



Extreme drought impacts have been underestimated in grasslands and shrublands globally

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Edited by Susan Harrison, University of California, Davis, CA; received June 12, 2023; accepted October 6, 2023

Climate change is increasing the frequency and severity of short-term (~1 y) drought events-the most common duration of drought-globally. Yet the impact of this intensification of drought on ecosystem functioning remains poorly resolved. This is due in part to the widely disparate approaches ecologists have employed to study drought, variation in the severity and duration of drought studied, and differences among ecosystems in vegetation, edaphic and climatic attributes that can mediate drought impacts. To overcome these problems and better identify the factors that modulate drought responses, we used a coordinated distributed experiment to quantify the impact of short-term drought on grassland and shrubland ecosystems. With a standardized approach, we imposed ~a single year of drought at 100 sites on six continents. Here we show that loss of a foundational ecosystem function—aboveground net primary production (ANPP)—was 60% greater at sites that experienced statistically extreme drought (1-in-100-y event) vs. those sites where drought was nominal (historically more common) in magnitude (35% vs. 21%, respectively). This reduction in a key carbon cycle process with a single year of extreme drought greatly exceeds previously reported losses for grasslands and shrublands. Our global experiment also revealed high variability in drought response but that relative reductions in ANPP were greater in drier ecosystems and those with fewer plant species. Overall, our results demonstrate with unprecedented rigor that the global impacts of projected increases in drought severity have been significantly underestimated and that drier and less diverse sites are likely to be most vulnerable to extreme drought.

climate extreme | Drought-Net | International Drought Experiment | productivity

Most terrestrial ecosystems are impacted to some degree by drought, defined meteorologically as an anomalous period of low precipitation relative to normal (1). While droughts vary widely with respect to severity, duration, and spatial extent, multi-year drought events that incur catastrophic ecological, economic, and societal impacts tend to capture the lion's share of the attention by scientists and the public (e.g., the 1930's US Dust Bowl, ref. 2; the 2000 to 2003 US Southwest drought, ref. 3; the 2012 to 2016 California Drought, The authors declare no competing interest.

This article is a PNAS Direct Submission.

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This article contains supporting information online at https://www.pnas.org/lookup/suppl/doi:10.1073/pnas. 2309881120/-/DCSupplemental.

Published January 8, 2024.

ref. 4; the 2001 to 2009 Millennium Drought in Australia, ref. 5; the 2015 to 2017 drought in Cape Town, South Africa, ref. 6). Yet, globally most droughts are *short-term*, lasting ~1 y in duration (7). Because short-term droughts are so numerous, they can cause substantial loss of ecosystem functioning at local, regional, and global scales (8, 9). As Earth's climate continues to change, short-term droughts that are statistically extreme in intensity (e.g., rare with respect to the long-term climate record, ref. 10) will become more common (11, 12), with 1-in-100-y droughts potentially happening every 2 to 5 y (7). Indeed, evidence of such drought intensification already exists for some regions (13). Unfortunately, because of the historic rarity of extreme drought, we have limited, and primarily anecdotal, estimates of the magnitude of their ecological consequences.

Knowledge of how short-term extreme drought may alter ecosystem functioning is particularly important for grasslands and shrublands. These ecosystems cover more than 40% of the ice-free terrestrial land surface (14, 15) and are found in every region of the globe (15). Grasslands and shrublands are characterized by high variability and frequent deficits in precipitation (16), and thus, are expected to be the most vulnerable to climate change (17). Moreover, grasslands and shrublands store more than 30% of the global stock of carbon (15) and contribute significantly to variability in global terrestrial carbon sinks (18) and atmospheric CO₂ concentrations (19). Thus, grassland and shrubland ecosystems can be expected to cause greater variation in global carbon cycling with intensifying droughts in the future.

Fortunately, many drought experiments have been conducted in grasslands and shrublands, relative to other ecosystems (e.g., forests, ref. 1), and a consensus has emerged based on recent meta-analyses of these studies. These meta-analyses show the expected-that drought has negative impacts on multiple aspects of ecosystem functioning, particularly those functions related to C cycling (e.g., productivity)-but also that considerable variation in terrestrial ecosystem responses is observed among studies (20, 21). Much of this variation could be caused by differences in the magnitude and duration of the droughts (or alteration in precipitation) imposed among the experiments included in these meta-analyses (20, 21). Although most droughts imposed are not statistically extreme, Wang et al. (21) showed that magnitude and duration were important factors for determining variation in ecosystem responses to experimental alterations in precipitation. They found a linear decrease in ecosystem functioning with greater reductions in precipitation; but, over time, productivity became less responsive to altered precipitation (21). One might conclude from this analysis that the effects of droughts, when imposed at statistically (i.e., historically) extreme levels, would result in even further declines in function with the greatest effects manifested in the short term. However, such extreme reductions in precipitation are uncommon in experiments (10); instead, precipitation reductions are for the most part within the range of nominal variability of a particular ecosystem. Consequently, we lack the critical understanding of how grassland and shrubland ecosystems will respond to a future where historically extreme droughts will become the norm rather than the exception.

Here we report results from the first-of-its-kind coordinated distributed experiment the International Drought Experiment or IDE—designed to impose a statistically extreme, short-term (~1 y, *Materials and Methods*) drought across grassland and shrubland sites globally, using a common methodology (22). At the time of analysis, IDE consisted of 44 sites that experimentally imposed a historically extreme, 1-in-100-y drought treatment for at least a full growing season. The IDE network also provided an additional 56 sites imposing a less severe drought treatment, one that was not extreme by our definition but rather within the range of historic variability (hereafter referred to as nominal drought; Fig. 1 and *Materials and Methods*). These 100 sites were arrayed across six continents and spanned broad climatic (Fig. 1 and *SI Appendix*, Table S2) and edaphic gradients (23). At all sites, we measured annual aboveground net primary production (ANPP, *Materials and Methods*), a foundational component of the global carbon cycle, as a metric of drought-induced loss of ecosystem functioning in these grasslands and shrublands.

The results from this globally distributed experiment allowed us to 1) quantify the effects of short-term extreme drought on ANPP and determine if this effect differed between grassland and shrubland ecosystems globally, 2) compare the effects of extreme drought to less severe, nominal (or non-extreme) drought on ANPP, and 3) broadly assess factors potentially contributing to variation in ecosystems' responses to both extreme and nominal drought. We expected to observe a significant loss in ANPP with extreme drought, and that this loss in ANPP would be greater in grasslands vs. shrublands, consistent with past studies (24–26). Furthermore, we expected that extreme drought would suppress ANPP substantially more than nominal drought. We also expected that the extreme drought effects would differ from those derived in previous meta-analyses of

Significance

Drought has well-documented societal and economic consequences. Climate change is expected to intensify drought to even more extreme levels, but because such droughts have been historically rare, their impact on ecosystem functioning is not well known. We experimentally imposed the most frequent type of intensified drought-one that is ~1 y in duration—at 100 grassland and shrubland sites distributed across six continents. We found that loss of aboveground plant growth, a key measure of ecosystem function, was 60% greater when short-term drought was extreme (≤1-in-100-y historical occurrence). This drought-induced loss in function greatly exceeds previously reported losses for grasslands and shrublands, suggesting that the global impacts of projected increases in drought severity have been substantially underestimated.



Fig. 1. Geographic extent and climate space encompassed by the IDE. The 100 grasslands (green circles) and shrublands (brown triangles) included in the analysis spanned six continents (*A*) and broad gradients of mean annual temperature and MAP (*B*). Closed symbols denote sites (n = 44) that experienced statistically extreme 1-in-100-y drought (i.e., below average annual precipitation during the experiment year). Open symbols denote IDE sites (n = 56) that experienced nominal (not statistically extreme) drought (i.e., average or above-average annual precipitation during the experiment year). Photos: Shown are drought shelters at representative sites on each continent. Drought shelters were designed to exclude a fixed proportion of each rainfall event from the plots below. The proportion excluded was selected to impose a 1-in-100-y drought for each site during years with average annual precipitation (based on long-term precipitation records, see *Materials and Methods* for details; see *SI Appendix*, Table S2 for site codes).

experimental drought results (20, 21, 27), given that past meta-analyses are subject to publication bias (28) and that their effect sizes were based on studies that varied widely in the type, duration, and magnitude of the drought imposed (29–31).

Results and Discussion

Consistent with most previous research, drought experimentally imposed over ~1 y (<2 y), whether nominal or extreme in magnitude, reduced ANPP relative to ambient conditions (Fig. 1 and *SI Appendix*, Table S3). For those sites that experienced extreme drought, ANPP was reduced on average by ~35% overall; by ~38% and 21% for grasslands and shrublands, respectively (Fig. 2*A*). Thus, across the 44 sites that experienced extreme drought, grasslands incurred greater losses in ANPP than shrublands, consistent with previous studies (24–26), though this difference was not statistically significant (*SI Appendix*, Table S4). For those sites that experienced nominal drought, ANPP was suppressed by ~21%, much less (>half) than in sites experiencing extreme drought, and there was a smaller difference between grasslands and shrublands in these nominal drought responses (Fig. 2*A*). This suggests that ANPP of grasslands and shrublands responds similarly to drought unless droughts are extreme, in which case these ecosystems are more likely to diverge in their average response to even a single year of drought.

The 95% CIs for mean reductions in ANPP due to nominal (non-extreme) droughts, and for grasslands and shrublands separately, overlapped with the range of mean effects reported in recent meta-analyses of drought experiments (Fig. 2*A*). This equivalence between the mean effects of nominal droughts in the IDE network and past meta-analyses occurred despite the wide variety of experimental protocols for imposing drought treatments included in the meta-analyses. In contrast, the effect of extreme drought on ANPP was well outside the range of these past reported effects, with the reduction in ANPP more than 1.5-fold greater. Thus, our results suggest that past studies have underestimated the ecosystem effects of statistically extreme droughts—the droughts of the future.

While it is reassuring that there is similarity among the full suite of sites in our study (n = 100), the subset of sites subjected to nominal drought (n = 56), and the mean effect sizes of meta-analyses, there are several reasons to view the IDE estimates as being more robust. First, by including results from all sites (including those



Fig. 2. Response of ANPP to a standardized 1-y drought for 100 IDE sites. (*A*) Mean drought response for sites that experienced extreme drought or nominal conditions for: all sites, grasslands, and shrublands (*SI Appendix*, Tables S3 and S4). The gray bar indicates the range of ANPP loss from –19 to –12.6% found in Song et al. and Wang et al. (20, 21), respectively. Drought response is calculated as: ln(average ANPP_{DROUGHT}/average ANPP_{CONTROL}); 0 (black dashed line) represents no effect of drought, and negative numbers indicate less ANPP in drought vs. control plots. (*B*) Mean drought response for each site ordered from negative (*Top*) to positive (*Bottom*). Site codes and corresponding site information are listed in *SI Appendix*, Table S2. Shown are 95% CIs for mean site-level drought responses. *Indicates site with CI that was omitted for clarity because it exceeds the *x* axis scale.

with no evidence of a drought effect, Fig. 2*B*), we eliminated the long-standing issue of publication bias affecting meta-analysis effect sizes (i.e., bias towards significant results, (28, 32)). In addition, although the statistical power was relatively low for detecting drought effects at individual sites—a concern of many global change experiments (32)—the large number of IDE sites, almost twice as many as included in Song et al. (20) and Wang et al. (21), provided broader and in some cases denser geographic coverage.

Furthermore, the standardized experimental design and sampling protocols we used ensured drought treatments were imposed, and responses assessed comparably, across all sites. Thus, variation in our dataset should be attributable to ecological differences among sites and not methodological differences inherent in meta-analyses. By reducing methodological differences, we anticipated that site-to-site variability would be reduced in this coordinated, distributed experiment (22, 23), at least relative to previous studies. Contrary to that expectation, a surprising amount of variation was still observed in ANPP responses to both extreme and nominal drought across sites (Fig. 2*B*). While 79 sites experienced the expected losses in ANPP, 21 sites were insensitive to the 1-y extreme drought, i.e., control and treatment ANPP means differed by <1% or were slightly higher in treatment plots, suggestive of high resistance to short-term drought (Fig. 2*B*). Variation in ecosystem

responses was also observed with nominal drought, with 39 of 56 sites experiencing a loss in ANPP, but 17 sites displaying high drought resistance. Thus, individual IDE sites still differed in their responsiveness to both extreme and nominal drought despite the use of common protocols. This begs the question: What factors are contributing to the large variation in drought response among sites?

Determinants of Variation in Ecosystem Response to Drought.

Although there are myriad factors that may contribute to site-level variation in the ANPP responses observed, we focused on seven key abiotic and biotic variables that were reliably available for >75% of the IDE sites. These included mean annual precipitation (MAP), previous year's precipitation (relativized by MAP), historic variability in precipitation (expressed as the interannual coefficient of variation of MAP), aridity index (AI), soil texture, plant species richness, and the dominant plant growth-form (expressed as proportion of graminoids) of the ecosystem (*Materials and Methods* and *SI Appendix*, Table S5).

Past empirical studies have indicated that the factors above may underpin variations in ANPP responses among sites to drought. For example, evidence suggests that drier and more arid sites (low MAP and AI) tend to be more sensitive to drought than wetter or less arid sites (33–36). But, in addition to MAP (or AI), historic variability in precipitation also may contribute to variation in drought responses (37, 38). In this case, sites that experience higher year-to-year variation in MAP are expected to be less sensitive to extreme drought, a product of these ecosystems being adapted to large interannual variations in precipitation. Research also suggests that legacy effects of the previous year's precipitation may play an important role in determining plant productivity, such that responses lag behind the increases or decreases in precipitation from the previous year (39–41). Finally, the inverse soil texture hypothesis (42) proposes that plants growing in coarsetextured (sandy) soils should experience less water stress than plants growing in fine-textured soils in relatively arid ecosystems, with the opposite pattern for ecosystems with higher precipitation. This interaction between soil texture and MAP is expected to be amplified with drought, but this prediction has rarely been tested (43). With IDE spanning a broad range of edaphic conditions (23), we provide one of the first tests of the inverse soil texture hypothesis on drought responses.

In addition to the abiotic factors listed above, plant species richness has been shown to influence the magnitude of ecosystem response to drought, with more species-rich communities being more resistant to drought than less species-rich communities (44). There is also abundant evidence that growth forms differ in their sensitivity to drought, with grasses and grass-like plants (i.e., graminoids) typically more sensitive to water deficits than woody plants (45, 46) or forbs (47). Given that the IDE sites represent a range of plant species richness and graminoid abundance (*SI Appendix*, Table S5), we evaluated the relationship between average plot-level species richness and abundance of graminoids and the magnitude of the drought response observed.

Three of these potential sources of variation in drought response had statistical support in the IDE dataset. We found weak evidence (P = 0.08) for MAP and moderate evidence (P = 0.02) for aridity (low values indicate lower plant water availability or more arid sites) being related to magnitude of drought responses (*SI Appendix*, Table S6). Drier sites (lower MAP or greater aridity) experienced greater losses in productivity than wetter grassland and shrublands. This finding matches studies demonstrating that production losses at more arid sites are greater in response to drought (32, 35–37) and supports the Huxman-Smith model (35) of greater sensitivity of ANPP to interannual variation in precipitation (and dry years) in more arid sites. Finally, as demonstrated previously (44), there was moderate evidence for more species-rich sites being more resistant to a loss in productivity than less rich sites (P = 0.04).

In contrast, we found no evidence that previous year's precipitation (as relativized by MAP), CV in MAP, percent sand (a key component of soil texture), or the proportion graminoids explained the variation in drought responses observed across all sites (Fig. 3 and *SI Appendix*, Table S6). We also found no evidence for an interactive effect between MAP and percent sand on drought response (*SI Appendix*, Table S7), and thus no support for the inverse soil texture hypothesis affecting differential drought sensitivity.

The Importance of Drought Severity. As indicated above, drought magnitude or severity (i.e., % reduction in precipitation relative to the control) was an important predictor of ecosystem response to drought in the recent meta-analysis by Wang and et al. (21). Because the passive approach to imposing drought employed with IDE (Fig. 1; *Materials and Methods*) relies on ambient precipitation levels, the actual amount of precipitation that was reduced with the drought treatment at each site (with respect to the long-term record, i.e., MAP) varied with the amount of annual precipitation received during the year of the experiment. We used this variation in drought severity to determine if differences in drought responses could be explained simply by the magnitude of drought imposed (calculated



Fig. 3. Effect of abiotic and biotic factor on drought response. Relationships between 1-y drought responses across IDE sites and (A) MAP, (B) previous year's precipitation (relativized by MAP), (C) interannual percent coefficient of variation (CV) of MAP, (D) natural log of the AI, (E) percent sand, (F) average proportion of graminoids, and (G) average richness of plant species. Information on abiotic and biotic characteristics for each site can be found in *SI Appendix*, Table S5. Model results are summarized in *SI Appendix*, Table S6. Drought response is calculated as: In(average ANPP_{DROUGHT}/average ANPP_{CONTROL}); 0 (black dashed line) represents no effect of drought, and negative numbers indicate less ANPP in drought vs. control plots. Lines are shown only for significant relationships. Shaded area represents the 95% CI. *Previous year's precipitation included the precipitation in the 365 to 730 d preceding the biomass collection date and was relativized by MAP.



Fig. 4. Relationship between drought severity and drought response. A linear mixed effects model found strong evidence for a negative effect of increasing drought severity (becoming more negative) on drought response for 1 y of drought across all sites (intercept = -0.11; slope = 0.53; P = 0.009; adjusted R² = 0.06). Model results for the effects of drought severity on drought response for extreme and nominal sites are summarized in the Main Text. Shaded area represents the 95% CI. Drought response is calculated as: In(average ANPP_{DROUGHT}/average ANPP_{CONTROL}). For drought responses, 0 represents no effect of drought, negative numbers indicate less ANPP in drought vs. control plots. Drought severity is calculated as: (Precip_{DROUGHT}-MAP)/MAP; MAP = mean annual precipitation. Because ambient precipitation during the experiment year determines the severity of the imposed drought, positive drought severity can occur during anomalously wet years when plots beneath drought shelters also experience above average precipitation. The open symbols denote those IDE sites (n = 56) where ambient precipitation was above average, and thus the imposed drought was not statistically extreme (1-in-100 y). Closed symbols denote those sites (n = 44) with average or below average annual precipitation during the experiment year. All of these IDE sites experienced statistically extreme drought. The filled orange square denotes the mean drought response for sites experiencing extreme drought whereas the open orange square is the mean for sites that experienced less severe drought. Note that there was no relationship between drought severity and drought response when only those sites that experienced extreme drought are considered. The red dashed lines provide visual guides for 50% and 75% reductions in ANPP.

as drought severity = (Precipitation_{DROUGHT}-MAP)/MAP). As expected, we found strong evidence (P = 0.009) that increasing (more negative) drought severity led to larger reductions in ANPP when examined across all 100 sites (Fig. 4). Further, the amount of variability explained doubled when drought severity, MAP, previous year's precipitation, CV in MAP, percent sand, and proportion of graminoids were included together in the model (*SI Appendix*, Table S8); though, drought severity remained the only significant factor in the model, underscoring the primary importance of this metric in determining the magnitude of the drought response.

One concern with passively reducing ambient precipitation is that in particularly dry years, differences in ecosystem responses between drought and control treatments tend to be minimized (48). In other words, if ambient precipitation is well below average, ANPP would be expected to be low even in control treatments and further reductions in precipitation with the drought treatment may not cause any additional appreciable reductions in ANPP. Thus, the difference between the drought and control treatments would be small, resulting in an effect size close to zero. This would give the appearance of the site being highly resistant to drought. We examined whether this was a possibility for sites with a large drought severity index (45% reduction in precipitation) and yet a drought response close to zero. We found that this phenomenon may indeed be responsible for the high resistance observed for these few sites (n = 4), but for most sites that exhibited resistance to drought (n = 17), this potential experimental phenomenon

could be dismissed, suggesting that other factors are contributing to high resistance of these sites to a single year of extreme drought.

With the above analyses, we were able to evaluate how climate, soil texture, vegetation structure, and drought severity broadly influence drought response. However, given that drought severity was the primary determinant of variation in drought response, other factors that we were unable to include in our analyses are likely contributing to observed high variability in ecosystem response to extreme drought. Moving forward, a key challenge is to determine what these other factors are and identify those ecosystem attributes (e.g., soil fertility, root:shoot ratios, plant community composition, plant-microbe interactions, etc.) that may strongly influence resistance to a single year of extreme drought. Measurement of these factors at the site level and inclusion of these factors in future analyses will be crucial for predicting and mitigating the impacts of extreme drought as climate changes.

Despite the uncertainty of what may be determining variation in drought response in grasslands and shrublands globally, our analysis suggests that overall, ANPP declines as a linear function of increasing drought severity (Fig. 4). In other words, there was no evidence for catastrophic or nonlinear losses in ANPP when single-year droughts become statistically extreme. However, the results from this globally distributed drought experiment do indicate that losses in ANPP are greater than previously expected when drought is historically extreme. With climate change, droughts are not only expected to become more extreme, but also more frequent and longer in duration. It remains unknown what effects these aspects of intensified drought may have when overlaid with greater losses in ANPP with increased drought severity.

In conclusion, given that many ecosystems, particularly grasslands and shrublands, experience substantial interannual variability in precipitation (16, 49), it is not surprising that short-term precipitation reductions that are not statistically extreme would result in only "moderate" (~20%) losses in productivity. Even such moderate responses are likely to have important implications for the global carbon cycle and the wildlife, livestock, and human populations that rely on plant production. Of greater concern, however, is that grassland sites and grassland and shrubland sites combined experienced a magnified loss of function (more than 1.8- and 1.5-fold greater reduction, respectively) when drought was statistically extreme for ~1 y. Clearly, with climate change increasing drought intensity and frequency (50), and given that effects can linger long after drought ends, even more substantial impacts on the global carbon cycle can be expected. Indeed, reductions in ANPP exceeding 35% are not often observed in moderate droughts of longer duration (24, 51), and a recent study suggests that experimental droughts may underestimate the magnitude of ANPP loss by more than half when compared to naturally-occurring droughts (52). Thus, results from our distributed experimental approach reveal that extreme droughts are likely to substantially slow C sequestration in grasslands and shrublands, surpassing predictions from past meta-analyses (20, 53) and experiments (9, 52). Finally, the underlying cause of the striking range in ecosystem responses to short-term extreme drought, from highly resistant to highly vulnerable, remains unresolved. Results from our globally distributed and standardized drought experiments demonstrated little to no evidence for key factors typically thought to drive ecosystem variability in response to drought: CV of MAP, previous year's precipitation, soil texture, and proportion of graminoids. Yet, we found strong support for MAP/aridity and plant species richness being at least partially predictive of ANPP response to extreme drought. If traditionally invoked variables do not explain most of the cross-site variation in responses to intensified droughts, we must rethink our measurements and experiments to allow us to identify

other underexplored factors. Understanding the determinants of differences in both short- and longer-term drought vulnerability will provide critical insight into both the mitigation potential and adaptative capacity of ecosystems in a future where today's extremes become the norm rather than the exception.

Materials and Methods

The IDE. IDE was initiated in 2013 as part of the Drought-Net Research Coordination Network, funded by the US NSF. Drought-Net is a global network of researchers committed to understanding how terrestrial ecosystems respond to extreme drought. For all network investigators, we provided and continue to provide standard experimental protocols on the DroughtNet website (droughtnet.weebly.com). Sites must follow these protocols for their data to be included in our analyses. At the time of analysis, 141 sites had joined our network, but data from only 100 of the sites (Fig. 1*A* and *SI Appendix*, Table S2) had been submitted and/or met our criteria for inclusion in this analysis (*SI Appendix*, Table S9). These 100 sites were well distributed across gradients of MAT and MAP (Fig. 1*B*) and represented two ecosystem types: grasslands and shrublands. Most (>90%) of the sites were dominated by perennials. Furthermore, like most grasslands and shrublands globally, all the sites had some history of management (*SI Appendix*, Table S10), but only 13 sites that we know of were actively mowed (n = 6), burned (n = 5), or grazed (n = 2).

The target for the IDE drought treatment was a statistically extreme, 1-in-100-y drought imposed year-round. To achieve the target level of extremeness, each site's treatment magnitude (reduction in precipitation) was based on the past 100 y of climate data from the site or 100 y of interpolated data from the Terrestrial Precipitation Analysis tool (54). Because precipitation history and variability are unique to each site, this approach allowed us to target the common level of statistical extremeness by allowing the proportional reduction in precipitation to vary across sites (55 and *SI Appendix*, Fig. S1). This contrasts with the alternative approach of imposing a fixed reduction in precipitation (e.g., 50%), which can result in very different levels of extremeness across sites (55).

The target level of extreme drought was imposed at each site using infrastructure that is commonly used in short-statured ecosystems (56). The infrastructure consisted of two or more open-sided shelters, each a minimum of $2 \text{ m} \times 2 \text{ m}$, with roofs that were partially covered with transparent strips of plastic (either V-shaped or corrugated). The percentage roof coverage was dictated by the target level of precipitation reduction (Fig. 1). This shelter design has been shown to have minimal effects on microclimate (57-59), while matching key characteristics (e.g., number of consecutive dry days, size of events, number of events) of naturally occurring extreme dry years across a range of ecosystems (55). For those sites (n = 9) with both control plots (no infrastructure) and plots with an infrastructure control (i.e., structures that mimic the shading of shelters but allow rain to pass through), there was no evidence that ANPP was affected by the shelter infrastructure (mean difference = 5.25%; 95% CI = -6.39%, 18.33%; t-value = 0.89; df = 33; P = 0.38). This suggests an absence of significant nontarget effects of the drought shelters, which has also been demonstrated in other experiments (57-59).

We also chose the shelter infrastructure to impose drought because it is highly cost-effective and can be consistently deployed across a range of short-statured ecosystems, making it amenable for use in a coordinated distributed experiment (22). However, because we manipulate precipitation passively, the target level of drought extremeness may or may not be achieved in any given year, depending on ambient precipitation amount (48). For example, an above-average year of precipitation will result in drought treatments that are less severe, while a below-average precipitation year will result in an even more extreme drought than the target level. In total, 44 of the 100 IDE sites received average or below-average precipitation in the first year of the treatment, and thus imposed the target statistically extreme, 1-in-100-y drought (*SI Appendix*, Fig. S2). The remaining 56 sites received above-average precipitation, and thus imposed a non-extreme ("nominal") drought, which was within the range of historic variability.

To be included in this analysis, sites needed to collect peak live aboveground biomass as an estimate of annual ANPP (60). While we recognize that there are numerous other ecosystem processes that can be impacted by drought, ANPP was selected because it can be comparably estimated and readily standardized across sites. Moreover, ANPP is a low-cost measurement that requires much less investment of time than other measures of ecosystem functioning–a crucial feature of response variables in CDEs and other successful experimental networks, such as the Nutrient Network (22, 61). ANPP was estimated either destructively and/or non-destructively using methods appropriate for the particular ecosystem as cited in Fahey and Knapp (62), with herbaceous-dominated sites encouraged to follow the Nutrient Network's protocols (https://nutnet.org/). We relied on investigators to use their expertise in determining the most appropriate methods–either destructive or non-destructive–to estimate ANPP for their study system. Sites then separated ANPP estimates into live and dead before further classifying live biomass by growth form (graminoid, grass, forb, woody, etc.) and submitting all estimates in grams of dry biomass per m². Standing dead biomass could be separated into current and previous year's growth where appropriate.

Drought Response Metric. For each site, we calculated relative drought response as the ratio of average ANPP in the drought plots compared to average ANPP in the control plots, as a metric of ecosystem response to imposed drought. Specifically, we adapted equations from Smith et al. (63) and Kreyling et al. (64) to define relative drought response as: In(ANPP_{DROUGHT}/ANPP_{CONTROL}).

Drought Severity Metric. We calculated the actual severity of the drought that was imposed during the year of precipitation manipulation using meteorological data that was either collected 1) on-site (site-submitted) or 2) from a nearby weather station (mean distance = 10.2 km). The nearby weather station data was obtained either from the Global Historical Climatology Network (GHCN, 65) or the Climate Hazards Group InfraRed Precipitation with Stations (CHIRPS, 66). When data from GHCN, CHIRPS, or local weather stations were not available, we used site-submitted annual precipitation estimates. We first calculated the amount of precipitation each site received in the 365 d preceding collection of ANPP. We then used this precipitation calculation, the site-reported percent reduction in precipitation imposed with the drought treatment, and days the drought shelters were in place prior to the collection of ANPP to estimate the total amount of precipitation reduced at each site. To qualify as 1 y of drought, a site's shelters needed to be in place for at least one full growing season (within 1 wk of 120 treatment days prior to harvest), but less than 2 y (within 1 wk of 650 d). For the drought severity metric, we compared the estimated precipitation received in drought plots to MAP (mm) as reported by each site, and calculated the deviation from this number: (Precip_{drought} - MAP)/MAP.

Abiotic and Biotic Factors. We included in our analysis seven abiotic and biotic factors that have been hypothesized to be important in influencing variability in drought response among sites (SI Appendix, Table S5). These factors included: MAP (mm), previous year's precipitation (mm), historical variability in MAP (interannual CV), AI, soil texture, proportion of graminoids, and site richness. As indicated above, we used MAP as reported by each site. We used either site-submitted, GHCN, or CHIRPS data to calculate how much precipitation each site received in the 365 to 730 d preceding the ANPP harvest and then relativized this value by site MAP. We refer to this as previous year's precipitation in our analyses. Historical variability of precipitation was estimated using the average coefficient of variation for the 30 y of precipitation from each site using GHCN station data. The AI (an estimate of plant moisture availability), was calculated as the MAP divided by potential evapotranspiration (67); Trabucco et al. (67) uses estimates from World Clim v2 (68) to calculate these values. Lower AI values indicate lower plant water availability. For the interannual coefficient of variation in MAP, we pulled data from the Multi-Source Weighted-Ensemble Precipitation tool (69).

As a proxy for soil texture, we used a weighted mean for percent sand in the top 100 cm of soil for each site (n = 96) available in the ISRIC World Soil Information (70), which yielded information for 96 sites. We used this global dataset because site-level data was only available for 27 of the sites. However, for those 27 sites we conducted a Pearson correlation test between site-derived values and the global dataset to confirm whether the global dataset measures were accurate, which we found to be the case (r = 0.67, P < 0.001). To calculate the proportion of graminoids, we used 77 sites that submitted ANPP data for each functional group. For each plot at a site, we divided the ANPP of graminoids by the total ANPP (all functional groups added together). We then averaged these proportions across plots for each site. For richness, we used plant species composition data submitted by 68 of the 100 sites at the time of analysis. We only used data from control plots and averaged richness at the plot level for each site. **Statistical Analyses.** For all statistical analyses, we used the language of evidence (71) to describe our results as an alternative to typical statistical significance testing (i.e., using a significance cut-off of *P*-value ≤ 0.05). With this approach, we ascribe *P*-values as following Muff et al. (71): >0.1 = little or no evidence, between 0.05 and 0.10 = weak evidence, between 0.01 and 0.05 = moderate evidence, and ≤ 0.01 = strong evidence.

To test whether ANPP in drought plots differed from that in control plots, we used six one-sided *t* tests in R (version 4.0.2, 72), identifying whether responses were significantly less than 0. We chose to use one-sided *t* tests given that out a priori hypothesis was for drought to result in a reduction in ANPP (rather than either a decrease or increase, which would be relevant for an expectation of increased ANPP with drought). However, to examine whether results differed based on one-sided vs. two-sided *t* tests, we conducted two-sided *t* tests for the six comparisons and found that this did not affect the results, except in the case of the shrubland nominal drought test (one-sided *P*-value = 0.07, two-sided *P*-value = 0.15). We tested sites with nominal and extreme droughts separately, and tested ecosystem types together and separately (*SI Appendix*, Table S3). To test for differences in grassland and shrubland responses to drought across all sites, we used linear regression (*SI Appendix*, Table S4).

To examine whether the seven abiotic and biotic factors described above explained any variation in the drought response observed, we used linear mixed effects models for all 100 sites, or separately for sites that experienced extreme or nominal drought. We built separate models that tested the following explanatory variables: ecosystem type (grassland or shrubland), MAP (mm), previous-year's precipitation (mm, relativized by MAP), average coefficient of variation (%) for MAP, AI (scaled using the natural log), average percent sand, average proportion of graminoids, and average plot-level richness. We also explored the inverse soil texture hypothesis using a linear mixed effects model (site set as a random effect) to test how the interaction of average percent sand and MAP affected drought response.

We used a Pearson correlation to test collinearity among drought severity (proportion), length of drought (days), and a categorical variable for whether sites had ambient precipitation equal to or above (group 1) or below (group 2) MAP. We set drought severity as our fixed effect as it integrates the length of drought (r = -0.59, P < 0.001), deviation in ambient precipitation from MAP (r = 0.60, P < 0.001), and a site's drought shelter design into a single variable. To test the nature of the relationship between drought response and various factors (drought severity, MAP, previous year's precipitation, aridity, coefficient of variation, and plant community richness) we compared both linear and non-linear models (asymptotic regression and general additive model with a spline function set to 3) using AIC. We did not find evidence for a non-linear response (*SI Appendix*, Table S11) and proceeded with building linear models using the Imer function (package stats) in R. We also built a multiple linear regression to see whether drought severity combined with MAP, previous year's precipitation, proportion graminoids, CV, and average percent sand could further explain variation in drought response.

Data, Materials, and Software Availability. Derived data are provided in *SI Appendix*, Table S5. All code and derived data have been deposited in Dryad (73). All other data are included in the manuscript and/or *SI Appendix*.

ACKNOWLEDGMENTS. We are grateful to our IDE collaborators who established and maintained the IDE experiments, collected field data, and shared their data with the IDE community (SI Appendix, Table S1). We thank A. Tatarko for assisting with data management. We also thank the landowners that made the IDE possible. Research support was provided by the following: US National Science Foundation (NSF) Research Coordination Network grant to M.D.S., O.E.S., and R.P.P. (DEB-1354732); US Department of Agriculture's National Institute of Food and Agriculture (USDA-NIFA) Postdoctoral Fellowship grant to K.D.W. (2020-67034-31898); USDA-NIFA Conference Grant to M.D.S. (2020-67019-31757); US Geological Survey (USGS) John Wesley Powell Center for Analysis and Synthesis grant to M.D.S., S.L.C., and S.M.M.; USGS grant to M.D.S. (G21AC10266-00); Global Drought Synthesis Group grant to M.D.S., K.D.W., P.W., O.E.S., L.A.G. funded by the NSF Long-term Ecological Research Network Office and the National Center for Ecological Analysis and Synthesis, University of California-Santa Barbara; National Key Research and Development Program of China (2022YFE0128000; 2021YFD2200405); and National Natural Science Foundation of China (32061123005, 32071627, 31930078, 31971461). Funding for specific experimental sites within this synthesis paper came from the USDA

Forest Service Rocky Mountain Research Station and the USDA Agricultural Research Service, and the findings and conclusions are those of the authors and should not be construed to represent any official USDA determination of policy. Any use of trade, firm, or product names is for descriptive purposes only and does not imply endorsement by the U.S. Government. Additional acknowledgements include: NSF Long Term Research in Environmental Biology (LTREB DEB 1754106 and 2326482) to Arizona State University and Long Term Ecological Research (LTER) program to New Mexico State University (DEB-2025166); Sevilleta and Central Arizona-Phoenix Long-Term Ecological Research Programs (NSF DEB-1655499 ad DEB-1832016); The Institute for the Study of Ecological and Evolutionary Climate Impacts supported sites at University of California-Santa Cruz; USDA Agriculture and Food Research Initiative (AFRI) Physiology of Agricultural Plants Program, Grant #2017-67013-26191; US Department of Energy (DOE) Environmental System Science Program (DE-SC0021980); Federal Ministry of Education and Research (grant 031B1067C); European Research Council [ERC Grant agreement 647038 (BIODESERT)] and Generalitat Valenciana (CIDEGENT/2018/041); USGS Ecosystems Mission Area; CONICET and Universidad de Buenos Aires; Tarbiat Modares University; Department of Biology at Kansas State University and NSF LTER program to Kansas State University (DEB-144048); Austrian Science Fund, Austrian Academy of Sciences and Austrian Research Promotion Agency; Utah Agricultural Experimental Station; Grants from the US NSF Long-Term Ecological Research Program (LTER) including DEB-1234162 and DEB-1831944. Further support was provided by the Cedar Creek Ecosystem Science Reserve and the University of Minnesota; US Bureau of Land Management (Grant No. L16AS00178); NSERC Discovery Grants to J.F.C., E.G.L., and J.L.W.; a joint strategic grant from the Alberta Livestock and Meat Agency (now Alberta Ministry of Agriculture and Forestry) and Emissions Reduction Alberta; CNPg/FAPEG - PELD-PNE (Site 13); DOE Office of Science Early Career Award DE-SC0015898; German Research Foundation (DFG), Grant number DU1688/1-1; iDiv and sDiv, the Synthesis Centre of iDiv (FZT 118, 202548816) and the Gottfried Wilhelm Leibniz Prize (Ei 862/29-1), both granted by the DFG; F.M.F. and D.H. received a PhD scholarship by Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES), Ministry of Education, Federal Government of Brazil; Australian Research Council (DP210102593); Texas A&M Savanna Long-Term Research and Education Initiative, Sid-Kyle Foundation, and Sonora Research Station personnel; Staatsbosbeheer (Dutch State Forestry Service) for giving permission to use the protected nature area at Rhijnauwen; Colorado Agricultural Experiment Station; National Research, Development and Innovation Fund (Fund) of Hungary (112576, 129068); City of Boulder Open Space & Mountain Parks Funded Research Program, Garden Club of America, and USDA National Institute of Food and Agriculture Predoctoral Fellowship (Project Accession Number, 1019166); German Federal Government (BMBF) through the SPACES initiative ("Limpopo Living Landscapes" project-grant 01LL1304D; "SALLnet" project-grant 01LL1802C); Post-doctoral fellowship of CAPES-Brasil, Programa CsF; PGI UNS 24/ZB81; Swiss NSF, grants 149862, 185110; Environment Research and Technology Development Fund (JPMEERF15S11420) of the Environmental Restoration and Conservation Agency of Japan; The Teshio Research Forest of Hokkaido University provided in situ support; German Academic Exchange Programme (DAAD)-SPACES scholarship for short term visit to Germany 2015 to 2017; Australian Research Council (DP150104199; DP190101968); Alexander von Humboldt Foundation (AvH; grant 33000351); The University of Winnipeg, In-kind support provided by Manitoba Beef and Forage Initiatives Inc.; Catalan Government grants SGR 2921-1333, the Spanish Government grant PID2022-140808NB-I00, and the Fundación Ramón Areces grant CIVP20A6621; PI-IUNRN 40-C-873; the French government IDEX-ISITE initiative 16-IDEX-0001 (CAP 20-25); Brazilian National Research Council (CNPq grant 307689/2014-0) and Fundação de Amparo à Pesquisa do Estado do Rio Grande do Sul (FAPERGS grant 17/2551-0001106-6); Post-doctoral fellowship of Inter-American Institute for Global Change Research (IAI)CRN3005, which is supported by the US NSF (Grant GEO-1128040); Great Western Woodlands Supersite, part of Australia's Terrestrial Ecosystem Research Network, and thanks to the Department of Biodiversity, Conservation and Attractions Western Australia for hosting the site and assistance with construction of shelters; US Department of Agriculture–National Institute of Food and Agriculture award 2019-68012-29819; National Council of Scientific and Technological Development (CNPq, grant number 310340/2016-0); Research Funds for ICBR (1632021023); Israel Ministry of Science and Technology; National Natural Science Foundation of China (31870456); National Research Foundation, Grant No: 116262; DFG Priority Program SPP-1803 "EarthShape: Earth Surface Shaping by Biota" (TI 338/14-1

and -2), and the German Ministry of Education and Research (BMBF); Generalitat Valenciana, Project R2D-RESPONSES TO DESERTIFICATION (CIPROM/2021/001); NORWEGIAN RESEARCH COUNCIL MILJØFORSK project 255090 (LandPress: Land use management to ensure ecosystem service delivery under new societal and environmental pressures in heathlands); Australian Research Council (DP150104199; DP190101968, DP210102593); Centre for Integrative Ecology, Deakin University; The Hermon Slade Foundation, Australia; Estonian Research Council (PRG609); Research Station Bad Lauchstädt of the Helmholtz-Centre for Environmental Research-UFZ, Germany; National Research Foundation Grant CSRU180504326326; and European Commission (GYPWORLD, H2020-MSCA-RISE-777803) and Spanish Government (Querpin PID2021-126927NB-I00).

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