Small-scale water deficits after wildfires create long-lasting ecological impacts

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Abstract

Ecological droughts are deficits in soil–water availability that induce threshold-like ecosystem responses, such as causing altered or degraded plant-community conditions, which can be exceedingly difficult to reverse. However, ‘ecological drought’ can be difficult to define, let alone to quantify, especially at spatial and temporal scales relevant to land managers. This is despite a growing need to integrate drought-related factors into management decisions as climate changes result in precipitation instability in many semi-arid ecosystems. We asked whether success in restoration seedings of the foundational species big sagebrush (Artemisia tridentata) was related to estimated water deficit, using the SoilWat2 model and data from >600 plots located in previously burned areas in the western United States. Water deficit was characterized by: (1) the standardized precipitation-evapotranspiration index (SPEI), a coarse-scale drought index, and (2) the number of days with wet and warm conditions in the near-surface soil, where seeds and seedlings germinate and emerge (i.e. days with >5 cm deep soil water potential >−2.5 MPa and temperature above 0 °C). SPEI, a widely used drought index, was not predictive of whether sagebrush had reestablished. In contrast, wet-warm days elicited a critical drought threshold response, with successfully reestablished sites having experienced seven more wet-warm days than unsuccessful sites during the first March following summer wildfire and restoration. Thus, seemingly small-scale and short-term changes in water availability and temperature can contribute to major ecosystem shifts, as many of these sites remained shrubless two decades later. These findings help clarify the definition of ecological drought for a foundational species and its imperiled semi-arid ecosystem. Drought is well known to affect the occurrence of wildfires, but drought in the year(s) after fire can determine whether fire causes long-lasting, negative impacts on ecosystems.

Introduction

Moisture deficit is one of the key ways that climate variability affects ecological processes, populations, community composition and success or failures of land management interventions, such as restoration of native vegetation. However, there is little guidance on how to determine when water deficit becomes ecologically impactful and thus constitutes ‘ecological drought’ (Slette et al 2019, Zang et al 2019). Drought has traditionally been defined as a deficit in precipitation relative to evapotranspiration (i.e. meteorological drought), shortage of water pools on earth (i.e. hydrological drought), or water deficits affecting human needs (i.e. agricultural or socioeconomic drought; Wilhite and Glantz 1985, Crausbay et al 2017). Ecological drought is a deficit in water availability that causes ecosystems to cross adaptive
thresholds, resulting in major changes in their structure and function (Crausby et al 2017, Snyder et al 2019). Even though the duration and magnitude of a water deficit causing threshold-like change must be known for a particular ecosystem or community in order to designate and predict ecological drought, there are few examples of ecological drought being resolutely (i.e. highly-resolved to a specific time, soil-depth, and soil–water availability) quantified in ways that are spatially and temporally robust and thus, generalizable.

‘Drought’ has traditionally been identified by meteorological or hydrological drought indices such as Palmer Drought Severity Index (PDSI), Standardized Precipitation Index (SPI), and more recently, by the Standardized Precipitation Evapotranspiration Index (SPEI) (Alley 1984, Beguería et al 2010, Vicente-Serrano et al 2012). Vicente-Serrano et al (2012) empirically compared these three indices to one another and determined that SPEI was more flexible on shorter time scales than PDSI (i.e. monthly) and it could better identify drought impacts than SPI, owing to SPEI’s inclusion of water outputs in evapotranspiration. However, quantifying ecological drought using site-level condition data (e.g. edaphic properties, vegetation structure, etc) from meteorological/hydrological drought indices can result in mischaracterizing the severity and impact of ecological drought.

Drought indices have been found to be correlated with certain coarse-scale shifts in ecosystem productivity, such as annual net primary productivity (ANPP) at the regional scale, which has provided understanding of climate effects on seasonal plant responses (Ruppert et al 2015). However, temporary reductions in ecosystem productivity do not constitute threshold-like changes if they are reversible, which is often the case when vegetation composition remains intact. Therefore, these types of changes do not qualify as ecological drought. In contrast, loss of dominant species in a plant community does constitute a threshold change if the loss is permanent or long-term (Bestelmeyer et al 2018, Law et al 2018). Drought indices do not relate well at the individual plot or local community scale, and thus are poor predictors of within-season plant responses and demographic processes such as germination, establishment, or mortality, which can lead to restructuring of plant communities. To better define and understand ecological drought, we must first identify the ecologically-relevant and biophysically meaningful metrics of water deficit that lead to demographic success or failure of a given plant species.

One biophysically relevant metric that directly relates to water deficit and plant survival is water potential. This metric provides a concise means of quantifying the availability of water to plants and other organisms (Campbell and Norman 1998), especially when it is measured at specific places and times where plants have critical interactions with soil and air (Breshears et al 2009, Roundy et al 2018). Using soil–water potential to define and quantify ecological drought should facilitate detection of water availability thresholds that influence key plant demographic transitions. Several studies have addressed threshold responses to variability in soil–water potential at the individual plant-, plot-, or site-level (e.g. Adams et al 2009, Breshears et al 2009, Roundy et al 2018), but few of these studies addressed threshold responses at the population-level (i.e. whole-stand) or across a species’ range.

Ecological droughts may be most likely to induce rapid change in plant communities recovering from disturbance, whether the recovery is natural or augmented by restoration. In certain ecosystems, plant germination and initial seedling survival are both sensitive to water deficits. These demographic bottlenecks can limit post-restoration plant establishment, particularly in drylands such as sagebrush steppe (James et al 2011, 2019, Kildisheva et al 2016). Restoration seedings in sagebrush ecosystems often fail in part due to soil–water deficits (i.e. ecological drought) during critical germination or emergence periods (Gornish et al 2015, Germino et al 2018). However, few studies have quantified the daily soil-surface conditions that most likely affect the demographic success of seeds and seedlings (but see Roundy et al 2018, James et al 2019 for examples involving grass species). Soil-surface moisture can become uncoupled from weather station-measured rain, atmospheric vapor pressure deficit, and bulk soil volumetric water content due to solar and terrestrial radiation balances at the soil surface—especially during the spring plant-emergence period. Nonetheless, current approaches for assessing demographic responses to soil–water availability tend to use climate variables such as annual evapotranspiration, seasonal water supply, or climatic water deficit (Dilts et al 2015). To date, no studies have attempted to define or quantify ecological drought, which could be a major driver in restoration failure or success.

Weather and ‘drought’ impacts on restoration are of particular interest in western North America, where only about half of the ~1000 000 km² once dominated by big sagebrush (Artemesia tridentata Nutt) and perennial bunchgrass communities remains. Sagebrush populations or stands have been steadily extirpated when wildfires, induced by exotic annual grasses, burn repeatedly and at increasingly larger extents (Brooks et al 2004, Balch et al 2013). Without major intervention, the remaining sagebrush steppe is at risk of ecosystem degradation and conversion to annual grasslands. Managers of these landscapes often seed after fire with mixtures of sagebrush and other shrubs, perennial grasses, and forbs to stabilize the soil, compete with exotic annual grasses, and rehabilitate wildlife habitat (Pilliard et al 2017b). These restoration efforts and investments have been substantial in recent decades due to increased fire frequencies, with hundreds of thousands of hectares seeded annually at large
costs (Pilliod et al 2017b). Thus, understanding the drivers that lead to seeding success is a priority for land management organizations in dryland ecosystems as outlined in the US Integrated Rangeland Fire Management Strategy (http://integratedrangelandfiremanagementstrategy.org/).

Here, we use a data set of sagebrush occupancy, SPEI, and simulations of soil–water potential from 627 plots that were seeded after wildfires across the Great Basin over the last ~40 years (figure 1) to investigate the influence of ecological drought on restoration outcomes. We asked whether successful and unsuccessful sagebrush seedings (i.e. plots with and without sagebrush years to decades after seeding) could be related to either the SPEI index or to soil–water potential in the specific times and locations (soil surfaces) where seeds and seedlings emerge and face strong selection and mortality. We focus on the month of March, when sagebrush typically germinate and emerge, and when soil water is typically at or below soil–water potential thresholds of −2.5 MPa and soil temperatures are typically above 0°C (Meyer and Monsen, 1992, DiChristina and Germino 2006, Schlaepfer et al 2014). To further guide our attempt to quantify ecological drought we hypothesized that: (1) SPEI, a commonly used drought index, would be a useful predictor of plot-level big sagebrush establishment, which we infer based on its presence years after fire and seeding, and (2) soil–water potentials, modeled from weather data, provide a more robust means of quantifying the magnitude and duration of water deficit that constitutes ecological drought for sagebrush.

Methods

Plot locations and sampling

Sagebrush abundance was measured between 2011 and 2016 on 627 plots across the Great Basin (figure 1). Each plot burned once between 1976 and 2008 prior to our sampling. Plots were seeded with big sagebrush in the fall or winter after fire, as documented in federal agency records contained in the Land Treatment Digital Library (Pilliod and Welty 2013). Of the 627 plots, 452 were placed in randomly selected burned-treated areas and the other 175 were placed in sites identified by the Bureau of Land Management cooperators as having successful post-restoration sagebrush establishment. We used an additional 127 random plots that burned once between 2009–2015 and were subsequently seeded during the following fall or winter with big sagebrush for statistical model validation.

Three 50 m AIM style belt transects were used to quantify sagebrush density in each plot (Herrick et al 2005), although the analysis here focuses on sagebrush occupancy (present or not detected). The central start point was randomly determined and the three transects proceeded away from this central point at an event 120° separation for a length of 50 m each. The width of the belt transects was variable to provide a balance of sampling time allocated per plot relative to the relative abundance of sagebrush. Transects were set at 6 m wide by default, but widths were reduced to 4 m, 2 m, or 1 m if observers expected to detect more than 20, 50, or 70 individual plants per transect, respectively.

Figure 1. Map of 627 sagebrush sampling sites in the western United States of America (a), distribution of elevation (stacked histogram), mean annual precipitation (MAP) and mean annual temperature (MAT) (b). Sites with sagebrush present are in blue and sites where sagebrush was not detected are in orange.
**SPEI calculation**

Pre-emergence drought severity was estimated at every plot using the SPEI (Vicente-Serrano et al. 2010, Beguería and Vicente-Serrano 2013) for the periods including 1, 2, 3, and 6 months prior to seedling emergence, which was assumed to be March. SPEI is calculated on a monthly basis from a climatic water balance that incorporates moisture deficit or surplus across a user-defined time scale and is adjusted to a log-logistic probability distribution. The SPEI averaging periods were determined from analyses of SPEI computation lengths, soil depths and corresponding $R^2$ values (unpublished data). SPEI was calculated at each plot using the ‘SPEI’ package in R (Beguería and Vicente-Serrano 2013), using values of precipitation and potential evapotranspiration extracted from the University of Idaho Gridded Surface Meteorological Dataset (gridMET; Abatzoglou 2013) for 1979–2014 at a 4 km resolution. Because the meteorological data are only available from 1979–2014, we were unable to use the 6 earliest burned plots for this analysis; hence our sample size was $N = 621$ for the SPEI-based model.

**Ecohydrological modeling**

Soil–water availability was simulated from measured and estimated inputs for each plot using SOILWAT2 (Version 3.2.2, Schlaepfer and Murphy 2018), a daily time-step, multiple soil layer, process-based, simulation model of ecosystem water balance (Schlaepfer et al. 2012, Bradford et al. 2014). SOILWAT2 uses weather data and soil properties (i.e. texture) at each soil layer, and monthly vegetation (live and dead biomass, litter, and active root profile) to simulate daily water balance processes. We used the Livneh (1915–2008, 6 km resolution) and PRISM (2009–2015; 4 km resolution) daily weather datasets to extract daily temperature (minimum and maximum) and precipitation, as well as the monthly normals of cloud cover, wind speed and relative humidity, which were downscaled to daily values for input into SOILWAT2 (Livneh, 2014), PRISM Climate Group 2017). Livneh data were used for the 627 model development plots and PRISM data were used for the 127 validation plots so that weather data corresponded with respective post-fire time periods. Plot specific soil properties and characteristics were obtained for each plot through a combination of field sampling and use of the SSURGO national database where field sampling data were not available (Soil Survey Staff 2017). The field sampling for soil data consisted of recording soil depth and determining soil texture for each plot (Barnard et al. 2019). We analyzed soil texture at 0–10 cm depth for 335 plots in this study using the Bouyoucos hydrometer method including peroxide pre-treatment and extended settling time for detection of clays. At each plot four soil subsamples were taken for analysis, which were then averaged together for input into SoilWat2. Nearly all sagebrush establishment after fire results from seeds that germinate during spring (March) in the year after fire, when adequate surface moisture and suitable air and soil temperatures coincide and cover of other plant species is still absent or scarce (Schlaepfer et al. 2014, Germino et al. 2018). Our analyses focused on this time period (March) and assumed 0% cover of other plant species. We also focused on the thin, near-surface soil-layer (0–5 cm) where sagebrush seeds germinate and emerge.

**Analyses**

All analyses and figures were completed in RStudio (R Core Team 2019). SPEI is calculated on a monthly basis, meaning that the calculation considers at least the previous 30 d. Thus, to determine if sagebrush occurrence (presence or not detected) could be explained by variation in SPEI among the plots, we needed to consider the duration for which SPEI was calculated prior to the presumed March germination and emergence period. We ran a binomial generalized linear mixed model (GLMM) to relate sagebrush occupancy (binary response variable) to the SPEI values (continuous, fixed effects predictor variable) for each SPEI monthly calculation (1, 2, 3, or 6 month(s)), with plot ID as a random effect ($\alpha = 0.05$; lme4 R package, Bates et al. 2015).

The second analysis was a logistic regression GLMM of the 627 plots to relate sagebrush occupancy (response variable) to March near-surface soil–water potentials, March soil temperatures, and elevation (explanatory variables). Soil–water potential and soil temperature were each averaged across March to provide mean March values for each plot. We included plot ID as a random effect in the binomial logistic regression GLMM ($\alpha = 0.05$; lme4 R package). Results from this model were used to predict the probability of sagebrush establishment using the 127-plot validation dataset.

In the final analysis, we used a gamma log-link function GLMM to assess the relationship between simulated soil–water potentials (absolute values of SWP as the response variable) and the fixed effects of sagebrush emergence (binary, present or not detected), soil temperature, and elevation (both scaled to improve model convergence), plus the random effects of plot ID and Julian calendar day in March for which soil water data were simulated. The plot ID and calendar day factors were included to explain variability across space and time. We parameterized the GLMM with data from all 627 plots and then validated the model using our 127 validation plots and the same gamma log-link GLMM structure. Due to the complex model covariance structure we could not predict the probability of each validation plot being occupied using the original GLMM, however were able to assess whether the GLMM model estimates, levels of significance, and model residuals were similar when
parameterized with validation plots (Wegner and Olden 2012).

We quantified the duration of ecological drought using two different approaches. First, we calculated the mean temperature and water availability for each day in March, separately for plots that were occupied by sagebrush and for plots where sagebrush was not detected. Second, for each plot, we tallied the number of days with soil temperatures above an \textit{a priori} threshold of 0 °C and soil water potentials less than an \textit{a priori} threshold of −2.5 MPa. These \textit{a priori} thresholds were based on thermal requirements for germination (Meyer and Monsen 1992) and based on xylem water potentials that cause net photosynthesis to be zero or less in sagebrush seedlings establishing in the first-year post-fire (DiChristina and Germino 2006).

We quantified ecological drought for the 627 model development plots as well as the 127 validation plots and examined the difference in number of days between the two data sets.

**Results**

SPEI did not differ between plots based on sagebrush occupancy, regardless of whether SPEI was calculated using values from 1, 2, 3, or 6 m prior to the March post-fire (figure 2).

Sagebrush was present at 69% of the 627 plots used to parameterize the models and at 57% of the 127 validation plots. In the logistic regression model, mean March soil–water availability was not a significant predictor of sagebrush occurrence (figure 3). However, increases in soil temperature significantly reduced the probability of sagebrush establishment (figure 3; \(P = 0.002\)) and sites with higher elevation had significantly greater probability of sagebrush occurrence (figure 3; \(P < 0.001\)). Moreover, we found that \textit{daily} soil–water availability at the soil-surface during the narrow March window following each post-fire seeding was a strong predictor of sagebrush occupancy (figure 4; \(P < 0.001\); table 1). Increased soil temperatures reduced soil–water availability, whereas increases in elevation increased soil–water availability (table 1; \(P < 0.001\) and \(P < 0.001\) respectively). Running the same gamma log-link GLMM with the 127 validation plots resulted in similar coefficients and model residuals as were obtained for the full model using the 627 plots. Plots occupied by sagebrush had greater soil–water availability than unoccupied plots (figure 4; \(P = 0.046\); table 1), and soil temperature was negatively related to soil–water availability (figure 4; \(P < 0.001\); table 1). Elevation was not a significant predictor in the gamma log-link GLMM parameterized with the 127 validation plots, though elevation was a significant factor in the GLMM parameterized with the 627 plots. The explained variance improved for the 127 validation plots compared to the 627 plots (\(R^2 = 0.35\), \(R^2 = 0.13\) respectively; table 1).

Of the 627 plots, those with sagebrush present had an average of 23 d during the March following fire and seeding in which mean daily water availability and temperatures of soil were \(-2.5\) MPa and \(>0\) °C, respectively, compared to an average of 16 d for plots where sagebrush was not detected (\(\Delta = 7\) d). In contrast, analysis of our 127 validation plots revealed that
Figure 3. Probability of sagebrush seedling establishment as a function of monthly soil–water availability, soil temperature and elevation. Gray region in the 95% CI and points show data used to fit the model.

Figure 4. Mean differences in water potential (lower panels and lower half of top panels) and temperature (top of upper panels) in the soil surface (0–5 cm) in the year following fire and seeding, for sites that did or did not have sagebrush during observations made in 2011–2016) (blue and orange, respectively). The green box indicates the critical germination and emergence period in March, which is zoomed into in the top panels. Two water potential thresholds are highlighted, including the nominal permanent wilting point (−1.5 MPa), and the point at which photosynthesis in seedlings ceases (−2.5 MPa). In the upper panels, solid lines represent soil–water potentials and dashed lines are soil temperatures. The two panels on the left represent 627 plots, while the two right panels represent 127 plots used for validating statistical models.
plots where sagebrush was present averaged 26 d meeting these criteria, compared to an average of 24 d for plots where sagebrush was not detected ($\Delta = 2$ d).

**Discussion**

Defining and then quantifying ecological drought requires relating soil–water availability to plant responses, and here we demonstrate this using a large dataset of post-fire seeding outcomes for big sagebrush in the Great Basin. Counter to expectations, we found that early spring SPEI did not predict post-fire and restoration sagebrush occupancy (figure 2). Thus, we conclude that SPEI is likely unsuited to quantify ecological drought for our application. However, in support of our second hypothesis, daily soil–water availability during March, a direct measure of water availability for plants, was useful in quantifying ecological drought as it pertains to sagebrush in the Great Basin (figure 4, lower panel). Daily soil–water availability combined with soil temperature was important for understanding when ecological drought conditions were met (figure 4 upper panel), whereas similar metrics, measured on a monthly-scale, were not predictive (figure 3). To our knowledge, this is the first time that daily soil–water availability and soil temperature have been used to identify the duration (i.e. less than 7 d) of water deficit capable of leading to a demographic bottleneck of sagebrush establishment, ultimately qualifying as an ecological drought. The influence of short-term ecological drought on sagebrush restoration success has clear implications for understanding how climate change may affect sagebrush population dynamics in the Great Basin (Shriver et al. 2019).

Ecological droughts are often thought of in terms of months or years rather than days. However, more frequent, short-term droughts can have profound negative impacts on plant survival (Adams et al. 2009). Although we assessed sagebrush occupancy years later, our data suggest that initial sagebrush germination and emergence success is dictated by daily water availability and temperatures of soil in the first March post-fire. Our results are similar to those of James et al. (2019), who found increased soil temperatures reduce the establishment of seedlings in the Great Basin (figure 3). Similarly, James et al. (2019) and Young et al. (2019) also did not find a correlation between a climatic drought variable, monthly climatic water deficit (CWD = potential evapotranspiration/actual evapotranspiration), and seedling establishment (figure 2). Climatic variables (i.e. those based on monthly or yearly timeframes) may be too coarse for predicting seedling establishment because of the narrow temporal windows associated with establishment. This suggests that daily environmental conditions are needed to understand seedling bottlenecks.

In the Great Basin, soil water-availability during early spring is critical for native plant species to germinate, emerge, and establish. Water deficits during this critical time period results in decreased establishment success (James et al. 2011, Roundy et al. 2018). In the near-future (i.e. 2050), climate conditions are

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**Table 1.** Two generalized mixed-effects models: (1) logistic regression GLMM with sagebrush presence as the response variable; (2) Gamma log-link GLMM with soil–water availability as the response variable. The gamma log-link GLMM has two datasets present: 1) original data that had 627 plots, and 2) validation data that had 127 plots. To compare effectiveness of the GLMM an $R^2$ value is listed. Model variable significance is represented through bolded values ($\alpha = 0.05$).

| GLMM Type                  | Response Variable | Estimate | Std. error | P value | $R^2$ |
|----------------------------|-------------------|----------|------------|---------|-------|
| **Logistic regression GLMM** | Response variable: sagebrush presence |          |            |         |       |
|   | (Intercept)        | 1.49     | 0.21      | < 0.001  |       |
|   | Soil–water availability | 0.04     | 0.05      | 0.402   |       |
|   | Soil temperature   | −0.14    | 0.04      | 0.002   |       |
|   | Elevation          | 0.74     | 0.15      | < 0.001 |       |
| **Gamma log-link GLMM**    | Response variable: soil–water availability |          |            |         | 0.13  |
|   | (Intercept)        | −0.71    | 0.20      | < 0.001  |       |
|   | Sagebrush Present  | −0.67    | 0.19      | < 0.001  |       |
|   | scaled Soil Temperature | 0.70    | 0.01      | < 0.001  |       |
|   | scaled Elevation   | −0.45    | 0.09      | < 0.001  |       |
| **Validation data (N = 127)** |                      |          |            |         | 0.35  |
|   | (Intercept)        | −0.50    | 0.27      | 0.064    |       |
|   | Sagebrush Present  | −0.68    | 0.34      | 0.046    |       |
|   | scaled Soil Temperature | 0.56    | 0.03      | < 0.001  |       |
|   | scaled Elevation   | −0.30    | 0.17      | 0.076    |       |
predicted to include increases in daytime minimum temperatures by 2 °C as well as increases in winter precipitation in the form of rain (Palmquist et al 2016, Brabec et al 2017, Snyder et al 2019). Our results suggest that a 2 °C increase in daytime minimum temperatures would cause a ~13% reduction in sagebrush establishment probability (figure 3). An increase in springtime temperatures would also dry soil surfaces faster, potentially leading to establishment failure (current study; Shriver et al 2018) if increased temperatures are not accompanied by increased March precipitation. Warmer and wetter winter conditions will also favor exotic-annual grass establishment (Concilio et al 2013, Compagnoni and Adler 2014), which can increase wildfire prevalence in the Great Basin (Abatzoglou and Kolden 2011, Bradley et al 2016, Pilliod et al 2017a).

Climate change and specifically drought will impact both fire occurrence and perhaps more critically, ecosystem recovery, especially reestablishment of foundational perennial species, in the Great Basin. Germination and emergence will likely occur earlier in the late winter or early spring, potentially shifting the ‘window’ of establishment when water-availability and temperature conditions are each more suitable. However, it is difficult to predict if the temporal ‘window,’ during which suitable temperature and water availability overlap for seedling establishment, will become longer or shorter. Warmer winter conditions could eliminate snow from the soil surface, causing more, or stronger, freeze-thaw events that reduce seedling establishment because of frozen soils and a lack of available soil–water (Gornish et al 2015). Thus, restoration of semi-arid landscapes may become even more necessary (because of reduced natural re-establishment) and more difficult because of climate change, though improvements in weather forecasting can help mitigate the impacts on restoration investments (Hardegree et al 2018). Understanding how ecological droughts impact different plant life stages and species is critical for determining restoration potential for sites. We suggest that ecologists evaluate the use of soil–water availability in appropriate spatio-temporal scales, especially at time periods when critical demographic transitions occur, as a means to define and quantify ecological drought. This approach gives mechanistic insights into how plant populations and communities will change because of soil–water deficits.

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Data availability

The data that support the findings of this study are openly available at doi:10.5066/P9LDKQE2.

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