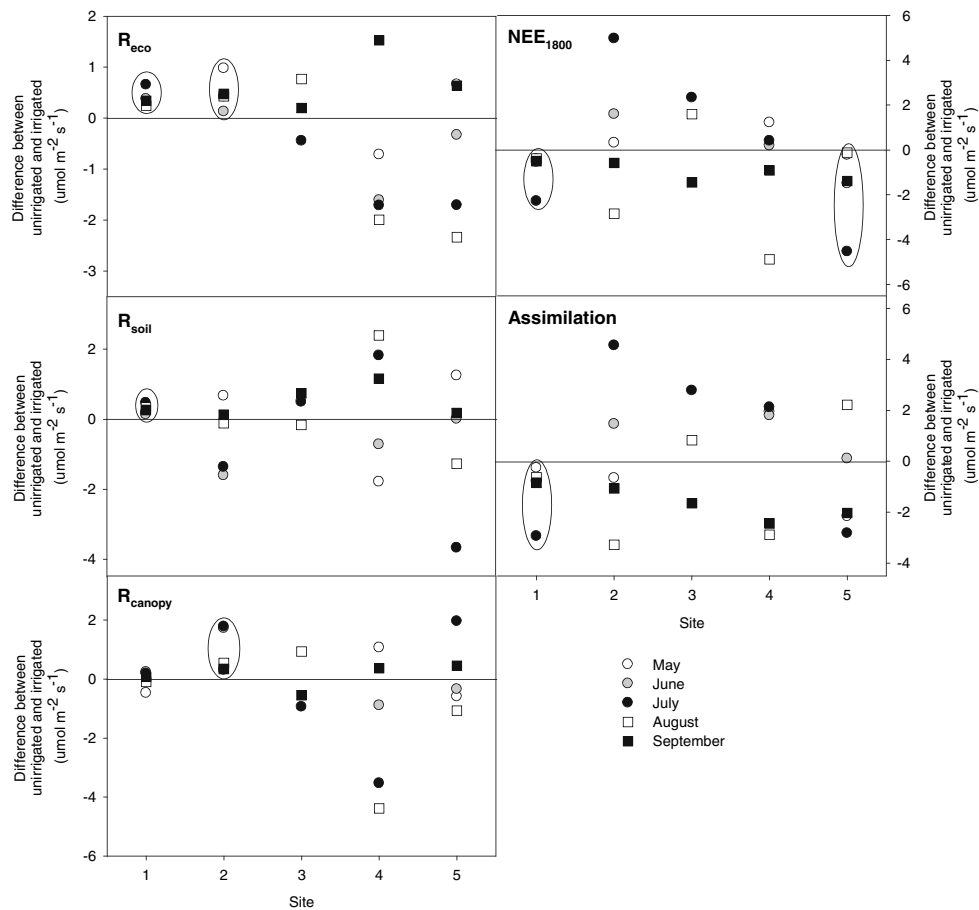


Fig. 4 Differences in net ecosystem CO₂ exchange (NEE_{1800}), gross assimilation (Assimilation), and ecosystem (R_{eco}), soil (R_{soil}) and canopy (R_{canopy}) respiration between the irrigated and unirrigated plots at the five study sites. For R_{eco} , R_{soil} , and R_{canopy} , positive differences indicate that fluxes were higher at the irrigated compared to the unirrigated plots. For NEE_{1800}

and Assimilation, negative values indicate higher rates at the irrigated plots. Ellipses denote significant treatment effects on ecosystem CO₂ fluxes at a specific site based on paired *t*-tests ($\alpha = 0.1$). Site 1 = Crystal Bench, Site 2 = Specimen Ridge, Site 3 = Canyon, Site 4 = Mammoth Hot Springs, Site 5 = Norris

could be related to differences in soil (e.g., chemical and physical properties, hydrology) and vegetation (e.g., water use efficiencies, total biomass, vegetation composition, density of plants) properties among the five locations (e.g., Silletti and Knapp 2001; Farmer et al. 2003). Irrigation increased soil moisture the most at sites 1 and 2, which had the sandiest soils among the sites. Sand-rich soils are known to have higher rates of percolation and lower water retention compared to clay rich soils (e.g., English et al. 2005). Consequently, the regular water additions in our study appeared to have especially helped to maintain higher soil moisture levels in grasslands where water from natural rainfall events likely drained quickly through the soil. The reason for finding a positive response of CO₂ fluxes to

water additions at the wettest (site 5), but not at the two sites (3 and 4) located in the middle of our gradient, is less clear. Potential explanations include the composition and structure of the vegetation growing at this site. The plant species growing at site 5, where the available resources were greatest among the sites, likely were stronger competitors than the dominating plant species found at sites 3 and 4 (Grime 1979). Consequently, these competition-adapted plants may have been more responsive to even small changes in resource availability. Further, the contrasting responses detected between sites 3/4 and site 5 could be related to differences in the number of individual plants growing per m² grassland (plant density; site 3: ~1,650 tillers m⁻²; site 4: ~2,050 tillers m⁻²; site 5:

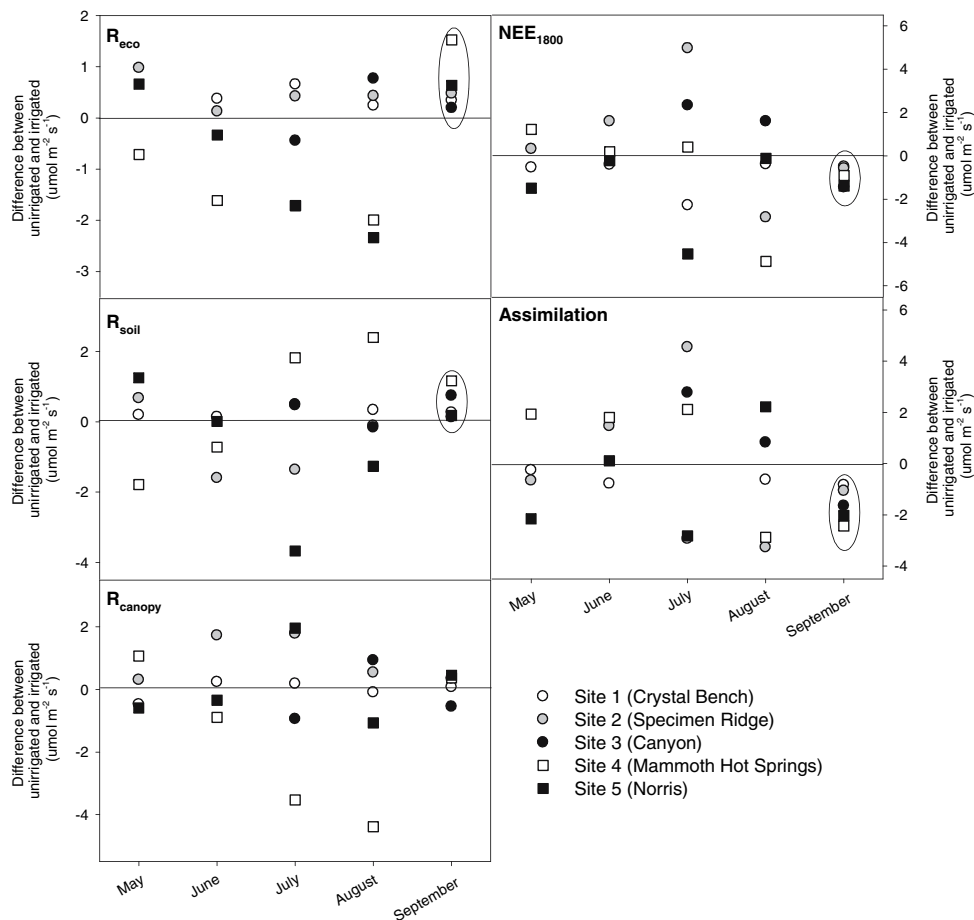


Fig. 5 Differences in net ecosystem CO_2 exchange (NEE_{1800}), gross assimilation (Assimilation), and ecosystem (R_{eco}), soil (R_{soil}) and canopy (R_{canopy}) respiration between the irrigated and unirrigated plots over the course of the growing season. For R_{eco} , R_{soil} , and R_{canopy} , positive differences indicate that

fluxes were higher at the irrigated compared to the unirrigated plots. For NEE_{1800} and Assimilation, negative values indicate higher values at the irrigated plots. Ellipses denote significant treatment effects on ecosystem CO_2 fluxes in a specific month, based on paired t -tests ($\alpha = 0.1$)

$\sim 2,700$ tillers m^{-2} ; unpublished data). Knapp and Smith (2001) suggested that vegetation types with low numbers of plants per area would respond less to increases in soil moisture compared to vegetation types with high density of plants (i.e., the meristem limitation hypothesis). Corroborating evidence for Knapp and Smith's (2001) hypothesis comes from Yahdjian and Sala (2006), who reported differences in the response of grassland to irrigation related to plant density when treating grassland with water additions after several years of experimental drought in Patagonia (Argentina). In their study, plant density decreased with increasing drought level, which resulted in a reduced response to subsequent water additions (Yahdjian and Sala 2006).

When considering the response in ecosystem CO_2 exchanges of all our sites during the growing season, we found no statistically significant difference in fluxes until late in the season. In September all the irrigated plots had significantly higher NEE_{1800} , Assimilation, R_{eco} and R_{soil} compared to the unirrigated plots. Consequently, in September, water-limitation occurred at all the sites. The higher fluxes at the irrigated plots late in the growing season likely originated from delayed senescence of plants at the irrigated compared to the unirrigated plots. Less dead biomass was observed on the irrigated than on the unirrigated plots (A. Risch, personal observations). Similar to our study, increased soil moisture led to longer plant survival and later peak biomass in pot

experiments (Mamolos et al. 2001; Yang et al. 2001). Irrigation-delayed senescence also was found in a field experiment with corn in New Zealand (Stone et al. 2001). Higher R_{soil} at the irrigated plots in September might have been caused by greater root and soil microbial respiration. For example, Liu and Li (2005) found increased root respiration under well watered compared to water-limited conditions in a greenhouse experiment with wheat. When irrigating the high arctic semi-desert of Greenland, Illeris et al. (2003) found the water addition led to higher soil microbial activity. A similar late-season increase in R_{eco} by irrigation, along with an absence of an early-season response, as in our study, was reported by Chimner and Welker (2005) in a mixed grass prairie in Wyoming, USA.

Our results showed a strong heterogenous response of grassland to water additions during a growing season of approximately average precipitation: the sites at the driest and wettest ends of the topo-edaphic gradient responded positively to irrigation, but no irrigation effect was detected at sites located in the middle of the gradient. At the dry end, the positive responses likely were associated with soil properties, while plant attributes may have caused the increases in ecosystem CO_2 fluxes at the wettest end of the gradient. The regular water additions also had an effect on ecosystem CO_2 fluxes at all sites in September, by delaying plant senescence and potentially increasing soil microbial activity. Thus, it is likely that years of above-average amounts of precipitation would lead to increases in net C gains late in the season, which in turn could have a positive effect on ecosystem C cycling in the following year. Further, if long-term trends in precipitation change, as predicted for many regions of North America (e.g., Cubasch et al. 2001), soil moisture gradients and the distribution of plant species will likely shift across the landscape. Such changes, in turn, could affect how this heterogeneous landscape responds to variable climate.

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