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

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Stress gradients and biodiversity: monoculture vulnerability drives stronger biodiversity effects during drought years

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Abstract. Climate change will increase the likelihood and severity of droughts into the future. Although diversity may buffer plant communities against the negative effects of drought, the mechanisms underlying this pattern remain unclear. Higher-diversity plant communities may have a higher likelihood of including more drought-resistant species that can compensate for drought-sensitive species (“insurance effects”). Alternatively, higher-diversity communities may alter environmental conditions and improve performance of even drought-sensitive species. Here we planted nonleguminous forbs and grasses into monocultures and four- and eight-species mixtures, and measured species and plot productivity every year from 2000 to 2010. We found that six of our eight species were suppressed when growing in monoculture during dry years. These same species were unaffected by drought when growing in higher-diversity mixtures. Because of this poor performance in monoculture (not insurance effects), the biodiversity productivity relationship was strongest during the driest years. If biodiversity ameliorates hot/dry conditions and therefore improves performance of drought-sensitive species during periods of low rainfall, this may mean biodiversity can be used as a tool to protect individual species from drought conditions.

Key words: BEF; climate change; ecosystem function and services; microclimate amelioration; species-specific responses.

INTRODUCTION

Rapidly changing climatic conditions are increasing the global frequency and severity of extreme weather events such as drought (Trenberth 2011, Dai et al. 2018). Globally, the area of dry lands may increase by up to 10% by 2100, because of both decreased precipitation and increased evapotranspiration with climate warming (Sherwood and Fu 2014). Regionally, severe drought risk may increase by up to 60% in some areas of Europe by 2100, and growing-season soil moisture is projected to decrease by 2–3% in the next 20 yr (Ruostenoja et al. 2017).

Drought often negatively affects ecosystems: drought can decrease agricultural yields and ecosystem carbon sequestration (Ciais et al. 2005) and make ecosystems more vulnerable to subsequent disturbances (Niinemets 2010). However, biodiversity can buffer ecosystems against the loss of productivity and other ecosystem functions (Tilman and Downing 1994, Isbell et al. 2015;

but see Pfisterer and Schmid 2002, Vogel et al. 2012, Craven et al. 2016). The ability of biodiversity to buffer ecosystems against stressful events is often considered a consequence of so-called “insurance effects”; higher-diversity ecosystems are more likely to contain species that are capable of persisting and sometimes thriving in severe environmental conditions; these species can then compensate for losses experienced by more vulnerable species (Yachi and Loreau 1999, Hector et al. 2010, Hautier et al. 2014). An assessment of the insurance hypothesis thus requires a comparison of species-specific responses to drought: some drought-sensitive species may perform poorly during a drought, regardless of the diversity or identity of the species around them. Some drought-tolerant species may perform well during a drought, and their enhanced performance during drought may compensate for the poor performance of drought-sensitive neighbors in higher-diversity plant communities.

Importantly, experimental results from the facilitation literature demonstrate that drought may also increase the importance and strength of direct interspecific facilitative interactions between plants (e.g., the stress gradient hypothesis, Bertness and Callaway 1994). In the

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context of biodiversity ecosystem functioning (BEF) research, the stress-gradient hypothesis predicts that drought-sensitive species may be buffered against climate extremes when growing in (1) higher-diversity or higher biomass plant communities, and (2) near drought-resistant neighbors (Caldeira et al. 2001, Wright et al. 2014, 2015). This is because higher-diversity communities can be more productive and thus provide greater shade, cooler air temperatures, increased relative humidity, increased likelihood of including deep-rooted species, and increased surface soil moisture at the community level (Wright et al. 2014, Steinauer et al. 2015, Cowles et al. 2016). These buffering effects can reduce water losses for drought-sensitive species (Wright et al. 2015), and this may allow individual drought-sensitive species to persist during drought, depending on the diversity and identity of their neighbors.

Furthermore, resource partitioning theory suggests that intraspecific competition should be greater than interspecific competition for many species (HilleRisLambers et al. 2012). Although conspecific neighbors likely compete for resources in overlapping ways, heterospecific neighbors may have complementary root distributions (but see Barry et al. 2020), alternative physiological strategies to combat water stress (Fotelli et al. 2000), or alternative energy allocation patterns that reduce water loss aboveground (e.g., lower specific leaf area of leaves). If intraspecific competition outweighs interspecific competition, this could buffer species from strong competition for soil water during drought.

If individual species are buffered from negative environmental conditions in higher-diversity communities (but not in lower-diversity communities, e.g., Fig. 1), this may also help explain how higher-diversity communities are protected from productivity losses during drought. This mechanism has not been given full attention in the biodiversity-stability literature thus far. Here, we use data from a 10-yr biodiversity experiment in Wageningen, the Netherlands to assess the following two hypotheses: (1) BEF relationships are stronger during periods of low rainfall and (2) stronger BEF relationships during periods of low rainfall are related to individual species performing better when growing near heterospecific neighbors than when growing near conspecific neighbors. This may be due to either interspecific facilitation or alleviation of intraspecific competition.

MATERIALS AND METHODS

Field experiment

The Wageningen Biodiversity Experiment was established in 2000 and ended in 2010 at Wageningen University in The Netherlands (Van Ruijven and Berendse 2003, 2010, Van Ruijven et al. 2003). The experiment is characterized by a cool temperate climate with an average annual temperature of $10.2^{\circ} \pm 0.7^{\circ}\text{C}$ and approximately 847.3 mm of rainfall annually (Bakker et al.

2016). The experiment was established using four grass species (*Agrostis capillaris* L., *Anthoxanthum odoratum* L., *Festuca rubra* L., and *Holcus lanatus* L.) and four forb species (*Centaurea jacea* L., *Leucanthemum vulgare* Lamk., *Plantago lanceolata* L., and *Rumex acetosa* L.). We planted 3-week-old seedlings into equally spaced 12 seedling \times 12 seedling grid cells within 1 \times 1 m plots (144 seedlings per plot). The plots were assigned to one of four diversity levels in six replicated blocks. Each block contained eight monocultures (one for each species), four mixtures of two and four species, and one eight-species mixture (17 plots per block and 102 total experimental plots). Species in two- and four-species plots were assigned randomly, such that the same two- or four-species mixtures were never replicated (in order to maximize the number of species combinations examined). All plots were watered regularly during the first 3 months of growth to ensure proper establishment. After this period, we stopped watering plots. Species composition of plots was maintained by hand weeding. All plots were surrounded with a buffer zone that was sown with a mixture of grass species that were not in the species pool used in our experiment. These walkways were mown regularly.

Plot biomass was measured annually by clipping all plant material to 2.5 cm above the soil surface in August of each year. Biomass was sorted to species, dried for at least 48 h at 70°C , and weighed. To avoid edge effects, only the central 60 \times 60 cm of plot biomass were used for analyses.

Rainfall measurements

Precipitation data were obtained from the Haarweg meteorological station in Wageningen, located approximately 2 km from the experimental site (Tank et al. 2002). Precipitation was recorded in millimeters and summed per month over the course of the growing season (April–August) from 2000 to 2010. Average precipitation per month was then averaged for each growing season and used as a covariate in our analyses. Importantly, this 10-yr period covered a wide range of rainfall conditions. This included the severe drought that occurred in Europe in July 2003, with temperatures up to 6°C greater than long-term averages and precipitation at 50% below the long-term average (Ciais et al. 2005). This also included the 2006 drought that was characterized by just 47 mm of precipitation from June to July 2006 (compared to a long-term average of 140 mm) and had strong effects on plant growth in this experiment (Van Ruijven and Berendse 2010).

Biomass/performance indices

We assessed the strength of the BEF effect by comparing biomass per square meter (g/m^2) in each diversity level in each year. To assess species-specific growth in a given year at a given level of plant diversity in

comparison with long-term average growth (suppression or enhancement), we calculated: $VI = B_{ijk} - \bar{B}_{iR}$ where VI is the vulnerability index, B_{ijk} is the biomass of species i in plot j in year k , and $-\bar{B}_{iR}$ is the average biomass of species i for that (R) richness level (1, 2, 4, or 8 species). Thus, we could assess how the performance of each individual species in a given plot in a given year (with a given amount of precipitation) deviated from how that species usually performed in that level of biodiversity. For example, a VI greater than 0 indicated that an individual species was growing better than average and a VI less than 0 indicated that the species was growing worse than average. We also calculated: $VI_{prop} = B_{ijk}/\bar{B}_{iR}$ to assess proportional responses of individual species in comparison with their long-term average performance. A VI_{prop} greater than 1 indicated better than average performance, whereas a value between 0 and 1 indicated worse-than-average performance.

We used our absolute VI index to understand species-specific patterns underlying the standard BEF relationship. We used our proportional VI index to understand relative species-specific patterns underlying a *proportional* BEF relationship. In particular, we were interested in whether some species perform well regardless of the identity or diversity of their neighbors. We were also interested in whether some species grow proportionally more during drought and thus compensate for losses experienced by drought-sensitive neighbors. Finally, our species-specific metrics can assess whether some species are less sensitive to the drought depending on the identity and diversity of their neighbors.

Statistical analyses

We assessed how the absolute and proportional BEF relationship changed as a function of annual precipitation using a mixed-effects model that included plot biomass (or proportional biomass) as the response variable, continuous fixed effects for year of study, species richness, annual precipitation (April–August), and the interaction between species richness and precipitation. We also included a random effect for plot nested in block to account for repeated measurements of the same plots over time and the blocked spatial layout of the experiment. We also included a random effect for year of study (discrete variable) to account for random variation in performance of plots from year to year (each year has its own y-intercept) that is not associated with precipitation or continuous changes in performance over (PlotBiomass ~ Sp Rich × Precipitation + year, random = Block/plot, random = year, e.g., Davidson et al. 1984). We also tested several other slightly more complicated error covariance structures (plot and block nested in year vs. plot nested in year and plot nested in block) and chose this error structure because it yielded the lowest Akaike information criterion (AIC_c) scores. Finally, we also analyzed all of our results using a Type III ANOVA design (main results reported here are Type II ANOVA,

Type III ANOVA results in Appendix S1: Table S1). There is an ongoing debate in the statistical community about which approach is more appropriate for unbalanced data, thus we report on both here (Langsrud 2003). Importantly, the two approaches do not affect the interactions that we are concerned with for the purposes of this study.

We also binned the precipitation measurements into three groups: low (51–57 mm rainfall), medium (61–74 mm rainfall), and high (76–92 mm rainfall). We used these three groups as this allowed for a near-equal sample size within each binned category ($n = 4$ yr in low rainfall and medium rainfall, $n = 3$ yr in the high rainfall group). We also included species richness as a categorical variable in this model to be able to use post hoc Tukey tests to compare between all binned groups (e.g., monocultures in rainy years vs. high-diversity plots in dry years) and to determine whether differences in BEF relationships were due to monoculture suppression during periods of low rainfall or due to higher diversity enhancement during periods of high rainfall. All other aspects of the model structure for this analysis were identical to the model described above. Overall results from this analysis were not different from the continuous variable model, and thus we just report on the results from the Tukey tests here.

To assess how individual species biomass was enhanced or suppressed by annual precipitation (continuous) and modified by community diversity (continuous), we used a mixed effects model with VI (or VI_{prop}) as the response variable, continuous fixed effects for year of study, species richness, annual precipitation (April to August), the interaction between species richness and precipitation, and a random effect for year and plot nested within block (VI ~ Sp Rich × Precipitation, random = Block/plot, random = year).

RESULTS

Ambient precipitation at the site ranged from 51 mm per month during the 2010 growing season to 91 mm per month during the 2007 growing season with no significant correlation between time and growing-season precipitation (Fig. 2, $R = -0.31$). Growing season precipitation did not have a strong *overall* effect on biomass production ($\chi^2_{1,9} = 3.24$, $P = 0.07$, Table 1). Year of study had a significant negative effect on overall biomass production ($\chi^2_{1,9} = 8.13$, $P = 0.0004$, Table 1) indicating reduced performance over time. Year of study also had a significant negative effect on biomass production of *Festuca rubra*, *Holcus lanatus*, *Plantago lanceolata*, and *Rumex acetosa* (Table 2).

We found that the strength of the BEF relationship increased during drier years (51–57 mm precipitation class, SR × precipitation interaction, Table 1, Fig. 3a), and this corresponded with lower monoculture performance in the drier years paired with no significant changes in higher diversity mixture performance in any

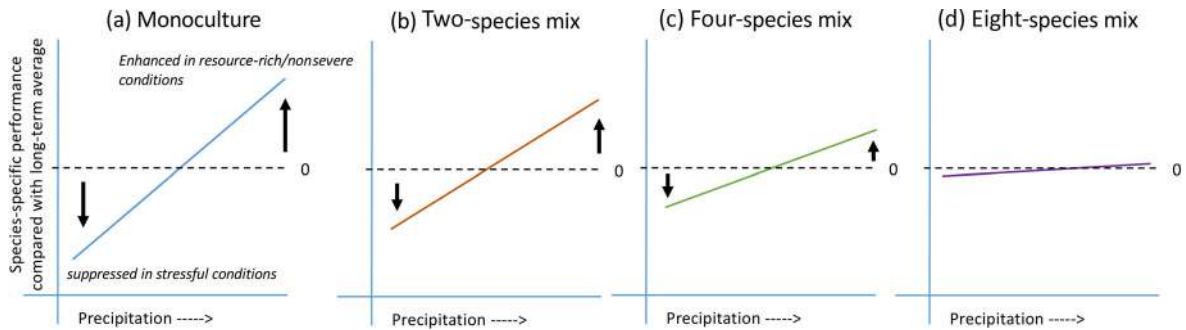


FIG. 1. Conceptual model of the monoculture vulnerability hypothesis. Some drought-sensitive species may be strongly suppressed when growing alone in monoculture during dry years (a). These same species may thrive in monoculture during wet years (a). If biodiversity provides some kind of buffer against fluctuations in precipitation (drought conditions), these same species should not respond strongly to fluctuations in precipitation when they are growing in higher-diversity mixtures (b–d).

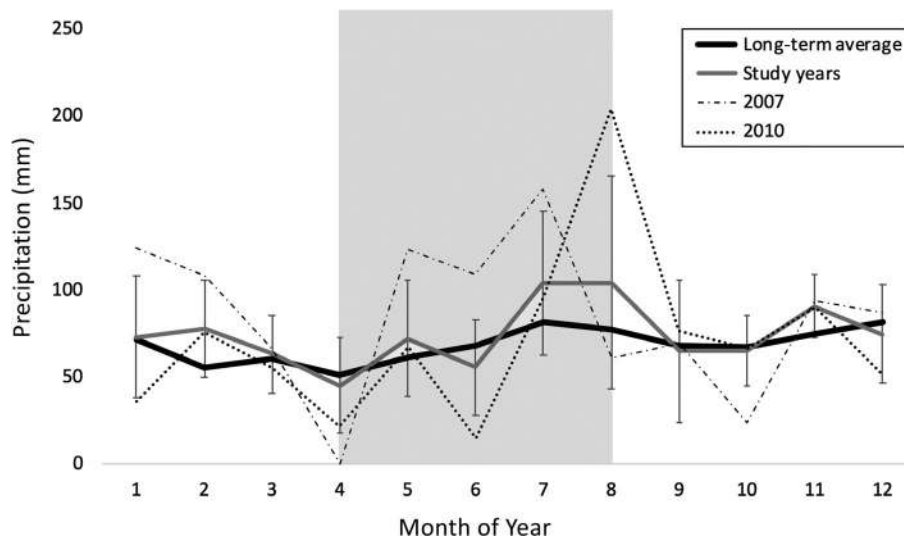


FIG. 2. Precipitation patterns over the course of this study in comparison with the long-term average (1951–2012). The data presented here cover the 2000–2010 growing season (April–August, shaded gray). The study years were overlapping with long-term average trends. The wettest year of the study was 2007, and the driest year of the study was 2010.

years (Fig. 3a). We also found that rainfall strongly affected our proportional community biomass measures. When we compared annual proportional biomass production in each plot with long-term averages, we found that monocultures grew 25% less than average during dry years and 30% greater than average during wet years (SR \times precipitation interaction, Table 1, Fig. 3b). Higher-diversity mixtures did not grow more or less than average in wet or dry years (Fig. 3b).

In terms of species-specific changes in growth (VI index), we found that six out of the eight species performed poorly when growing in monoculture during dry years but not wet years (*Agrostis capillaris*, *Anthoxanthum odoratum*, *Festuca rubra*, *Holcus lanatus*, *Leucanthemum vulgare*, and *Plantago lanceolata*). The first year of the experiment was also the second wettest year of the experiment, and thus these species were likely affected

by both precipitation effects and increased performance in monoculture in their first year (Fig. 4). However, even after accounting for the effects of time in our statistical model, there was still an interaction between precipitation and the species richness of the plot. In other words, these species were unaffected by precipitation when growing in higher-diversity mixtures but strongly affected by precipitation when growing alone in monoculture (SR \times precipitation interactions, Table 2, Fig. 4). The other two species (*Rumex acetosa* and *Centaurea jacea*) were not directly affected by precipitation in any of the diversity mixtures.

In terms of proportional growth responses (VI_{prop}), we found that two of these species were negatively affected by drought when growing in monoculture but not when growing in mixture (similar to absolute VI analyses). We found that one species (*Holcus lanatus*)

TABLE 1. We used a mixed-effects model to assess the role of year of the study (Year), planted species richness (Sp Richness), precipitation during the growing season (April–August) and the interaction between species richness and precipitation on biomass production in our plots.

Factor	Absolute BEF					Proportional BEF				
	Estimate	Standard error	df	ChiSq	P value	Estimate	Standard error	df	ChiSq	P value
Year	−8.83	3.10	1, 9	8.13	0.004*	−0.06	0.02	1, 9	9.48	0.002*
Species richness	36.1	7.11	1, 100	32.0	<0.0001*	0.16	0.04	1, 100	0.00	1.000
Precipitation	1.96	0.81	1, 9	3.24	0.07	0.01	0.005	1, 9	3.24	0.07
Species richness × precipitation	−0.24	0.09	1, 1009	6.73	0.0095*	−0.002	0.0005	1, 1009	19.7	<0.0001*

Notes: We included a random effect for plot nested within block to account for repeated measurements taken over time and spatial autocorrelation of measurements taken within closer proximity to one another at the field site (blocked design). We also included a random effect for year of study to account for random year-to-year variation in the plots that is not associated with precipitation or continuous change over time. Significant results are marked with an asterisk.

TABLE 2. We used a mixed-effects model to assess the role of year of the study (Year), planted species richness (Sp Richness) and precipitation during the growing season (April–August) and the interaction between species richness and precipitation on vulnerability index for each individual species in our plots.

	Year			Species richness			Precipitation			Species richness × precipitation		
	df	ChiSq	P value	df	ChiSq	P value	df	ChiSq	P value	df	ChiSq	P value
<i>Agrostis capillaris</i>	1, 9	1.97	0.16	1, 28	0.00	1.00	1, 9	5.89	0.02*	1, 289	49.4	<0.0001*
<i>Anthoxanthum odoratum</i>	1, 9	2.71	0.10	1, 27	0.00	1.00	1, 9	9.93	0.002*	1, 279	9.55	0.002*
<i>Centaurea jacea</i>	1, 9	0.46	0.50	1, 28	0.00	1.00	1, 9	0.57	0.45	1, 289	0.78	0.38
<i>Festuca rubra</i>	1, 9	12.7	0.0003*	1, 29	0.00	1.00	1, 9	3.44	0.06	1, 299	20.5	<0.0001*
<i>Holcus lanatus</i>	1, 9	4.63	0.03*	1, 28	0.00	1.00	1, 9	2.48	0.12	1, 289	33.5	<0.0001*
<i>Leocanthemum vulgare</i>	1, 9	0.10	0.32	1, 28	0.00	1.00	1, 9	0.17	0.68	1, 289	18.7	<0.0001*
<i>Plantago lanceolata</i>	1, 9	11.7	0.0006.*	1, 28	0.00	1.00	1, 9	4.18	0.04*	1, 289	29.1	<0.0001*
<i>Rumex acetosa</i>	1, 9	25.3	<0.0001*	1, 28	0.00	1.00	1, 9	8.19	0.004*	1, 289	0.14	0.70

Notes: Vulnerability index was assessed as biomass production in a given year in comparison with average annual biomass production for that species. We included a random effect for plot nested within block to account for repeated measurements in the same plots over time, autocorrelation of multiple measurements taken on different species within a single plot at each sampling point, and the spatially nested nature of plots within larger blocks at our field site. We also included a random effect for random year-to-year variation in the plots that is not associated with precipitation or continuous change over time. Because our index is normalized to each species richness level, we (a priori) expect a perfect fit for each species as a function of species richness. Interactions between species richness and precipitation indicate biologically meaningful results. Significant results are marked with an asterisk.

grew proportionally more in high-rainfall years, but more strongly in higher-diversity plots (indicating compensation for other species that may have been affected negatively), and we found that one species (*Centaurea jacea*) grew proportionally more in drought years, but mostly just in higher-diversity plots (Appendix S1: Table S2 and Fig. S1).

DISCUSSION

Here we confirm past results that higher-diversity communities can better resist productivity changes caused by drought (Isbell et al. 2015). Importantly, we also demonstrate, for the first time, that this can

be driven by a buffering of drought-sensitive species responses when these species are grown in higher-diversity mixtures. Specifically, we found that six of our eight grassland species were sensitive to annual precipitation, but only when growing in monoculture. These same individual species, when growing in higher-diversity mixtures, during these same dry years, were unaffected by annual changes in precipitation. This may be due to alleviation of intraspecific competition. This result is also consistent with past evidence that diversity may moderate environmental conditions and make the microclimate more hospitable for sensitive species (Caldeira et al. 2001, Wright et al. 2015, 2017, Barry et al. 2018). We call this the “monoculture

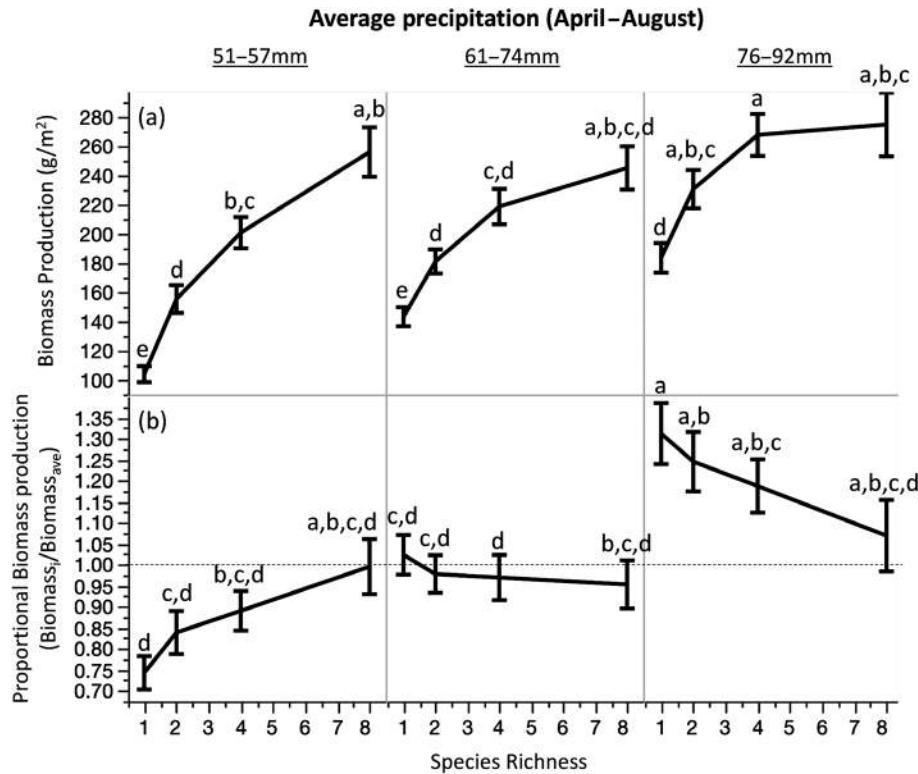


FIG. 3. We assessed the effects of species richness on biomass production every year from 2000 to 2010. We also assessed how species richness and precipitation affected proportional biomass production in year *i* compared with average biomass production for that species richness level across all 11 yr of the experiment. The strength of the effect of species richness on biomass production was strongest in dry years (51–61 mm rainfall). Proportional biomass production was also strongly affected by growing-season precipitation. Error bars represent standard error of the mean. Letters indicate significant differences according to post hoc Tukey tests to assess multiple comparisons between species diversity levels at different levels of precipitation.

vulnerability hypothesis”; some species are sensitive to drought conditions when growing in monoculture but are buffered from these negative effects when growing in higher-diversity mixtures.

BEF relationships are stronger during periods of low rainfall

Climate change is expected to increase the severity and frequency of drought in the future, and past work has demonstrated that biodiversity may be used as a tool to mitigate some of the most negative effects of drought. However, the majority of past work on the role of biodiversity during drought has focused on insurance effects: higher-diversity communities are more likely to resist drought at the community level because of a higher likelihood of containing drought-resistant species (Naehm and Li 1997). In other words, drought-resistant species can compensate at the community level for drought-sensitive species in higher-diversity mixtures (Hector et al. 2010). These so-called insurance effects have been suggested to lead, at least partially, to the maintenance of biomass production in mixtures during drought, but average monoculture productivity declines (Tilman and Downing 1994, Isbell et al. 2015).

A recent meta-analysis by Isbell et al. (2015) demonstrated consistent patterns in ecosystem responses to natural drought across 46 different experiments: higher-biodiversity plant communities were more resistant to changes in biomass production during drought conditions. Further, during moderate drought (drought events occurring between once every 4 yr or once per decade), monocultures were suppressed strongly during the drought, whereas higher-diversity mixtures maintained normal levels of biomass production (Isbell et al. 2015). Our findings provide additional mechanistic insight by focusing on individual species: individual species were sensitive to drought conditions when growing in monoculture, but these same species were insensitive to the drought when growing in higher-diversity mixtures.

Interestingly, these results differ from past assessments of drought effects during experimental drought manipulations (e.g., rainfall exclusion). In fact, a recent meta-analysis of experimental drought demonstrated that higher-diversity communities responded *more* negatively to experimental drought than lower-diversity communities (Craven et al. 2016). It is thus far unclear why drought responses differ between experimental manipulations (Craven et al. 2016) and natural variation in drought conditions (Isbell et al.

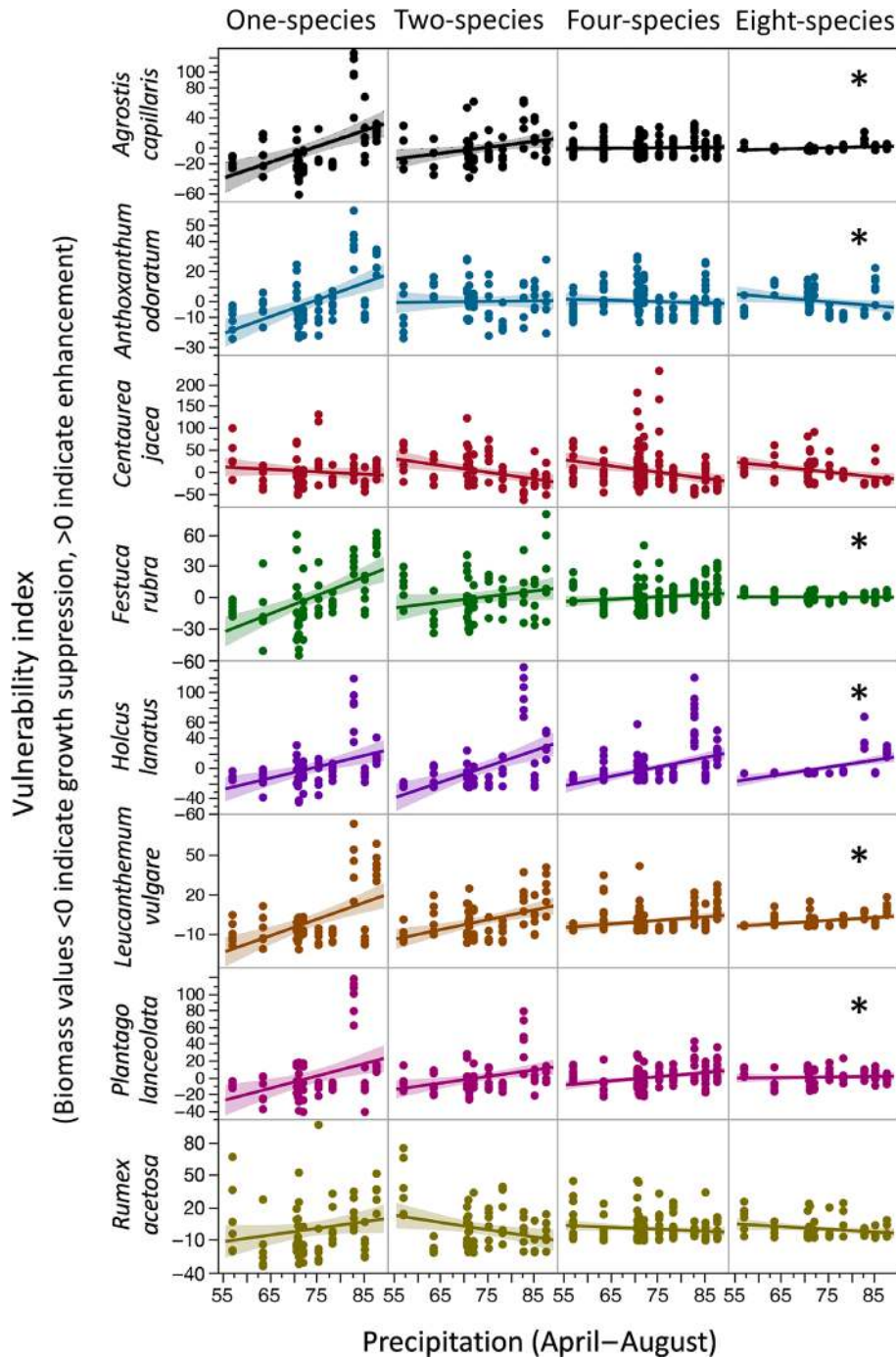


Fig. 4. We measured the vulnerability index for each species in each year of the experiment by comparing species-specific biomass production in that year and compared it with average biomass production for that species over time. We found a significant interaction between growing-season precipitation and species richness of the plot for six of eight species: *Agrostis capillaris*, *Anthoxanthum odoratum*, *Festuca rubra*, *Holcus lanatus*, *Leucanthemum vulgare*, and *Plantago lanceolata* (indicated with an asterisk). Shaded regions indicate 95% confidence intervals.

2015). One possibility that is relevant for the current study is a difference in how drought is applied between experimental droughts (rainfall is removed) vs. natural droughts (rainfall is reduced AND

evaporative demand is increased). Future work should assess these facilitation indices in response to experimental manipulations of drought via careful manipulation of soil moisture and increased evaporative

demand (see Vicca et al. 2012 and Kreyling et al. 2017 for discussion of associated issues).

Insurance effects

One forb species (*Centaurea jacea*) in our analysis appears to be compensating slightly for relative losses experienced by drought-sensitive species in higher-diversity plots during dry years. In drought years, this species grew approximately 1.5 times more than it did in an average year. However, insurance effects do not appear to be the main driver of the maintenance of high productivity over time in higher-diversity plots (only one of eight species showed such a pattern). Furthermore, if insurance effects drove the sustained biomass production of high-diversity communities during drought, we would expect to see decreased biomass production in drought-sensitive species at all diversity levels. We did not observe this effect, suggesting alternative mechanisms underlie our results.

Facilitation and alleviation of competition in higher-diversity mixtures

Six of our eight species performed poorly when growing in monoculture during dry years but not wet years (*Agrostis capillaris*, *Anthoxanthum odoratum*, *Festuca rubra*, *Holcus lanatus*, *Leucanthemum vulgare*, and *Plantago lanceolata*). These same species were unaffected by drought when growing in higher-diversity mixtures (in terms of absolute biomass production). Both *Leucanthemum vulgare* and *Plantago lanceolata* followed these same patterns when assessed in terms of proportional biomass production. Cong et al. (2014) demonstrated that these same six species have proportionally more root biomass in the shallowest root horizon in comparison with the two species that did not follow these trends (*Centaurea jacea* L., and *Rumex acetosa* L.). Shallow rooting depth may be one of several factors contributing to drought sensitivity of these six species. Past work at a temperate North American grassland has shown that higher-diversity microclimates are cooler and more humid, and have lower vapor pressure deficit and higher surface level soil moisture than lower-diversity communities (Wright et al. 2015, Cowles et al. 2016). This has been shown to lead to decreased mortality for some species (Wright et al. 2013), reduced water stress during periods of drought (Caldeira et al. 2001, Wright et al. 2014), and increased growth of sensitive species growing in higher-diversity mixtures during periods of drought (Wright et al. 2015). These known changes in microclimate conditions in higher-diversity communities, paired with our data on species-specific responses to drought in higher-diversity communities, indicate the important role that biodiversity-microclimate effects may play in explaining stronger BEF relationships during dry years.

Although strong interspecific facilitation may help explain these results, alleviation of strong intraspecific

competition could also result in similar patterns. Lower-diversity communities may experience greater competition for soil water during dry years. If intraspecific competition for soil moisture outweighs interspecific competition during these dry periods (e.g., niche complementarity effects, HilleRisLambers et al. 2012), individual species may experience alleviation of intense intraspecific competition when growing in higher-diversity plots. Although we cannot rule this out as a contributing mechanism, a recent meta-analysis by Barry et al. (2020) found that spatial resource partitioning for belowground resources is quite rare in 21 experimental manipulations of grassland biodiversity around the world. Although competition for soil moisture is inevitably occurring, it is likely paired with well-known aboveground effects of diversity on evaporative demand. Higher-diversity communities offer both alleviation of intense intraspecific competitions, and unique microclimatic conditions that reduce water losses.

Finally, monoculture suppression has been demonstrated in BEF research in the past, but this work has mainly focused on the suppressive role of species-specific pests and pathogens (Hendriks et al. 2013). Species-specific pests and pathogens accumulate in monocultures over time and suppress monoculture performance. These density-dependent effects are diluted in higher-diversity mixtures, where herbivory by species-specific pests and pathogens is reduced (Mommer et al. 2018). Although this type of monoculture suppression may also be occurring in our experiment, we would not expect it to increase in drought years when pathogen densities are likely relatively low (unless low pathogen load is paired with disproportionately high pathogen sensitivity). We would also expect this effect to be more tightly tied to increased pathogen suppression over time, as opposed to peaks in suppression tied to particular years.

CONCLUSION AND FUTURE PERSPECTIVES

Recent work has found consensus that higher-diversity plant communities resist against biomass losses during moderate natural droughts (Isbell et al. 2015). Although the vast majority of past studies have pointed at the importance of insurance effects to explain these biodiversity patterns, we present clear evidence that protection of even drought-sensitive species may also underlie these results. This is an important distinction to make. Whereas insurance effects emphasize the *probability* of including drought-resistant species in mixture, our results are less reliant on drought-resistant species and point to the emergent properties of biodiversity in higher-diversity mixtures. In fact, higher-diversity mixtures can protect even drought-sensitive species from the negative effects of drought. The theoretical and management distinctions between these two mechanisms are important. For example, if abiotic facilitation is responsible for biodiversity-stability patterns during droughts, biodiversity may be one method by which to maintain

populations of sensitive or rare species in the face of climate change. Furthermore, insurance effects and facilitation are likely driven by different environmental factors. For example, although insurance effects may not be directly related to evaporative demand, facilitation is stronger and more important when evaporative demand increases (via microclimate amelioration, Wright et al. 2015). Future experiments that manipulate the role of biodiversity during drought should be careful to assess all aspects of drought in order to capture the multiple mechanisms that likely underlie the ability of biodiversity to buffer ecosystems against drought.

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