



Seasonal variation in carbon dioxide exchange over a Mediterranean annual grassland in California

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Abstract

Understanding how environmental variables affect the processes that regulate the carbon flux over grassland is critical for large-scale modeling research, since grasslands comprise almost one-third of the earth's natural vegetation. To address this issue, fluxes of CO₂ (F_c , flux toward the surface is negative) were measured over a Mediterranean, annual grassland in California, USA for 2 years with the eddy covariance method.

To interpret the biotic and abiotic factors that modulate F_c over the course of a year we decomposed net ecosystem CO₂ exchange into its constituent components, ecosystem respiration (R_{eco}) and gross primary production (GPP). Daytime R_{eco} was extrapolated from the relationship between temperature and nighttime F_c under high turbulent conditions. Then, GPP was estimated by subtracting daytime values of F_c from daytime estimates of R_{eco} .

Results show that most of carbon exchange, both photosynthesis and respiration, was limited to the wet season (typically from October to mid-May). Seasonal variations in GPP followed closely to changes in leaf area index, which in turn was governed by soil moisture, available sunlight and the timing of the last frost. In general, R_{eco} was an exponential function of soil temperature, but with season-dependent values of Q_{10} . The temperature-dependent respiration model failed immediately after rain events, when large pulses of R_{eco} were observed. Respiration pulses were especially notable during the dry season when the grass was dead and were the consequence of quickly stimulated microbial activity.

Integrated values of GPP, R_{eco} , and net ecosystem exchange (NEE) were 867, 735, and -132 g C m^{-2} , respectively, for the 2000–2001 season, and 729, 758, and 29 g C m^{-2} for the 2001–2002 season. Thus, the grassland was a moderate carbon sink during the first season and a weak carbon source during the second season. In contrast to a well-accepted view that annual production of grass is linearly correlated to precipitation, the large difference in GPP between the two seasons were not caused by the annual precipitation. Instead, a shorter growing season, due to late start of the rainy season, was mainly responsible for the lower GPP in the second season. Furthermore, relatively higher R_{eco} during the non-growing season occurred after a late spring rain. Thus, for this Mediterranean grassland, the timing of rain events had more impact than the total amount of precipitation on ecosystem GPP and NEE. This is because its growing season is in the cool and wet season when carbon uptake and respiration are usually limited by low temperature and sometimes frost, not by soil moisture.

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1. Introduction

Understanding how gross primary production (GPP) and ecosystem respiration (R_{eco}) respond to

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environmental variables and how these carbon flux components vary on daily, seasonal, annual and interannual temporal scales are prerequisites for understanding what controls and modulates net ecosystem exchange (NEE) of ecosystems and biomes (Law et al., 1999). Since carbon fluxes of biomes respond differently to environmental forcing variables and their perturbations, it is essential to characterize the physiological responses for each major biome.

It is also imperative to assess the relationship between GPP and canopy structure over a variety of ecosystems. Such data are needed to validate indices that are being evaluated with reflectance data measured by instruments on satellite-based remote sensing platforms—normalized difference vegetation index (NDVI)—to upscale carbon assimilation fluxes to regional, continental and global scales (Running et al., 1999; Turner et al., 1999; Myneni et al., 2002; Wylie et al., 2003).

At present, most long-term carbon flux studies have focused on various temperate conifer and broadleaved (deciduous and evergreen) forests, tropical and boreal forests. Only a few long-term carbon flux studies have been conducted over grasslands (Meyers, 2001; Flanagan et al., 2002; Suyker et al., 2003; Gilmanov et al., 2003), even though grassland is the largest of the four major natural biomes (Sims and Risser, 2000).

Grasslands growing in temperate continental areas are perennial and their growing season is limited to the warm summer. Consequently, temperature, precipitation, photosynthesis and respiration are in phase with one another. From a functional perspective, biogeographic conditions and natural selection have interacted to produce grasslands where a significant fraction of species use the C₄ photosynthetic pathway (Ham and Knapp, 1998; Kim and Verma, 1990; Ehleringer et al., 1997).

While both perennial and annual grasslands exist, the majority of carbon flux studies over grasslands have been restricted to continental regions. Mediterranean grasslands, in contrast, are functional during the winter and early spring and are dead during the summer, as temperature is out phase with precipitation (Bartolome, 1979; Chiariello, 1989). Consequently, the length of its growing season depends on the timing of the beginning and end of the rainy season. And because the growing season is during the cool, wet winter Mediterranean regions consist mainly of annual

C₃ species (Huenneke and Mooney, 1989; Ehleringer et al., 1997).

At the ecosystem scale, grasslands can be either net carbon sources or sinks. Depending on precipitation, more carbon uptake occurs during wet years and vice versa (Flanagan et al., 2002; Meyers, 2001; Suyker et al., 2003). These physiological results, based on the eddy covariance method, are consistent with numerous ecological studies, which have shown that above-ground net primary production (ANPP) of grasslands growing in the continental region of North America is correlated linearly with annual precipitation (Sims and Singh, 1978; Webb et al., 1978; Sala et al., 1988; Pardo et al., 1999; Lauenroth and Sala, 1992; Knapp and Smith, 2001).

Studies on the micrometeorological and ecophysiological control on the carbon exchange of annual grasslands in California are rare (Barbour and Minnich, 2000). Only Valentini et al. (1995) has published measurements of carbon and water use of Californian grassland, but their study period was not on a continuous basis for a full year. Furthermore, it was specific to grass growing on serpentine soil near the Pacific coast and in an urban air-shed area.

Several arguments can be forwarded to articulate a need for long-term carbon flux measurements over annual, C₃ grassland ecosystems in California. First, they are model ecosystems for studying the environmental perturbation of ecosystem carbon fluxes. Specifically, they experience modulations of temperature, rainfall, soil moisture and leaf area index across a wider range of environmental conditions than are observed over mesic and temperate ecosystems that are more commonly studied by sites across the AmeriFlux and FLUXNET networks (e.g. Baldocchi et al., 2001; Law et al., 2002). This unique feature provides us with the ability to define the light-, temperature- and soil moisture-response functions for carbon uptake and ecosystem respiration better. Secondly, because the grasslands experience distinct wet and dry periods they provide us with a better ability to quantify how carbon gain and water loss compete as the soil moisture pool is depleted.

In this paper, we report a 2-year carbon flux study over California grassland. The main objective is to quantify NEE of the grassland and its partitioning into GPP and R_{eco} using eddy covariance technique. Specifically, we examine the seasonal and interannual

variability in carbon exchange in term of biotic and environmental drivers. Three specific questions that are addressed with our data are: (1) How do the two major components of carbon balance (i.e. GPP and R_{eco}) respond to changes in grass phenology, soil moisture and temperature? (2) How does the seasonal distribution and total amount of precipitation affect GPP and R_{eco} ? (3) What is the interannual variation of NEE, GPP and R_{eco} of an annual grassland? Information on water and energy exchange from this grassland is reported separately (Baldocchi et al., 2003).

2. Materials and methods

2.1. Site description, climate, soil and grass species

The research site is a grazed grassland opening in a region of oak/grass woodland. It was established in October 2000 as part of the AmeriFlux network (Law et al., 2003). It is located on the foothills of the Sierra Nevada Mountains, and is about 35 km southeast of Sacramento (38°24.400 N, 120°57.044 W, and 129 m a.s.l.).

The soil is an Exchequer very rocky silt loam (Lithic xerorthents). It contains 30% sand, 57% silt and 13% clay. Its bulk density at surface layer (0–30 cm) is around $1.43 \pm 0.10 \text{ g cm}^{-3}$ ($n = 27$). Total nitrogen and carbon content of the soil were about 0.14 and 1.39%, respectively. The soil profile is about 0.5 m deep, and overlays fractured rock.

The climate at the site is Mediterranean type with clear days, high temperatures, and virtually no rainfall during the summer. In contrast, the winter is relatively cold and wet. The mean annual temperature was 16.2 °C during 2001 and 566 mm of precipitation fell. These values are close to climatic means, determined over 30 years from a nearby weather station at Ione, CA (mean air temperature is 16.3 °C and mean precipitation is 559 mm).

The grassland is dominated by cool-season C_3 annual species. More than 95% of species composition at the site, are *Brachypodium distachyon* L., *Hypochaeris glabra* L., *Trifolium dubium* Sibth., *Trifolium hirtum* All., *Dichelostemma volubile* A., and *Erodium botrys* Cav. The maximum grass height in the peak growth period (late April to early May) could reach up to $0.55 \pm 0.12 \text{ m}$ ($n = 25$).

2.2. Flux measurement

The flux of CO_2 over the grassland was measured continuously with an eddy covariance system. The fetch from all directions is more than 200 m. Calculations with a footprint model (Schmid, 1994) indicated that the fetch was well within the flux footprint. The flux system, which was mounted at 2 m above the ground, consisted of a 3D sonic anemometer (Model 1352, Gill Instruments Ltd., Lymington, England) and an open-path and fast response infrared gas analyzer (IRGA, Li 7500, Li-Cor Inc., Lincoln, NE, USA). The anemometer and the IRGA provide digital output of the fluctuations in three wind components (w , u , v), sonic temperature (T_s), water vapor (ρ_v) and CO_2 density (ρ_c). The raw data from each 30 min period were recorded at the rate of 10 Hz into separate files on a laptop computer. Spectra and co-spectra computations show that these sampling rates are adequate for measuring fluxes with this configuration.

The IRGA was swapped out every month with a newly calibrated one. The CO_2 signal of the IRGA was calibrated against gas mixtures in air that were referenced to standards prepared by NOAA's Climate Monitoring & Diagnostics Laboratory (NOAA/CMDL). The span for the water vapor was calibrated with a dew point generator (model Li-610, Li-Cor Inc., Lincoln, NE, USA). Zeros for both CO_2 and water vapor channels were calibrated with 99.99% nitrogen gas after passing through soda lime and magnesium perchlorate. Calibration results showed that the cumulative deviations for zero drift and span change for both CO_2 and water vapor channels over a period of one full year were less than 3 and 1%, respectively. Thus, shifts of zero and span over a month period can be considered as insignificant.

The eddy flux system was powered by eight 12 VDC deep cycle batteries that were charged by eight solar panels (Model SP75, Siemens) in wintertime and six panels in the rest time of a year. The system used approximately 3 A at 12 V.

2.3. Meteorology, soil and other parameter measurements

Along with the flux measurements with the eddy covariance technique, standard meteorological and soil parameters were measured continuously with an array

of sensors. Photosynthetic photon flux density, Q_p , reflected Q_p , and R_n were measured with a quantum sensor (Kipp and Zonen PAR-Lite, Delft, Holland) and a net radiometer at the height of 2.5 m (Kipp and Zonen, Delft, Holland), respectively. Precipitation was measured with a tipping-bucket rain gauge (Texas Electronics, Texas). Air temperature and relative humidity at the height of 2.5 m were measured with a shielded and aspirated sensor (HMP-35 A, Vaisala, Helsinki, Finland). Soil temperature at the depths of 0.02, 0.04, 0.08, 0.16 and 0.32 m were measured with three multiple-level thermocouple sensors. Soil volumetric water content was measured with frequency-domain reflectometer probe (ML2x, Delta-T Devices, Burwell, Cambridge, UK) at depths of 0.05, 0.10 and 0.20 m. Profiles of soil moisture (at the depth of 0–0.15, 0.15–0.30, 0.30–0.45, and 0.45–0.60 m) were also measured weekly with a time-domain reflectometers (Moisture Point, model 917, Environmental Sciences Inc., Vancouver, Canada) over much larger sample area. Soil heat flux was obtained by averaging the output of three heat flux plates (model HFP-01, Hukseflux Thermal Sensors, Delft). They were buried 0.01 m below the surface and were randomly placed within a few meters of the flux system. All channels from meteorological and soil sensors, except the rain gauge, were scanned every 5 s with data-loggers (CR10X or CR23X, Campbell Scientific Inc., Logan, UT, USA), and then 30 min mean data were stored. The 30 min mean data were retrieved by the laptop computer used for the eddy covariance measurement.

Leaf area index (LAI) was determined at intervals of 2–4 weeks. Grass was harvested from four sample plots (0.25 m × 0.25 m) in the prevailing direction of eddy covariance flux system. Then leaves were separated from the stem and their areas were measured with a leaf area meter (Li-Cor 3100, Lincoln, NE, USA).

2.4. Data processing and flux computation

Standard micrometeorological software was used to compute flux covariances from the raw data. Computations included spike removal, coordinate rotation, and application of standard gas laws. In practice, the analytical method used to measure CO₂ has an impact on the computation of the flux covariance. When CO₂ is

measured with an infrared spectrometer one samples molar density, ρ_c (moles per unit volume). In principle, changes in molar density can occur by adding molecules to or removing them from a controlled volume or by changing the size of the controlled volume, as is done when pressure, temperature and humidity change in the atmosphere. By measuring the eddy flux covariance in terms of molar density, the net flux density of CO₂ (F_c) across the atmosphere–biosphere interface can be expressed as

$$F_c = \overline{w\rho_c} = \overline{w'\rho'_c} + \overline{w}\overline{\rho_c} \quad (1)$$

The first term on the right-hand side of Eq. (1) is the covariance between the fluctuations of vertical velocity (w') and CO₂ density (ρ'_c). The second term is the product of the mean vertical velocity (\overline{w}) and CO₂ density ($\overline{\rho_c}$). The mean vertical velocity is non-zero and arises from air density fluctuations (Webb et al., 1980). The magnitude of \overline{w} is too small (<1 mm s⁻¹) to be detected by sonic anemometry, so it is usually computed on the basis of temperature (T) and humidity density (ρ_v) fluctuations using the Webb–Pearman–Leuning (1980) algorithm:

$$F_c = \overline{w'\rho'_c} + \frac{m_a}{m_v} \frac{\overline{\rho_c}}{\overline{\rho_a}} \overline{w'\rho'_v} + \left(1 + \frac{\overline{\rho_v}m_a}{\overline{\rho_a}m_v}\right) \frac{\overline{\rho_c}}{\overline{T}} \overline{w'T'} \quad (2)$$

Other variables in Eq. (2) are the molecular weights of air, m_a , and water vapor, m_v , dry air density, ρ_a . From Eq. (2), positive F_c represents the ecosystem losing carbon via respiration, whereas negative indicates the net carbon gain via photosynthesis.

To assess the accuracy of the eddy covariance measurements, we analyzed linear regressions between the sum of latent heat (LE), sensible heat (H) and soil heat flux (G) vs net radiation (R_n):

$$LE + H + G = a_0 + aR_n \quad (3)$$

At the beginning of this study, we were able to attain a high degree of closure of the surface energy balance. During 2000, the intercept, slope and r^2 were 5.5, 0.94 and 0.93, respectively. As the experiment continued into 2001 and 2002 we observed degradation in the degree of energy balance closure. However, we did not observe any systematic changes from upper envelopes of seasonal trends in net radiation, sensible heat flux and latent heat flux. Close inspection of the

data showed that the long-term degradation in closing the surface energy balance was an artifact of measuring soil heat flux within a cow-proof enclosure. The enclosure had more plant matter and detritus than the surrounding paddock and this caused the magnitude and amplitude of soil heat flux to lower with time. When the change in heat storage term above soil heat flux plates was taken into account, the regression, defined by Eq. (3), exceeded 0.90 for both year 2001 and 2002 (Baldocchi et al., 2003).

2.5. Methods for gap filling, low U^* correction and estimating GPP and R_{eco}

For long-term and continuous field measurements, missing data is unavoidable due to malfunction of the instrumentations or power failure. From the 2-year measurements, we missed about 17.3% of data in 2001 and 8.2% in 2002. Another 13.5% of data points were rejected in 2001 and 10.9% in 2002 due to the bad statistics of certain turbulent variables, e.g. Reynolds stress, skewness and kurtosis of w and ρ_c signals, or off normal scales of T_s , ρ_v , or ρ_c . In order to obtain the information on the annual sum of the carbon flux data, we used the following procedure to fill missing and bad data. For small blocks (less than an hour) of missing data, simple interpolation method was used. Larger blocks of missing data during the growing season were filled by using a rectangular hyperbolic light-response function (Falge et al., 2001):

$$F_c = \frac{F_{\text{max}} \alpha Q_p}{\alpha Q_p + F_{\text{max}}} + R_{\text{eco}} \quad (4)$$

where Q_p ($\mu\text{mol m}^{-2} \text{s}^{-1}$) is incident photosynthetically active radiation, F_{max} ($\mu\text{mol m}^{-2} \text{s}^{-1}$) the maximum CO_2 flux at infinite light, α the apparent quantum yield, and R_{eco} the respiration from soil and plants; we tried other types of light-response functions, and results showed those functions did not provide a better fit to Eq. (4).

Nighttime missing data were filled by using the following exponential relationship between F_c from periods of high turbulence when the friction velocity (U^*) was higher than 0.1 m s^{-1} and soil temperature at depth of 0.04 m (T_{soil}):

$$F_c = b_0 \exp(bT_{\text{soil}}) \quad (5)$$

where b_0 and b are two empirical coefficients, from which Q_{10} can be estimated as

$$Q_{10} = \exp(10b) \quad (6)$$

The reason we chose the soil temperature at the depth of 0.04 m in this analysis is that it reduced the sources of variation most for regression of Eq. (5) than that at other depths. The soil temperature function (Eq. (5)) was also used to extrapolate daytime ecosystem respiration. During the non-growing summer period, all missing, rejected data were filled or replaced by using Eq. (5).

We also corrected F_c for the CO_2 storage term before gap filling to avoid double counting (Aubinet et al., 2002). Since we did not have CO_2 profile data, the CO_2 storage term was estimated only based on the one point CO_2 concentrations from the open-path IRGA of the eddy covariance system. This approach might be an acceptable estimation, since the height of our flux system was only at 2 m high, and because the storage term sums to zero on daily and annual time scales (Baldocchi, 2003).

GPP was obtained by subtracting R_{eco} from NEE (Barr et al., 2002; Flanagan et al., 2002), which required us to evaluate R_{eco} for day and nighttime periods. Because of the strong seasonality in grass phenology and in soil moisture, we evaluated the respiration temperature coefficient, Q_{10} , using short, 5-daytime windows, for the fast transition periods. During the non-growing summer season, when soil moisture changed little, we used 1-month time window.

3. Results and discussion

3.1. Weather conditions and canopy structure

In order to study the carbon exchange between the grassland and the atmosphere, it is necessary to first understand the seasonality of key environmental variables. Seasonal variations in daily maximum (T_{max}) and minimum (T_{min}) air temperature, daily precipitation (PPT), volumetric soil water content (θ_v), reflectivity of Q_p (γ), and LAI are presented in Fig. 1.

There were pronounced variations in temperature during the growing season. T_{max} ranged from moderate levels ($\sim 10^\circ\text{C}$) in the winter and early spring to extreme levels ($>40^\circ\text{C}$) by early summer during the

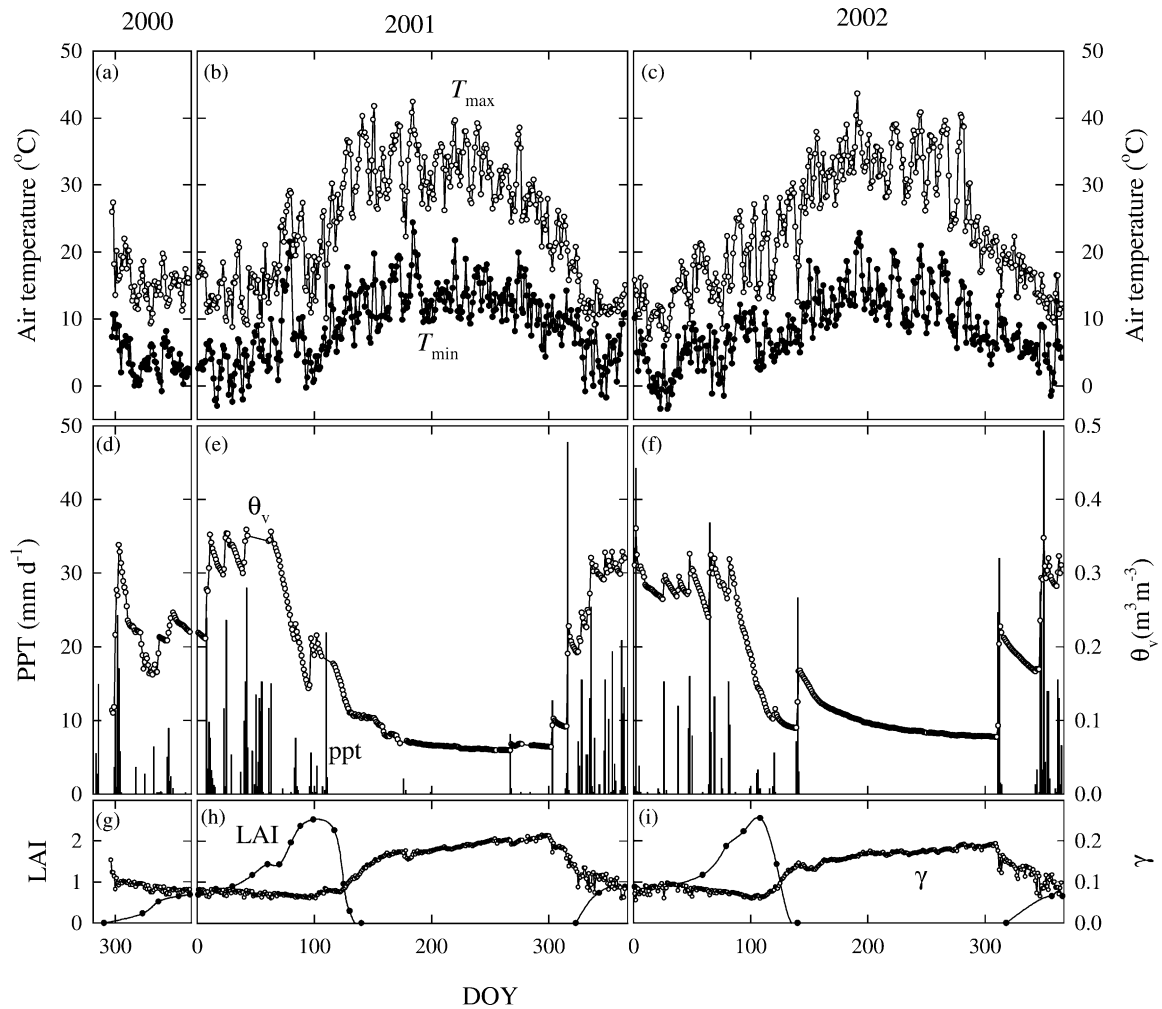


Fig. 1. Seasonal variation of major meteorological variables over the course of the study. Data shown in the top three panels ((a)–(c)) are maximum (T_{max}) and minimum air temperature (T_{min}). Daily total precipitation (PPT) and averaged volumetric soil water moisture (θ_v) from the depth of 0.05, 0.10, and 0.20 m are shown in the middle three panels ((d)–(f)). Leaf area index (LAI) and Q_p albedo (γ) were also shown ((g)–(i)). γ was Q_p -weighted averaged when PAR > 100 $\mu\text{mol m}^{-2} \text{s}^{-1}$.

grass senescence period. The wide temperature range encountered by the ecosystem over the course of a day is also worth noting. In summer temperature it was well over 25 °C. Even in winter and early spring it could be as large as 15 °C (Fig. 1b and c).

Due to the Mediterranean climate, about 98% of precipitation fell during the wet seasons (i.e. from October to April). Annual precipitation during 2001 and 2002 was 567 and 494 mm, respectively. The precipitation in both years was not significantly different from the normal, which was 559 mm. Winter rainfall

provided enough moisture to recharge the soil profile. During the wet season, θ_v (weighted-average from 0.05, 0.10, and 0.20 m) was around 0.30 m³ m⁻³ or higher. When the wet season was over, θ_v declined within a month to a low steady state value in the range of 0.07–0.10 m³ m⁻³ (Fig. 1e and f). Vigorous growth of grass, combined with a shallow soil profile, relatively light soil texture, and high evaporative demand, contributed this rapid depletion of soil moisture. This large seasonal variation in θ_v , along with the wide range of diel and seasonal temperature, provides us a

unique opportunity to better define temperature- and moisture-response curves of carbon flux of the ecosystem.

Timing of rainfall is also important. In the second season there was a large storm just after the grass senescence; the storm produced 37 mm precipitation over the period of 3 days (19–21 May 2002). Because transpiration ceased after the grass died and soil evaporation was chronically low, θ_v was consistently $0.02\text{--}0.04\text{ m}^3\text{ m}^{-3}$ higher during the summer non-growing season in 2002 than in the previous year. This had a great impact on the soil respiration, as we will demonstrate below.

Because of the shallow rooting depth, the grass phenology closely followed soil moisture, which in turn depended on the rainfall. Grass seed germination normally occurs in the fall, 1 week after a major rain event with total precipitation of at least 15 mm (Bartolome, 1979; Chiariello, 1989). One such rain event occurred on 10 October for the 2000–2001 growing season, 1 month earlier as compared to 11 November for the second season. As a result, the grass germinated almost 1 month earlier in the first season. After germination, due to low soil and air temperature and occasional frosts, the grass underwent a period of slow vegetative growth in the wintertime as we can see from LAI data in both seasons (Fig. 1g–i). In the spring, warming temperatures, longer day length, and ample soil moisture accelerated the grass growth to a peak period with the maximum LAI of around 2.5 for the two seasons. The peak growth period did not last very long. Shortly after the wet season was over and most of the available soil moisture was utilized, the grass quickly senesced—around DOY140 for both seasons—leaving dormant grass seeds to transmit biological activity across the dry and hot summer.

Fig. 1 also illustrates the seasonal variations in Q_p albedo (γ), which was determined with an upward- and downward-facing quantum sensors. To avoid low solar angles when γ was not truly representative of the canopy surface, γ data were Q_p -weighted average when incident Q_p values were higher than $100\text{ }\mu\text{mol m}^{-2}\text{ s}^{-1}$. The seasonal change in γ was almost like the mirror image of variations in LAI, showing that γ decreased gradually as canopy size developed. Whereas during the fast senesced phase in the late spring when LAI plummeted to zero in just 3 weeks, γ jumped up correspondingly. Such strong

negative relationship between γ and LAI suggests that γ could be used as an alternative parameter to quantify the canopy activity when LAI information is not available. In fact, a study from a central Oklahoma tallgrass prairie showed that sometimes γ was even a better parameter than LAI to describe transitional periods of vegetation activity, including the end of peak growth, the entire pre-grow and post-burn periods (Burba and Verma, 2001).

3.2. Response of nighttime F_c to temperature

As we described above, nighttime gap filling were based on the soil temperature function (i.e. Q_{10} relation, Eq. (5)) of the R_{eco} obtained under strong turbulence conditions ($U^* > 0.1\text{ m s}^{-1}$). To avoid the confounding effects of grass phenology and soil moisture on the temperature function (Luo et al., 1996; Xu and Qi, 2001a; Flanagan et al., 2002), we developed a specific response curve for different time windows, sometimes as short as 5 days, during which both LAI and θ_v changed little. Fig. 2a shows three examples of temperature function from DOY129 to DOY139 (late senescence), DOY180 to DOY230 (non-growing season), and DOY347 to DOY365 (winter, LAI < 1.0) in 2001. The exponential function of Eq. (5) described the relationships between soil temperature at 0.04 m and R_{eco} very well. From Eq. (6), Q_{10} was estimated to be 2.5, 2.2, and 2.1 for the three examples, respectively.

High Q_{10} values from the present study were observed in wet season when grass was very active, whereas low Q_{10} values were mostly from non-growing summertime, a reflection of the different temperature sensitivities for autotrophic and heterotrophic respiration and the turnover times of the multiple carbon pools. High temperature sensitivity may include the direct physiological effect of temperature on root and microbial activities and the indirect effect related to photosynthetic assimilation and carbon allocation to roots (Davidson et al., 1998). Evidence for the indirect effect of photosynthesis on autotrophic respiration comes from a series of recent studies (Bremer et al., 1998; Höglberg et al., 2001; Bowling et al., 2002). During the dry season, the main portion of ecosystem respiration comes from the more recalcitrant carbon material, which has been shown to have low temperature sensitivity (Liski et al., 1999). Thus, in modeling

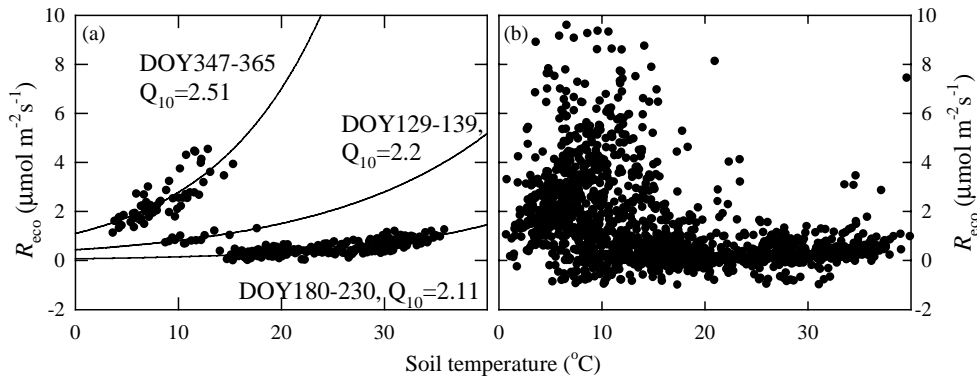


Fig. 2. Response of ecosystem respiration (R_{eco}) to change in soil temperature at the depth of 4 cm. Data were from 2000 to 2001 season, and were 3 h binned. Data in (a) were three examples from different growth periods each with similar LAI and soil moisture content. While in (b), data across the whole season were presented.

long-term ecosystem respiration, one should account for how Q_{10} varies over the season with changes in soil moisture, temperature and phenology.

The measured values of Q_{10} were close to the mean value deduced from a global survey of soil respiration (Raich and Schlesinger, 1992). A wider variation in Q_{10} values has been reported in the literature as soil moisture drops (Davidson et al., 1998; Xu and Qi, 2001a) or as soil temperature increases (Tjoelker et al., 2001). For example, Xu and Qi (2001b) reported that Q_{10} varied from 1.0 to 2.2 from a study conducted at a Sierra Nevada forest. Davidson et al. (1998) reported that Q_{10} for soil respiration ranged from 3.4 to 5.6 for different study sites in a temperate mixed hardwood forest. Considering variations in soil moisture, Reichstein et al. (2002b) reported a significant decline in Q_{10} (from 0.5 to 2.6) when the soil moisture dropped for three Mediterranean evergreen sites.

When R_{eco} was plotted against soil temperature over the whole season (Fig. 2b), no single temperature function was found to describe the variations in R_{eco} . The maximum R_{eco} occurred when the soil temperature was around 7–13 °C, soil moisture was ample, and the grass was experiencing its peak growth period. R_{eco} declined to less than $1.0 \mu\text{mol m}^{-2} \text{s}^{-1}$ during the non-growing season when soil temperature was above 15 °C and θ_v was less than $0.10 \text{ m}^3 \text{ m}^{-3}$. These results clearly indicates that the potential to enhance ecosystem respiration, by temperature during the dry season, is countered the inhibition due to severe drought.

3.3. Response of daytime F_c to incident radiation

Because photosynthesis is driven by light, we first examine how F_c responds to changes in Q_p . Because the structure and function of the grassland changes appreciably over the year we examine this relation for four different phenoseasons (Fig. 3). In general, the rectangular hyperbolic function can be used to describe the relationship between Q_p and F_c with success. Except for the fourth period, which was from the senescent phase (Fig. 3d), no clear light saturation of F_c was observed, even at LAI of less than 1.0. Based on the statistical analysis by using Eq. (4), the regression coefficients indicated that changes in Q_p accounted for more than 80% of the variations in F_c . The F_{max} for the first three growing periods (Fig. 3a–c) was -10.8 , -24.6 , and $-40.2 \mu\text{mol m}^{-2} \text{s}^{-1}$, respectively, and α was 0.014, 0.023, and 0.024. The quantum yield was well within the range of published data for C_3 grasses (Ruimy et al., 1995), and was very close to the value from other eddy covariance studies over temperate C_3 grassland (e.g. Flanagan et al., 2002). The low α at early season was most likely due to small canopy size, low temperature, and immature leaves. At the end of the senescence phase (DOY130–140, Fig. 3d), F_c increased along with Q_p in the early morning, and then showed a trend of decrease as Q_p further increased. This result is probably due to enhanced ecosystem respiration with increasing in temperature.

Seasonal changes in the diurnal patterns of CO_2 exchange can provide insights on Q_p and LAI interact

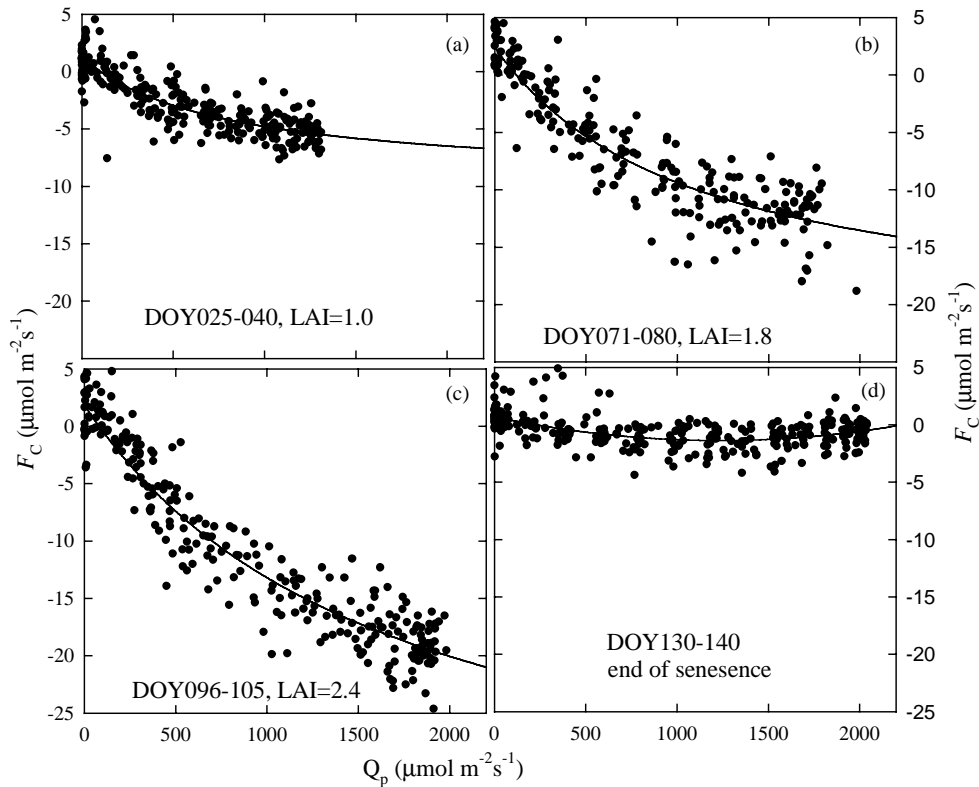


Fig. 3. Examples of light-response curves of CO₂ flux (F_c) at different growth stages from 2001 to 2002 growing season. Fitted curves are rectangular hyperbolic as described in Eq. (4), except in (d) which is quadratic polynomial. The growth stage and its leaf area (LAI) are also given.

to control photosynthesis and respiration. Mean diurnal courses of F_c and Q_p from different growth period are presented in Fig. 4, to illustrate this point; data were binned from five consecutive days to reduce the sampling error. Both examples were from sunny days with ample soil moisture. One was from wintertime during DOY031–035 in 2001 when LAI was 1.0, and the other from the peak growth period during DOY101–105 in 2002 with LAI of 2.2. The diurnal course of F_c depended mainly on LAI and Q_p , showing a similar temporal pattern to the Q_p curves. The maximum F_c for the two cited periods, -9.0 and $-19.5 \mu\text{mol m}^{-2} \text{s}^{-1}$, respectively, was centered around noontime. There was no consistent difference in the relationship between F_c and Q_p between morning and afternoon, indicating unlimited soil moisture conditions for grass carbon uptake. The observed values of F_c for this C₃ annual grassland was in the mid-

dle range of the values reported from other grassland studies. For example, Valentini et al. (1995) observed the maximum rate of CO₂ uptake to range between -6 and $-8 \mu\text{mol m}^{-2} \text{s}^{-1}$ over a serpentine grassland in California. In contrast, much higher maximum rates of CO₂ uptake (between -30 and $-40 \mu\text{mol m}^{-2} \text{s}^{-1}$) have been reported from more productive perennial grasslands, which contain C₄ species (Kim and Verma, 1990; Suyker and Verma, 2001; Dugas et al., 1999). Also noted from Fig. 4 is that nighttime R_{eco} was much higher for the peak growth stage than for the early season, reflecting importance of photosynthetic activity on the ecosystem respiration.

3.4. GPP in relation to LAI and R_{eco}

Over the course of the growing season, day-to-day variations in GPP, on sunny days, were highly

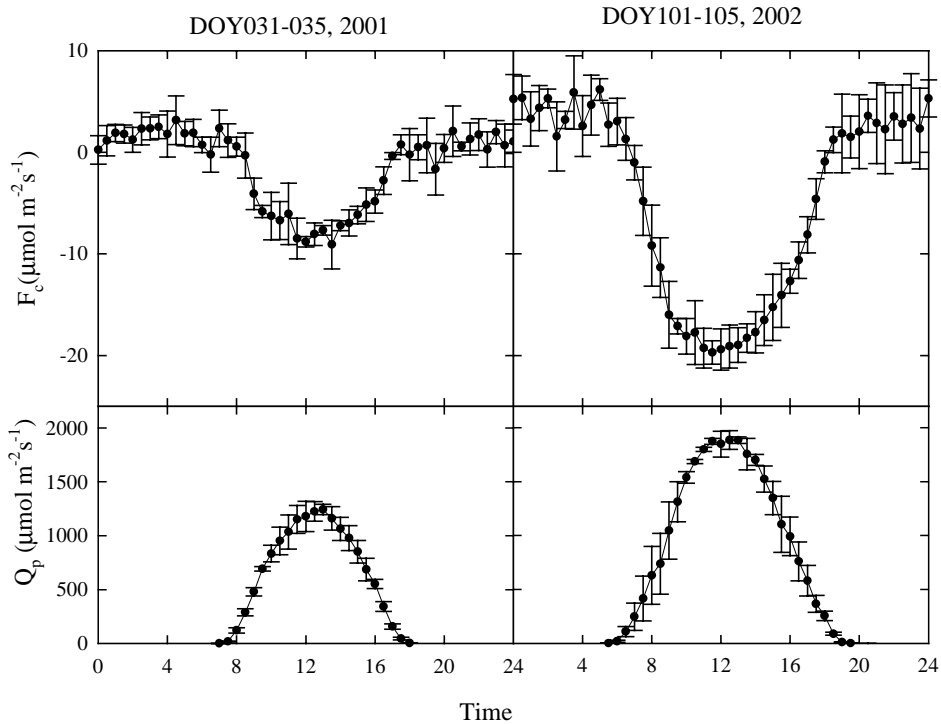


Fig. 4. Examples of 5-day binned diurnal variations in CO_2 flux (F_c) and photosynthetic active radiation (Q_p) growth periods. DOY031–035, 2001 and DOY101–105, 2002. LAI was around 1.0 and 2.2, respectively. Error bars represent the standard deviation.

correlated with variations in LAI (Fig. 5). For this annual grassland, over 84% of the variance in GPP was explained by changes in LAI. The remaining 16% of the variance was due to variations in other weather/soil variables, such as vapor pressure deficit, temperature, direct and diffuse radiation and soil moisture. In general, GPP increased by about 3.9 g C m^{-2} per day for each incremental increase in LAI. The linear relationship we observed is consistent with other grassland studies (Saigusa et al., 1998; Flanagan et al., 2002), but the slope we report is smaller than the one reported by Flanagan et al. (2002) for a continental grassland ($7\text{--}9 \text{ g C m}^{-2}$ per day per LAI).

Evidence exists in the literature that plant respiration can be scaled from plant photosynthesis, but only a few studies have presented information on GPP and ecosystem respiration (Law et al., 2002). When R_{eco} was plotted against GPP, we found that there was a strong linear relation ($r^2 = 0.78$, Fig. 6). This result indicates that R_{eco} was more closely related to the canopy photosynthetic activity than to tempera-

ture. These results are in agreement with a number of recent studies that have demonstrated a close linkage between the photosynthesis activity and respiration. For example, based on carbon flux data from 18 sites across European forests, Janssens et al. (2001) found that productivity of forests overshadows temperature in determining soil and ecosystem respiration. A study by Högberg et al. (2001) in a boreal pine forest in Sweden showed that up to 37% decrease in soil respiration was detected within 5 days after the stem bark of pine trees was girdled. Thus, the exponential function for ecosystem respiration (Eq. (5)) holds for a limited time period when LAI and soil moisture are similar. Therefore, in simulating R_{eco} across the whole season, the impact of canopy photosynthesis activity must be taken into account (Janssens et al., 2001). Most recent results have shown that there is a time lag for R_{eco} to respond to the change in photosynthesis (Ekbald and Högberg, 2001; Bowling et al., 2002), making it more challenging to model ecosystem respiration.

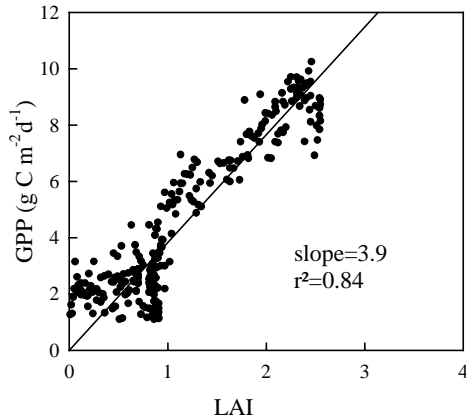


Fig. 5. The relationship of daily total gross primary production (GPP) and leaf area index (LAI). Data were obtained from the whole study period. GPP data from sunny days only and the linear regression was forced to pass through zero.

3.5. Influence of rain events on R_{eco}

We observed large pulses of R_{eco} immediately after individual rain events, especially during the summer dry period, when the grass was dead. Examples from year 2001 are presented in Fig. 7. The first rain event occurred late evening on DOY267 and lasted

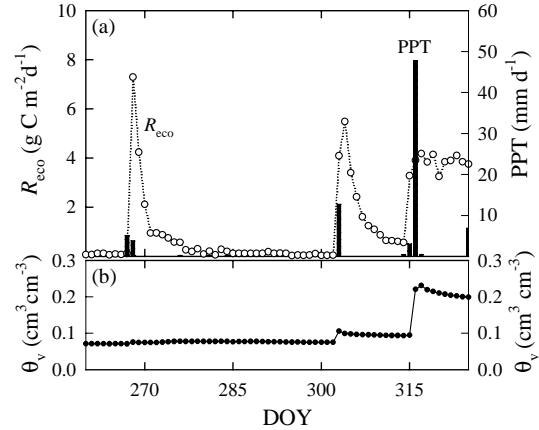


Fig. 7. Examples of influence of rain events on the ecosystem respiration (R_{eco}) from year 2001. Data in (a) are the daily total R_{eco} and precipitation (PPT) and in (b) are volumetric soil moisture content (θ_v) as averaged from the depth of 0.05, 0.10 and 0.20 m.

until early in the morning on the next day. The total precipitation was only 8.9 mm, and soil moisture did not even show any detectable increase (Fig. 7b). R_{eco} on DOY268 suddenly increased to 7.29 g C m^{-2} per day from the background of 0.12 g C m^{-2} per day on the previous day. Then in just 3 days, R_{eco} plummeted to 0.95 g C m^{-2} per day on DOY271. After the second rain event (12.7 mm rainfall), the R_{eco} again greatly increased from 0.05 g C m^{-2} per day on DOY302 to 5.49 g C m^{-2} per day on DOY304. Then, R_{eco} showed an exponential decrease with time, which was most likely associated with a fast decrease in soil moisture, as soil evaporation is a function of the inverse of the square root of time after a wetting event (Denmead, 1984). With the third rain event, which occurred on DOY315–316 with total precipitation of 51.8 mm, R_{eco} increased to 3.28 g C m^{-2} per day and did not show the sharp decrease as the previous two rain events. This could be due to relative large amount of precipitation for that event as shown with consistently high θ_v . It is worth noting that the peak value of R_{eco} after each rain event showed a gradual decrease. We attributed this observation to the facts that soil temperature became colder and labile carbon pool size at soil surface became smaller.

The sustained rates of respiration after rain were not due to prolonged physical displacement of air by the rain. Both laboratory experiments (Birch, 1958;

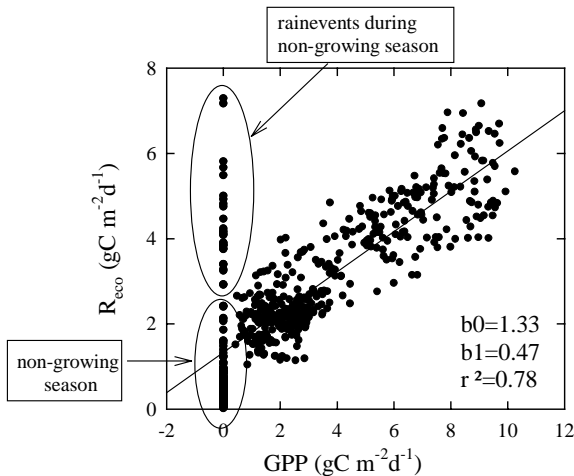


Fig. 6. The linear relation between daily gross primary production (GPP) and ecosystem respiration (R_{eco}). Data were compiled from the whole study period. Data points from rain events and non-growing seasons were excluded in the linear regression analysis.

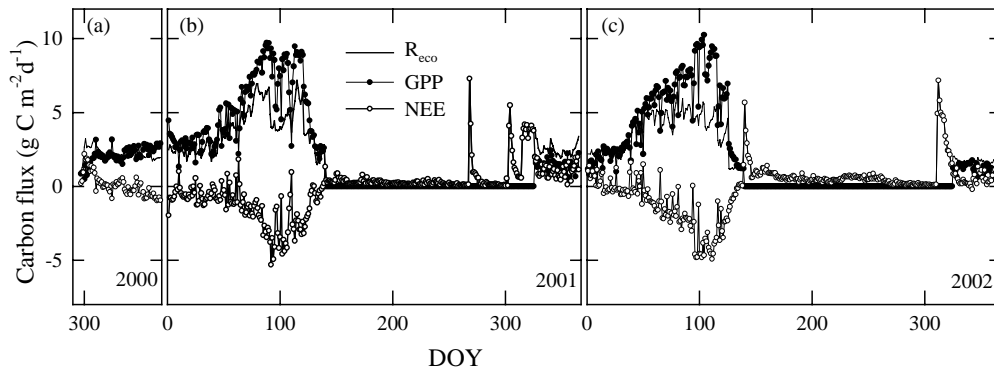


Fig. 8. Seasonal pattern of daily total ecosystem respiration (R_{eco}), gross primary production (GPP), and net ecosystem exchange (NEE) over the course of this study from DOY296 (22 October 2000) to the end of the year 2002.

Griffiths and Birch, 1961; Orchard and Cook, 1983) and field measurements (Liu et al., 2002; Rey et al., 2002) show that microbial respiration is able to turn on quickly (within hours) with the addition of water, and respiration rate ceases as soon as the soil layer dries.

3.6. Seasonal variations and cumulative GPP, R_{eco} , and NEE

Fig. 8 illustrates the seasonal variations in daily GPP, R_{eco} , and NEE over the course of this study. Due to the earlier rain in the winter of 2000, the ecosystem started uptaking carbon around DOY297 and showed a slight increase over the wintertime. R_{eco} was almost unchanged after grass germination, and was very close to GPP, resulting in NEE close to zero. While in the winter of 2001, the grass germinated about 1 month later as compared with the previous year because of late start of the rain. As a result, GPP was consistently lower than R_{eco} , causing positive NEE for the whole winter. In the spring, 2 years showed quite similar pattern of seasonal variations in GPP, R_{eco} , and NEE. Both GPP and R_{eco} gradually increased in January and February, and NEE was slightly negative. Then as the temperature warmed up and day length increased, GPP and R_{eco} accelerated in March and April, but with GPP at higher rate, making the ecosystem a strong carbon sink. The daily maximum NEE reached up to -4.8 g C m^{-2} per day. As the soil dried out in early summer, GPP and R_{eco} plummeted to near zero around DOY140. In summer, after grass senescence, the grassland continuously lost carbon via soil respira-

tion, but at very low rate due to the low soil moisture content.

We observed quite different rates of R_{eco} in the summer between the 2 years. R_{eco} in 2001 was in the range of $0.1\text{--}0.3 \text{ g m}^{-2}$ per day, as compared to that of $0.3\text{--}1.2 \text{ g m}^{-2}$ per day during the second season (Fig. 8b and c). This contrast in R_{eco} was probably caused by the difference in soil moisture. As already described in Fig. 1f, just after grass senescence there was a large storm with a total amount of 37 mm rainfall occurred during DOY139–141 2002. This late spring rain was a source of moisture left in the soil profile. It allowed microbes to decompose carbon substrate at a higher rate and caused the ecosystem to lose more carbon during the dry summer as compared with the previous year. This event can be a source of interannual variability in NEE, as will discuss below.

The maximum values of GPP, R_{eco} , and NEE from the two seasons were very similar, which were around 10.1 , 6.5 , and -4.8 g m^{-2} per day, respectively. For comparison, we cite a study conducted over a temperate, C_3 grassland near Alberta, Canada. Flanagan et al. (2002) reported similar seasonal maximums of 8.2 , 4.0 and -5.0 for GPP, R_{eco} , and NEE during the wet year. Higher values of maximum daily NEE have been reported in other studies of tallgrass, prairies in Kansas and Oklahoma; i.e. -6.3 g C m^{-2} per day by Kim et al. (1992), -7.6 g C m^{-2} per day by Dugas et al. (1999), and -6.0 g C m^{-2} per day by Suyker and Verma (2001). Those higher values may be attributed to larger canopy size and the dominant warm-season C_4 species.

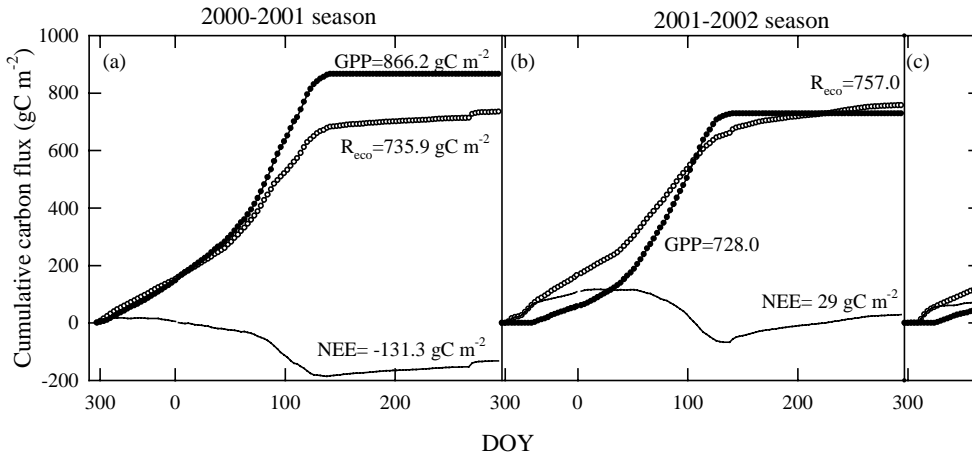


Fig. 9. Cumulative gross primary production (GPP), ecosystem respiration (R_{eco}), and net ecosystem exchange (NEE) over the two seasons.

The cumulative GPP, R_{eco} and NEE over the two seasons (2000–2001 and 2001–2002) were presented in Fig. 9. Since the growing season for the grass is across two calendar years, we did not compute cumulative GPP, R_{eco} and NEE over the calendar year. Instead, we computed those cumulative parameters based on the growing season of the grass, which is defined as the period from the first major rain event that produced enough precipitation (>15 mm) for the grass seeds germination to such similar rain event in the next year. As shown in Fig. 9, GPP, R_{eco} and NEE were 867, 735, and -132 g C m^{-2} for 2000–2001 season, and 729, 758 and 29 g C m^{-2} for 2001–2002 season, respectively. Based on the cumulative NEE data, the grassland was a moderate carbon sink for the first season, while it is a weak source for the second season. As compared to the second season, the ecosystem fixed more carbon, yet respired less during the first season. To better understand this intriguing finding,

we divided each grass season into three periods, winter growth, spring growth, and summer non-growth periods (Table 1). Winter growth period was defined as the time from the first major rain event to the 31 December. Spring growth period was defined as the time from 1 January to the senescence of the grass, summer non-growth period from the grass senescence to the beginning of the next season. During the winter period of the first season, due to the earlier start of the grass germination, GPP was 148 g C m^{-2} , as compared that of only 58 g C m^{-2} in the same period of the second season. The ecosystem fixed almost 90 g C m^{-2} more carbon than that in the second season, even though precipitation was much less in the first season. Despite the large difference in GPP, the total respiration was comparable for the two seasons (153 and 166 g C m^{-2}). During the spring growth period, both GPP and R_{eco} between the two seasons did not show large difference (Table 1). But during the

Table 1
Comparison of precipitation (ppt), ecosystem respiration (R_{eco}), gross primary production (GPP), and net ecosystem carbon exchange (NEE) at different growth periods for the two seasons

Growth period	2000	2001			2002			2000–2001 season	2001–2002 season	Over the whole course of study
	winter	Spring	Summer	Winter	Spring	Summer	Winter			
ppt (mm)	120	311		255	255		239	431	510	1180
R_{eco} (g C m^{-2})	153	528	54	166	498	94	124	735	758	1617
GPP (g C m^{-2})	148	719		58	671		49	867	729	1645
NEE (g C m^{-2})	5	-191	54	108	-173	94	75	-132	29	-28

summer periods, quite different results were observed. Because of late spring rain in the second season after the grass senescence, the soil moisture content was slightly higher than the previous year. As a result of better soil moisture condition, more ecosystem respiration with total carbon loss of 94 g C m^{-2} occurred, as compared to that of 54 g C m^{-2} during the first season. So for the second season late start of growing season combined with better soil moisture in the summer made the ecosystem a weak carbon source. Overall, the system lost 29 g C m^{-2} for the second season, whereas it gained 132 g C m^{-2} carbon in the first season. Across the whole study period from October 2000 to the end of 2002, the total carbon uptake was slightly more than the carbon loss via respiration (Table 1).

In the literature, it is well established that annual grassland productivity (normally based on ANPP) is well correlated with annual precipitation across the North American grasslands (Sims and Singh, 1978; Sala et al., 1988). On the temporal level, grassland ANPP has also been shown to have the strongest and positive response to interannual variations in precipitation (Knapp and Smith, 2001). Furthermore, carbon flux measurements with eddy covariance technique also support this well-established view. For examples, from a 3-year eddy covariance measurement over northern temperate grassland in Canada, Flanagan et al. (2002) found that GPP and aboveground biomass production were positively correlated well with annual precipitation. Another flux study conducted over a tallgrass prairie in Oklahoma showed that precipitation is a dominating factor in controlling the NEE (Suyker et al., 2003). From the present study, even with only two-season data, one can still see clearly that the timing of the rain events played much stronger role than total precipitation in determining the GPP and NEE, by influencing the length of the growing season and the ecosystem respiration. This is probably because the growing season of this Mediterranean grassland was confined in the winter and early springtime when potential evapotranspiration was low and usually soil moisture was not limiting factor for carbon uptake. Hence, the widely accepted view of the positive relationship between grass production and precipitation may not hold for Mediterranean types of grassland. The impact of the length of growing season on NEE was also observed from

a synthesis based on the carbon flux data across different temperate deciduous forests (Baldocchi et al., 2001).

3.7. Impact of U^* correction on estimate of annual NEE

As we know, the reliability of nighttime F_c measured with eddy covariance method under low turbulent conditions has been questioned (e.g., Aubinet et al., 2002; Massman and Lee, 2002), and there is potential systematic bias on underestimating respiration at night. Among others, poor turbulent mixing, horizontal advection or drainage has been identified as main causes responsible for the underestimation of carbon flux (Massman and Lee, 2002). In Fig. 10a, we present a relation between U^* and nighttime 0.5 h F_c . To avoid effects from other variables on F_c , data were from the peak growth period (DOY070–120) of 2001. Overall trends shown in the dataset is that F_c was strongly underestimated under low U^* condition. Also there appears to exist a U^* threshold which was around 0.1 m s^{-1} . The lower nighttime F_c under calm conditions has been reported in other studies over forest sites (Goulden et al., 1997; Aubinet et al., 2002) and from grassland sites (Suyker and Verma, 2001; Flanagan et al., 2002). Normally, the data from those conditions have often been corrected to estimate the annual NEE (Falge et al., 2001). One of the most common procedures is to develop a temperature-dependent correction function of F_c obtained during periods of high turbulence when friction velocity (U^*) is above a certain threshold (Aubinet et al., 2002; Flanagan et al., 2002).

It is clear that under low U^* conditions half-hour measurements of F_c could be significantly underestimated. How different thresholds in the process of U^* correction would affect the estimated annual NEE is less clear, however. Data shown in Fig. 10b illustrate that as U^* threshold increased from 0.0 to 0.1 m s^{-1} the estimated NEE became less negative or more positive. Then further increase in U^* threshold only slightly affected the estimated NEE. The differences of estimated NEE between uncorrected (i.e. U^* threshold = 0 m s^{-1}) and corrected at the threshold of 0.1 m s^{-1} were only 44 and 34 g C m^{-2} for two seasons, which was near the upper bound of uncertainty of NEE with the eddy covariance technique

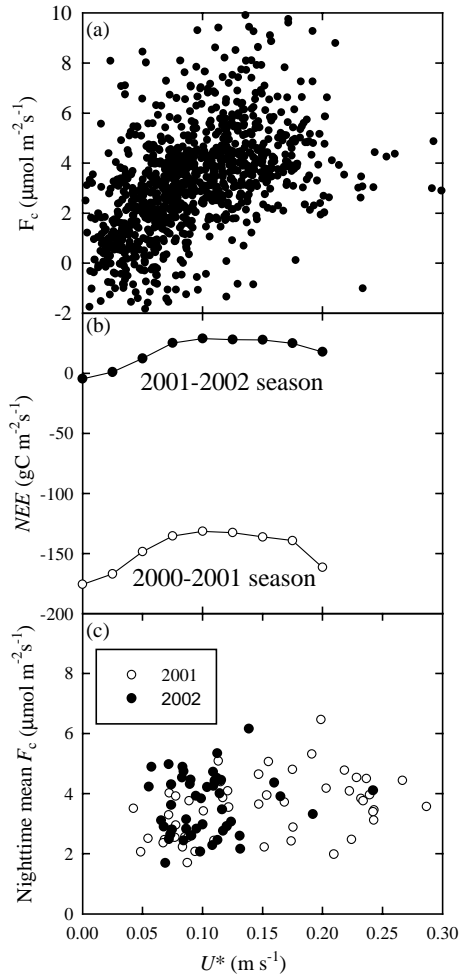


Fig. 10. (a) Influence of friction velocity (U^*) on nighttime 0.5 h CO_2 flux (F_c). (b) Impact of different U^* threshold on estimated annual ecosystem carbon exchange (NEE). (c) Influence of nighttime mean U^* on mean F_c . To avoid possible confounding effect of soil moisture and grass phenology on ecosystem respiration, only data from the peak growth period (DOY070–120) were included in this analysis of (a) and (c).

(Moncrieff et al., 1996; Goulden et al., 1996a,b). Thus, the impact of using different U^* threshold on annual NEE was much less significant than that on the 0.5 h F_c data. This seems indicating that major portion of respired CO_2 from the ecosystem during very stable conditions might not leave the system. On the other hand, the storage term could not entirely account for the decrease in F_c (data not shown). So we

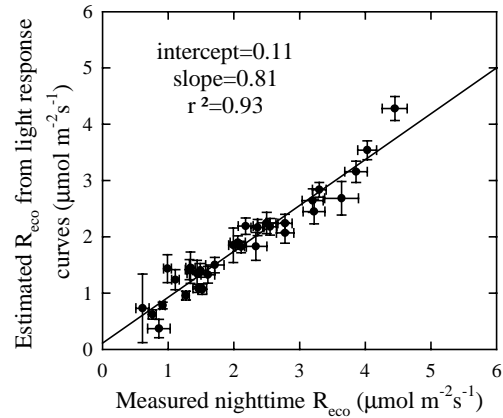


Fig. 11. Ecosystem respiration (R_{eco}) determined by using Eq. (4) as compared to the mean of measured nighttime ecosystem respiration.

argue that most of respired CO_2 might accumulate at microtopographic troughs under stable condition, and it could flush out and pick up by the eddy covariance system when the air became unstable. Two lines of evidence can be used to support this argument. One is data presented in Fig. 10c, which showed a poor correlation between mean nighttime F_c and U^* . Another evidence is from the analysis of R_{eco} using light-response function (Eq. (4)) as shown in Fig. 11. R_{eco} estimated from light-response curves was correlated very well with the nighttime mean F_c . Linear regression (with the mean nighttime F_c as independent variable) yielded a slope of 0.81 and intercept of $0.11 \mu\text{mol m}^{-2} \text{s}^{-1}$ ($r^2 = 0.93$), demonstrating the reliability of nighttime F_c data. So from this analysis, low U^* correction for our site may be not needed to produce accurate annual sums. A similar slope of 0.78 (Suyker and Verma, 2001) or even higher slope of 0.94 (Falge et al., 2002) has been reported from studies over a wide range of functional types, including tropic, Mediterranean, temperate and boreal forests, and C_3 and C_4 grassland. Some studies have shown that estimated annual NEE was insensitive to U^* threshold (Barford et al., 2001), while others shown that the sensitivity was very high (Barr et al., 2002). Thus, U^* correction could be site specific, depending on the characteristics of terrain and vegetation, and instrument heights (Aubinet et al., 2002; Massman and Lee, 2002).

4. Conclusions

We presented data from a CO₂ flux study over annual grassland in California using the eddy covariance method. Results showed that seasonal trends of both GPP and R_{eco} followed closely the change in LAI. R_{eco} followed the exponential function of soil temperature with season-dependent Q_{10} values. We observed large R_{eco} pulses immediately after rain events, especially those in the non-growing season. During the non-growing seasons from mid-May to September, despite high temperature, severe drought inhibited R_{eco} to a very low level (less than 1.0 g C m⁻² per day). Nighttime low U^* was showed to have a limited impact on estimated annual NEE and nighttime mean F_c , suggesting that drainage may not occur under stable condition. Integrated GPP, R_{eco} , and NEE were 867, 735, and -132 g C m⁻², respectively, for 2000–2001 season, and 729, 758, and 29 g C m⁻² for 2001–2002 season. Higher GPP in the first season was mainly attributed to earlier start of the growing season. Thus, change in length of the growing season due to time of onset of rain was mainly responsible for the variations of annual GPP. So from two-season data, for this Mediterranean grassland, the start of the wet season and the timing of rain event had larger impact than total precipitation on GPP and R_{eco} , and consequently on NEE. More studies are needed to understand how GPP, R_{eco} , and NEE of this Mediterranean grassland respond to a wider spectrum of changes in rainfall patterns due to El Niño and La Niña climate events.

Observations of respiratory pulses after rain events are becoming more common as more investigators report carbon fluxes from Mediterranean ecosystems (Reichstein et al., 2002a,b). The next generation of soil respiration models will need to assess these pulses and the rate they decrease as the upper soil dries, in addition to considering the basal effects of soil temperature and deep soil moisture. To do so we need better information on the physiological mechanisms that describe how soil microbes respond to sudden increase in moisture (Halverson et al., 2000).

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