



Precipitation legacy effects on dryland ecosystem carbon fluxes: direction, magnitude and biogeochemical carryovers

W. Shen¹, G. D. Jenerette², D. Hui³, and R. L. Scott⁴

¹Center for Ecology and Environmental Sciences, South China Botanical Garden, the Chinese Academy of Sciences, Guangzhou, 510650, China

²Department of Botany and Plant Sciences, Center for Conservation Biology, University of California Riverside, Riverside, CA 92521, USA

³Department of Biological Sciences, Tennessee State University, Nashville, TN 37209, USA

⁴USDA Agricultural Research Service, Southwest Watershed Research Center, Tucson, AZ 85719, USA

Correspondence to: W. Shen (shenweij@scbg.ac.cn)

Received: 5 June 2015 – Published in Biogeosciences Discuss.: 1 July 2015

Revised: 25 December 2015 – Accepted: 7 January 2016 – Published: 21 January 2016

Abstract. The precipitation legacy effect, defined as the impact of historical precipitation (PPT) on extant ecosystem dynamics, has been recognized as an important driver in shaping the temporal variability of dryland aboveground net primary production (ANPP) and soil respiration. How the PPT legacy influences whole ecosystem-level carbon (C) fluxes has rarely been quantitatively assessed, particularly at longer temporal scales. We parameterized a process-based ecosystem model to a semiarid savanna ecosystem in the southwestern USA, calibrated and evaluated the model performance based on 7 years of eddy-covariance measurements, and conducted two sets of simulation experiments to assess interdecadal and interannual PPT legacy effects over a 30-year simulation period. The results showed that decreasing the previous period/year PPT (dry legacy) always increased subsequent net ecosystem production (NEP) whereas increasing the previous period/year PPT (wet legacy) decreased NEP. The simulated dry-legacy impacts mostly increased subsequent gross ecosystem production (GEP) and reduced ecosystem respiration (R_e), but the wet legacy mostly reduced GEP and increased R_e . Although the direction and magnitude of GEP and R_e responses to the simulated dry and wet legacies were influenced by both the previous and current PPT conditions, the NEP responses were predominantly determined by the previous PPT characteristics including rainfall amount, seasonality and event size distribution. Larger PPT difference between periods/years resulted in larger legacy impacts, with dry legacies fostering more C

sequestration and wet legacies more C release. The carryover of soil N between periods/years was mainly responsible for the GEP responses, while the carryovers of plant biomass, litter and soil organic matter were mainly responsible for the R_e responses. These simulation results suggest that previous PPT conditions can exert substantial legacy impacts on current ecosystem C balance, which should be taken into account while assessing the response of dryland ecosystem C dynamics to future PPT regime changes.

1 Introduction

Drylands play an important role in global carbon (C) cycle and future C sequestration (Houghton et al., 1999; Asner et al., 2003), as they cover 30–45 % of the earth's land surface (Asner et al., 2003; Reynolds et al., 2007), store about 15 % of the global soil organic carbon (Schlesinger, 1991), and represent 30–35 % of terrestrial net primary production (Field et al., 1998). Driven by sporadic precipitation (PPT) and nonlinear biological responses, dryland C fluxes are especially variable across time and space (Maestre et al., 2012; Collins et al., 2014), making the prediction of dryland C budgets a challenging task (Jenerette et al., 2012). Moreover, climate models predict that the intra- and interannual PPT variability may be further intensified in dryland regions with longer drought durations and more large-sized events (Solomon et al., 2007; Diffenbaugh et al., 2008; Cook and

Seager, 2013). Further, sequences of wet years followed by sequences of dry years and vice versa are also increasingly likely (Peters et al., 2012; Sala et al., 2012). Understanding the response of dryland ecosystem C fluxes to PPT variation is, therefore, important to characterizing the global C cycle and predicting how future PPT regime changes will affect dryland C balance.

As a measure of ecosystem C balance, net ecosystem production (NEP) has a value that is positive when an ecosystem accumulates C and negative when an ecosystem loses C. Dryland NEP is closely tied to current-year PPT amount, with wetter-than-average years being a C sink, drier-than-average years being a C source, and years with average rainfall being C neutral (Flanagan et al., 2002; Hastings et al., 2005). Additionally, at seasonal scales, the distribution of PPT in addition to the total amount can have large influences on ecosystem production (Porporato et al., 2004; Katul et al., 2007). At interannual scales a PPT legacy effect, defined as the impact of past PPT conditions on the current structure and functioning of ecosystems (Lauenroth and Sala, 1992; Sala et al., 2012; Monger et al., 2015), has also been found to play an important role in shaping the temporal variability of dryland ecosystem C fluxes (Knapp et al., 2002; Huxman et al., 2004a, b; Heisler and Weltzin, 2006; Sala et al., 2012; Ogle et al., 2015). For example, Hasting et al. (2005) attributed the C sink status of a desert shrub ecosystem in the early spring of 2002 to the above-average rainfall in the late fall of 2001. Scott et al. (2009) and Hamerlynck et al. (2013) found that a cool-season (December–April) drought was followed by an unusually large net C loss during the following warm monsoon season (July–September) in a semiarid savanna and a semi-desert grassland. Moreover, the savanna ecosystem has recently been a net C source, and one hypothesized but untested explanation is due to an increase in current respiration of organic C that accumulated in the preceding wetter decade (Scott et al., 2009). While these studies reveal the existence of PPT legacy effects on NEP at the seasonal scale, only a few studies have quantitatively assessed the contribution of PPT legacy to the temporal variability of dryland NEP at interannual and interdecadal timescales (Williams and Albertson, 2006), mainly because it is methodologically difficult to separate the past and current PPT impacts on C fluxes with the limited observational data (Sala et al., 2012), and there is a general lack of field manipulative experiments to address the PPT legacies at these scales (Reichmann et al., 2013a).

Much of our current understanding of the PPT legacy effects on dryland C fluxes is based on aboveground net primary production (ANPP). A number of studies have documented that dryland ANPP is not only linearly related to current-year PPT but also closely related to the PPT amount and seasonality several months to years before (Lauenroth and Sala, 1992; Oesterheld et al., 2001; Huxman et al., 2004c). For example, field studies have found a positive legacy impact where ANPP is higher than expected if pre-

ceded by a wetter year, or lower than expected if preceded by a drier year (Jobbagy and Sala, 2000; Oesterheld et al., 2001; Wiegand et al., 2004; Sherry et al., 2008; Sala et al., 2012). Proposed mechanisms explaining such observed positive PPT legacy effects on ANPP mainly involve the structural carryovers between years, which can be leaf and root biomass (Oesterheld et al., 2001); the composition of species differing in rooting depth and phenology (Paruelo et al., 1999; Jobbagy and Sala, 2000); or the density of seeds, tillers and plant individuals (Oesterheld et al., 2001; Yahdjian and Sala, 2006; Reichmann et al., 2013a). Alternatively, a negative legacy effect occurs when production is lower than expected if preceded by a wet period or higher than expected if preceded by a dry period (Jenerette et al., 2010). A negative PPT legacy effects may be influenced more by biogeochemical carryovers that influence the resource availability to respond to current PPT (Evans and Burke, 2013; Reichmann et al., 2013b), whereby increased growth in response to a higher PPT can reduce the available nutrients (e.g., nitrogen, N) for the following period and vice versa. Although various mechanisms have been proposed for the PPT legacy impacts on ANPP, few of them have been rigorously tested, and the key underlying mechanisms still remain poorly understood (Sherry et al., 2008; Williams et al., 2009; Sala et al., 2012; Monger et al., 2015).

Soil respiration (R_s), as a major component of ecosystem C efflux, has also been found to have lagged responses to PPT variations (Huxman et al., 2004b; Sponseller, 2007; Ma et al., 2012; Cable et al., 2013). This is particularly true at the event scale; after a period of drought, a rainfall event can result in a pulse of CO_2 efflux that may be orders of magnitude larger than that before the event and then decline exponentially for a few days to weeks (Xu et al., 2004; Jenerette et al., 2008; Borken and Matzner, 2009; Cable et al., 2013; Oikawa et al., 2014). At a seasonal scale, Vargas et al. (2010) found no lags between R_s and soil moisture across 13 vegetation types, including four grasslands; however, Hamerlynck et al. (2013) presented longer-term ecosystem flux data that suggest seasonal drought legacies affect ecosystem respiration (R_e) in a semi-desert grassland in southeastern AZ, USA. They posited that the increased C substrate availability resulting from the previous cool-season drought-induced plant mortality was responsible for the higher R_e in the following monsoon season. However, very few studies have been devoted to understanding the PPT legacy impacts on dryland respiration at greater than seasonal timescales.

In this study, we conducted simulation experiments with a widely used dryland ecosystem model, Patch Arid Land Simulator (PALS; Kemp et al., 1997, 2003; Reynolds et al., 2004; Shen et al., 2009), to analyze the PPT legacy effects on ecosystem-level C fluxes including NEP, gross ecosystem production (GEP), and R_e . The PALS model was built on the pulse-reserve concept (Noy-Meir, 1973) and had been used to analyze the impacts of antecedent moisture conditions and the lagged responses of different plant functional

types (FTs) in three North American deserts at the rainfall event scale (Reynolds et al., 2004). We parameterized, calibrated, and evaluated the model based on the long-term eddy-covariance-measured fluxes in a semi-desert savanna ecosystem in the southwestern USA (Scott et al., 2009) to analyze the PPT legacy effects at interannual and interdecadal scales. Specifically, we addressed the following two questions. First, what are the direction and magnitude of ecosystem C flux responses to dry and wet legacies? We expected that the PPT legacy impacts would occur over annual and decadal scales in correspondence to PPT fluctuations at these scales, and the dry- and wet-legacy impacts would differ in direction and magnitude. Second, how are the direction and magnitude of PPT legacy effects related to the PPT characteristics of both the previous and the current year/period? We expected that greater variability in PPT would lead to corresponding increases in legacy effect. For PPT characteristics, we were interested not only in the annual and seasonal PPT amount but also in between-event interval and event size distribution, since these variables are widely recognized key PPT features to dryland ecosystems (Porporato et al., 2004; Katul et al., 2007; Shen et al., 2008a).

2 Methods

2.1 Model description

PALS is a process-based ecosystem model that consists of four modules: atmospheric forcing, a water cycling and energy budget, plant production and respiration, and soil organic matter (SOM) decomposition and heterotrophic respiration (R_h). The four modules are interactively linked by the cycling of C, N, and H_2O through the atmosphere–plant–soil continuum. The PALS model explicitly considers seven plant FTs commonly found in the North American warm deserts: evergreen shrub, deciduous shrub, perennial forb, perennial C_3 and C_4 grasses, and native and exotic C_3 annual grasses (Reynolds et al., 1997; Shen et al., 2009). Since the detailed model structure and mechanistic relationships have been presented in several publications (Kemp et al., 1997, 2003; Reynolds et al., 1997, 2000, 2004; Gao and Reynolds, 2003; Shen et al., 2005, 2008a, b, 2009), here we briefly describe the four modules and refer to the specific literature for detailed description.

The atmospheric driving force module reads in data for atmospheric driving variables (e.g., atmospheric $[CO_2]$, N deposition rate, daily maximum and minimum air temperatures, PPT, relative humidity, and solar radiation) and, based on these driving variables, calculates other important variables such as vapor pressure deficit (VPD), which directly influences stomatal conductance and indirectly influences soil temperature, SOM decomposition and soil respiration. Calculations of VPD and soil temperature can be found in Eqs. (2)–(7) in Shen et al. (2005).

The water cycling and energy budget module mainly calculates soil water contents at six layers, the rates of water infiltration into and percolation out of a layer, and water losses via evaporation and transpiration from different layers. Water infiltration and percolation rates of a layer are determined by the effective PPT reaching the soil surface, previous water content, and the water holding capacity as a function of soil texture (Shen et al., 2005). Soil evaporation is determined by soil water availability and energy available in the two top soil layers (10 cm in depth). Water uptake by plants is partitioned among the soil layers according to the proportion of roots in each layer for all plant FTs (Kemp et al., 1997; Shen et al., 2008b). Canopy transpiration is calculated by using the energy budget and the canopy stomatal resistance (Reynolds et al., 2000; Gao and Reynolds, 2003).

The plant production and respiration module mainly simulates phenology, primary production, growth and maintenance respiration, photosynthate allocation, and litterfall of each plant FT. Three major phenophases (i.e., dates of germination, leafing, and dormancy) are determined in PALS based on the observed dates, air temperature, and PPT (Shen et al., 2009). Primary production for each FT is calculated based on the leaf area, potential net photosynthetic rate, stomatal conductance, leaf N content modifier, and the difference between intercellular and atmospheric $[CO_2]$. The plant photosynthesis rate is estimated as a product of stomatal conductance and the partial pressure gradient between atmospheric and intercellular $[CO_2]$. The stomatal conductance is calculated as an exponential function of leaf water potential that decreases linearly with atmospheric vapor deficit (see Eqs. (10)–(14) in Shen et al., 2005). Photosynthate is allocated to different plant organs (leaf, stem, and root) using fixed allocation ratios after subtracting the maintenance respiration, which is estimated as a function of live biomass, basal respiration rate, and modifiers of temperature and plant water potential (Shen et al., 2008a). Growth respiration is calculated based on the growth yield coefficient and the net photosynthate used for growth (Shen et al., 2008a). Litterfall amount is mainly determined as a function of observed dormancy dates, maximum air temperature and drought conditions (Shen et al., 2008a, 2009).

The SOM decomposition and heterotrophic respiration module simulates the decomposition of metabolic and structural litter material; SOM in active, slow and passive pools; and CO_2 emissions associated with these decomposition processes (Kemp et al., 2003; Shen et al., 2009). The SOM decomposition rate or heterotrophic rate is calculated as a first-order kinetic rate with a decomposition coefficient multiplied by the pool size and the temperature and moisture scalars (see Eqs. (A4)–(A11) in Shen et al., 2009). In addition, this module also simulates the dynamics of soil mineral N pool by using N mineralization and atmospheric deposition as the major inputs, and plant N uptake and leaching loss as the major outputs. Among these the N mineralization and plant uptake processes are modeled in more detail while the rates

of the other processes are basically assigned with empirical constant values. The N mineralization processes are directly coupled to litter and SOM decomposition processes and are calculated as a product of the C flow rates and the C / N ratio of the corresponding litter or SOM pools (Parton et al., 1993; Kemp et al., 2003). The plant N uptake is a product of water transpiration and N concentration in soil solution (see Eq. 8 in Shen et al., 2008b).

2.2 Model parameterization

For this study, we modified and parameterized PALS to represent an upland mesquite savanna ecosystem in the Santa Rita Experimental Range (SRER; 31.8214° N, 110.8661° W, elevation 1116 m), about 45 km south of Tucson, AZ, USA. Soils at this site are a deep sandy loam (Scott et al., 2009), and the mean groundwater depth likely exceeds 100 m (Barron-Gafford et al., 2013). PPT was therefore considered as the only source of water input into the system. Based on the vegetation composition (Scott et al., 2009), there were five major plant FTs included in PALS: shrub (e.g., *Prosopis velutina*), subshrub (e.g., *Isocoma tenuisecta*), C₄ perennial grass (e.g., *Digitaria californica*), perennial forb (e.g., *Ambrosia psilostachya*), and C₃ annual grass, among which the velvet mesquite shrub with average height of ca. 2.5 m accounted for ~35 % of the total canopy cover and other FTs (mainly perennial grasses) accounted for ~22 % (Scott et al., 2009). Therefore, we derived the site-characteristic parameters for the two major FTs (shrub and perennial grass) from previous studies carried out in SRER, with those for the other FTs being adopted from a generic parameter data set for the PALS model to be used in the North American warm deserts (Reynolds et al., 2004; Shen et al., 2005). These site-specific parameters mainly included plant-related parameters (e.g., canopy cover, C allocation ratio, rooting distribution ratio, and the initial values of living and dead plant biomass pools) and soil-related parameters (e.g., soil chemical and physical properties, C / N ratios, decomposition rates, and initial values of the litter and SOM pools). The values of these parameters are provided in Table S1 in the Supplement, with the cited literature also being listed below the table.

For the climatic variables used to drive the PALS model, we compiled a 30-year meteorological data set that included daily PPT, maximum and minimum air temperatures (T_{\max} and T_{\min}), relative humidity (RH), and total solar radiation (S_{rad}) from 1981 to 2010. The T_{\max} , T_{\min} , RH, and S_{rad} data from 1981 to 1990 were observations from the Tucson weather station (about 50 km north of the mesquite savanna site and lower elevation) and obtained by accessing Arizona Meteorological Network online data (AZMET: <http://ag.arizona.edu/azmet>). The remaining 20 years (1991–2010) of T_{\max} , T_{\min} , RH and S_{rad} data were observations from the Kendall Grassland meteorological site (about 85 km east of the mesquite savanna site and slightly higher elevation) and obtained by accessing Southwest Watershed Research Cen-

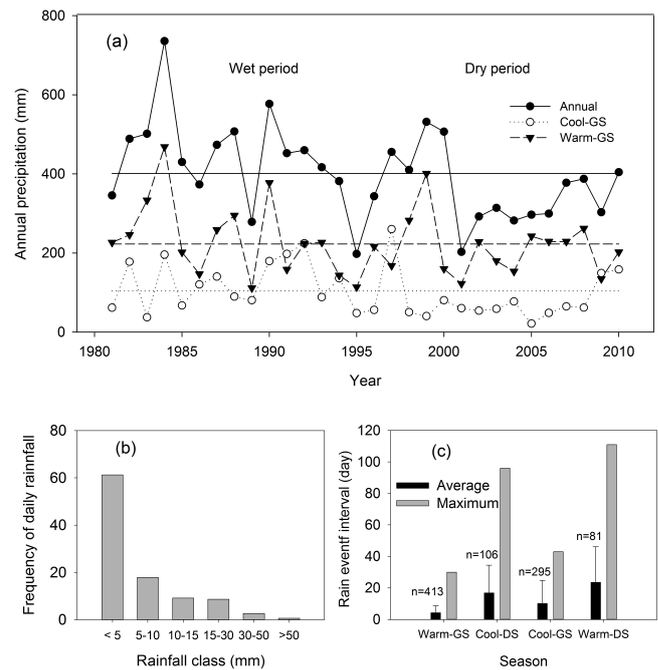


Figure 1. Precipitation characteristics in the 30 years (1981–2010) at the Santa Rita mesquite savanna site. **(a)** Annual and seasonal precipitation amount; **(b)** frequency distribution of daily rainfall; **(c)** mean and maximum between-event interval (BEI). Horizontal lines within **(a)** indicate mean annual and seasonal precipitation. The warm growing season (warm-GS) is from July through September, the cool dry season (cool-DS) from October to November, the cool growing season (cool-GS) from December through March, and the warm dry season (warm-DS) from April through June. Error bars in panel **(c)** represent standard deviations, and n is the number of rain event pairs used to calculate the between-event interval in the 30 years.

ter (SWRC) online data (<http://www.tucson.ars.ag.gov/dap/>). The 30-year PPT data were observations from the Santa Rita watershed rain gage no. 5 (1.5 km from the site) and obtained also from the SWRC online data access. These different sources of meteorological data were adjusted based on the 7 years (2004–2010) of meteorological data obtained from the AmeriFlux eddy-covariance flux tower at the mesquite savanna site (US-SRM; see Fig. S1 in the Supplement). Lastly, we used the AZMET and SWRC data from 1981 to 2003 plus the flux tower data from 2004 to 2010 to drive the model.

Since our simulation experiment was based on the manipulations of the 30-year (1981–2010) PPT data, we report the PPT characteristics here in more detail. In the past 30 years, the mean annual PPT (MAP) amount was 401 mm at the site, slightly greater than the long-term (1937–2007) mean of 377 mm (Scott et al., 2009). These 30 years were divided into two periods: a wet period from 1981 to 1994 with a MAP of 449 mm and a dry period from 1995 to 2010 with a MAP of 347 mm (Fig. 1a). For the analysis of PPT legacy effects at interdecadal scale, the wet period was treated as the previ-

ous period and the dry period as the current period. For the analysis of PPT legacy effects at interannual scale, the annual scale was defined as being from July through June of the next year. To analyze the relationship between PPT legacy effects and seasonal rainfall characteristics, each year was further divided into four seasons (with their mean rainfall in parentheses): the main warm growing season from July to September (warm-GS, 224 mm), the cool dry season from October to November (cool-DS, 48 mm), the minor cool growing season from December to March (cool-GS, 104 mm), and the warm dry season from April to June (warm-DS, 26 mm). At the site, as in many other dryland regions (Sala et al., 1992; Heisler-White et al., 2008), most rainy days have only light-rainfall events. About 80 % of daily rainfall was < 10 mm, with medium- to large-sized events (10–50 mm) accounting for about 20 % and only 10 events larger than 50 mm in the 30 years (Fig. 1b). The no-rain-day duration between events (hereafter between-event interval or BEI) was ~ 5 days on average in the warm-GS and ~ 10 days in the cool-GS (Fig. 1c). The average BEI was ~ 17 days in the cool-DS and 24 days in the warm-DS, but there could be no rain for 3 months in these dry seasons (Fig. 1c).

2.3 Model calibration and evaluation

After model parameterization, we calibrated the model based on 4 years (2004–2007) of CO_2 and H_2O flux data monitored using the eddy-covariance technique at the savanna site. Detailed descriptions of instrumentation, sensor heights and orientations, and data-processing procedures for the eddy-covariance data can be found in Scott et al. (2009). During model calibration, we mainly adjusted the parameter values of photosynthate allocation ratios, live biomass death rates, and SOM decomposition rates to achieve a best fit between modeled and observed GEP and R_e , since these parameters have been identified as the most sensitive and uncertain ones (e.g., photosynthate allocation ratios) in influencing the modeled ecosystem carbon fluxes (Shen et al., 2005). The model performed well in capturing the seasonal variation patterns of actual evapotranspiration (AET), GEP, R_e , and NEP in the 4 calibration years (Fig. S2), with larger C fluxes during the warm-GS than in the other seasons. At the annual scale, simulated AET, GEP, and R_e explained over 60 % of the variations in the observations (Fig. 2, left panels), but the correlation between the simulated and observed NEP was very weak (Fig. 2d). This was mainly because the model substantially overestimated GEP (120 g C m^{-2} simulated vs. 52 g C m^{-2} observed) in the cool-GS of 2006 (Fig. S3b). Further explanations on the possible causes of the GEP overestimation in 2006 shall be provided later in the Discussion section. If the data of this year were excluded, the explanatory power for annual NEP could reach 74 %. Since our goal was to use an empirically plausible model to understand the long-term temporal variations in ecosystem fluxes, we consider the calibration results acceptable.

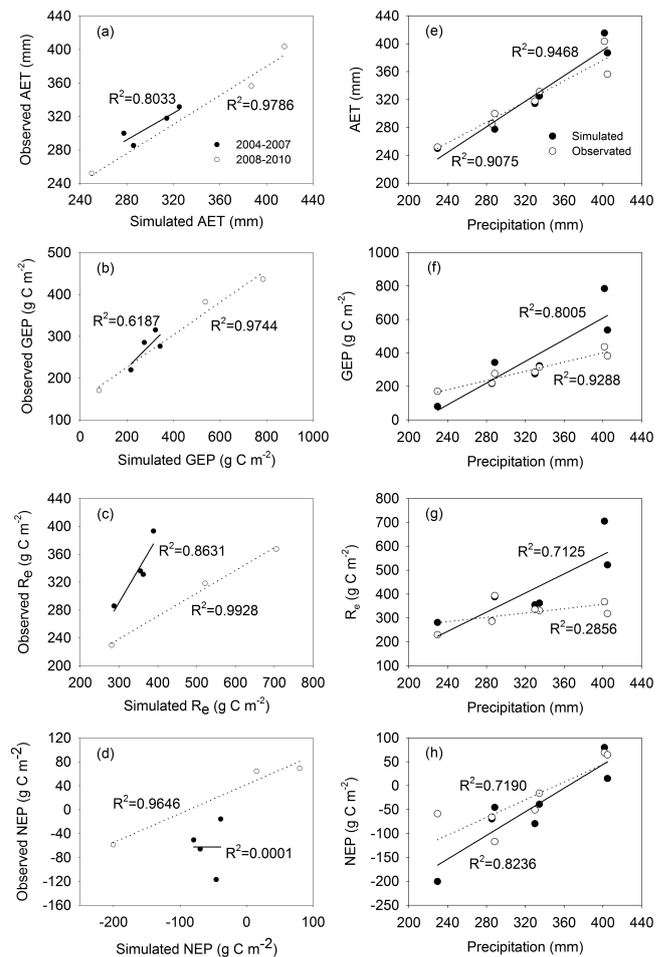


Figure 2. Comparison of the model-simulated water and carbon fluxes with the eddy-covariance observations at the mesquite savanna site. Left panels show the comparison between the modeled and observed fluxes in 4 calibration (2004–2007; solid dots) and 3 validation years (2008–2010; open dots). Right panels show the relationships of the simulated (solid dots) and observed (open dots) fluxes with precipitation in the 7 years (2004–2010). R^2 is the coefficient of determination describing the proportion of the variance in measured fluxes explained by the model for the left panels or that explained by precipitation for the right panels. AET represents actual evapotranspiration, GEP gross ecosystem production, R_e total ecosystem respiration, and NEP net ecosystem production.

The model performance was further evaluated by assessing the degree of correlation between the PALS-simulated and flux-tower-measured C and H_2O fluxes from 2008 through 2010, which were not used for model calibration. The coefficients of determination (R^2), which describe the proportion of the variance in measured data explained by the model, were all larger than 0.9 in the 3 validation years (2008–2010; Fig. 2, left panels). These evaluation results indicate that the model was capable of capturing the temporal variability of observed fluxes at the annual scale. Furthermore, we also analyzed the relationships between the ob-

served and simulated fluxes with the corresponding current-year PPT to see how the flux variations were explained by current-year PPT under baseline conditions (i.e., the PPT variations shown in Fig. 1). The explanatory power (R^2) for both the observed and simulated fluxes were mostly over 70 % (Fig. 2, right panels), which further indicates that the model is capable of capturing the impacts of PPT variability on ecosystem fluxes. The following simulation experiments were therefore designed to discriminate the contributions by previous- and current-year PPT impacts.

2.4 Simulation experiments

We designed two sets of simulation experiments to examine the interdecadal and interannual PPT legacy effects. To analyze the interdecadal legacy effects, we first changed the PPT of the 14-year previous period (1981–1994) by 0, ± 10 , ± 30 , ± 50 and ± 80 % (multipliers of existing daily PPT amounts in the record) while keeping the 16-year current-period (1995–2010) PPT unchanged. After these manipulations, the average PPT of the previous period ranged from 93 mm, corresponding to the 80 % decrease, to 837 mm, corresponding to the 80 % increase. This design detects how changes in previous-period PPT influence the current-period C fluxes and the associated C pool dynamics. On top of each previous period PPT manipulation level, we further changed the current-period PPT by 0, ± 10 , ± 30 , ± 50 , and ± 80 %, which resulted in the average current-period PPT varying from 69 to 621 mm. This design detects how changes in the current-period PPT influence the legacies resulting from changes in the previous-period PPT. As a result, we conducted 73 simulation runs, corresponding to the 73 combinations of the above previous- and current-period PPT manipulations (9 previous PPT levels times 8 current PPT levels plus 1 baseline run).

To analyze the interannual legacy, we changed the PPT of each individual year by ± 30 % while keeping the PPT of the subsequent years unchanged. This design resulted in 54 simulation runs (27 years from 1981 to 2007 times 2 PPT manipulation levels) and illustrates the effects of changes in the PPT of the previous 1 year on the C fluxes and resource pools of the current year(s). After a 30 % PPT change, annual PPT ranged from 162 to 925 mm in the 27 years, which was large enough to cover the PPT interannual variation at the study site. Another consideration of using 30 % as the PPT manipulation level was that future projected annual PPT variation in dryland regions will be -30 to $+25$ % (Bates et al., 2008; Maestre et al., 2012).

2.5 Data analysis

The legacy effect was quantified as the C flux (or resource pool size) of the current period/year after PPT changes in the previous period/year minus that without PPT changes in the previous period/year. As an example, the following equation

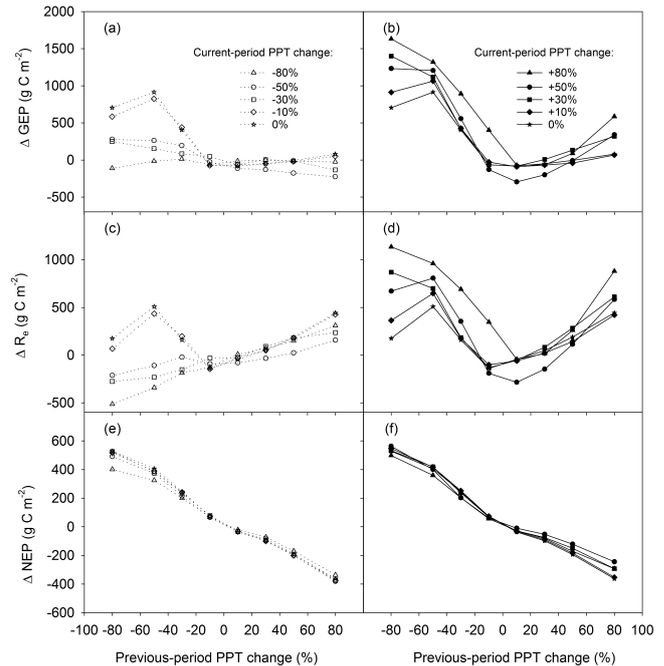


Figure 3. Interdecadal legacy effects of changing the previous-period (1981–1994) precipitation on the cumulative carbon fluxes of the current period (1995–2010). Interdecadal legacy effects on carbon fluxes (e.g., ΔNEP) are calculated as the difference between the current-period flux with previous-period PPT changes and that without previous-period PPT changes. Dashed lines with open symbols represent different levels of decreasing the current-period precipitation (left panels). Solid lines with filled symbols represent increasing the current-period precipitation (right panels).

calculates the legacy effect of increasing the previous-period PPT by 30 % on the current-period NEP:

$$\text{Legacy}_{\text{NEP}} = \Delta\text{NEP} = \text{NEP}_{\text{PPT}+30\%}^{\text{CP}} - \text{NEP}_{\text{PPT}+0\%}^{\text{CP}}, \quad (1)$$

where $\text{NEP}_{\text{PPT}+30\%}^{\text{CP}}$ is the cumulative NEP throughout the current period (1995–2010) under a 30 % previous-period (1981–1994) PPT increase; $\text{NEP}_{\text{PPT}+0\%}^{\text{CP}}$ is the cumulative NEP throughout the current period with no previous-period PPT change (i.e., the baseline PPT conditions shown in Fig. 1). This method directly quantifies whether changes in PPT of the previous period will impose a positive, a negative, or no legacy effect on the C fluxes (or resource pools) of the current period. For simplicity, hereafter we refer to the legacy effect resulting from the decreased previous-period/year PPT as the dry legacy and that resulting from the increased previous-period/year PPT as the wet legacy. Spearman correlation analysis was used to detect the relationships between legacy effects and PPT characteristics, including PPT amount, BEI, and the number of large (≥ 10 mm) vs. small (< 10 mm) events at yearly and seasonal scales. The correlation analysis was performed in SPSS 16.0 (Chicago, IL, USA).

3 Results

3.1 Interdecadal legacy

Changes in PPT of the previous period (1981–1994) imposed obvious legacy impacts on the C fluxes of the current period (1995–2010). The direction of the simulated interdecadal dry and wet legacies on GEP and R_e was dependent upon the direction of both the previous- and current-period PPT changes. When the current-period PPT was reduced (Fig. 3, left panels), the simulated dry legacies mostly increased the current-period GEP (Fig. 3a) but decreased R_e (Fig. 3c); whereas wet legacies imposed little impacts on the current-period GEP (Fig. 3a) but mostly increased R_e (Fig. 3c). When the current-period PPT was enhanced (Fig. 3, right panels), both the dry and wet legacies mostly increased GEP and R_e (Fig. 3b, d). Regardless of current-period PPT changes, NEP always increased with dry legacies and decreased with wet legacies (Fig. 3e, f), indicating a consistent negative NEP response to PPT legacies.

The simulated absolute magnitude of the PPT legacy influence on ecosystem C fluxes (i.e., GEP, R_e , and NEP) generally increased with the absolute magnitude of changes in the previous-period PPT (Figs. 3, 4). Increasing the current-period PPT generally amplified the legacy effects compared to decreasing the current-period PPT (comparing the left to the right panels of Fig. 3). The magnitude of the PPT legacies was also significantly correlated with the PPT difference between the current and previous period (Δ PPT, equals to the current-period PPT minus the previous-period PPT; Fig. 4). If the previous period was wetter than the current period (i.e., Δ PPT < 0 or a wet-to-dry period transition), the legacy effect on R_e was negatively correlated with Δ PPT (Fig. 4c) but that on NEP was positively correlated with Δ PPT (Fig. 4e), indicating more current-period C release after a wetter previous period. In contrast, if the previous period was drier than the current period (i.e., Δ PPT > 0 or a dry-to-wet period transition), the correlations were all positive for GEP, R_e and NEP (Fig. 4, right panels), indicating more current-period C sequestration after a drier previous period.

The resource pool dynamics were also shaped by the alterations in the previous- and current-period PPTs. We only showed the 30 % decrease and increase in the previous- and current-period PPT (i.e., 4 out of 72 pairs of PPT change combinations) as representative examples in Fig. 5, because the major response patterns for the other paired combinations were similar. The PPT legacy impacts generally lasted for about 6–8 years for plant biomass, litter mass and soil water content (SWC), and much longer for soil organic matter (SOM) and soil mineral N (N_{soil} ; Fig. 5). Based on the resource pool responses in the early 1–2 years (i.e., 1995 and 1996) of the current period, the dry legacies decreased biomass, litter and SOM (Fig. 5a–f), but positively impacted N_{soil} (Fig. 5g–h). Contrastingly, the wet legacies increased biomass, litter and SOM (Fig. 5a–f) but negatively impacted

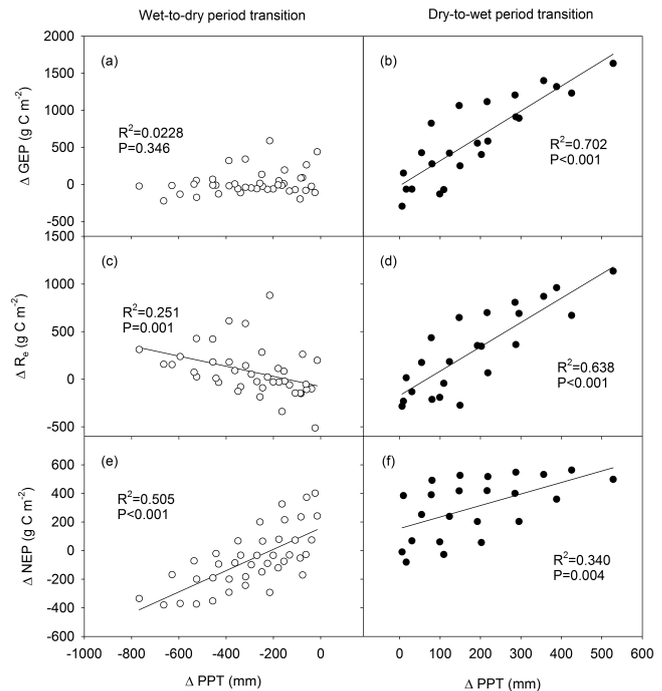


Figure 4. Spearman correlations of interdecadal precipitation legacy effects with the precipitation difference between periods (Δ PPT). Interdecadal Δ PPT is calculated as the mean PPT of the current period (1995–2010) minus that of the previous period (1981–1994). Interdecadal legacy effects on carbon fluxes (e.g., Δ NEP) are calculated as the difference between the current-period flux with previous-period PPT changes and that without previous-period PPT changes. Sample size is 41 for the wet-to-dry period transition (left panels) and 23 for the dry-to-wet period transition (right panels). GEP represents gross ecosystem production, R_e ecosystem respiration, and NEP net ecosystem production. R^2 is the coefficient of determination, and P is probability.

N_{soil} (Fig. 5g–h). Similar to the influences on C fluxes, increasing the current-period PPT (Fig. 5, right panels) amplified the PPT legacy impacts on biomass and litter (Fig. 5a–d), and hastened the recovery rates of SOM and N_{soil} to their baseline levels (Fig. 5e–h).

3.2 Interannual legacy

At the interannual scale, a 30 % decrease or increase in PPT could have legacy impacts on ecosystem C cycling lasting for 2–12 years (Fig. 6a–b). Notably, the direction of GEP and R_e responses to decreasing or increasing previous-year PPT could be positive or negative (Fig. 6c–f). The dry- or wet-legacy effects on these two fluxes were variable; idiosyncratic; and, in some cases, large at this timescale. However, the simulated dry legacies mostly increased NEP (Fig. 6g), whereas the simulated wet legacies mostly decreased NEP (Fig. 6h), which was similar to legacy responses at the interdecadal scale (Fig. 3e–f).

Table 1. Spearman correlation coefficients between interannual legacy effects and precipitation characteristics. Significant correlations are indicated with * for $0.01 < P \leq 0.05$ and ** for $P \leq 0.01$ (2-tailed; $n = 27$).

Precipitation characteristics	Dry legacy (previous-year PPT -30%)			Wet legacy (previous-year PPT $+30\%$)		
	ΔGEP	ΔR_e	ΔNEP	ΔGEP	ΔR_e	ΔNEP
Previous-year PPT characteristics						
Yearly rainfall	0.134	0.033	0.0270	-0.324	-0.180	-0.374
Warm-GS rainfall	0.303	0.072	0.519**	-0.430*	-0.065	-0.579**
Warm-GS BEI	-0.069	0.137	-0.399*	-0.075	0.053	-0.262
Warm-GS NE > 10 mm	0.329	0.067	0.636**	-0.535**	-0.227	-0.619**
Current-year PPT characteristics						
Yearly rainfall	0.278	0.162	0.484*	-0.466*	-0.600**	-0.224
Cool-GS rainfall	0.528**	0.338	0.495*	-0.277	-0.331	-0.218
Yearly BEI	-0.512**	-0.285	-0.686**	0.359	0.352	0.255
Cool-GS BEI	-0.519**	-0.286	-0.510**	0.151	0.088	0.214
Yearly NE > 10 mm	0.331	0.178	0.512**	-0.567**	-0.583*	-0.398*
Cool-GS NE < 10 mm	0.614**	0.577**	0.398*	-0.105	-0.075	-0.128
PPT difference (ΔPPT) between current and previous year						
Yearly rainfall	0.088	-0.135	0.466*	0.078	-0.088	0.252
Warm-GS rainfall	-0.059	-0.042	0.074	0.206	-0.096	0.326
Cool-GS rainfall	0.326	0.048	0.374*	0.248	0.160	0.209

Abbreviations: PPT: precipitation; GEP: gross primary production; R_e : ecosystem respiration; NEP: net ecosystem production; GS: growing season; BEI: between-event interval; NE: number of rainfall events.

The correlation analysis showed that not only rainfall amount but also BEI and event size distribution could influence the magnitude of the simulated dry and wet legacies (Table 1). The warm-GS PPT of a previous year was significantly correlated with the dry legacies for NEP and the wet legacies for GEP and NEP (Table 1). On the other hand, the cool-GS PPT of a current year influenced the dry and wet legacies for C fluxes, but not all of them were statistically significant (Table 1). These results indicate that the legacies were mainly generated in the warm-GS of a previous year, but the current-year cool-GS PPT conditions could influence the C flux responses to the previous-year legacies. Unlike at the interdecadal scale (Fig. 4), our correlation analysis showed that only the dry legacies for NEP had significant correlations with the PPT difference (ΔPPT) between 2 consecutive years or cool-GSs (Table 1), indicating that the larger the PPT difference between a previous dry year and a current wet year, the greater the legacy impacts on NEP imposed by the previous dry year.

To analyze the interannual PPT legacy impacts on the dynamics of resource pools (i.e., biomass, litter, SOM, N_{soil} , and SWC), 2 wet years (1983 and 1994) and 2 dry years (1986 and 1995) were chosen as examples (see Fig. 1a). The simulated dry legacies reduced biomass, litter and SOM but increased N_{soil} and SWC in the first current year (Fig. 7). In contrast, the simulated wet legacies imposed the opposite direction of impacts on the five resource pools (Fig. 7). The simulated PPT legacy impacts on the resource pools could

also last for several years, and the direction and magnitude of the legacy impacts in the following years could differ from those in the first year as described above. For example, increasing the PPT of 1995 by 30% caused a positive legacy impact on the biomass of the first following year (i.e., 1996; Fig. 7b), but it became negative in the later following years (e.g., in 1998; Fig. 7b), further indicating that current-year PPT conditions can influence the direction and magnitude of previous-year PPT legacies.

4 Discussion

4.1 Direction and magnitude of the simulated PPT legacies

Through this simulation analysis we demonstrated that previous PPT could impose substantial legacy impacts on current ecosystem C fluxes at interannual and interdecadal timescales. Notably, our simulation results support the hypothesis proposed for our study site (Scott et al., 2009) that the accumulated SOM during the previous wet period contributed to the net C release from the ecosystem during the current dry period. This specific test illustrates a major finding from our simulation study of a negative PPT legacy effect on NEP; i.e., decreasing previous PPT increased current NEP, whereas increasing previous PPT decreased current NEP (Figs. 3, 6). Increasing prior PPT (wet legacy) led to limited changes in GEP but consistently increased R_e .

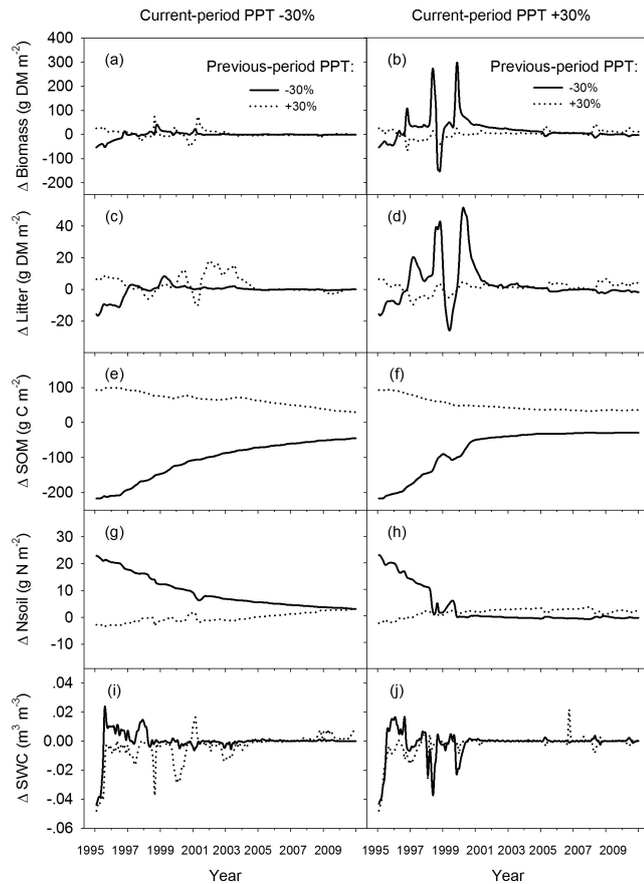


Figure 5. Interdecadal precipitation legacy effects on the resource pool dynamics. Left panels show the resource pool responses under a 30 % decrease, while right panels show those under a 30 % increase in the precipitation (PPT) of the current period from 1995 to 2010. Legacy effects on pool size (e.g., Δ Biomass) are quantified as the difference between the current-period pool size with previous-period PPT change and that without previous-period PPT change. Dashed lines represent a 30 % decrease, while solid lines represent a 30 % increase in the PPT of the previous period from 1981 to 1994. SOM represents soil organic matter, N_{soil} soil mineral nitrogen, and SWC soil water content.

Decreasing prior PPT (dry legacy) led to more variable effects for both GEP and R_e that were strongly conditioned on current-period PPT such that increasing current PPT was associated with increases in the dry-legacy effect. Overall, the effects on GEP were larger than R_e for reduced prior PPT and smaller for increased prior PPT, which resulted in a consistent negative PPT legacy on NEP regardless of current PPT. The complexity in the legacy effects on ecosystem C cycling we show here are in part influenced by the contrasting PPT legacy responses of C uptake and emission and their distinct interactions with current PPT distributions.

In projecting future dryland C dynamics, the effects of PPT legacies increase the complexity of ecosystem responses to PPT variability. One consistent interaction between legacy

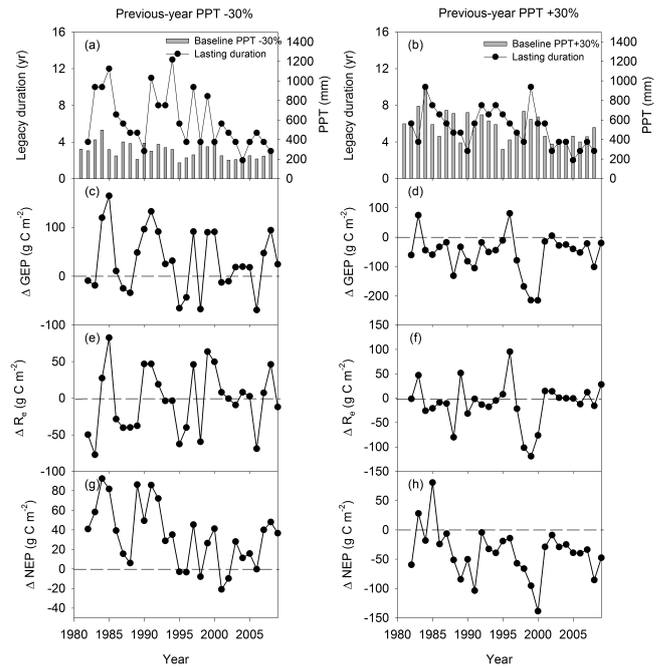


Figure 6. Interannual precipitation legacy effects on the ecosystem carbon fluxes. (a) and (b) show the lasting duration of dry (left panels) and wet (right panels) legacies, respectively. The legacy lasting duration is quantified as the number of years during which the legacy impacts on ecosystem fluxes exist after a previous-year PPT change. (c) through (h) show the responses of gross ecosystem production (GEP), ecosystem respiration (R_e) and net ecosystem production (NEP) to dry (left panels) and wet (right panels) legacies. Bars in the background of (a) and (b) represent the previous-year PPT amount after a 30 % decrease and increase, respectively.

and current PPT effects was that larger between-period PPT differences could result in larger legacy effects (Fig. 4), which is in agreement with what has been found in some field studies. For example, the magnitude of drought legacy on ANPP is proportional to the severity of the drought (Yahdjian and Sala, 2006; Swemmer et al., 2007), and dry- or wet-year legacies on ANPP are linearly related to the PPT difference between years (Sala et al., 2012; Reichmann et al., 2013a). Our simulation analysis detected that not only annual PPT amount but also finer-scale PPT characteristics such as GS rainfall, BEI, and event size could be important in determining the interannual-scale PPT legacy effects (Table 1). These simulation results suggest that PPT legacy effects may play a more important role in shaping the temporal variability of dryland ecosystem C fluxes under the projected increase in future PPT variability (Solomon et al., 2007; Cook and Seager, 2013) but that their characterization remains a challenge.

The influence of PPT legacies on dryland ecosystem C balance may strongly interact with other sources of variability in dryland C balance, including current-year PPT (Flanagan et al., 2002; Hastings et al., 2005), growing-season length (Xu and Baldocchi, 2004; Ma et al., 2007), seasonal

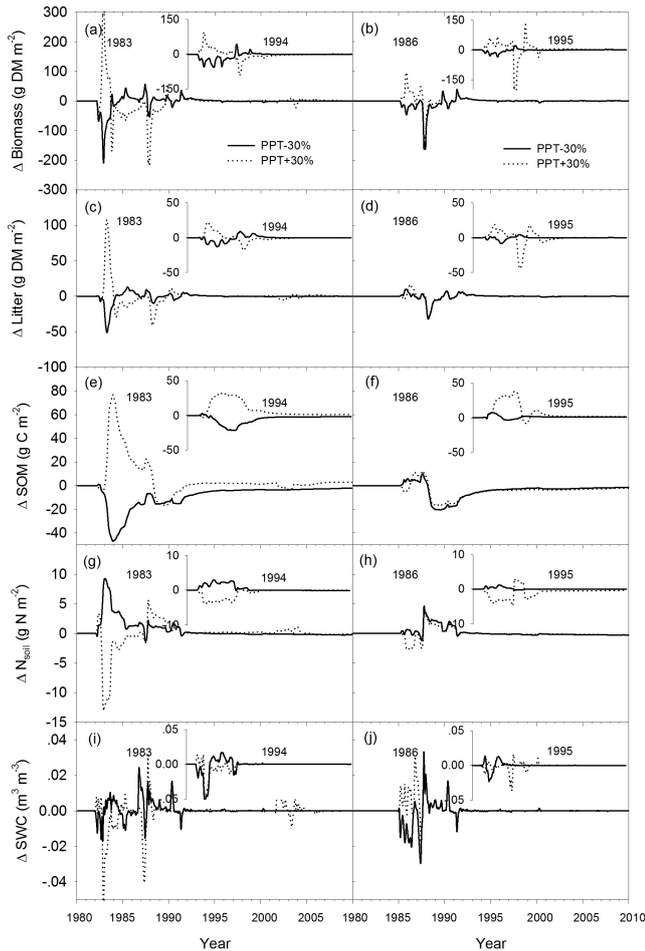


Figure 7. Interannual precipitation legacy effects on resource pool dynamics. Left panels show the legacy effects on pool dynamics in 2 representative wet years, while right panels for 2 representative dry years. Legacy effects on pool size (e.g., Δ Biomass) are quantified as the difference between the current-year pool size with previous-year PPT change and that without previous-year PPT change. Solid lines represent a 30 % decrease, while dashed lines represent a 30 % increase in the previous-year precipitation (PPT). SOM represents soil organic matter, N_{soil} soil mineral nitrogen, and SWC soil water content.

drought (Scott et al., 2009, 2010; Hamerlynck et al., 2013), and other factors such as temperature and vegetation composition (Hui et al., 2003; Hamerlynck et al., 2010; Barron-Gafford et al., 2012; Scott et al., 2014). These interactions are shown by several examples from our simulations. While PPT was wetter than normal in 1987 (537 mm), the NEP was $-85 \text{ g C m}^{-2} \text{ yr}^{-1}$ (a C source), due to the negative wet-legacy impacts on NEP from several previous wet years before (1982–1985; see Fig. 6h). PPT was nearly normal in 2008 (402 mm), but the simulated NEP was $80 \text{ g C m}^{-2} \text{ yr}^{-1}$ (a C sink), due to the positive dry-legacy impacts on NEP from several previous dry years (2002–2007; see Fig. 6g). Our findings of substantial PPT legacy effects are consis-

tent with a recent analysis of 14 years (1997–2011) of eddy-covariance measurements, where Zielis et al. (2014) reported that inclusion of previous year's weather (PPT and temperature) into the linear predicting models for NEP increased the explained variance to 53 % compared to 20 % without accounting for previous year's weather, indicating that previous year's weather also played an important role in determining the C balance of the subalpine spruce forest in Switzerland. Although response patterns generated from this simulation study compared well with previous field observations, there exists no field study that, to our knowledge, provides a similarly comprehensive analysis of PPT legacies. The simulation experimental design of this study provides helpful insights into designing field manipulative experiments to further test the modeled patterns by focusing on contrasting wet and dry legacies, separating ecosystem production and decomposition, and exploring the difference in prior and current PPT on the magnitude of the PPT legacy effect.

4.2 Potential mechanisms of the modeled PPT legacies

There are three basic mechanisms explaining why PPT legacy impacts can occur in a model system like PALS. First, the rate of C fluxes is a function of not only various environmental factors (e.g., PPT and temperature) but also the pool size itself. For example, soil heterotrophic CO_2 efflux (R_h) rate is a product of the decomposition coefficient, two scalar functions accounting for temperature and moisture influences, and also the size of the SOM pool (Kemp et al., 2003; Shen et al., 2009). Change in the SOM pool size from previous PPT thereby affects current R_h . Second, different C pools have different turnover rates that determine whether biogeochemical materials (e.g., biomass or SOM) can be carried over. If the material produced in a previous year has a turnover rate of less than 1 year, it will not be carried over to the next year to form a legacy impact as explained in the first mechanism. In addition, the turnover rates of different C pools also determine legacy duration. For example, SOM pools in the model have relatively slower turnover rates than biomass pools (Shen et al., 2005, 2008b), thus resulting in the longer-lasting legacy impacts on SOM than on biomass or litter pools (Figs. 5 and 7). Third, the interactions between C fluxes and resource (e.g., N or water) availability also determine the direction and magnitude of legacy effects. For example, N carryover as a legacy of a prior dry period (Fig. 5g, h) can impose impacts on the current-period GEP only when the current-period PPT is not so limiting (Fig. 3b). These are the general mechanisms explaining the occurrence of the modeled PPT legacies from a systems perspective. Below we discuss more specifically the major patterns and the responsible biogeochemical carryovers found in this study.

An intuitive first explanation of the simulated wet legacies would be the carryover of water. However, in most cases soil water carryover did not occur because the wet legacies on SWC were mostly negative or close to zero at the beginning

of the current period/year (Figs. 5i–j; 7i–j). Soil water carryover was therefore not the major contributor to the modeled PPT legacy effects at interdecadal and interannual scales. This simulation result corroborates field studies that have shown that carryover of water across long temporal scales is rare in dryland ecosystems, because the rainy growing seasons or wet years are often separated by dry dormant seasons or dry years resulting in short residence times (Oesterheld et al., 2001; Reichmann et al., 2013a; Scott et al., 2014).

The carryover of soil N (N_{soil}) is mainly responsible for the modeled GEP responses. In the PALS model, the photosynthetic rate is linearly related to N availability if plant N demand is not fulfilled (Reynolds et al., 2004; Shen et al., 2005). Therefore, the enhanced N_{soil} from dry legacies (Figs. 5g, h and 7g, h) generated mostly positive responses of GEP (Figs. 3a, b and 6c). The simulated dry legacies increased N_{soil} mainly through suppressed plant growth (e.g., the reduced biomass and litter production shown in Figs. 5 and 7) that limited N uptake, which is consistent with the results of many field measurements that N_{soil} accumulates under drought conditions (Reynolds et al., 1999; Yahdjian et al., 2006; Yahdjian and Sala, 2010; de Vries et al., 2012; Evans and Burke, 2013; Reichmann et al., 2013b). Although diverse mechanisms of inorganic N accumulation during dry periods have been proposed in field studies – such as the diffusion restriction of N ions in thin water films of dry soil, the reduced N immobilization by microbial growth and plant uptake, and the reduced N loss from the soil via leaching (Yahdjian et al., 2006) – our simulation results suggest that reduced plant uptake may be the main contributor to the N_{soil} accumulation during dry periods. Given the accumulated N_{soil} as a dry legacy, how ecosystem C fluxes such as GEP respond to this dry legacy may be influenced by current PPT conditions. When current PPT conditions were favorable (e.g., the increasing current-period PPT treatment shown in Fig. 3b and the relatively wet years shown in Fig. 6c), GEP mostly increased with a dry legacy (or the accumulated N) because both N and H_2O availabilities were favorable for plant growth (or GEP). Contrastingly, when current PPT conditions were unfavorable (e.g., the decreasing current-period PPT treatment shown in Fig. 3a and the relatively dry years shown in Fig. 6c), the GEP responses could be reduced because of the constrained plant growth and the reduced biomass in previous dry years (see Figs. 5c and 7b).

Similarly, the mostly negative responses of GEP to wet legacies (see Figs. 3a, b and 6d) can be explained by the reduced N_{soil} (Figs. 5g, h and 7g, h). The decrease of N_{soil} with increasing PPT in the PALS model is mainly attributed to the increases in plant N uptake and the N leaching loss that is calculated as a linear function of PPT amount (Shen et al., 2005). Similar to our simulation results, several field studies found that N uptake increases and N_{soil} decreases under wet conditions in dryland ecosystems (McCulley et al., 2009; McCalley and Sparks, 2009; Yahdjian and Sala, 2010; Reichmann et al., 2013b). However, contrary to our model assump-

tion that N leaching loss is greater in wet than in dry years, some recent field studies have reported that the N leaching loss actually is higher in dry than in wet years or at wet sites (McCulley et al., 2009; Evans et al., 2013; Reichmann et al., 2013b; Homyak et al., 2014), resulting in a more “open” N cycle under drier conditions. If these recent field study results are also true for our semi-desert savanna ecosystem, the model assumption could potentially cause an overestimation of N_{soil} carryover effects as shown in Figs. 3 and 6. Further studies are needed to discriminate the relative contributions of different N processes (e.g., plant uptake, microbial immobilization and mineralization, denitrification, ammonia volatilization, and leaching) to the dynamics of soil inorganic N pools. Nevertheless, this simulation analysis highlights the importance of interactions between N and H_2O availabilities in creating the legacy impacts of PPT and in shaping the temporal variability of dryland ecosystem C fluxes.

The carryover of organic material (biomass, litter and SOM) is mainly responsible for the modeled R_e responses. In the PALS model, the autotrophic (R_a) and heterotrophic (R_h) respiration rates are linearly related to the size of biomass, litter and SOM pools (Kemp et al., 2003; Shen et al., 2008a, 2009). The previous wet condition benefited biomass, litter and SOM accumulation (Figs. 5 and 7), which resulted in the mostly positive wet-legacy impacts on R_e (Figs. 3c, d and 6f). Conversely, the dry legacy decreased these pools (Figs. 5 and 7) and therefore resulted in the mostly negative dry-legacy impacts on R_e (Figs. 3c, d and 6e). Contrary to our simulation results that dry legacies are mostly negative on SOM and R_h , some field studies suggest that the labile C resulting from litter decomposition in a dry season may stimulate R_h in the following wet season (Jenerette et al., 2008; Scott et al., 2009; Ma et al., 2012). This is likely because the labile soil C pool in the PALS model only accounts for $\sim 3\%$ of the total SOM and has a very short residence time (1.7 year; see Table S1); small amounts of seasonal labile C carryover therefore may not exert obvious legacy impacts on the total SOM pool size and R_h across interannual and interdecadal scales. These results imply that the PPT legacy effects differ in direction and magnitude, depending on the type of C fluxes under consideration, the type of legacies (i.e., dry vs. wet), and the temporal scale of analysis.

Several lines of future research will likely be needed to continue improving the understanding of ecosystem legacy dynamics. Structural shifts in vegetation composition such as woody-plant encroachment (Potts et al., 2008; Scott et al., 2014), exotic-species invasion (Hammerlynck et al., 2010; Scott et al., 2010), and changes in microbial communities (de Vries et al., 2012; Evans and Wallenstein, 2012; Collins et al., 2014) may also interact with the biogeochemical processes to shape the PPT legacy effects on the temporal variability of dryland C fluxes. Furthermore, we need to better understand the legacy effects of extreme events such as the cool-GS drought in 2006 (see Fig. 1a) so that these important events can be adequately simulated. This cool-GS

drought may have caused increased plant mortality as reported for a semi-desert grassland near our study site (Scott et al., 2010; Hamerlynck et al., 2013), but that is poorly represented in the model and may have caused the overestimation of the modeled GEP in comparison with the observation (see Fig. S3b). Finally, our approach that uses a highly resolved process model provides information complementary to contrasting analytical approaches that evaluate ecosystem responses to statistical rainfall regimes (Rodrigo-Iturbe et al., 2006; Katul et al., 2007; Porporato and Rodríguez-Iturbe, 2013). Improvement of these alternative modeling approaches is needed to understand both general and specific ecosystem responses to changing PPT regimes at temporal scales from events to decades.

5 Conclusions

We learned through this simulation analysis that (1) previous PPT conditions can impose substantial legacy impacts on the C balance of dryland ecosystems, with dry legacies fostering more current C sequestration and wet legacies causing more current C release; (2) the responses of ecosystem C fluxes to the simulated dry and wet legacies are mostly opposite in direction and asymmetrical in magnitude, with dry legacies being greater for GEP than for R_e and wet legacies being greater for R_e than for GEP; (3) the carryover of N_{soil} is mainly responsible for the GEP responses, and the carryovers of biomass, litter and SOM are mainly responsible for the R_e responses; and (4) the simulated PPT legacy effects can last for several years even with a 1-year PPT change, and therefore the direction and magnitude of interannual PPT legacy effects are less predictable than interdecadal ones. These simulation results suggest that dryland ecosystems such as these in the southwestern USA may emit more C that was sequestered in the past into the atmosphere with the predicted drying trend in the region (Seager et al., 2007; Solomon et al., 2007). The temporal variability of ecosystem C fluxes may be further intensified in the region due to the increasing PPT variability and the associated legacy impacts.

The Supplement related to this article is available online at doi:10.5194/bg-13-425-2016-supplement.

Acknowledgements. We thank the USDA Agricultural Research Service (ARS), Southwest Watershed Research Center (SWRC) and the Arizona Meteorological Network (AZMET) for making their meteorological data open to access. W. Shen acknowledges the financial support from the Natural Science Foundation of China (31130011, 31425005 and 31290222), the Major State Basic Research Development Program (973 Planning Program 2011CB403206), and the Natural Science Foundation of Guangdong Province, China (S2012020011084). Partial funding for the

Santa Rita mesquite savanna eddy-covariance site is provided by the US Department of Energy AmeriFlux Office (grant DE-AC02-05CH11231).

Edited by: C. A. Williams

References

- Asner, G. P., Archer, S., Hughes, R. F., Ansley, R. J., and Wessman, C. A.: Net changes in regional woody vegetation cover and carbon storage in Texas Drylands, 1937–1999, *Glob. Change Biol.*, 9, 316–335, 2003.
- Barron-Gafford, G. A., Scott, R. L., Jenerette, G. D., Hamerlynck, E. P., and Huxman, T. E.: Temperature and precipitation controls over leaf- and ecosystem-level CO_2 flux along a woody plant encroachment gradient, *Glob. Change Biol.*, 18, 1389–1400, 2012.
- Barron-Gafford, G. A., Scott, R. L., Jenerette, G. D., Hamerlynck, E. P., and Huxman, T. E.: Landscape and environmental controls over leaf and ecosystem carbon dioxide fluxes under woody plant expansion, *J. Ecol.*, 101, 1471–1483, 2013.
- Bates, B. C., Kundzewicz, Z. W., Wu, S., and Palutikof, J. P. (Eds.): *Climate Change and Water*. Technical Paper of the Intergovernmental Panel on Climate Change, IPCC Secretariat, Geneva, 2008.
- Borken, W. and Matzner, E.: Reappraisal of drying and wetting effects on C and N mineralization and fluxes in soils, *Glob. Change Biol.*, 15, 808–824, 2009.
- Cable, J. M., Ogle, K., Barron-Gafford, G. A., Bentley, L. P., Cable, W. L., Scott, R. L., Williams, D. G., and Huxman, T. E.: Antecedent conditions influence soil respiration differences in shrub and grass patches, *Ecosystems*, 16, 1230–1247, 2013.
- Collins, S. L., Belnap, J., Grimm, N. B., Rudgers, J. A., Dahm, C. N., D’Odorico, P., Litvak, M., Natvig, D. O., Peters, D. C., Pockman, W. T., Sinsabaugh, R. L., and Wolf, B. O.: A multiscale, hierarchical model of pulse dynamics in arid-land ecosystems, *Annu. Rev. Ecol. Evol. S.*, 45, 397–419, 2014.
- Cook, B. I. and Seager, R.: The response of the North American Monsoon to increased greenhouse gas forcing, *J. Geophys. Res.-Atmos.*, 118, 1690–1699, 2013.
- de Vries, F. T., Liiri, M. E., Bjornlund, L., Setälä, H. M., Christensen, S., and Bardgett, R. D.: Legacy effects of drought on plant growth and the soil food web, *Oecologia*, 170, 821–833, 2012.
- Diffenbaugh, N. S., Giorgi, F., and Pal, J. S.: Climate change hotspots in the United States, *Geophys. Res. Lett.*, 35, L16709, doi:10.1029/2008GL035075, 2008.
- Evans, S. E. and Burke, I. C.: Carbon and nitrogen decoupling under an 11-year drought in the shortgrass steppe, *Ecosystems*, 16, 20–33, 2013.
- Evans, S. E. and Wallenstein, M. D.: Soil microbial community response to drying and rewetting stress: does historical precipitation regime matter?, *Biogeochemistry*, 109, 101–116, 2012.
- Field, C. B., Behrenfeld, M. J., Randerson, J. T., and Falkowski, P.: Primary production of the biosphere: Integrating terrestrial and oceanic components, *Science*, 281, 237–240, 1998.
- Flanagan, L. B., Wever, L. A., and Carlson, P. J.: Seasonal and interannual variation in carbon dioxide exchange and carbon balance

- in a northern temperate grassland, *Glob. Change Biol.*, 8, 599–615, 2002.
- Gao, Q. and Reynolds, J. F.: Historical shrub-grass transitions in the northern Chihuahuan Desert: modeling the effects of shifting rainfall seasonality and event size over a landscape gradient, *Glob. Change Biol.*, 9, 1475–1493, 2003.
- Hamerlynck, E. P., Scott, R. L., Moran, M. S., Keefer, T. O., and Huxman, T. E.: Growing season ecosystem and leaf-level gas exchange of an exotic and native semiarid bunchgrass, *Oecologia*, 163, 561–570, 2010.
- Hamerlynck, E. P., Scott, R. L., and Barron-Gafford, G. A.: Consequences of cool-season drought-induced plant mortality to Chihuahuan Desert grassland ecosystem and soil respiration dynamics, *Ecosystems*, 16, 1178–1191, 2013.
- Hastings, S. J., Oechel, W. C., and Muhlia-Melo, A.: Diurnal, seasonal and annual variation in the net ecosystem CO₂ exchange of a desert shrub community (*Sarcocaulis*) in Baja California, Mexico, *Glob. Change Biol.*, 11, 927–939, 2005.
- Heisler, J. L. and Weltzin, J. F.: Variability matters: towards a perspective on the influence of precipitation on terrestrial ecosystems, *New Phytol.*, 172, 189–192, 2006.
- Heisler-White, J. L., Knapp, A. K., and Kelly, E. F.: Increasing precipitation event size increases aboveground net primary productivity in a semi-arid grassland, *Oecologia*, 158, 129–140, 2008.
- Homyak, P. M., Sickman, J. O., Miller, A. E., Melack, J. M., Meixner, T., and Schimel, J. P.: Assessing nitrogen-saturation in a seasonally dry Chaparral watershed: Limitations of traditional indicators of N-saturation, *Ecosystems*, 17, 1286–1305, 2014.
- Houghton, R. A., Hackler, J. L., and Lawrence, K. T.: The US carbon budget: Contributions from land-use change, *Science*, 285, 574–578, 1999.
- Hui, D. F., Luo, Y. Q., and Katul, G.: Partitioning interannual variability in net ecosystem exchange into climatic variability and functional change, *Tree Physiol.*, 23, 433–442, 2003.
- Huxman, T. E., Cable, J. M., Ignace, D. D., Eilts, J. A., English, N. B., Weltzin, J., and Willimas, D. G.: Response of net ecosystem gas exchange to a simulated precipitation pulse in a semi-arid grassland: the role of native versus non-native grasses and soil texture, *Oecologia*, 141, 295–305, 2004a.
- Huxman, T. E., Snyder, K. A., Tissue, D., Leffler, A. J., Ogle, K., Pockman, W. T., Sandquist, D. R., Potts, D. L., and Schwinning, S.: Precipitation pulses and carbon fluxes in semiarid and arid ecosystems, *Oecologia*, 141, 254–268, 2004b.
- Huxman, T. E., Smith, M. D., Fay, P. A., Knapp, A. K., Shaw, M. R., Loik, M. E., Smith, S. D., Tissue, D. T., Zak, J. C., Weltzin, J. F., Pockman, W. T., Sala, O. E., Haddad, B. M., Harte, J., Kock, G. W., Schwinning, S., Small, E. E., and Willimas, D. G.: Convergence across biomes to a common rain-use efficiency, *Nature*, 429, 651–654, 2004c.
- Jenerette, G. D., Scott, R. L., and Huxman, T. E.: Whole ecosystem metabolic pulses following precipitation events, *Funct. Ecol.*, 22, 924–930, 2008.
- Jenerette, G. D., Scott, R. L., and Huete, A. R.: Functional differences between summer and winter season rain assessed with MODIS-derived phenology in a semi-arid region, *J. Veg. Sci.*, 21, 16–30, 2010.
- Jenerette, G. D., Barron-Gafford, G. A., Guswa, A. J., McDonnell, J. J., and Villegas, J. C.: Organization of complexity in water limited ecohydrology, *Ecohydrology*, 5, 184–199, 2012.
- Jobbagy, E. G. and Sala, O. E.: Controls of grass and shrub above-ground production in the Patagonian steppe, *Ecol. Appl.*, 10, 541–549, 2000.
- Katul, G. A., Proporato, A., and Oren R.: Stochastic dynamics of plant-water interactions, *Annu. Rev. Ecol. Evol. S.*, 38, 767–791, 2007.
- Kemp, P. R., Reynolds, J. F., Pachepsky, Y., and Chen, J. L.: A comparative modeling study of soil water dynamics in a desert ecosystem, *Water Resour. Res.*, 33, 73–90, 1997.
- Kemp, P. R., Reynolds, J. F., Virginia, R. A., and Whitford, W. G.: Decomposition of leaf and root litter of Chihuahuan desert shrubs: effects of three years of summer drought, *J. Arid Environ.*, 53, 21–39, 2003.
- Knapp, A. K., Fay, P. A., Blair, J. M., Collins, S. L., Smith, M. D., Carlisle, J. D., Harper, C. W., Danner, B. T., Lett, M. S., and McCarron, J. K.: Rainfall variability, carbon cycling, and plant species diversity in a mesic grassland, *Science*, 298, 2202–2205, 2002.
- Lauenroth, W. K. and Sala, O. E.: Long-term forage production of North-American shortgrass steppe, *Ecol. Appl.*, 2, 397–403, 1992.
- Ma, S. Y., Baldocchi, D. D., Xu, L. K., and Hehn, T.: Inter-annual variability in carbon dioxide exchange of an oak/grass savanna and open grassland in California, *Agr. Forest Meteorol.*, 147, 157–171, 2007.
- Ma, S. Y., Baldocchi, D. D., Hatala, J. A., Detto, M., and Yuste, J. C.: Are rain-induced ecosystem respiration pulses enhanced by legacies of antecedent photodegradation in semi-arid environments?, *Agr. Forest Meteorol.*, 154, 203–213, 2012.
- Maestre, F. T., Salguero-Gomez, R., and Quero, J. L.: It is getting hotter in here: determining and projecting the impacts of global environmental change on drylands Introduction, *Philos. T. R. Soc. B.*, 367, 3062–3075, 2012.
- McCalley, C. K. and Sparks, J. P.: Abiotic gas formation drives nitrogen loss from a desert ecosystem, *Science*, 326, 837–840, 2009.
- McCulley, R. L., Burke, I. C., and Lauenroth, W. K.: Conservation of nitrogen increases with precipitation across a major grassland gradient in the Central Great Plains of North America, *Oecologia*, 159, 571–581, 2009.
- Monger, C., Sala, O. E., Duniway, M. C., Goldfus, H., Meir, I. A., Poch, R. M., Throop, H. L., and Vivoni, E. R.: Legacy effects in linked ecological-soil-geomorphic systems of drylands, *Front. Ecol. Environ.*, 13, 13–19, 2015.
- Noy-Meir, I.: Desert ecosystems: environment and producers, *Annu. Rev. Ecol. Evol. S.*, 4, 25–51, 1973.
- Oosterheld, M., Loreti, J., Semmartin, M., and Sala, O. E.: Inter-annual variation in primary production of a semi-arid grassland related to previous-year production, *J. Veg. Sci.*, 12, 137–142, 2001.
- Ogle, K., Barber, J. J., Barron-Gafford, G. A., Bentley, L. P., Young, J. M., Huxman, T. E., Loik, M. E., and Tissue, D. T.: Quantifying ecological memory in plant and ecosystem processes, *Ecol. Lett.*, 18, 221–235, 2015.
- Oikawa, P. Y., Gratz, D. A., Chatterjee, A., Eberwein, J. E., Allsman, L. A., and Jenerette, G. D.: Unifying soil respiration pulses, inhibition, and temperature hysteresis through dynamics of labile carbon and soil O₂, *J. Geophys. Res.-Biogeo.*, 115, 521–536, 2014.

- Parton, W. J., Scurlock, J. M. O., Ojima, D. S., Gilmanov, T. G., Scholes, R. J., Schimel, D. S., Kirchner, T., Menaut, J. C., Seastedt, T., Moya, E. G., Kamalrut, A., and Kinyamario, J. I.: Observations and modeling of biomass and soil organic-matter dynamics for the grassland biome worldwide, *Global Biogeochem. Cy.*, 7, 785–809, 1993.
- Paruelo, J. M., Lauenroth, W. K., Burke, I. C., and Sala, O. E.: Grassland precipitation-use efficiency varies across a resource gradient, *Ecosystems*, 2, 64–68, 1999.
- Peters, D. P. C., Yao, J., Sala, O. E., and Anderson, J. P.: Directional climate change and potential reversal of desertification in arid and semiarid ecosystems, *Glob. Change Biol.*, 18, 151–163, 2012.
- Porporato, A. and Rodriguez-Iturbe, I.: Ecohydrology bearings – invited commentary from random variability to ordered structures: a search for general synthesis in ecohydrology, *Ecohydrology*, 6, 333–342, 2013.
- Porporato, A., Daly, E., and Rodriguez-Iburbe, I.: Soil water balance and ecosystem response to climate change, *Am. Nat.*, 164, 625–632, 2004.
- Potts, D. L., Scott, R. L., Cable, J. M., Huxman, T. E., and Williams, D. G.: Sensitivity of mesquite shrubland CO₂ exchange to precipitation in contrasting landscape settings, *Ecology*, 89, 2900–2910, 2008.
- Reichmann, L. G., Sala, O. E., and Peters, D. P. C.: Precipitation legacies in desert grassland primary production occur through previous-year tiller density, *Ecology*, 94, 435–443, 2013a.
- Reichmann, L. G., Sala, O. E., and Peters, D. P. C.: Water controls on nitrogen transformations and stocks in an arid ecosystem, *Ecosphere*, 4, 1–17, 2013b.
- Reynolds, J. F., Virginia, R. A., and Schlesinger, W. H.: Defining functional types for models of desertification, in: *Plant Functional Types: Their Relevance to Ecosystem Properties and Global Change*, edited by: Shugart, T. M. and Woodward, F. I., Cambridge University Press, Cambridge, 1997.
- Reynolds, J. F., Virginia, R. A., Kemp, P. R., de Soyza, A. G., and Tremmel, D. C.: Impact of drought on desert shrubs: Effects of seasonality and degree of resource island development, *Ecol. Monogr.*, 69, 69–106, 1999.
- Reynolds, J. F., Kemp, P. R., and Tenhunen, J. D.: Effects of long-term rainfall variability on evapotranspiration and soil water distribution in the Chihuahuan Desert: A modeling analysis, *Plant Ecol.*, 150, 145–159, 2000.
- Reynolds, J. F., Kemp, P. R., Ogle, K., and Fernandez, R. J.: Modifying the “pulse-reserve” paradigm for deserts of North America: precipitation pulses, soil water, and plant responses, *Oecologia*, 141, 194–210, 2004.
- Reynolds, J. F., Stafford Smith, D. M., Lambin, E. F., Turner, B. L., Mortimore, M., Batterbury, S. P. J., Downing, T. E., Dowlatabadi, H., Fernandez, R. J., Herrick, J. E., Huber-Sannwald, E., Jiang, H., Leemans, R., Lynam, T., Maestre, F. T., Ayarza, M., and Walker, B.: Global desertification: Building a science for dryland development, *Science*, 316, 847–851, 2007.
- Rodriguez-Iturbe, I., Isham, V., Cox, D. R., Manfreda, S., and Porporato, A.: Space-time modeling of soil moisture: Stochastic rainfall forcing with heterogeneous vegetation, *Water Resour. Res.*, 42, W06D05, doi:10.1029/2005WR004497, 2006.
- Sala, O. E., Lauenroth, W. K., and Parton, W. J.: Long-term soil-water dynamics in the shortgrass steppe, *Ecology*, 73, 1175–1181, 1992.
- Sala, O. E., Gherardi, L. A., Reichmann, L., Jobbagy, E., and Peters, D.: Legacies of precipitation fluctuations on primary production: theory and data synthesis, *Philos. T. R. Soc. B.*, 367, 3135–3144, 2012.
- Schlesinger, W. H.: *Biogeochemistry: An Analysis of Global Change*, Academic Press Inc, San Diego, CA, 1991.
- Scott, R. L., Jenerette, G. D., Potts, D. L., and Huxman, T. E.: Effects of seasonal drought on net carbon dioxide exchange from a woody-plant-encroached semiarid grassland, *J. Geophys. Res.-Biogeol.*, 114, G04004, doi:10.1029/2008JG000900, 2009.
- Scott, R. L., Hamerlynck, E. P., Jenerette, G. D., Moran, M. S., and Barron-Gafford, G. A.: Carbon dioxide exchange in a semidesert grassland through drought-induced vegetation change, *J. Geophys. Res.-Biogeol.*, 115, G03026, doi:10.1029/2010JG001348, 2010.
- Scott, R. L., Huxman, T. E., Barron-Gafford, G. A., Jenerette, G. D., Young, J. M., and Hamerlynck, E. P.: When vegetation change alters ecosystem water availability, *Glob. Change Biol.*, 20, 2198–2210, 2014.
- Seager, R., Ting, M. F., Held, I., Kushnir, Y., Lu, J., Vecchi, G., Huang, H. P., Harnik, N., Leetmaa, A., Lau, N. C., Li, C. H., Velez, J., and Naik, N.: Model projections of an imminent transition to a more arid climate in southwestern North America, *Science*, 316, 1181–1184, 2007.
- Shen, W. J., Wu, J. G., Kemp, P. R., Reynolds, J. F., and Grimm, N. B.: Simulating the dynamics of primary productivity of a Sonoran ecosystem: Model parameterization and validation, *Ecol. Model.*, 189, 1–24, 2005.
- Shen, W. J., Jenerette, G. D., Hui, D. F., Phillips, R. P., and Ren, H.: Effects of changing precipitation regimes on dryland soil respiration and C pool dynamics at rainfall event, seasonal and interannual scales, *J. Geophys. Res.-Biogeol.*, 113, G03024, doi:10.1029/2008JG000685, 2008a.
- Shen, W. J., Wu, J. G., Grimm, N. B., and Hope, D.: Effects of urbanization-induced environmental changes on ecosystem functioning in the Phoenix metropolitan region, USA, *Ecosystems*, 11, 138–155, 2008b.
- Shen, W. J., Reynolds, J. F., and Hui, D. F.: Responses of dryland soil respiration and soil carbon pool size to abrupt vs. gradual and individual vs. combined changes in soil temperature, precipitation, and atmospheric [CO₂]: a simulation analysis, *Glob. Change Biol.*, 15, 2274–2294, 2009.
- Sherry, R. A., Weng, E. S., Arnone, J. A., Johnson, D. W., Schimel, D. S., Verburg, P. S., Wallace, L. L., and Luo, Y. Q.: Lagged effects of experimental warming and doubled precipitation on annual and seasonal aboveground biomass production in a tallgrass prairie, *Glob. Change Biol.*, 14, 2923–2936, 2008.
- Solomon, S., Qin, D., Manning, M., Chen, Z., Marquis, M., Averyt, K. B., Tignor, M., and Miller, H. L. (Eds.): *Climate Change 2007: The Physical Science Basis*, Cambridge University Press, Cambridge, 2007.
- Sponseller, R. A.: Precipitation pulses and soil CO₂ flux in a Sonoran Desert ecosystem, *Glob. Change Biol.*, 13, 426–436, 2007.
- Swemmer, A. M., Knapp, A. K., and Snyman, H. A.: Intra-seasonal precipitation patterns and above-ground productivity in three perennial grasslands, *J. Ecol.*, 95, 780–788, 2007.

- Vargas, R., Baldocchi, D. D., Allen, M. F., Bahn, M., Black, T. A., Collins, S. L., Yuste, J. C., Hirano, T., Jassal, R. S., Pumpanen, J., and Tang, J. W.: Looking deeper into the soil: biophysical controls and seasonal lags of soil CO₂ production and efflux, *Ecol. Appl.*, 20, 1569–1582, 2010.
- Wiegand, T., Snyman, H. A., Kellner, K., and Paruelo, J. M.: Do grasslands have a memory: Modeling phytomass production of a semiarid South African grassland, *Ecosystems*, 7, 243–258, 2004.
- Willimas, C. A. and Albertson, J. D.: Dynamical effects of the statistical structure of annual rainfall on dryland vegetation, *Glob. Change Biol.*, 12, 777–792, 2006.
- Williams, C. A., Hanan, N., Scholes, R. J., and Kutsch, W.: Complexity in water and carbon dioxide fluxes following rain pulses in an african savanna, *Oecologia*, 161, 469–480, 2009.
- Xu, L. K. and Baldocchi, D. D.: Seasonal variation in carbon dioxide exchange over a Mediterranean annual grassland in California, *Agr. Forest Meteorol.*, 123, 79–96, 2004.
- Xu, L. K., Baldocchi, D. D., and Tang, J. W.: How soil moisture, rain pulses, and growth alter the response of ecosystem respiration to temperature, *Global Biogeochem. Cy.*, 18, GB4002, doi:10.1029/2004GB002281, 2004.
- Yahdjian, L. and Sala, O. E.: Vegetation structure constrains primary production response to water availability in the Patagonian steppe, *Ecology*, 87, 952–962, 2006.
- Yahdjian, L. and Sala, O. E.: Size of precipitation pulses controls nitrogen transformation and losses in an arid Patagonian ecosystem, *Ecosystems*, 13, 575–585, 2010.
- Yahdjian, L., Sala, O. E., and Austin, A. T.: Differential controls of water input on litter decomposition and nitrogen dynamics in the Patagonian Steppe, *Ecosystems*, 9, 128–141, 2006.
- Zielis, S., Etzold, S., Zweifel, R., Eugster, W., Haeni, M., and Buchmann, N.: NEP of a Swiss subalpine forest is significantly driven not only by current but also by previous year's weather, *Biogeosciences*, 11, 1627–1635, doi:10.5194/bg-11-1627-2014, 2014.