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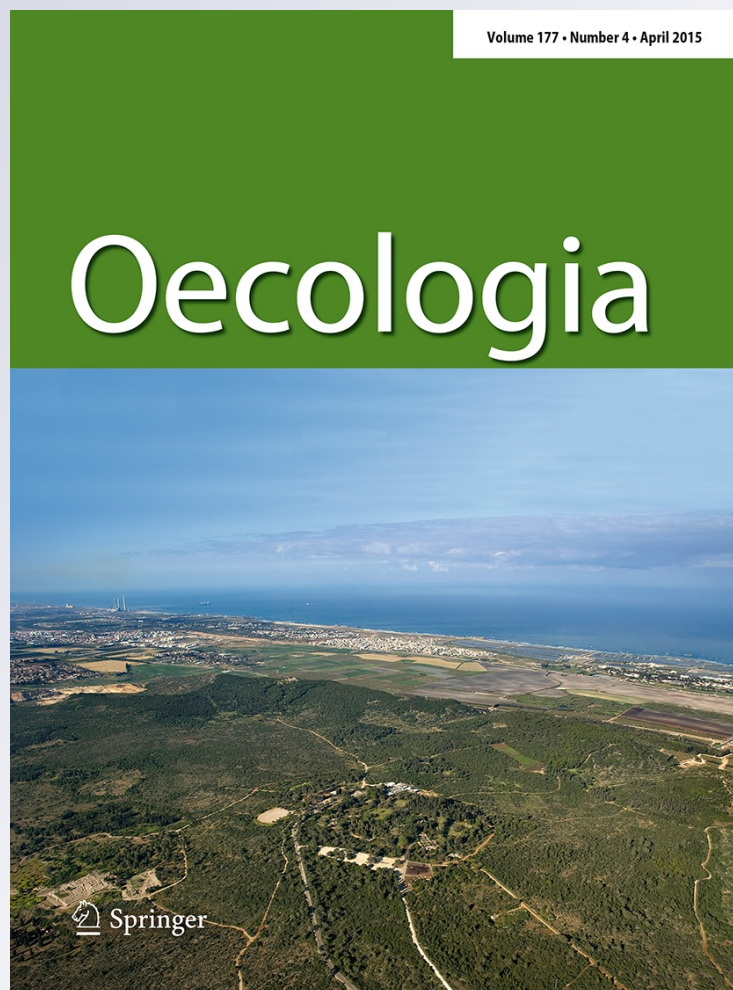
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# Climatic controls of aboveground net primary production in semi-arid grasslands along a latitudinal gradient portend low sensitivity to warming

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**Abstract** Although climate models forecast warmer temperatures with a high degree of certainty, precipitation is the primary driver of aboveground net primary production (ANPP) in most grasslands. Conversely, variations in temperature seldom are related to patterns of ANPP. Thus forecasting responses to warming is a challenge, and raises the question: how sensitive will grassland ANPP be to warming? We evaluated climate and multi-year ANPP data (67 years) from eight western US grasslands arrayed along mean annual temperature (MAT; ~7–14 °C) and mean annual precipitation (MAP; ~250–500 mm) gradients. We

used regression and analysis of covariance to assess relationships between ANPP and temperature, as well as precipitation (annual and growing season) to evaluate temperature sensitivity of ANPP. We also related ANPP to the standardized precipitation evaporation index (SPEI), which combines precipitation and evapotranspiration to better represent moisture available for plant growth. Regression models indicated that variation in growing season temperature was negatively related to total and graminoid ANPP, but precipitation was a stronger predictor than temperature. Growing season temperature was also a significant parameter in more complex models, but again precipitation was consistently a stronger predictor of ANPP. Surprisingly, neither annual nor growing season SPEI were as strongly related to ANPP as precipitation. We conclude that forecasted warming likely will affect ANPP in these grasslands, but that predicting temperature effects from natural climatic gradients is difficult. This is because, unlike precipitation, warming effects can be positive or negative and moderated by shifts in the  $C_3/C_4$  ratios of plant communities.

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## Introduction

Of the predicted climatic changes forecast to occur due to anthropogenic disruption of the global climate system, the most certain of these is for increased atmospheric temperatures (IPCC 2013). Moreover, there is abundant evidence that warming is already occurring across much of the globe (Rummukainen 2012). While virtually every biome may be impacted by warming, the degree of impact is likely to

vary, and it is therefore essential to understand how sensitive different ecosystems are to temperature change. For grassland ecosystems, precipitation has long been considered the dominant climatic driver of ecosystem function over much of the globe (Harpole et al. 2007; Merbold et al. 2009; Jung et al. 2011; Zhang et al. 2014). This has been repeatedly demonstrated for aboveground net primary production (ANPP) in site-based observational studies (Sala et al. 1992; Briggs and Knapp 1995; Jobbágy et al. 2002; Derner et al. 2008), studies across broad geographic gradients (Webb et al. 1978; Knapp and Smith 2001; Reed et al. 2009; Guo et al. 2012), and in a number of experiments in grassland ecosystems (Yahdjian and Sala 2006; Sherry et al. 2008; Cherwin and Knapp 2012). Indeed, at large spatial scales, mean annual precipitation (MAP) may account for 90 % of the variation in ANPP (Sala et al. 1988).

Temperature on the other hand is seldom statistically related to spatial or temporal variation in ANPP in grasslands (Sala et al. 1988; Del Grosso and Parton 2008; Guo et al. 2014) except perhaps at global scales (e.g., Whittaker 1975; Frank and Inouye 1994; Gang et al. 2013). In experiments with warming treatments, results have been mixed with some studies showing positive responses (Lin et al. 2010), others negative responses (Niu et al. 2008, 2011; Cantarel et al. 2013), while others found little response (Fay et al. 2011; Xu et al. 2013) to increased temperatures. In one of the few studies demonstrating temperature effects on grassland ANPP across natural climatic gradients, Epstein et al. (1997) reported negative effects of temperature on ANPP in the central US. Although regression coefficients were not strong (average  $r^2 \sim 0.2\text{--}0.3$ ) temperature effects were greater in drier (<600 mm MAP) than more mesic grasslands. Similarly, Wu et al. (2011) in a meta-analysis of warming experiments reported that warming often had a negative effect on productivity unless additional precipitation was added. These and other studies suggest that the primary effect of higher temperatures in many grassland ecosystems may be indirect through negative effects on water balance (Peñuelas et al. 2007; Xu et al. 2012; Dulamsuren et al. 2013).

Grasslands cover approximately a third of the earth's land surface and provide vital ecosystem services that include the sequestration of C in the soil, forage production and habitat critical for many species (Lal 2004; Chou et al. 2008). Because many of these grassland ecosystem services depend on productivity (often measured as ANPP), it is important to better understand the drivers of spatial and temporal patterns of productivity. ANPP is an important integrative variable of ecosystem function as well as a key component of the global C cycle. Given the high degree of certainty in forecasted increases in air temperatures, insight into how grassland ANPP may be affected is critical. If effects of increasing temperatures are primarily manifest

through negative impacts on water balance, semi-arid and arid grasslands should be among the most sensitive ecosystems to this consequence of warming. For example, Huxman et al. (2004) inferred that ecosystems with low precipitation inputs should be the most responsive to changes in water availability, and Knapp and Smith (2001) reported that grasslands were more responsive to precipitation variability than most other ecosystem types in North America. Studies on other continents have found similar patterns (Li et al. 2011; Sala et al. 2012).

Recently, De Frenne et al. (2013) advocated the use of natural gradients of climatic variation to assess ecological responses to climate change, and we adopted this approach to assess the temperature sensitivity of ANPP in semi-arid grasslands. Our goal was to assess sensitivity from a spatial perspective by relating variation in ANPP to temperature variables across a range of sites, as has been done in past ANPP precipitation studies (Sala et al. 1988; Epstein et al. 1997; Huxman et al. 2004; Verón et al. 2005). The analysis was based on data from eight semi-arid native grasslands with collectively 67 years of ANPP data. These sites occurred along a western US climatic gradient in which both mean annual temperature (MAT) and MAP varied two-fold. We predicted that if there were direct effects of variation in temperatures on ANPP then either a positive or negative effect of higher temperatures on ANPP would be detected. Positive effects could result from lengthening the growing season, which would be particularly important along the northern end of the gradient. Negative effects could result from exceeding the thermal optima of the dominant plants, which would likely be more important at the southern end of the gradient. Similarly, if the effects of increasing temperatures were indirect then a negative effect of temperature on ANPP would also be detected, but this would be the result of the effects of soil drying. To further examine this indirect effect, we also incorporated the standardized precipitation evaporation index (SPEI) (Vicente-Serrano et al. 2010) into our analysis as an independent variable. The SPEI adjusts precipitation inputs by estimates of evapotranspiration (ET; largely driven by temperature) and thus explicitly includes the potential negative effects of temperature on water balance (Vicente-Serrano et al. 2010). Our expectation was that combining precipitation inputs with the negative effects of temperature on water balance would yield a model that explained a greater variation in ANPP than precipitation or temperature alone.

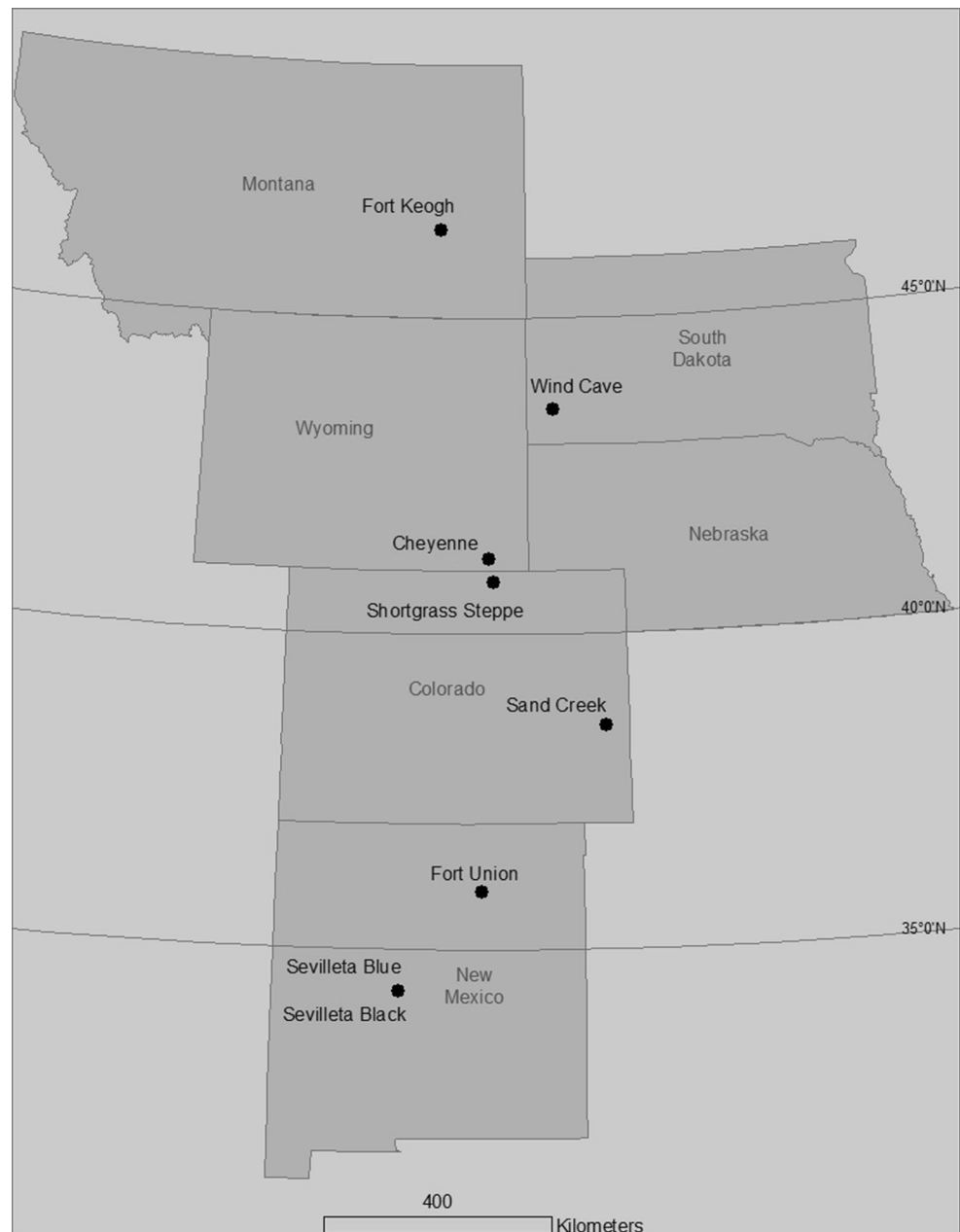
## Materials and methods

Annual net primary production (ANPP;  $\text{g/m}^2$ ) precipitation and temperature data (annual and growing season, 1 April to 30 September) were compiled from eight

semi-arid grassland sites located in the Great Plains region of the US (Fig. 1; Table 1). We defined the “growing season” for all grasslands as 1 April–30 September, even though growing season length is variable across this latitudinal gradient. However, analyses with shorter growing seasons in the north and longer seasons in the south only marginally altered the results reported below. Thus, we opted to maintain a consistent growing season period for all grasslands. Across these sites, 30-year MAT varied ~twofold, from 7.8 to 14.2 °C, as did MAP (250–500 mm). Distances between sites usually exceeded 100 km except for the two sites at the southern end of the gradient. These were located at the Sevilleta Long-term

Ecological Research (LTER) site and both were included because the dominant grass species differed between sites (*Bouteloua gracilis* vs. *Bouteloua eriopoda*). Data availability for ANPP varied from 3 to 16 years at individual sites (Table S1), and our primary focus was on relating patterns of ANPP and temperature across this 1,600-km latitudinal gradient, although temporal variation was included, consistent with past studies that have assessed the sensitivity of ANPP to precipitation in grasslands (Sala et al. 1988; Huxman et al. 2004; Wu et al. 2011; Jobbágy et al. 2002; Peñuelas et al. 2007). Soils varied from sandy loams to clay (Table 1), and as expected for the central US, grassland communities were dominated by

**Fig. 1** Locations of eight grasslands included in this study. Additional information for each site can be found in Tables 1 and S1



**Table 1** Site characteristics of the eight grassland sites along a latitudinal gradient of semi-arid grasslands in the western US

Site	MAP (mm)	GS.Precip. (mm)	MAT (°C)	GS.Temp. (°C)	C <sub>3</sub> :C <sub>4</sub> <sup>a</sup> (%)	Soil type
Fort Keogh	316.0	246.1	7.83	16.9	87:13	Silty clay loam <sup>b</sup>
Wind Cave	499.1	378.7	8.4	15.8	75:25	Sandy loam <sup>c</sup>
Cheyenne	404.9	306.8	8.1	14.9	59:41	Fine-loamy <sup>d</sup>
SGS	389.7	290.6	8.4	15.2	42:58	Sandy loam <sup>e</sup>
Sand Creek	393.9	312.7	10.9	19.0	14:86	Clay <sup>f</sup>
Fort Union	427.0	317.3	9.9	16.2	25:75	Sandy clay loam <sup>f</sup>
Sevilleta blue	256.0	178.8	14.2	21.6	22:78	Sandy loam/clay loam <sup>g</sup>
Sevilleta black	256.0	178.8	14.2	21.6	20:80	Sandy loam/clay loam <sup>g</sup>

Climate data are from the National Oceanic and Atmospheric Administration (NOAA) ([www.ncdc.noaa.gov/cdoweb/datasets](http://www.ncdc.noaa.gov/cdoweb/datasets)). More detailed information for each site can be found in the references provided after the soil type description

MAP Mean annual precipitation, GS.Precip. growing season precipitation, MAT mean annual temperature (all based on 30 years of data); SGS shortgrass steppe

<sup>a</sup> Determined using aboveground net primary production (ANPP) data collected from this study, and calculated as the proportion of C<sub>3</sub> and C<sub>4</sub> biomass (g/m<sup>2</sup>) relative to the total

<sup>b</sup> Heitschmidt et al. (2005)

<sup>c</sup> Smith (2012)

<sup>d</sup> Dijkstra et al. (2012)

<sup>e</sup> Lauenroth and Burke (2008)

<sup>f</sup> Cherwin and Knapp (2012)

<sup>g</sup> Muldavin et al. (2008)

C<sub>3</sub> plants in the northern sites grading to C<sub>4</sub> dominance in the southern sites (Teeri and Stowe 1976; Epstein et al. 1997; Table 1). Sites were not burned or grazed by livestock during the years that ANPP data were collected.

#### ANPP estimates

Methods used for estimating ANPP differed among sites. In the six northern sites, ANPP was estimated by harvesting peak or end of season biomass, sorting by species, then drying and weighing. Plot sizes varied from 0.1 to 0.25 m<sup>2</sup> and the number of harvested plots was >10/year at each site except at Cheyenne, where  $n = 5$ . ANPP estimates from the two southern sites were based on non-destructive allometric methods in which volume estimates were made for individual plants and ANPP was estimated using species-specific equations (Muldavin et al. 2008). For each site, total ANPP and ANPP of specific functional types including graminoids (grasses and sedges), forbs, and C<sub>3</sub> and C<sub>4</sub> photosynthetic pathways were estimated. Woody plants were included when estimating total ANPP, but they were a minor component of ANPP at all sites, therefore they were not analyzed as a functional group.

Climate data (including annual temperature and precipitation, as well as growing season temperature and precipitation) were compiled from daily site weather records or nearby weather stations (<15 km from study site). Many parameters (e.g., maximum and minimum values) at many

temporal scales (daily, monthly, annually) were initially considered and explored, but growing season and annual values were the most informative. Further details on each site as well as ANPP sampling and climate data can be found in the references in Table 1 and in the Electronic Supplementary Material (Table S2).

#### Data analysis

We related patterns of ANPP to variation in temperature and precipitation in three ways. First, we combined the 67 years of data available from all sites with corresponding climatic data to evaluate simple and multiple linear regression models relating temperature and precipitation to ANPP. We focused on simple linear and multiple regressions, (procREG, SAS version 9.3; SAS, Cary, NC) initially in order to more directly compare our results with other regional scale analyses. Further, non-linear models did not provide additional explanatory power. Because the number of years of data varied among sites and those sites with the greatest number of years (the most northern and the two southern sites; Table S1) could dominate and bias relationships, we calculated site means for ANPP and climate data. This eliminated temporal variability and limited our statistical power to the number of sites (eight), but allowed us to determine if using all 67 years of data led to qualitatively different relationships from those based on site means. Second, the combined data set was analyzed with analysis of covariance (ANCOVA; proc MIXED)

**Table 2** Results for simple linear regression models relating climatic variables to ANPP (total, graminoid and the proportion of ANPP from C<sub>3</sub> plants)

Dependent variable	Parameter	Regression coefficient	<i>p</i> -value	<i>r</i> <sup>2</sup>
Total ANPP	Annual temperature	−5.32	0.019	0.08
	Annual precipitation	0.33	<0.0001	0.39
	Annual SPEI	31.86	<0.0001	0.28
	Growing season temperature	−7.67	0.002	0.14
	Growing season precipitation	0.31	<0.0001	0.31
	Growing season SPEI	31.65	<0.0001	0.30
Graminoid ANPP	Annual temperature	−6.38	0.0002	0.15
	Annual precipitation	0.31	<0.0001	0.43
	Growing season temperature	−8.30	0.001	0.20
	Growing season precipitation	0.30	<0.0001	0.40
Relative C <sub>3</sub> ANPP	Annual temperature	−0.07	<0.0001	0.43
	Growing season temperature	0.27	−0.06	<0.0001

Only significant relationships are shown. When temperature and precipitation variables were combined in multiple regression models, none were significant, nor were any regression models significant for forb ANPP

*SPEI* Standardized precipitation evaporation index (estimates site water balance by combining precipitation inputs with estimates of evapotranspirational losses)

models that included site as a fixed effect, along with temperature and precipitation variables and all interaction terms. Finally, we downloaded SPEI values (Vicente-Serrano et al. 2010, from [www.sac.csic.es](http://www.sac.csic.es)) based on annual and growing season periods for each site and related these to patterns of ANPP. SPEI includes an estimate of ET driven primarily by temperature, thus by comparing SPEI–ANPP relationships to precipitation–ANPP relationships, we could explicitly assess the negative and indirect effects of temperature on water balance and consequently ANPP.

All models were evaluated for total ANPP, graminoid, forb, and the proportion of ANPP comprising C<sub>3</sub> species, as dependent variables. The latter dependent variable (which ranged in magnitude from ~90 to <5 %) was included because relative abundance of photosynthetic types was less variable from year to year than absolute ANPP values. This allowed us to focus more on broad-scale climatic drivers of ANPP by photosynthetic pathway.

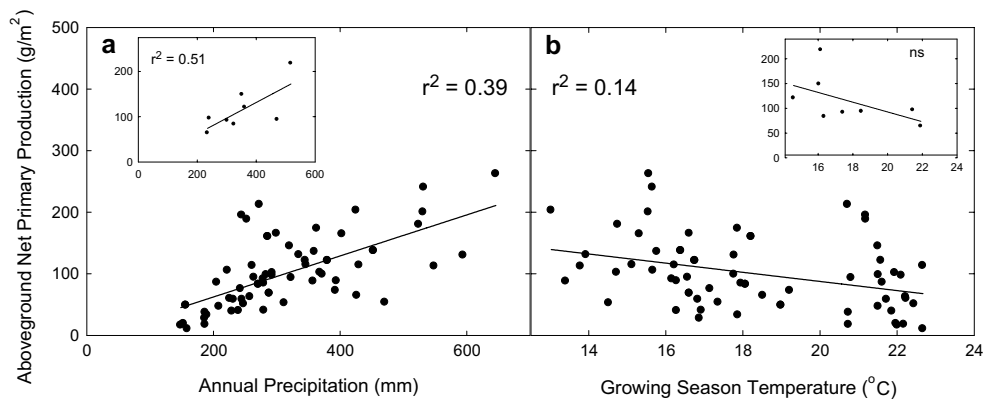
Akaike's information criterion (AIC) (Johnson and Omland 2004) was used to select those models that best fit the patterns of variation in ANPP (see Table S3, Supplementary Information for all candidate models). The AIC model selection procedure (SAS proc GLM select) compared each candidate model and assigned them an AIC value based on the each model's goodness of fit, taking into account the number of parameters by penalizing each model for additional terms. The model with the lowest AIC value was selected as the best model. To estimate the relative importance of particular model parameters, Akaike weights (*w*) were summed from those models that included the term of interest (Johnson and Omland 2004). On a scale

of 0–1, the parameters were assigned a weight based on model comparisons. Parameters in which *w* was near one were deemed the most important. We calculated *w* with MuMIn in R version 3.0.2 ([www.r-project.org](http://www.r-project.org)).

Finally, following Epstein et al. (1997), data were parsed into narrow ranges of precipitation (100- and 200-mm bins) to minimize variation in precipitation and increase the potential for temperature sensitivity to be manifest. Simple linear regression analyses were then conducted for those subsets of the data (*n* = 7–56 years depending on the particular range of precipitation binned).

## Results

Both annual and growing season precipitation were positively correlated with total ANPP in simple regression models (Table 2; Fig. 2) and accounted for 39 and 31 % of the variance in ANPP, respectively. In contrast, annual and growing season temperatures were negatively correlated with ANPP (Table 2; Fig. 2) and explained 8 and 14 % of the variance, respectively. When site means were used, precipitation–ANPP relationships remained statistically significant (*p* < 0.0001; Fig. 2, left inset), whereas temperature–ANPP relationships were not significant (although trends were similar; Fig. 2, right inset). The addition of temperature variables in multiple regression models did not improve the explanatory power of precipitation alone. However, despite temperature's relatively minimal predictive power in simple and multiple regressions, ANCOVA models selected by AIC included growing season temperature along with site, annual



**Fig. 2** Simple linear regression models that best fit patterns of aboveground net primary production (ANPP). **a** Relationship between annual precipitation (An.Precip.) and aboveground net primary production (ANPP) across eight grasslands (ANPP =  $-2.90 + 0.33 \times \text{An.Precip.}$ ). *Inset* Relationship based on site means of ANPP and precipitation

(ANPP =  $-7.32 + 0.35 \times \text{An.Precip.}$ ). **b** Relationship between ANPP and growing season temperature (GS.Temp.) across eight grasslands (ANPP =  $240.87 - 7.67 \times \text{GS.Temp.}$ ). *Inset* Relationship based on site means of ANPP and temperature. *ns* Not significant

**Table 3** Results from analyses of covariance relating climatic variables and site as a fixed effect to total ANPP and functional types

Selected models using AIC					
Dependent variable	Model	Parameters	AIC	<i>w</i>	<i>R</i> <sup>2</sup>
Overall ANPP	Site GS.Temp. An.Precip. GS.Temp. × An.Precip.		572.35		0.61
		Site		1.00	
		GS.Temp.		0.81	
		An.Precip.		1.00	
		GS.Temp. × An.Precip.		0.55	
Graminoid	Site GS.Temp. An.Precip. GS.Temp. × An.Precip.		544.24		0.63
		Site		1.00	
		GS.Temp.		0.58	
		An.Precip.		1.00	
		GS.Temp. × An.Precip.		0.20	
Forb	Site An.Precip. An.Precip. × Site		398.44		0.52
		Site		1.00	
		An.Precip.		1.00	
		An.Precip. × Site		0.99	
Relative C <sub>3</sub>	Site GS.Temp. An.Precip. An.Precip. × Site GS.Temp. × An.Precip.		-189.96		0.89
		Site		1.00	
		GS.Temp.		0.98	
		An.Precip.		1.00	
		An.Precip. × Site		0.98	
		GS.Temp. × An.Precip.		0.96	

Akaike's information criterion (AIC) was used to select models with the best fit and the least complexity. Simple and multiple regression models were included as candidate models, but none were selected. Models were selected for each of the functional types: total ANPP, graminoid, forb, and relative C<sub>3</sub> ANPP. Akaike weights (*w*) were used to assess how individual parameters improved the model selected

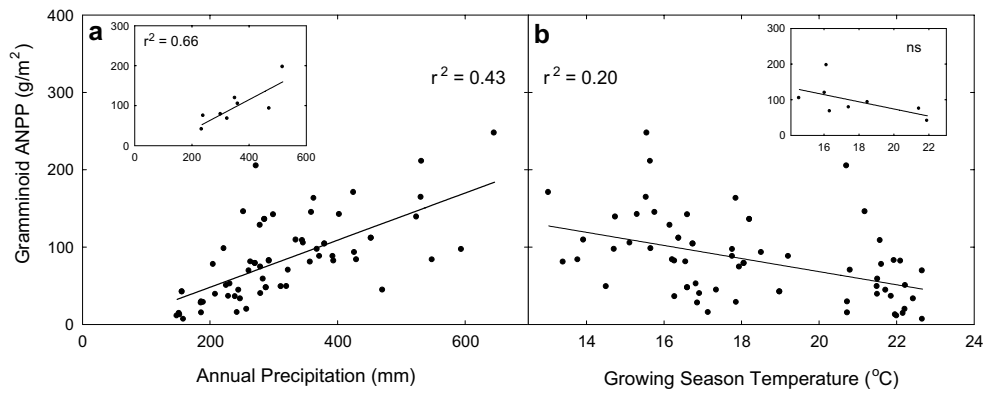
GS Growing season, An. annual, Temp. temperature, Precip. precipitation

precipitation and a growing season temperature × annual precipitation interaction term (Table 3). Although site and annual precipitation were the most influential parameters in the model (*w* = 1.0), growing season temperature

contributed substantially (*w* = 0.81), while the interaction term was the least important variable (*w* = 0.55; Table 3).

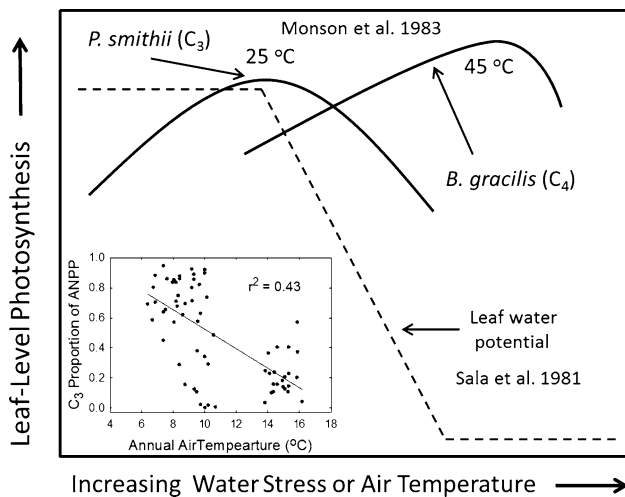
Graminoids comprised a large proportion of total ANPP at all sites and thus both annual and growing season





**Fig. 3** Simple linear regression models that best fit patterns of graminoid ANPP. **a** Relationship between An.Precip. and graminoid ANPP (Graminoid ANPP =  $-14.58 + 0.31 \times \text{An.Precip.}$ ). **b** Relationship between GS.Temp. and graminoid ANPP

(Graminoid ANPP =  $-233.20 - 8.30 \times \text{GS.Temp.}$ ). *Insets* Relationships based on site means for each of the eight grasslands (ANPP =  $-36.78 + 0.38 \times \text{An.Precip.}$ ). For abbreviations, see Fig. 2



**Fig. 4** Contrasting responses of leaf-level photosynthesis in two dominant semi-arid grassland grasses to alterations in temperature and increasing water stress. Along the north to south climatic gradient in this study (Fig. 1), there is a shift in the proportion of ANPP resulting from plants with the C<sub>3</sub> vs. C<sub>4</sub> photosynthetic pathways (*inset* relationship from this study; Table 1). The two dominant grasses along this gradient, *Pascopyrum smithii* and *Bouteloua gracilis*, both have broad photosynthetic response surfaces to varying temperatures and their respective temperature optima differ by 20 °C (Monson et al. 1983). Shifts in the abundance of C<sub>3</sub> and C<sub>4</sub> species along this gradient may moderate apparent temperature sensitivity of ANPP, whereas effects of temporal changes in temperature for both C<sub>3</sub> and C<sub>4</sub> grasses will depend upon whether temperatures are shifting towards or away from thermal optima. In contrast to temperature, responses of photosynthesis to water stress (*dashed line*) show a strong threshold response for all plant species [example shown is for *B. gracilis* (Sala et al. 1981)] and thus ANPP responses to changes in precipitation inputs are more likely to be consistently strong along the entire climatic gradient (spatially and temporally). Combined, spatial shifts in species-level traits and differences in the nature of physiological responses to change in water vs. temperature are key mechanisms explaining why precipitation but not temperature is a strong predictor of latitudinal variation in ANPP in western US semi-arid grasslands

precipitation were also strongly correlated with graminoid ANPP; annual precipitation again was the best predictor (Fig. 3; Table 2). In simple regression models, annual and growing season temperatures were highly correlated with graminoid ANPP, but similar to total ANPP, annual and growing season temperatures explained much less of the variability ( $r^2 = 0.15$  and  $0.20$  respectively; Fig. 3; Table 2). Similar to the results for total ANPP, when data were combined at the site level, relationships were statistically significant for precipitation but not temperature (Fig. 3, inset).

The best ANCOVA model for graminoid ANPP included the same parameters important for total ANPP (site, growing season temperature, annual precipitation, and a growing season temperature  $\times$  annual precipitation interaction term). For graminoids, site and annual precipitation were the most important model parameters, both with a weight of 1.0, whereas growing season temperature was less important ( $w = 0.58$ ; Table 3). In contrast to graminoids, forbs comprised a much smaller and more variable proportion of total ANPP among sites and as a result, no simple or multiple regression models with temperature or precipitation parameters were significant. The best ANCOVA model for forb ANPP included site, annual precipitation, and an interaction between these two terms (Table 3).

In contrast to absolute ANPP values, the proportion of ANPP from C<sub>3</sub> plants was strongly correlated with both growing season and annual temperature, but not with precipitation. In simple regressions, annual temperature was a better predictor of relative C<sub>3</sub> ANPP than growing season temperature ( $r^2 = 0.43$  and  $0.27$  respectively; Table 2), and was inversely related to C<sub>3</sub> production (Fig. 4). The best model selected using  $w$  included site, growing season temperature, annual precipitation and the interaction terms annual precipitation  $\times$  site and growing season temperature  $\times$  annual precipitation (Table 3).

Despite the combination of precipitation and temperature effects in SPEI values, simple regressions of growing season ( $r^2 = 0.30$ ) and annual SPEI ( $r^2 = 0.28$ ), although highly significant, explained less of the variation in ANPP than precipitation alone (Table 2). Furthermore, parsing the data into 100- and 200-mm ranges of precipitation after Epstein et al. (1997) did not improve relationships between ANPP and temperature variables along this gradient.

## Discussion

The goal of this study was to assess the temperature sensitivity of ANPP in semi-arid grasslands along a natural climatic gradient where temperature and precipitation both varied by twofold from north to south. Although growing season temperature was found to be negatively related to both total and graminoid ANPP with simple linear regression models, annual precipitation was a much stronger predictor of patterns of ANPP than temperature. Thus, our study was consistent with previous research (Lauenroth and Sala 1992; Li et al. 2011; Knapp et al. 2002; Vermeire et al. 2009; Sala et al. 1988), except that the amount of variance explained by precipitation was substantially less along this latitudinal gradient than in many of these other studies. Because temperature and precipitation were negatively correlated across these sites, inferences regarding temperature sensitivity are challenging. However, with models that accounted for site variation and interactions between temperature and precipitation, growing season temperature was still identified as a significant model parameter explaining variation in ANPP, albeit secondary to precipitation. A similar conclusion of weak but significant temperature effects in models dominated by water availability terms was reached in an analysis of tree growth in eucalypt forests arrayed along climatic gradients in Australia (Bowman et al. 2014).

Past studies (experimental and observational) have demonstrated negative effects of temperature on ANPP, and these have been argued to be indirect due to increased ET and reduced water availability to plants at higher temperatures (Epstein et al. 1997; Niu et al. 2008; Engel et al. 2009; Dulamsuren et al. 2013; Xu et al. 2013). We were unable to demonstrate such a negative statistical relationship between temperature and ANPP when we analyzed subsets of the data in which precipitation variation was restricted to narrow ranges (Epstein et al. 1997). Perhaps more surprising, however, was that simple linear regression models based on the SPEI explained less variance along this natural climatic gradient than precipitation alone. This suggests that across these eight semi-arid grasslands, the indirect effect of temperature on site water balance was not consistent with regard to effects of temperature on ANPP. This result and the inclusion

of significant precipitation  $\times$  temperature interaction terms in the best-fit models suggest that the relationship between temperature and ANPP is more complex than the precipitation–ANPP relationship (Vicente-Serrano et al. 2013).

Why is ANPP sensitivity to temperature difficult to detect along natural climate gradients?

The strong interdependency of temperature and precipitation in determining ecosystem function is well known (Rosenzweig 1968; Kardol et al. 2010; Frank and Inouye 1994; Rustad et al. 2001), yet at local to global scales, ecosystem function and structure are usually much better correlated with precipitation patterns than temperature (e.g., Sala et al. 1988; Del Grosso and Parton 2008). The exception to this generalization is in ecosystems with abundant water (Huxman et al. 2004; Kirwan et al. 2009). Below we explore several potential reasons why temperature sensitivity is difficult to detect and why as a result, predicting responses of ANPP to forecast warming is likely to be better informed by experiments than by using natural climatic gradients.

A fundamental difference between temperature and precipitation effects on plant and ecosystem processes is that for all but the most hydric ecosystems (Knapp et al. 2008), significant reductions in precipitation will always have a negative (or at best neutral) effect on ecosystem processes. This includes both leaf-level photosynthesis and ANPP (Fig. 4; Sala et al. 1981; Heitschmidt et al. 2005). In contrast, most C-gain processes at both the leaf and ecosystem levels have distinct thermal optima. Thus, alterations in temperature can have negative or positive impacts contingent upon temperatures shifting towards or away from thermal optima (Fig. 4). Such contingent positive or negative effects can occur on diurnal as well as seasonal time scales. Temperature impacts on C gain can also vary with soil moisture such that warm temperatures might positively affect ecosystem processes for several days after a substantial rain event, but have negative effects during dry periods when soil water is low (Niu et al. 2008; Zhou et al. 2008). Such contingent effects are likely reflected in the temperature  $\times$  precipitation interaction term in the models that explained variation in ANPP the best (Table 3).

In addition, precipitation tends to vary much more than temperature at interannual time scales. Based on long-term (30-year) climatic records for these eight sites, coefficients of variation (CV) for annual temperature were always less and often less than half the magnitude of the CVs for precipitation (Table S2). This pattern was also evident in the 67-year data set. Thus, even if ANPP was equally sensitive to alterations in temperature and precipitation, greater interannual variability in precipitation would increase the chance of detecting significant precipitation sensitivity relative to temperature.

Less interannual variation in temperature vs. precipitation may also lead to the strong correlation of MAT with the distribution of species with  $C_3$  and  $C_4$  photosynthetic pathways at regional to continental scales in grasslands (Teeri and Stowe 1976; Tieszen et al. 1979; Wittmer et al. 2010). Indeed in this study, precipitation better explained patterns of ANPP but variation in temperature explained more variance in the contribution of  $C_3$  species to ANPP. Greater stability in temperatures may allow species with different photosynthetic traits to align more strongly along temperature than precipitation gradients. This strong sorting of  $C_3/C_4$  photosynthetic pathways was clearly evident along the natural climatic gradient in this study (Table 1; Fig. 4). Differences in temperature optima of the dominant  $C_3$  and  $C_4$  grasses and their shift in abundance from north to south along a temperature gradient, combined with variable effects of alterations in temperature (Fig. 4), all likely contribute to low apparent temperature sensitivity of ANPP in this region and perhaps in many others.

Despite low sensitivity of ANPP to temperature along this natural climatic gradient, we are hesitant to conclude that these grasslands will be insensitive to forecasted warming. Indeed, because of low interannual variability in temperature, directional shifts in MAT by only a few degrees may lead to temperatures that routinely exceed historic levels (Mora et al. 2013), particularly for temperature extremes (Smith 2011). Further, because shifts in community composition and species distributions (including alterations in  $C_3/C_4$  composition) to directional changes in climate require more time than physiological responses of extant species (Smith et al. 2009; Vermeire et al. 2009), sensitivity of ANPP to future warming may lag the more immediate responses to changes in precipitation. Thus, although using natural climatic gradients for ecological climate change research may have many advantages (De Frenne et al. 2013), and such gradients have been particularly useful for providing insights into precipitation as a driver of ANPP, long-term experimentation may be required to better forecast ecosystem responses to warming (Knapp et al. 2012).

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