



# Non-native plant invasion after fire in western USA varies by functional type and with climate

Janet S. Prevéy · Catherine S. Jarnevich · Ian S. Pearse · Seth M. Munson · Jens T. Stevens · Kevin J. Barrett · Jonathan D. Coop · Michelle A. Day · David Firmage · Paula J. Fornwalt · Katharine M. Haynes · James D. Johnston · Becky K. Kerns · Meg A. Krawchuk · Becky A. Miller · Ty C. Nietupski · Jacquilyn Roque · Judith D. Springer · Camille S. Stevens-Rumann · Michael T. Stoddard · Claire M. Tortorelli

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**Abstract** Invasions by non-native plant species after fire can negatively affect important ecosystem services and lead to invasion-fire cycles that further degrade ecosystems. The relationship between fire and plant invasion is complex, and the risk of invasion varies greatly between functional types and across geographic scales. Here, we examined patterns and predictors of non-native plant invasion following fire across the western United States. We

specifically analyzed how the abundance of non-native plants after fire was related to fire characteristics and environmental conditions, such as climate, soil, and topography, in 26,729 vegetation plots from government networks and individual studies. Non-native plant cover was higher in plots measured after wildfires compared to prescribed burns or unburned plots. The post-fire cover of non-native species varied by plant functional type, and only the cover of short-lived (i.e., annual and biennial) forbs and short-lived C3 grasses was significantly higher in burned plots compared to unburned plots. Cool-season short-lived

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J. S. Prevéy (✉) · C. S. Jarnevich · I. S. Pearse  
U.S. Geological Survey, Fort Collins Science Center, 2150  
Centre Avenue, Building C, Fort Collins, CO 80526, USA  
e-mail: [jprevey@usgs.gov](mailto:jprevey@usgs.gov)

S. M. Munson  
U.S. Geological Survey, Southwest Biological Science  
Center, 2255 N. Gemini Dr., Flagstaff, AZ 86001, USA

J. T. Stevens  
USDA Forest Service Research and Development,  
Washington, DC, USA

J. T. Stevens  
Department of Biology, University of New Mexico,  
Albuquerque, NM, USA

K. J. Barrett · C. S. Stevens-Rumann  
Colorado Forest Restoration Institute, Department  
of Forest and Rangeland Stewardship, Colorado State  
University, Campus Mail 1472, Fort Collins, CO 80523,  
USA

J. D. Coop  
Western Colorado University, Gunnison, CO 81231, USA

M. A. Day  
USDA Forest Service, Rocky Mountain Research Station,  
Missoula Fire Sciences Lab, 5775 US Highway 10 W,  
Missoula, MT 59808, USA

D. Firmage  
National Park Service, 1 Zion Park Blvd, Springdale,  
UT 84720, USA

P. J. Fornwalt  
USDA Forest Service, Rocky Mountain Research Station,  
240 West Prospect Road, Fort Collins, CO 80526, USA

K. M. Haynes  
USDA Forest Service, Intermountain Region, 324 25th  
Street, Ogden, UT 84401, USA

grasses composed most of the non-native post-fire vegetation, with cheatgrass (*Bromus tectorum*) being the most recorded species in the dataset. Climate variables were the most influential predictors of the cover of non-native short-lived grasses and forbs after fires, with invasion being more common in areas with drier summers and a higher proportion of yearly precipitation falling in October through March. Models using future projected climate for mid (2041–2070) and end (2071–2100) of century showed a potential for increasing post-fire invasion risk at higher elevations and latitudes. These findings highlight priorities for mitigation, monitoring, and restoration efforts to reduce post-fire plant invasion risk across the western United States.

**Keywords** *Bromus tectorum* · Climate change · Disturbance · Exotic annual grasses · Invasive plants · Management · Wildfire

## Introduction

Invasions by non-native plant species following fire can have significant and lasting impacts on ecosystem services such as biodiversity, soil stability, and forage production (Hobbs and Huenneke 1992; Coates et al. 2016; Nagy et al. 2021). Non-native species may also alter trajectories in early successional communities, lead to the exclusion of native early seral species (Kulmatiski 2006), and have negative impacts on other biotic interactions (Mack et al. 2000; Coates et al. 2016). Non-native plants often invade native ecosystems after disturbances such as fire remove

established vegetation, which can increase the availability of limiting resources and benefit invasive species establishment (Davis et al. 2000; Davis and Pelser 2001; Shea and Chesson 2002; Daehler 2003). In some ecosystems, non-native plant invasions can also increase fire risk (Fusco et al. 2019), and create a positive invasion-fire feedback that can result in alternative, less diverse ecosystem states (D’Antonio and Vitousek 1992; Brooks et al. 2004; Davies et al. 2012).

Fire and invasion are both complex processes that are influenced by environmental conditions (climate, soil, topography), human disturbance (grazing pressure, proximity to roads and trails, and the overall intensity of disturbance), fire regime characteristics (past fire regime, departure from fire regime), and biological factors (pre-fire plant community characteristics, non-native propagule pressure, and potential invader characteristics). All these factors interact to influence how “invasible” a given ecosystem is after a fire (Keeley et al. 2003, 2011; Merriam et al. 2006; Perkins et al. 2011; Ellsworth et al. 2016; Williamson et al. 2020). For example, some shrubland ecosystems in western USA historically had long fire return intervals prior to European settlement, but now experience more frequent fires (Bukowski and Baker 2013), and this departure from past conditions can allow new species to invade (Keeley et al. 2011). Fire severity may also influence invasion, as higher severity may lead to greater disturbance, which can increase available resources for invasive plants (Fornwalt et al. 2010; Miller et al. 2010; Sherrill and Romme 2012). Invasion can increase landscape-scale fire spread and severity, which has been linked to tree mortality,

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J. D. Johnston  
College of Forestry, Oregon State University,  
Corvallis, CO, USA 97331

B. K. Kerns  
USDA Forest Service, Pacific Northwest Research Station,  
Corvallis Forestry Sciences Lab, Portland, OR, USA

M. A. Krawchuk  
Department of Forest Ecosystems and Society, College  
of Forestry, Oregon State University, 336 Richardson Hall,  
Corvallis, OR, USA 97331

B. A. Miller  
Blue Mountains Forest Partners, P.O. Box 595,  
Mt. Vernon, OR 97862, USA

T. C. Nietupski  
USDA Forest Service, 2150 Centre Avenue, Building A,  
Fort Collins, CO 80526, USA

J. Roque  
USDA Forest Service, Medicine Bow - Routt National  
Forests and Thunder Basin National Grassland, 2468  
Jackson St., Laramie, WY 82070, USA

J. D. Springer · M. T. Stoddard  
Ecological Restoration Institute, Northern Arizona  
University, Flagstaff, AZ 86011, USA

C. M. Tortorelli  
Department of Plant Sciences, University of California,  
Davis, CA, USA

a shift to non-forest vegetation types, and in some cases increase the potential for invasion (Coop et al. 2020; Kerns et al. 2020; Woolman et al. 2022; Tortorelli et al. 2023). However, to our knowledge there has been no research examining how fire severity may relate to plant invasions across large climate gradients and multiple ecosystem types.

To date, most of the negative impacts associated with the invasion of non-native plants after fire in North America have been in hotter, drier regions of the western part of the USA (D'Antonio and Vitousek 1992; Brooks 1999; Fusco et al. 2019; Wilder et al. 2021). Invasion by non-native plants can increase fine fuel loads and fuel continuity in historically sparsely vegetated and fuel-limited systems. Coupled with hotter and drier weather, invasion can increase the extent and frequency of both subsequent large destructive wildfires and subsequent invasions via a positive feedback (Brooks et al. 2004; Fusco et al. 2019; Wilder et al. 2021). Prescribed fire, which is often used to achieve restoration goals (Ditomaso et al. 2006; Calo et al. 2012), can similarly exacerbate plant invasions in some cases (Keeley et al. 2007; Sherrill and Romme 2012; Roundy et al. 2018). However, not all fires, or fire types, result in invasion by non-native species (Alba et al. 2015), some invasive species can reduce fire spread (Stevens and Beckage 2009), and not all non-native species become abundant after fires (Byers et al. 2002; Smith et al. 2008).

The non-native species that are successful after fires in semi-arid ecosystems are often fast-growing annuals that establish quickly and grow early after disturbances (Wolkovich and Cleland 2011), preempting newly available soil resources and outcompeting slower-establishing native species with more conservative growth strategies (Davis et al. 2000; Montesinos 2022). A changing climate in conjunction with an increase in the size and severity of fires at higher elevations and in more northern regions may lead to an increased risk of post-fire plant invasion in formerly resistant or resilient ecosystems (Concilio et al. 2013; Taylor et al. 2014; Lembrechts et al. 2016; Kerns et al. 2020). Increasingly, the risk of invasion by non-native grasses after disturbances at higher altitudes and in forested ecosystems has been recognized (Kerns et al. 2020; Smith et al. 2022). Forested, formerly invasion-resistant regions may not return to their historical forested state after high severity wildfires with climate change (Coop et al. 2020; Woolman

et al. 2022), and this can lead to increased potential for an even broader footprint of plant invasions (Franklin et al. 2006). Land managers are concerned with the higher potential for invasion risks associated with climate change and mega-fires (Williams 2013; USDA 2022), and thus need information to focus limited resources on managing potential invasions after large fires across broad spatial scales.

To deal with threats posed by increasing wildfire activity and intensity under warmer and drier conditions, managers and government officials are directing large amounts of money for prescribed fires and fuels treatments (see, e.g., USDA 2022). Consequentially, understanding the impacts of wildfires and prescribed fires on non-native plants at the landscape-scale is a high priority. Given the large number of interacting factors that can potentially determine post-fire invasion risk, it is difficult to accurately predict areas that are at the greatest risk of invasion after fires, both now and in the future (Beaury et al. 2020). To address the uncertainty associated with the interconnected risks of plant invasion and fire, we performed a quantitative data synthesis to explore how the abundance of non-native plants after fire varies in relation to fire characteristics and environmental conditions such as soils, topography, and climate, using vegetation data collected on the ground from across the western USA. We synthesized this large dataset to examine which types of invasive plants are the most successful invaders after fires and identify where post-fire plant invasions are most common.

We hypothesized the following: (1) Non-native plant species will differ in their successful establishment after fires based on their functional type and life history. Specifically, we hypothesized that non-native short-lived (annual and biennial) herbaceous species would be more abundant in burned plots compared to non-native long-lived herbaceous and woody species, because short-lived herbaceous species may respond to an increase in limited resources following disturbances more rapidly than slower growing plant species (Melgoza et al. 1990; Davis et al. 2000). (2) Non-native plant functional types that increase in cover following fires will be more abundant following wildfires compared with prescribed fires and will be most abundant in severely burned areas and areas that historically had long fire return intervals prior to European colonization of North America. We hypothesized this because

greater levels of disturbance and greater departures from historical conditions may increase invasion risk (Davis et al. 2000; Keeley et al. 2011). (3) Abundance of functional types that increase after fires will vary across climatic gradients. Specifically, we hypothesized that non-native plant functional types that benefit from fire will also be more abundant in regions with greater summer drought, and a greater proportion of yearly precipitation falling outside the traditional growing season, because quick growing and winter-annual non-native plants may more effectively capture limited pulses of resources after disturbances like fire and outcompete natives in these conditions (Melgoza et al. 1990; Davis et al. 2000; Prevéy and Seastedt 2014; Bradley et al. 2018). Strong correlations between climate and post-fire invasion may also indicate that invasion risk could change in the future as the climate changes.

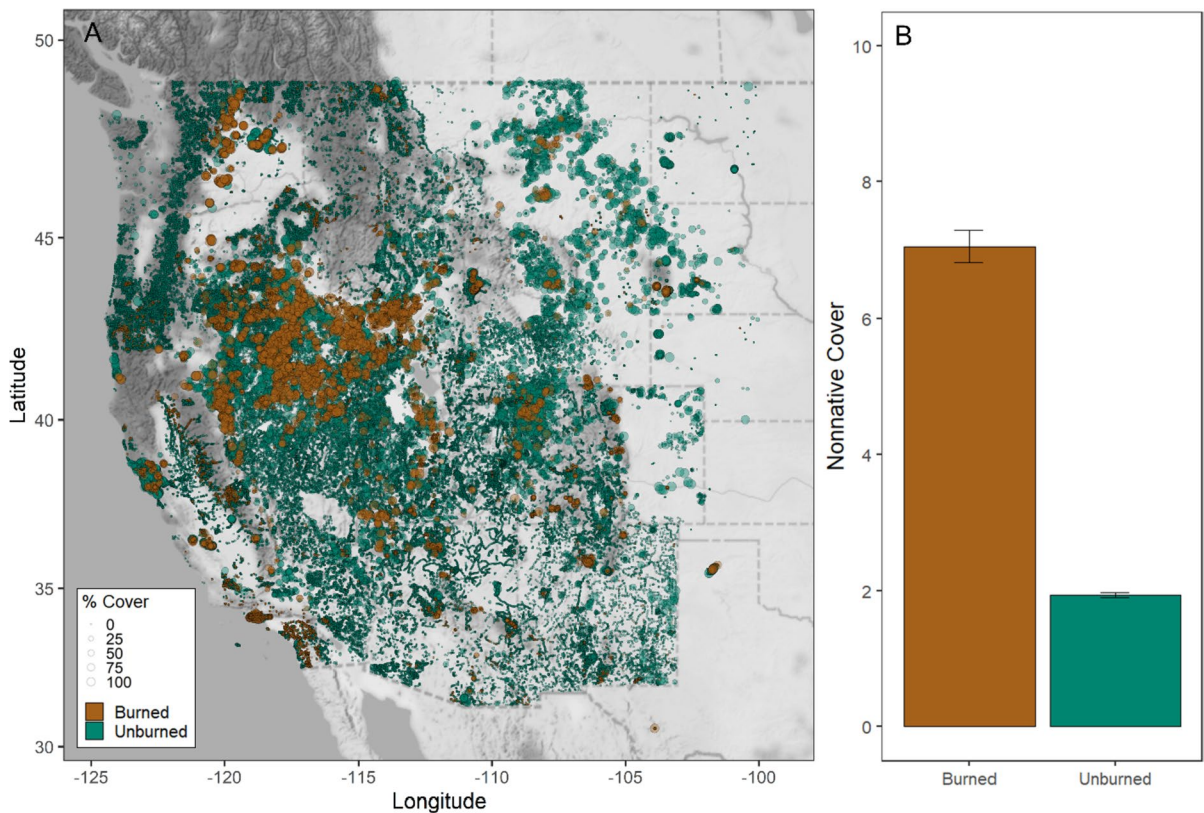
## Materials and methods

We compiled a large dataset of non-native plant abundance, fire characteristics, and environmental predictors across the western United States. We focused our analyses on vegetation plots in the Western USA, west of  $-100$  longitude, as there were too few vegetation plots measured after fires in the eastern part of the country to examine broad-scale patterns there. For these analyses, we combined data from two national-scale datasets of plant species cover data compiled from federal, state, county, and other sources: the Standardized Plant Community with Introduced Status Database (SPCIS; Petri et al. 2022) and the Public LANDFIRE Reference Database from the Landscape Fire and Resource Management Planning Tools program (LANDFIRE 2016a, b). Three of the main contributors of plot-level vegetation data to these databases were the Bureau of Land Management (BLM) Inventory and monitoring program, the National Park Service (NPS) Inventory and Monitoring plots, and U.S. Department of Agriculture (USDA) Forest Service (USFS) Forest Inventory and Analysis Program. The LANDFIRE reference database comes from county, state, and federal government sources that contributed plant cover data to the LANDFIRE project (LANDFIRE 2016a, b). To supplement this dataset, we added plot-level data from numerous targeted post-fire vegetation studies in the western USA

(Firmage and Ronsani 2021a, b, Table S1). These studies measured the absolute percent foliar cover of plant species within plots, or frequency of all plant species located at intervals along transects (hereafter, “plots”). We then used fire perimeter data from the combined wildland fire datasets for the United States and certain territories from 1800s to 2020 (Welty and Jeffries 2021), and fire perimeter and severity data from the Monitoring Trends in Burn Severity (MTBS) database, from 1980 through 2020 ([www.mtbs.gov/](http://www.mtbs.gov/)) to identify all plots that were measured up to 50 years following a fire and quantify the number of times plots had burned prior to measurement. We identified the historical mean fire return interval for all fires and historical fire regimes prior to European colonization of North America for each plot using derived layers from LANDFIRE (LANDFIRE 2016b). The resulting combined dataset includes cover data from 237,799 vegetation plots located west of  $-100$  longitude, measured between 1970 and 2020 (Fig. 1). Of these plots, 26,729 were burned from 1 to 50 years prior to vegetation measurements (Fig. 1). This dataset consists of plots measured at a single point in time; and thus results shown here infer impacts via a space-for-time substitution (Sofaer et al. 2018).

We summed the percent cover of non-native plants by plant functional type and photosynthetic pathway in each plot across the combined dataset, using the following categories: woody (trees and shrubs), short-lived forbs, long-lived forbs, C3 short-lived grasses, C3 long-lived grasses, and C4 grasses. There were very few non-native C4 grasses present in the dataset, so the short-lived and long-lived C4 categories were combined to allow for statistical comparisons of cover between burned and unburned plots. Vines were included in the forb categories. ‘Short-lived’ indicates annuals and biennials, and ‘long-lived’ refers to perennial plants. Non-native status and lifeform categories are based on the native status, duration, and growth habit information from the USDA Plants database (USDA and NRCS 2022) and photosynthetic pathway information was derived from Waller and Lewis (1979), Belmonte and de Agrasar (2002) and Bruhl and Wilson (2007). Plots with summed cover values for any functional type that totaled over 100% were converted to 100%.

To address our first hypothesis that some plant functional types would be more abundant after fire than others, we fit separate models with each



**Fig. 1** **A** Locations of all plots in the dataset. Brown points indicate plots that burned up to 50 years prior to vegetation measurements ( $n=26,729$ ), and green points indicate plots that were not identified as having burned within 50 years of measurements ( $n=211,070$ ). The size of points relates to the percent foliar cover of non-native plant species recorded at that

plot. **B** Average cover of all non-native plant species in plots burned up to 50 years prior to measurement, and unburned plots,  $\pm$  standard error. Map tiles for base map by Stamen Design, under CC BY 4.0. Data by OpenStreetMap, under ODbL

non-native functional type’s plot-level cumulative percent cover (hereafter ‘cover’) as response variables and burn status (burned or unburned) as the predictor variable. To examine our second hypothesis that abundance of non-native plant functional types following fires will depend on fire type and severity, we separately analyzed how fire type of the most recent fire (wildfire or prescribed fire), number of times burned over the past 50 years, time since the most recent fire, severity of the most recent fire based on burn severity mosaics from the MTBS database, historic fire return interval, and fire regime influenced the percent cover of non-native plants. Plots with fire types listed in MTBS as ‘unknown’ or ‘wildland fire use’ were grouped with the wildfire category. MTBS thematic burn severity classes were binned into three categories prior to analysis: ‘Low’, ‘Medium’, and

‘High’, based on changes in the Normalized Burn Ratio from LANDSAT images taken from before and after fires (Eidenshink et al. 2007). We performed separate analyses of thematic burn severity for plots located in forested versus grassland or shrubland land cover types according to the North American Environmental Atlas (NALCMS 2015). Fire regime analysis was restricted to categories with over 1000 observations. We tested for significant effects of all above variables using generalized linear mixed-effects models (glmm) in the statistical program R (R Core Team 2022). To accommodate the unbalanced nature of the data and the many plots where no non-native plants were found, we fit models with tweedie distributions using the glmmTMB package (Brooks et al. 2017). Tweedie distributions are generalized power law distributions that have been used

to model continuous data with many zeros (Dunn et al. 2018), and here provided a better fit and convergence than other distributions for our statistical models. For each analysis we also fit mixed-effects models only using data for each functional type in plots where those functional types were recorded (i.e., we removed plots with zero values for that specific functional type). We did this to account for potential biases in the way plot-level vegetation data were gathered. For example, if data on certain types of species were not measured in a plot, this would result in erroneous zero values. Additionally, a lack of cover of a non-native plant species may result from a lack of seed sources in an area, and not necessarily be indicative of responses to fire. Ecoregion, dataset, and survey year were considered random variables in models. Ecoregion categories were based on Level I Ecoregions of North America (Omernik and Griffith 2014), with the ‘North American Deserts’ category split into ‘Warm deserts’ and ‘Cold deserts’ to distribute the number of burned plots more evenly between regions (Fig. S1). We calculated pseudo  $R^2$  values for mixed effects models with continuous predictor variables using the DHARMA package (Hartig 2022), and we performed Tukey’s post hoc tests for multiple comparisons for models with categorical predictor variables (historic fire return interval and fire regime) using the multcomp package in R (Hothorn et al. 2008). Model assumptions, including lack of dispersion and normality of residuals, were checked visually in quantile–quantile plots, histograms of residuals, and plots of residuals versus fitted values.

To address our third hypothesis and examine how climate and other environmental factors influenced the post-fire invasion risk of functional types that benefited from fire, we modeled the abundance of different functional types using random forest models across the western USA. Random forest models were chosen for this portion of the analysis because they are robust to unbalanced, continuous data with non-normal distributions (Cutler et al. 2007; Mi et al. 2017), and have been found to perform well for spatial predictions of ecological phenomena (Evans et al. 2011). We used plot-level cover of non-native functional types in burned plots as response variables and included ecologically relevant bioclimatic variables derived from ClimateNA normals from 1981 to 2010 (Wang et al. 2016) as potential predictor variables. We also derived several precipitation variables

that we hypothesized would be important predictors of non-native plant success. Specifically, we calculated the total precipitation received over the growing season (April–September). We also determined the ratio of yearly precipitation received in winter (October–March), March, and spring (March–May). Our reasoning behind this is that some non-native species may be more successful in regions that receive more precipitation in winter or early spring and less over the traditional growing season (Prevéy and Seastedt 2014). We included a predictor variable calculating distance from roads and trails, as they can be vectors for the introduction and spread of invasive plant species (Gelbard and Belnap 2003). We also considered soil variables (% sand, % clay, bulk density, pH), and geography (slope, aspect, and continuous heat-insolation load index), as gradients in soils and topography are often associated with where plants grow on the landscape (Table 1). Preliminary correlation analyses were conducted to identify predictor variables that were strongly correlated with each other. We removed predictor variables that were correlated at > than 0.7 (Dormann et al. 2013), retaining the set of uncorrelated variables that had the most ecological relevance (Table 1). We also tested the effect of each predictor variable on non-native cover separately using glmms as described above to ensure the final selected variables were also those that explained a significant amount of variance in post-fire non-native plant cover individually. We performed ecoregion specific analyses for our six generalized ecoregions to test whether important predictors of post-fire invasion varied by ecoregion. We compared results to those from random forests trained on spatially thinned datasets that retained data from only 10,000 plots selected to be maximally distant from each other, and we compared results to random forest models only including data from plots where plant functional groups were present. We also compared results for random forests trained with data only from the most common non-native C3 short-lived grass in the dataset, cheatgrass (*Bromus tectorum*), and random forests trained with all non-native C3 short-lived grasses except cheatgrass to examine how these results reflect its influence and whether other annual grasses show similar responses. We fit random forest models using the randomForest package (Liaw and Wiener 2002) in R, and assessed accuracy using Pearson correlations between training and out of bag samples. We assessed spatial

**Table 1** All potential predictor variables considered for random forest models of post-fire non-native plant abundance across the western United States

Variable abbreviation	Variable name	Calculations/units	Source
<b>AHM</b>	<b>Annual heat-moisture index</b>	<b>(MAT + 10)/(MAP/1000)</b>	<b>Wang et al. (2016)</b>
April precip	April precipitation	mm	Wang et al. (2016)
August precip	August precipitation	mm	Wang et al. (2016)
BD	Bulk density of soil	kg/cubic-meter	Ramcharan et al. (2018)
bFFP	The day of the year on which the frost-free period (FFP) begins	DOY	Wang et al. (2016)
<b>CHILI</b>	<b>Continuous Heat-Load Index, based on latitude, aspect, and slope</b>	<b>Index between 0 and 1</b>	<b>USGS (2019)</b>
<b>Clay %</b>	<b>Percent clay content in top 5 cm of soil</b>	<b>%</b>	<b>Ramcharan et al. (2018)</b>
CMD	Hargreaves climatic moisture deficit	mm	Wang et al. (2016)
DD_0	degree-days below 0 °C	Number of days	Wang et al. (2016)
eFFP	The day of the year on which FFP ends	DOY	Wang et al. (2016)
EMT	Extreme minimum temperature over 30 years	°C	Wang et al. (2016)
Eref	Hargreaves reference evaporation	mm	Wang et al. (2016)
EXT	Extreme maximum temperature over 30 years	°C	Wang et al. (2016)
FFP	Frost-free period	Number of days	Wang et al. (2016)
<b>FRI</b>	<b>Historic fire regime interval</b>	<b>Mean number of years between fires</b>	<b>LANDFIRE (2016a, b)</b>
<b>GSP</b>	<b>April to September precipitation (mm)</b>	<b>(mm), Calculated from Climate NA monthly data</b>	<b>Wang et al. (2016)</b>
July precip	July precipitation	mm	Wang et al. (2016)
June precip	June precipitation	mm	Wang et al. (2016)
MAP	mean annual precipitation	mm	Wang et al. (2016)
<b>MAR</b>	<b>Mean annual solar radiation</b>	<b>(MJ m<sup>-2</sup> d<sup>-1</sup>)</b>	<b>Wang et al. (2016)</b>
Mar p	March precipitation	mm	Wang et al. (2016)
MAT	Mean annual temperature	°C	Wang et al. (2016)
May precip	May precipitation	mm	Wang et al. (2016)
MCMT	Mean coldest month temperature	°C	Wang et al. (2016)
MSP	May to September precipitation	mm	Wang et al. (2016)
MWMT	Mean warmest month temperature	°C	Wang et al. (2016)
MWP	October to March precipitation	mm	Wang et al. (2016)
Nitrogen	Nitrogen content	% weight	Ramcharan et al. (2018)
<b>Northness</b>	<b>Northness</b>	<b>Cosine (Aspect in radians)</b>	<b>USGS (2019)</b>
<b>PAS</b>	<b>Precipitation as snow between August in the previous year and July in the current year</b>	<b>mm</b>	<b>Wang et al. (2016)</b>
pH	Ph of soil	pH	Ramcharan et al. (2018)
<b>Remote</b>	<b>Remoteness from roads and trails</b>	<b>Cost surface based on travel times to nearest road or trail</b>	<b>US Census (2018)</b>
<b>Sand %</b>	<b>Percent sand in top 5 cm of soil</b>	<b>%</b>	<b>Ramcharan et al. (2018)</b>
Sept. precip	September precipitation		Wang et al. (2016)
SHM	summer heat-moisture index	((MWMT)/(MSP/1000))	Wang et al. (2016)
<b>Slope angle</b>	<b>Slope degrees</b>	<b>°</b>	<b>USGS (2019)</b>
SOC	Soil organic content	g/kg	Ramcharan et al. (2018)
TD	Temperature difference between MWMT and MCMT, or continentality	Temperature difference in °C	Wang et al. (2016)

**Table 1** (continued)

Variable abbreviation	Variable name	Calculations/units	Source
<b>WPR</b>	<b>% of annual precipitation received in Winter (Oct.-March)</b>	<b>%, Calculated from Climate NA monthly data</b>	Wang et al. (2016)

Variables in bold are the uncorrelated ( $<0.07$ ) predictors selected for final random forest models based on correlation analyses. Full citations for sources can be found in the Citations

autocorrelation of model results by visual inspection of residuals across plot locations in the western USA to ensure there were no spatial patterns in areas of over or under prediction.

To project potential climate-driven changes in post-fire invasion risk of functional types that benefited from fire, we used future projections of climate variables across the western USA to map predicted changes in post-fire risk for mid (2041–2070) and end (2071–2100) of century under a medium (SSP245) and a high (SSP585) greenhouse gas emission scenarios (Eyring et al. 2016; Wang et al. 2016). The SSP245 scenario is a medium radiative forcing scenario that predicts an additional radiative forcing of  $4.5 \text{ W m}^{-2}$  by 2100, and the SSP585 scenario represents the upper boundary of the range of climate change scenarios and predicts an additional radiative forcing of  $8.5 \text{ W/m}^2$  by the end of the twenty-first century (Eyring et al. 2016). We used downscaled future climate projections from a CMIP6 13-model ensemble (Adaptwest Project 2022; Wang et al. 2016, Mahony et al. 2022) to identify regions where climatic conditions matching those predicting current post-fire invasion risk may exist in the future.

## Results

Non-native plants were present in only 12% (28,391) of the 237,799 plots in the dataset. There were 667 non-native species identified across all plots in this dataset (Table S2). By non-native functional type, C3 short-lived grasses were present in 19,067 plots, C3 long-lived grasses were present in 6479 plots, short-lived forbs were present in 15,541 plots, long-lived forbs were present in 4873 plots, C4 grasses were present in 429 plots, and shrubs and trees were present in 1256 plots (Fig. S2).

Overall, burned plots had higher cover ( $z=47.13$ ,  $p<0.0001$ ) and species richness ( $z=46.10$ ,  $p<0.0001$ ) of non-native plants compared with

unburned plots (Table 2, Fig. 1). C3 short-lived grasses comprised most of the non-native post-fire vegetation (Fig. 2), and the C3 short-lived grass *Bromus tectorum* was the most common and abundant species in the dataset (Table 3, Table S2). Post-fire cover of non-native species varied by plant functional type, and only cover of non-native C3 short-lived grasses and short-lived forbs were significantly higher in burned plots compared to unburned plots when including all plots, as well as only the subset of plots where the non-native functional types were present (all  $p<0.0001$ , Table 2, Fig. 2).

Eleven percent (26,729) of the 237,799 plots in the dataset had burned within 50 years prior to measurement. Wildfire burned 25,661 plots; prescribed fires burned 1067 plots. Cover of non-native C3 short-lived grasses and short-lived forbs was higher following wildfires compared with prescribed burns, both in analyses including zero cover values, and when zero values were excluded (all  $p>0.05$ , Table 4, Fig. 3).

We were able to identify the total number of times a plot was burned prior to measurement for 18,306 of the 26,729 burned plots. Of these, 8932 of the plots burned once prior to measurement, 4828 burned twice, 2463 burned three times, 1161 burned four times, and 922 plots had burned 5 or more times prior to measurement. The number of times a plot had burned prior to measurement was weakly correlated with the abundance of C3 short-lived grasses and short-lived forbs (all  $p<0.06$ ; pseudo  $R^2=0.01$ , Table 4).

We identified the MTBS burn severity data class of the most recent fire for a subset of 6244 plots in the western USA that had burned after 1984 in fires larger than 1000 acres; 3831 of these plots were binned in the ‘low’ fire severity category, 1577 in the ‘medium’ category, and 836 plots in the ‘high’ category. There was no trend for higher cover of C3 short-lived non-native grasses or forbs with higher fire severity across all ecosystem types, or within either the shrubland/grassland or forested ecosystems (all  $p>0.07$ , pseudo



**Table 2** Estimates of fixed effects of fire status on non-native cover and species richness of the non-native plant functional types, for analyses with and without plots recording 0% cover of the functional type

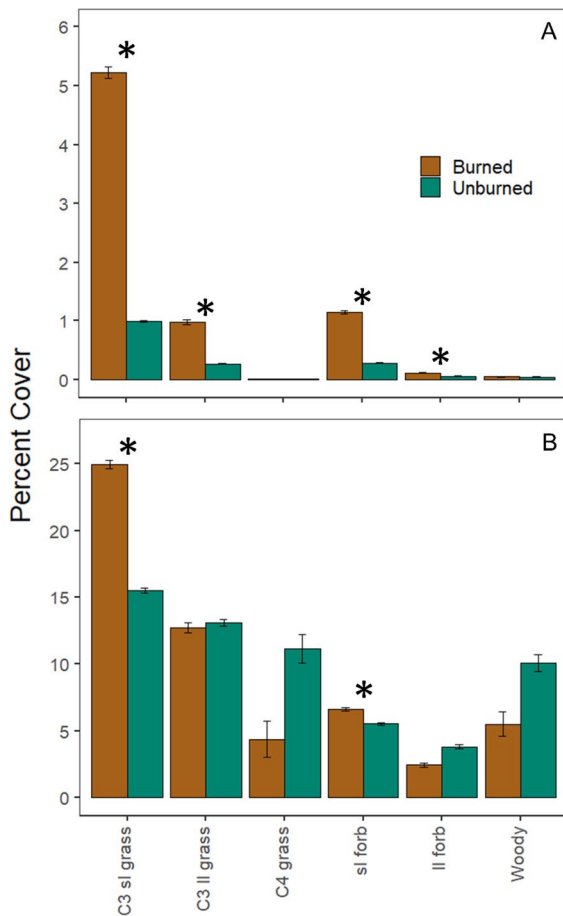
Non-native functional type	Fixed effect	Model type	Estimate	se	z	p	# Obs
Total non-native cover	Burned	All plots	1.57	0.03	48.21	<0.00001	237,799
Total non-native cover	Burned	Non-zero plots	0.60	0.02	27.49	<0.00001	28,391
Non-native sp. richness	Burned	All plots	0.85	0.02	46.35	<0.00001	237,799
Non-native sp. richness	Burned	Non-zero plots	0.21	0.01	19.76	<0.00001	28,391
c3 sl grass	Burned	All plots	1.64	0.04	46.15	<0.00001	237,799
c3 sl grass	Burned	Non-zero plots	0.57	0.02	24.67	<0.00001	19,067
c3 ll grass	Burned	All plots	1.86	0.07	25.00	<0.00001	237,799
c3 ll grass	Burned	Non-zero plots	-0.02	0.05	-0.52	0.60	6479
c4 grass	Burned	All plots	0.53	0.56	0.95	0.34	237,799
c4 grass	Burned	Non-zero plots	-0.18	0.28	-0.64	0.52	429
sl forb	Burned	All plots	1.54	0.04	38.19	<0.00001	237,799
sl forb	Burned	Non-zero plots	0.21	0.03	8.13	<0.00001	15,541
ll forb	Burned	All plots	1.09	0.08	12.95	<0.00001	237,799
ll forb	Burned	Non-zero plots	-0.01	0.06	-0.22	0.83	4873
Woody	Burned	All plots	0.74	0.34	2.20	0.06	237,799
Woody	Burned	Non-zero plots	-0.22	0.16	-1.36	0.17	1256

$R^2 < 0.15$ , Table 4). There was also no trend for a decrease in non-native C3 short-lived grass or short-lived forb cover based on the amount of time between when a fire occurred and when a plot was measured (all  $p > 0.07$ , pseudo  $R^2 < 0.009$ , Table 4). Both short-lived non-native functional types were more abundant in plots that historically experienced longer fire return intervals (all  $p < 0.03$ , Fig. 4), and were most abundant in fire regime category IV-A, which historically had fire return intervals from 36 to 100 years and experienced replacement fires greater than 66.7% of the time (all  $p < 0.05$ , Fig. 4, Table 4).

Random forest models that included data from all burned plots throughout the western USA showed that severity of drought, growing season precipitation, and the ratio of yearly precipitation received in winter were the most important variables predicting cover of the two functional types that benefited from fire in this dataset: non-native C3 short-lived grasses and non-native short-lived forbs (Tables 5 and 6, Fig. 5). These variables were also highly significant predictors of post-fire cover in single-variable glmms (Tables S3 and S4). The order of importance of the top two predictor variables did not differ between non-native C3 short-lived grasses and non-native short-lived forbs. The order of importance differed somewhat for random forests including only plots where lifeforms were

present, and models only including data from specific ecoregions (Tables S5 and S6). However, a climate variable was always ranked as the most important for each model type, with summer precipitation being the most important variable in most models (Tables S5 and S6). For non-native C3 short-lived grasses, non-native short-lived forbs, only *Bromus tectorum*, and all C3 short-lived grasses excluding *Bromus tectorum*, higher projected post-fire cover was predicted in semi-arid basins and steppes across the western USA for the current time period (Fig. 5). Areas projected to currently have higher post-fire cover of *B. tectorum* differed slightly from those projected to have higher cover of short-lived forbs or other C3 short-lived grasses, with higher predicted cover for *B. tectorum* overall, but relatively higher projected post-fire cover of short-lived forbs and other C3 grasses in the central and Mojave basins of California, and in the northern plains (Fig. 5).

Future predictions of invasion risk based on current relationships between post-fire cover of non-native C3 short-lived grasses and short-lived forbs and important environmental variables indicate that the risk of invasion may increase at higher elevations and latitudes, and in northern portions of the short-grass prairie (Figs. 6 and 7). Future projections show increased probabilities for more severe summer



**Fig. 2** Mean percent cover of non-native plant functional types in unburned plots and plots in the western United States that had burned in all fire types up to 50 years prior to measurement,  $\pm$  standard error. Asterisks indicate significant differences in mean cover by burn status. **A** Means of all cover response variables from all plots measured, including plots with 0% cover for the variable. **B** Means of all cover response variables only including plots where the target lifeform was present. Note the change in values on the y-axis. Maps of cover for each lifeform, along with sample sizes, are shown in Fig. S2

drought, as well as an increase in the percentage of total precipitation occurring in winter and early spring and falling as rain rather than snow. The higher SSP585 scenario projects a greater risk of invasion by mid-century with further increases in post-fire invasion risk at higher elevations and latitudes by the end of century, and the lower SSP245 scenario shows a more moderate increase in post-fire invasion risk that stabilizes by end of the century (Figs. 6 and 7).

## Discussion

Short-lived non-native plants were more abundant after wildfires than in unburned plots throughout the western USA, as predicted, which is in line with results from studies at smaller spatial scales (Alba et al. 2015; Williamson et al. 2020; Sofaer et al. 2022). Time since fire and fire severity did not significantly correlate with cover of non-native species, although satellite-derived measurements of fire severity may not be highly accurate across forest and shrubland ecosystems (Zhu et al. 2006; Storey et al. 2021). Short-lived non-native species were more abundant in burned areas with historically longer fire return intervals. Departure from historic fire regimes may create a pathway for invasion, as the plant species present in these ecosystems were adapted to disturbance regimes prior to European settlement, and the recent departure from these regimes may allow colonization by new species better evolved to deal with a higher disturbance frequency (Keeley et al. 2011; Brooks and Chambers 2011). This interpretation is reinforced by the relatively higher abundance of these non-natives following wildfire than prescribed fire (which is generally low severity in western forests), though we did not observe an effect of wildfire severity across the entire dataset.

In our dataset, short-lived C3 grasses were the most common non-native plants to establish after fires, with *Bromus tectorum* being the most abundant of these. Post-fire invasion of these non-native short-lived grasses was more likely in areas with low summer precipitation, higher annual heat-to-moisture ratios, and a higher proportion of yearly precipitation falling in winter and early spring. Currently, short-lived non-native C3 grasses are most invasive in semi-arid regions of western North America after fires. In these ecosystems, disturbance and departure from historical fire regimes (e.g. atypically severe or frequent fire) are important factors allowing for plant invasion (Lembrechts et al. 2016). Non-native species that can escape harsh conditions temporally (short-lived) and grow quickly (non-woody) can potentially thrive after disturbances, whereas the native species in these ecosystems often have a more conservative strategy to resist and survive harsh conditions (e.g. long-lived, slow growing, small leaves, thick cuticles, woody). In contrast, in moderate, wetter climates, many native species have evolved to grow quickly

**Table 3** The six most frequently encountered non-native species by functional group, the number of plots the species were identified in across the dataset, and the average percent cover of each species in plots where they were present

Lifeform	Species	# of Plots	% Cover
Short-lived C3 grasses	<i>Bromus tectorum</i>	14,213	18.6
	<i>Bromus arvensis</i>	2144	11.77
	<i>Bromus rubens</i>	1707	4.73
	<i>Bromus hordeaceus</i>	1371	6.28
	<i>Bromus diandrus</i>	1206	4.9
	<i>Bromus madritensis</i>	804	2.63
Long-lived C3 grasses	<i>Poa pratensis</i>	2213	8.71
	<i>Agropyron cristatum</i>	2113	14.37
	<i>Thinopyrum intermedium</i>	1447	7.59
	<i>Poa bulbosa</i>	897	9.38
	<i>Bromus inermis</i>	717	9.27
	<i>Phleum pratense</i>	512	4.37
C4 grasses	<i>Cynodon dactylon</i>	87	13.97
	<i>Eragrostis lehmanniana</i>	87	8.09
	<i>Pennisetum setaceum</i>	86	2.97
	<i>Eragrostis cilianensis</i>	68	1.97
	<i>Echinochloa crus-galli</i>	44	1.76
	<i>Setaria viridis</i>	33	4.13
Short-lived forbs	<i>Sisymbrium altissimum</i>	2062	4.73
	<i>Alyssum desertorum</i>	1951	6.53
	<i>Tragopogon dubius</i>	1820	0.83
	<i>Erodium cicutarium</i>	1753	4.48
	<i>Lactuca serriola</i>	1427	1.37
	<i>Melilotus officinalis</i>	1262	6.27
Long-lived forbs	<i>Taraxacum officinale</i>	2353	1.85
	<i>Cirsium arvense</i>	573	3.24
	<i>Rumex crispus</i>	359	0.9
	<i>Hypochaeris radicata</i>	301	1.86
	<i>Rumex acetosella</i>	248	2.13
	<i>Marrubium vulgare</i>	218	1.08
Shrubs and trees	<i>Tamarix spp.</i>	292	10.38
	<i>Nicotiana glauca</i>	189	0.46
	<i>Rubus armeniacus</i>	92	13.62
	<i>Elaeagnus angustifolia</i>	92	16.74
	<i>Genista monspessulana</i>	91	6.36
	<i>Rubus bifrons</i>	66	5.34

A full list of all non-native species found in the dataset is in Table S2

with abundant resources, and thus post-fire native competition and invasion resistance is higher (Chambers et al. 2019; Bekris et al. 2021). Higher cover of native species, and specifically native perennial species, has been shown to provide resistance to winter-annual and short-lived non-native plant invasions (Chambers et al. 2007; McGlone et al. 2011; Reisner et al. 2013; Williamson et al. 2020; Anthony and Germino 2023). As this synthesis shows, and is backed

by other research, the importance of climate and competition is especially apparent after disturbances (Taylor et al. 2014; Brummer et al. 2016).

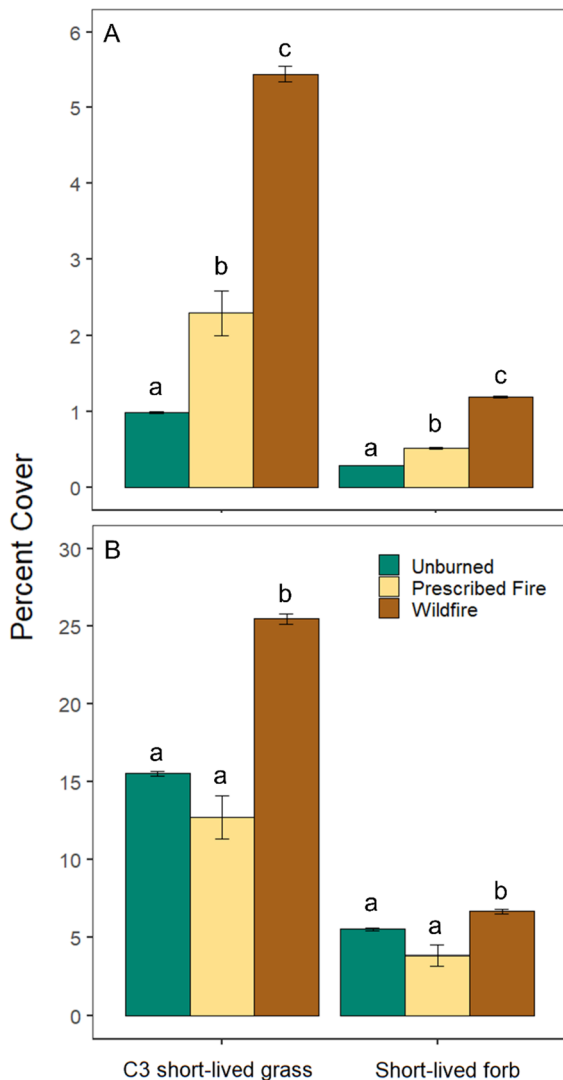
The non-native functional types that increased after fire, short-lived forbs and C3 grasses, were both strongly correlated with climate, and the regions where invasions following fire were predicted to be most likely were similar for both functional types. Furthermore, even though our results for short-lived

**Table 4** Estimates of fixed effects of fire type (unburned, wildfire, prescribed fire), the number of times a plot was burned in the last 50 years, the number of years since a fire occurred, the monitoring trends in burn severity (MTBS) fire severity classification, fire regime on percent cover of non-native functional types that increased following fires: short-lived grasses and short-lived forbs

Non-native functional type	Response values included	Fixed effect	Estimate	se	z	p	# obs
Short-lived grasses	All	Fire type (wildfire)	1.67	0.03	46.26	<0.00001	25,661
Short-lived grasses	All	Fire type (Rx fire)	0.97	0.16	6.22	<0.00001	1067
Short-lived grasses	Non-zero	Fire type (wildfire)	0.59	0.02	25.2	<0.00001	5373
Short-lived grasses	Non-zero	Fire type (Rx fire)	-0.001	0.11	-0.02	0.99	193
Short-lived grasses	All	# times burned	0.16	0.05	3.01	0.002	18,306
Short-lived grasses	Non-zero	# times burned	0.17	0.04	4.18	0.05	4610
Short-lived grasses	All	Time since fire	0.004	0.001	2.97	0.003	18,306
Short-lived grasses	Non-zero	Time since fire	0.002	0.001	1.67	0.09	5596
Short-lived grasses	All	Fire severity	-0.05	0.03	-1.74	0.08	6244
Short-lived grasses	Non-zero	Fire severity	0.04	0.02	1.36	0.17	3667
Short-lived grasses	All—forest	Fire severity	-0.50	0.42	-1.19	0.23	799
Short-lived grasses	All—grass/shrubland	Fire severity	-0.06	0.03	-1.79	0.07	5345
Short-lived grasses	All	FRI (36–100)	0.03	0.05	2.1	0.04	9881
Short-lived grasses	All	FRI (200+)	0.16	0.04	3.3	0.0009	8710
Short-lived grasses	All	Fire regime (I-C)	-1.83	0.09	*4.36	<0.00001	3206
Short-lived grasses	All	Fire regime (III-A)	-1.39	0.29	-3.55	0.0004	2303
Short-lived grasses	All	Fire regime (IV-A)	2.79	0.32	8.47	<0.00001	7578
Short-lived grasses	All	Fire regime (IV-B)	0.57	0.08	1.7	0.08	6509
Short-lived forbs	All	Fire type (wildfire)	1.56	0.04	38.53	<0.00001	25,661
Short-lived forbs	All	Fire type (Rx fire)	0.62	0.18	3.40	0.0007	1067
Short-lived forbs	Non-zero	Fire type (wildfire)	0.22	0.03	8.3	<0.00001	4463
Short-lived forbs	Non-zero	Fire type (Rx fire)	0.005	0.12	0.41	0.96	143
Short-lived forbs	All	# times burned	0.08	0.04	1.99	0.05	18,306
Short-lived forbs	Non-zero	# times burned	0.04	0.03	1.37	0.001	2681
Short-lived forbs	All	Time since fire	-0.005	0.002	-2.378	0.017, 0.35	18,306
Short-lived forbs	Non-zero	Time since fire	-0.0002	0.002	-0.113	0.91	4639
Short-lived forbs	All	Fire severity	0.06	0.05	1.20	0.23	6244
Short-lived forbs	Non-zero	Fire severity	0.12	0.04	3.22	0.06	2788
Short-lived forbs	All—forest	Fire severity	-0.23	0.36	-0.65	0.52	799
Short-lived forbs	All—grass/shrubland	Fire severity	0.01	0.05	0.27	0.78	5345
Short-lived forbs	All	FRI (36–100)	0.12	0.06	1.97	0.05	9881
Short-lived forbs	All	FRI (200+)	0.31	0.06	5.2	<0.00001	8710
Short-lived forbs	All	Fire regime (I-C)	-0.18	0.03	-1.51	0.13	3206
Short-lived forbs	All	Fire regime (III-A)	-0.14	0.01	-1.06	0.29	2303
Short-lived forbs	All	Fire regime (IV-A)	0.53	0.01	4.88	<0.00001	7578
Short-lived forbs	All	Fire regime (IV-B)	0.54	0.01	0.492	0.62	6509

C3 grasses were likely driven by relationships important for the dominant invasive *B. tectorum*, the spatial predictions from models trained on other C3 grasses excluding *B. tectorum* were similar to those from models trained on only *B. tectorum* cover data. These results highlight that semi-arid regions

receiving the majority of precipitation in the winter and early spring are susceptible to invasion after disturbances by short-lived non-native species that can germinate and grow quickly outside the historical growing season. Species traits are often the focus of studies of invasions (e.g. Drenovsky et al. 2012; Funk



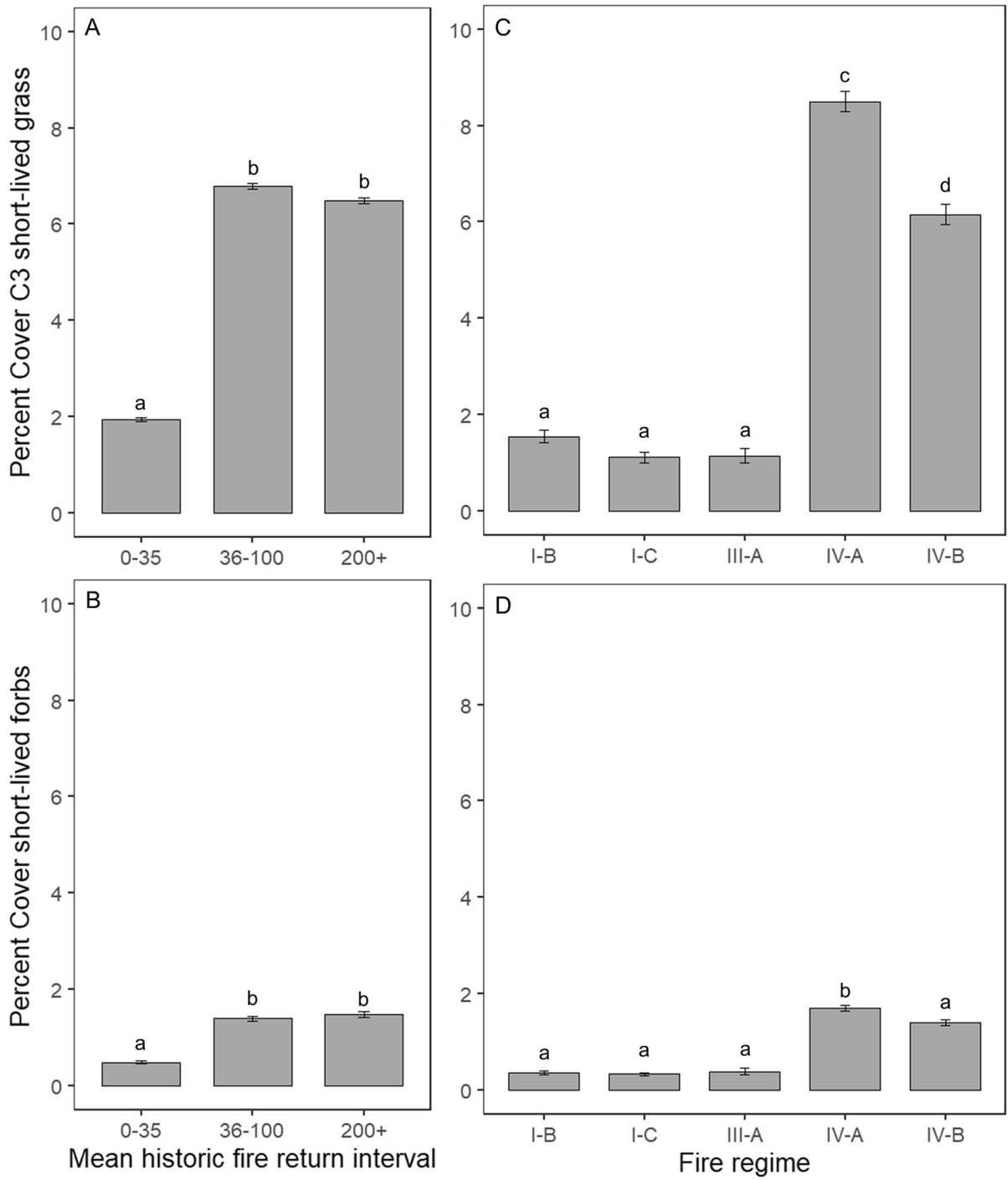
**Fig. 3** **A** Mean percent cover of fire-increasing non-native lifeforms in unburned plots ( $n=211,071$ ), plots that burned in a prescribed fire ( $n=1067$ ), and plots that burned in a wildfire ( $n=25,152$ ) up to 50 years prior to measurement,  $\pm$  standard error, including plots with 0% cover for the variable. **B** Means of all cover response variables only including plots where the target lifeform was present. Note the change in values on the y-axis. Different letters indicate significant differences in mean cover by fire type at  $p < 0.05$

et al. 2017), but abiotic factors also play an important role in determining where species become invasive (Davis et al. 2000). In regions with temporally limited resources, there may be a greater fluctuation in these resources after disturbances compared to regions with more moderate climates, and thus there

is a greater opportunity for invasion, perhaps regardless of the specific invader characteristics themselves (Davis et al. 2000). Highlighting the environmental conditions that influence ecosystem invasibility after disturbances can help focus management efforts to regions at the greatest risk following disturbance and with climate change.

Climate-driven changes in the timing of precipitation and drought may alter the locations where post-fire plant invasions for the species included in this dataset are most likely and most damaging (Bradley 2009; Taylor et al. 2014; Bradley et al. 2018). There is some evidence that these changes may already be occurring, as long-term monitoring plots from rangelands in western USA show an increase in the cover of annual plants across all regions over time, including in forested mountains and in the Northern Great Plains (Kleinhesselink et al. 2023). Future changes in invasion risk may impact decisions about prescribed fires, thinning operations, herbicide use, restoration planting, and where to increase post-disturbance monitoring for non-natives in higher elevation ecosystems (Brooks et al. 2004; Merriam et al. 2006; Keeley et al. 2011). Our results also highlight regions that may see a decrease in invasion risk by short-lived non-native C3 grasses and forbs in the future based on climate projections. Areas that are projected to see decreased risk of invasion are those that are projected to become very arid in the future, and these regions may become more resistant to invasion as there will be fewer resource pulses to enable non-native colonization following disturbances, potentially favoring the more conservative strategies of some native species (Zefferman et al. 2015). These results need to be interpreted with caution, but they are supported by other similar findings for individual non-native species (Bradley and Wilcove 2009; Allen and Bradley 2016). Targeting areas with a reduced projected invasion risk for restoration treatments could result in greater restoration success as non-natives become less climatically suited and thus less competitive with restored native plantings (Pilliod et al. 2021).

There are many critical caveats that should be considered when interpreting the results of our data synthesis. Perhaps most importantly, this dataset consists of temporally and spatially uneven samples consisting of a single post-fire measurement per plot; and thus results shown here infer impacts via a space-for-time substitution (Sofaer et al. 2018). For example, we



**Fig. 4** Average percent cover of functional types that increased following fire in burned plots. **A** non-native C3 short-lived grasses and **B** non-native short-lived forbs in binned categories based on LANDFIRE historic fire return intervals, 0–35 years ( $n=7702$ ), 36–100 years ( $n=9881$ ), and 200 years or greater ( $n=8710$ ). Average percent cover of **C** non-native C3 short-lived grasses and **D** non-native short-lived forbs by LANDFIRE fire regime classification: I-B is categorized as having percent replacement fire less than 66.7%, and a fire return interval of 6–15 years ( $n=3114$ ), I-C has 66.7% replacement fire, fire return interval of 16–35 years ( $n=3206$ ), III-A has less than 80% replacement fire, fire return interval of 36–100 years ( $n=2303$ ), IV-A has greater than 80% replacement fire, fire return interval of 36–100 years ( $n=7578$ ), IV-B has greater than 66.7% replacement fire, fire return interval of 101–200 years ( $n=6509$ ). Different letters denote significant differences at  $p < 0.05$

found that time since fire did not impact non-native plant cover. However, these data are from only a single point in time at each site, so we cannot accurately examine site-level cover changes over time. We also found no relationships between non-native cover and fire severity. Fire severity measurements for grass and shrubland ecosystems have been shown to be highly inaccurate (Zhu et al. 2006; Storey et al. 2021), and site-level analyses have shown that higher fire severity increases prevalence of some invaders in forests (Tortorelli et al. 2020). Additionally, the importance of climate variables as drivers of invasion risk likely results from the large scale of this analysis as well. Topography, soils, past management, and grazing history would likely become more important predictors of where non-native plants are abundant at finer scales (Pearson and Dawson 2003; Ellsworth et al. 2016). Additional data from long-term monitoring plots that are repeatedly measured over time, evenly dispersed across the landscape, and paired with accurate on-the-ground measures of fire severity across a range of habitat types will help refine predictions of post-fire invasion risk.

We did not examine native vegetation here, so we cannot directly infer impacts of competition, biotic resistance, or other plant interactions. This would have only been correlative with this data collection, as these data were taken at a single point in time, and time-series before and after fires would be needed for such analyses. We attempted to limit the influence of pre-existing vegetation in our analyses by only including “recently” burned (i.e., disturbed) plots. However, this type of analysis means we are unable to accurately identify how pre-fire plant community composition or vegetation type influences post-fire invasion patterns. There were also large data gaps that likely hinder the accurate assessment of post-fire invasion risk across the western USA. For example, we have very few data points from burned plots in southwestern hot desert ecosystems, where non-native long-lived grass species may be becoming more prevalent after fires (Brooks and Chambers 2011; Wilder et al. 2021). Additional monitoring data from burned areas in the southwestern region, as well as other under-sampled regions, will help provide more balanced and accurate predictions of post-fire invasion risk across the west. This synthesis is helpful in that it highlights regions where more data are needed.

Another important caveat with these analyses is that we sum the cover of all non-native species within a functional type together prior to evaluating how cover is influenced by environmental variables. This is partially due to the scarcity of data for any single species besides *Bromus tectorum* across the entirety of the western USA, but also stems from the goal to look at broad-scale patterns in invasion rather than replicate species-level distribution models and maps (e.g. Bradley 2009; Bradley et al. 2018; Sofaer et al. 2022). Individual species will respond in different ways to combinations of disturbance and climate (Tortorelli et al. 2020; Applestein and Germino

**Table 5** Random forest model results showing  $R^2$  and root-mean-squared-error (RSME) values for predictions from out-of-bag data not in the bootstrapped sample (oob), as well for the bootstrapped data (sample)

Model	$R^2$ oob	$R^2$ sample	RSME oob	RSME sample
Short-lived C3 grasses	0.44	0.89	11.63	5.80
Short-lived forbs	0.19	0.83	4.43	2.39

**Table 6** Importance values for the final set of predictor variables from random forest models of post-fire abundance of non-native C3 short-lived grasses and short-lived forbs

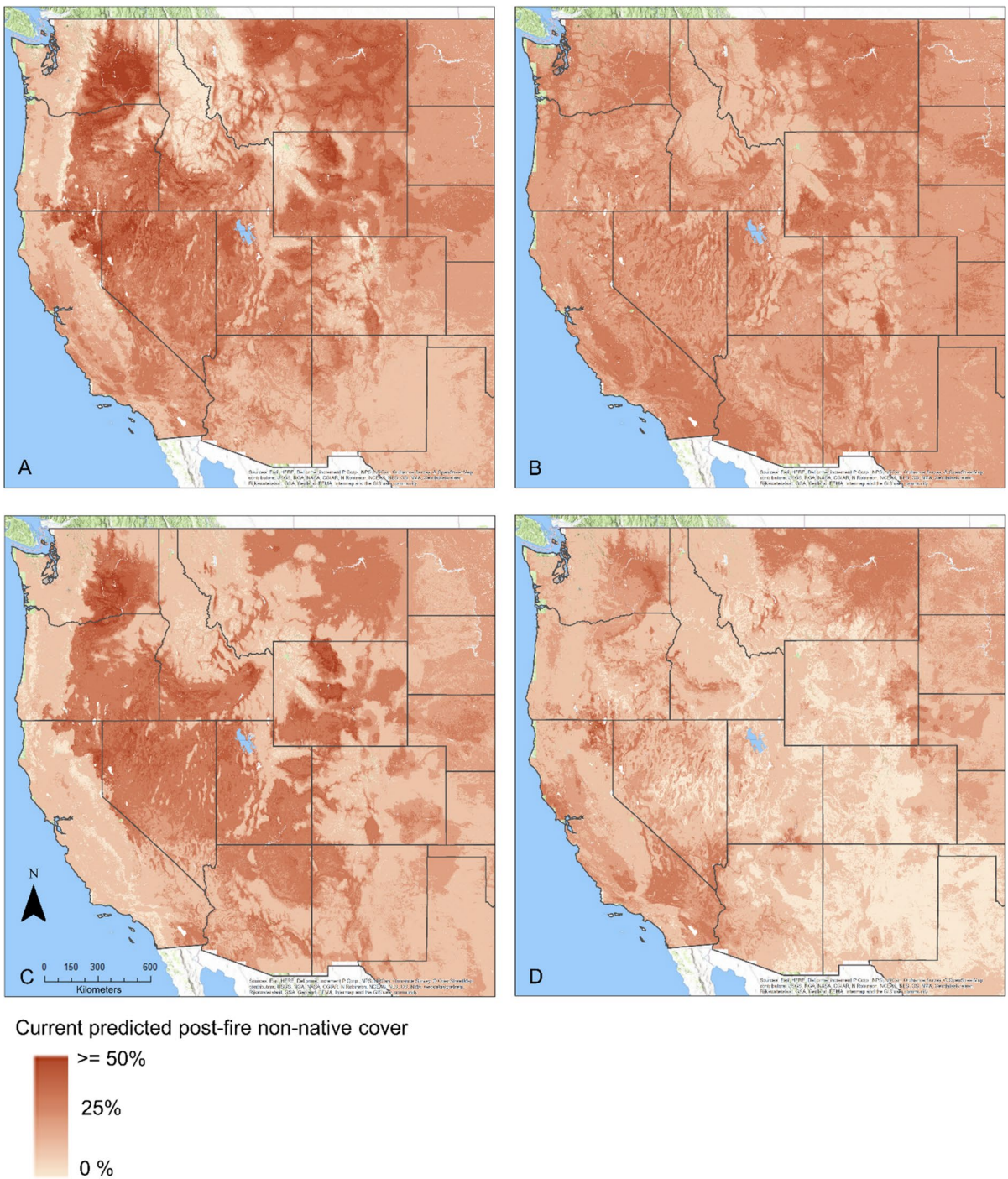
Lifeform	Variable	Permutation importance
C3 short-lived grasses	April-Sept. precipitation	17.80
	Winter precipitation ratio	15.50
	Precipitation as snow	15.16
	Annual heat-moisture index	13.78
	Mean annual radiation	8.65
	Historic fire return interval	8.24
	Remoteness	8.24
	% Sand	6.79
	% Clay	6.70
	CHILI	6.02
	Northness	5.84
	Short-lived forbs	April-Sept. precipitation
Winter precipitation ratio		6.89
Annual heat-moisture index		6.12
Precipitation as snow		5.97
MAR		3.36
Historic fire return interval		3.17
% Clay		3.09
% Sand		2.82
Remoteness		2.57
CHILI		1.73
Northness		1.66

Importance values show how much including the variable in the model increases model accuracy. Variables with higher importance values have greater predictive power. Descriptions of variable abbreviations can be found in Table 1

2022), and thus grouping multiple species together for analyses could obfuscate species-level patterns. In sagebrush steppe, *Ventenata dubia* has been shown to occupy a distinct niche from *Bromus tectorum* and *Taeniatherum caput-medusae* (Applestein and Germino 2022). Additionally, fire may not play as large of a role in enhancing invasion risk of specific species of short-lived non-natives compared to others (Ridder et al. 2021). For example, *Ventenata dubia* is currently a more successful invader at higher elevations than *Bromus tectorum*, and our combined climate predictions for all short-lived grasses may mask more rapid advances to higher elevations of *Ventenata dubia* with a changing climate (Tortorelli et al. 2020).

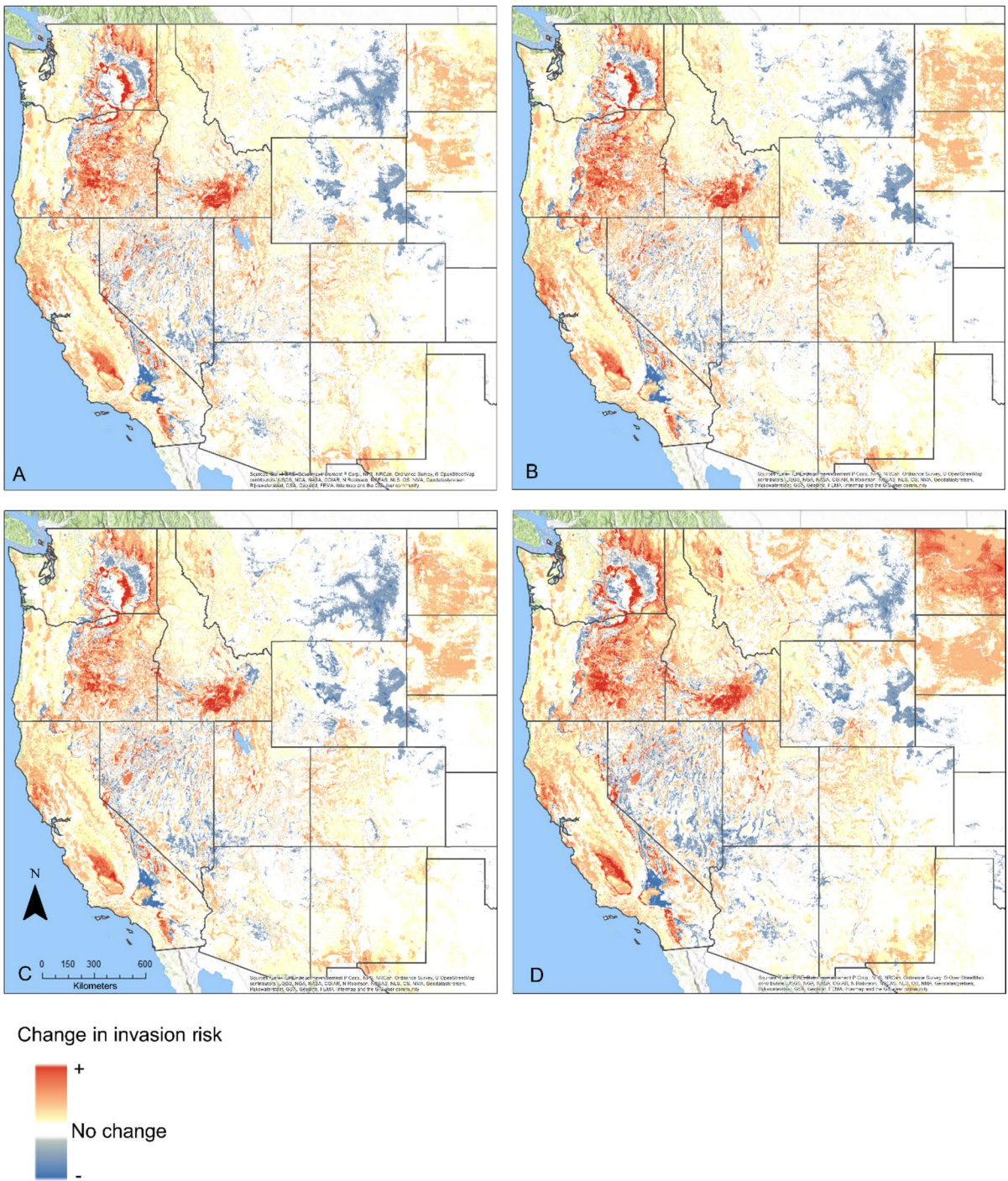
This synthesis is the first to pull extensively from multiple datasets to examine patterns in post-fire plant invasion risk at a subcontinental scale. We show that non-native invasion after fire in the western USA is overwhelmingly caused by short-lived plant species, and they are most abundant after fires in regions with dry summers and historically long fire return intervals. Our analyses clearly indicate a 'climate space' where non-native plant invasion is more likely after fire, specifically in regions with higher summer drought. This 'climate space' may change as the climate changes and summer droughts become more severe, so risk of post-fire plant invasions may change as well. The spatial projections of the results presented here can help identify hotspots where invasions are the most common following fire now and in the future. These spatial predictions provide an important step to help reduce the risk of





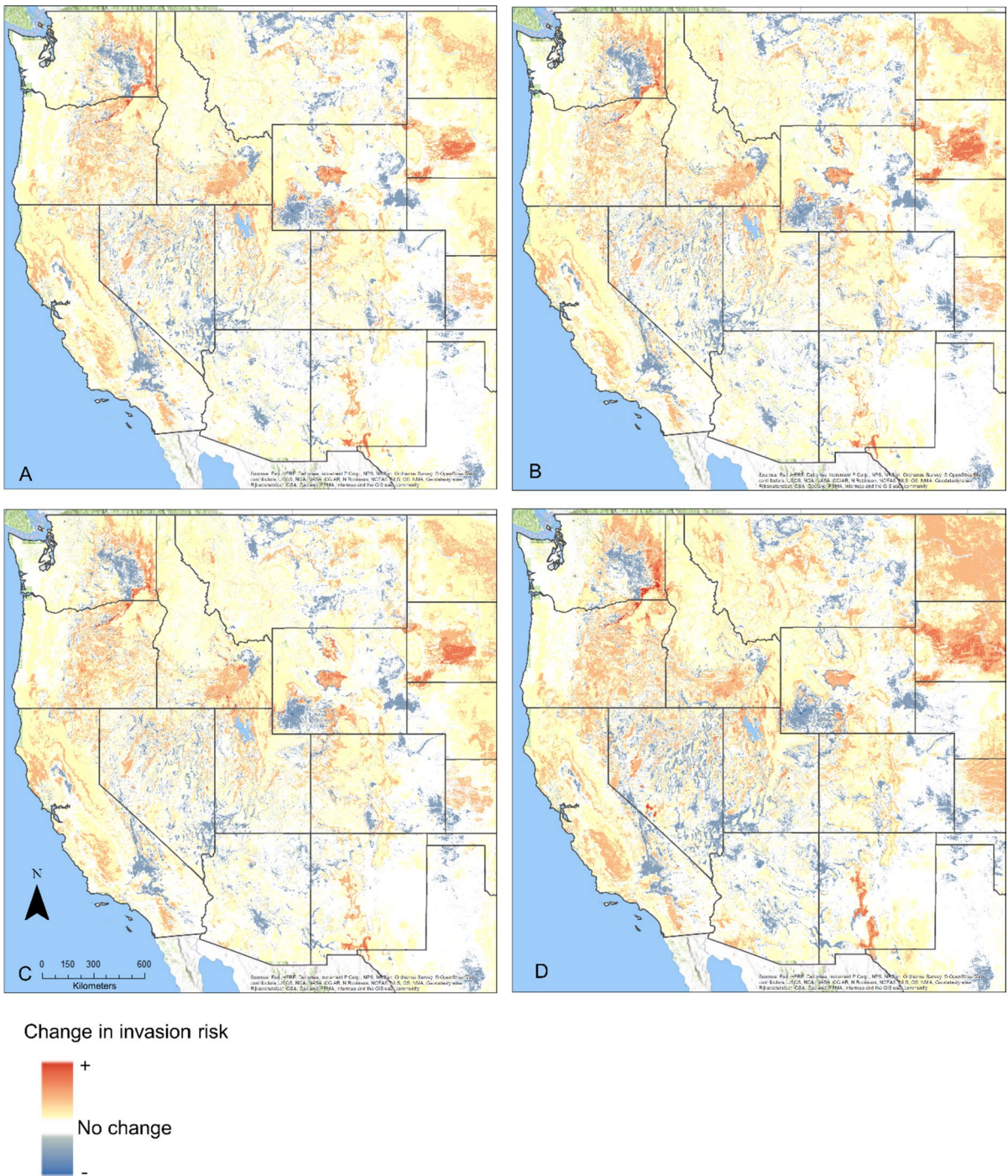
**Fig. 5** Projections of post-fire cover of non-native **A** C3 short-lived grasses, **B** short-lived forbs, **C** only *Bromus tectorum*, and **D** all other C3 short-lived grasses except *Bromus tectorum*

based on random forests using bioclimatic data for the current (1981–2010) time period. Base map from Esri and its licensors, copyright 2022



**Fig. 6** Projected changes in post-fire abundance of non-native C3 short-lived grasses for **A** 2041–2070 under a medium emissions climate scenario SSP245, **B** 2041–2070 under the high-

est emissions climate scenario SSP285, **C** 2071–2100 under SSP545, and **D** 2071–2100 under SSP585. Base map from Esri and its licensors, copyright 2022



**Fig. 7** Projected changes in post-fire abundance of non-native short-lived forbs for **A** 2041–2070 under a medium emissions climate scenario SSP245, **B** 2041–2070 under the high-

est emissions climate scenario SSP285, **C** 2071–2100 under SSP545, and **D** 2071–2100 under SSP585. Base map from Esri and its licensors, copyright 2022

invasion after wildfires by allowing for more targeted mitigation, monitoring, and restoration.

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**Author contributions** JSP, ISP, CSJ, conceived of the concept for this synthesis with input from SMM and JTS. KB, JDC, MAD, DF, PF, KH, JJ, BKK, MAK, BM, TN, JR, JDS, CSSR, MTS, and CT collected and contributed data to this synthesis. JSP led the writing of the manuscript and analysis of the data. All authors contributed to writing and edits of drafts of the manuscript and gave final approval for publication.

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**Data availability** Any unpublished plot-level plant cover data used in this manuscript will be made available upon request from the authors.

## Declarations

**Conflict of interest** The authors have no relevant financial or non-financial interests to disclose. Any use of trade, firm, or product names is for descriptive purposes only and does not imply endorsement by the U.S. Government.

## References

- AdaptWest Project (2022) Gridded current and projected climate data for North America at 1km resolution, generated using the ClimateNA v7.30 software. (T. Wang et al., 2022). Available at <https://adaptwest.databasin.org/>
- Alba C, Skálová H, McGregor KF et al (2015) Native and exotic plant species respond differently to wildfire and prescribed fire as revealed by meta-analysis. *J Veg Sci* 26:102–113. <https://doi.org/10.1111/jvs.12212>
- Allen JM, Bradley BA (2016) Out of the weeds? Reduced plant invasion risk with climate change in the continental United States. *Biol Conserv* 203:306–312. <https://doi.org/10.1016/j.biocon.2016.09.015>
- Anthony CR, Germino MJ (2023) Does post-fire recovery of native grasses across abiotic-stress and invasive-grass gradients match theoretical predictions, in sagebrush steppe? *Glob Ecol Conserv* 42:e02410. <https://doi.org/10.1016/j.gecco.2023.e02410>
- Applestein C, Germino MJ (2022) Patterns of post-fire invasion of semiarid shrub-steppe reveals a diversity of invasion niches within an exotic annual grass community. *Biol Invasions* 24:741–759. <https://doi.org/10.1007/s10530-021-02669-3>
- Beaury EM, Fusco EJ, Jackson MR et al (2020) Incorporating climate change into invasive species management: insights from managers. *Biol Invasions* 22:233–252. <https://doi.org/10.1007/s10530-019-02087-6>
- Bekris Y, Prevéy JS, Brodie LC, Harrington CA (2021) Effects of variable-density thinning on non-native understory plants in coniferous forests of the Pacific Northwest. *For Ecol Manag* 502:119699. <https://doi.org/10.1016/j.foreco.2021.119699>
- Belmonte MCB, de Agradar ZER (2002) Analysis of the patterns of distribution of photosynthetic pathways and representativity of the family Poaceae on Martín García Island, Río de La Plata, Buenos Aires, Argentina. *Flora Morphol Distrib Funct Ecol Plants* 197:351–360. <https://doi.org/10.1078/0367-2530-00051>
- Bradley BA (2009) Regional analysis of the impacts of climate change on cheatgrass invasion shows potential risk and opportunity. *Glob Change Biol* 15:196–208. <https://doi.org/10.1111/j.1365-2486.2008.01709.x>
- Bradley BA, Wilcove DS (2009) When invasive plants disappear: transformative restoration possibilities in the Western United States resulting from climate change. *Restor Ecol* 17:715–721. <https://doi.org/10.1111/j.1526-100X.2009.00586.x>
- Bradley BA, Curtis CA, Fusco EJ et al (2018) Cheatgrass (*Bromus tectorum*) distribution in the intermountain Western United States and its relationship to fire frequency, seasonality, and ignitions. *Biol Invasions* 20:1493–1506. <https://doi.org/10.1007/s10530-017-1641-8>
- Brooks ML (1999) Alien annual grasses and fire in the Mojave desert. *Madroño* 46:13–19
- Brooks ML, Chambers JC (2011) Resistance to invasion and resilience to fire in desert shrublands of North America. *Rangel Ecol Manag* 64:431–438. <https://doi.org/10.2111/REM-D-09-00165.1>
- Brooks ML, D'Antonio CM, Richardson DM et al (2004) Effects of invasive alien plants on fire regimes. *Bio-science* 54:677–688. [https://doi.org/10.1641/0006-3568\(2004\)054\[0677:EOIAPJ\]2.0.CO;2](https://doi.org/10.1641/0006-3568(2004)054[0677:EOIAPJ]2.0.CO;2)
- Brooks ME, Kristensen K, van Benthem KJ et al (2017) glmTMB balances speed and flexibility among packages for zero-inflated generalized linear mixed modeling. *The R journal* 9:378–400. <https://doi.org/10.3929/ethz-b-000240890>
- Bruhl J, Wilson K (2007) Towards a comprehensive survey of C3 and C4 photosynthetic pathways in Cyperaceae. *Aliso* 23:99–148. <https://doi.org/10.5642/aliso.20072301.11>
- Brummer TJ, Taylor KT, Rotella J et al (2016) Drivers of *Bromus tectorum* abundance in the Western North American sagebrush steppe. *Ecosystems* 19:986–1000. <https://doi.org/10.1007/s10021-016-9980-3>
- Bukowski BE, Baker WL (2013) Historical fire regimes, reconstructed from land-survey data, led to complexity and fluctuation in sagebrush landscapes. *Ecol Appl* 23:546–564. <https://doi.org/10.1890/12-0844.1>

- Byers JE, Reichard S, Randall JM et al (2002) Directing research to reduce the impacts of nonindigenous species. *Conserv Biol* 16:630–640. <https://doi.org/10.1046/j.1523-1739.2002.01057.x>
- Calo A, Shayna B, Scott J (2012) Integrated treatment with a prescribed burn and postemergent herbicide demonstrates initial success in managing cheatgrass in a northern Colorado natural area. *Nat Areas J* 32:300–304
- Chambers JC, Roundy BA, Blank RR et al (2007) What makes great basin sagebrush ecosystems invulnerable by *Bromus tectorum*? *Ecol Monogr* 77:117–145
- Chambers JC, Brooks ML, Germino MJ et al (2019) Operationalizing resilience and resistance concepts to address invasive grass-fire cycles. *Front Ecol Evol* 7:185. <https://doi.org/10.3389/fevo.2019.00185>
- Coates PS, Ricca MA, Prochazka BG et al (2016) Wildfire, climate, and invasive grass interactions negatively impact an indicator species by reshaping sagebrush ecosystems. *Proc Natl Acad Sci* 113:12745–12750. <https://doi.org/10.1073/pnas.1606898113>
- Concilio AL, Loik ME, Belnap J (2013) Global change effects on *Bromus tectorum* L. (Poaceae) at its high-elevation range margin. *Glob Change Biol* 19:161–172. <https://doi.org/10.1111/gcb.12032>
- Coop JD, Parks SA, Stevens-Rumann CS et al (2020) Wildfire-driven forest conversion in western North American landscapes. *Bioscience* 70:659–673. <https://doi.org/10.1093/biosci/biaa061>
- Cutler DR, Edwards TC Jr, Beard KH et al (2007) Random forests for classification in ecology. *Ecology* 88:2783–2792. <https://doi.org/10.1890/07-0539.1>
- Daehler CC (2003) Performance comparisons of co-occurring native and alien invasive plants: implications for conservation and restoration. *Annu Rev Ecol Evol Syst* 34:183–211. <https://doi.org/10.1146/annurev.ecolsys.34.011802.132403>
- D'Antonio CM, Vitousek PM (1992) Biological invasions by exotic grasses, the grass/fire cycle, and global change. *Annu Rev Ecol Syst* 23:63–87
- Davies GM, Bakker JD, Dettweiler-Robinson E et al (2012) Trajectories of change in sagebrush steppe vegetation communities in relation to multiple wildfires. *Ecol Appl* 22:1562–1577. <https://doi.org/10.1890/10-2089.1>
- Davis MA, Pelsor M (2001) Experimental support for a resource-based mechanistic model of invasibility. *Ecol Lett* 4:421–428. <https://doi.org/10.1046/j.1461-0248.2001.00246.x>
- Davis MA, Grime JP, Thompson K (2000) Fluctuating resources in plant communities: a general theory of invasibility. *J Ecol* 88:528–534. <https://doi.org/10.1046/j.1365-2745.2000.00473.x>
- Ditomaso JM, Brooks ML, Allen EB et al (2006) Control of invasive weeds with prescribed burning. *Weed Technol* 20:535–548
- Dormann CF, Elith J, Bacher S et al (2013) Collinearity: a review of methods to deal with it and a simulation study evaluating their performance. *Ecography* 36:27–46. <https://doi.org/10.1111/j.1600-0587.2012.07348.x>
- Drenovsky RE, Grewell BJ, D'Antonio CM et al (2012) A functional trait perspective on plant invasion. *Ann Bot* 110:141–153. <https://doi.org/10.1093/aob/mcs100>
- Dunn PK, Smyth GK (eds) (2018) Chapter 12: Tweedie GLMs. In: Generalized linear models with examples in R. Springer, New York, NY, pp 457–490
- Eidenshink J, Schwind B, Brewer K et al (2007) A project for monitoring trends in burn severity. *Fire Ecol* 3:3–21. <https://doi.org/10.4996/fireecology.0301003>
- Ellsworth LM, Wroblewski DW, Kauffman JB, Reis SA (2016) Ecosystem resilience is evident 17 years after fire in Wyoming big sagebrush ecosystems. *Ecosphere* 7:e01618. <https://doi.org/10.1002/ecs2.1618>
- Evans JS, Murphy MA, Holden ZA, Cushman SA (2011) Modeling species distribution and change using random forest. In: Drew CA, Wiersma YF, Huettmann F (eds) Predictive species and habitat modeling in landscape ecology: concepts and applications. Springer, New York, pp 139–159
- Eyring V, Bony S, Meehl GA et al (2016) Overview of the coupled model intercomparison project phase 6 (CMIP6) experimental design and organization. *Geosci Model Dev* 9:1937–1958. <https://doi.org/10.5194/gmd-9-1937-2016>
- Firmage D, and Ronsani KG (2021a) Bryce Canyon National Park FFI fire effects monitoring data
- Firmage D, Ronsani KG (2021b) Zion National Park FFI fire effects monitoring data
- Fornwalt PJ, Kaufmann MR, Stohlgren TJ (2010) Impacts of mixed severity wildfire on exotic plants in a Colorado ponderosa pine-Douglas-fir forest. *Biol Invasions* 12(8):2683–2695
- Franklin J, Spears-Lebrun LA, Deutschman DH, Marsden K (2006) Impact of a high-intensity fire on mixed evergreen and mixed conifer forests in the Peninsular Ranges of southern California, USA. *For Ecol Manag* 235:18–29. <https://doi.org/10.1016/j.foreco.2006.07.023>
- Funk JL, Larson JE, Ames GM et al (2017) Revisiting the Holy Grail: using plant functional traits to understand ecological processes. *Biol Rev* 92:1156–1173. <https://doi.org/10.1111/brv.12275>
- Fusco EJ, Finn JT, Balch JK et al (2019) Invasive grasses increase fire occurrence and frequency across US ecoregions. *PNAS* 116:23594–23599. <https://doi.org/10.1073/pnas.1908253116>
- Gelbard JL, Belnap J (2003) Roads as conduits for exotic plant invasions in a semiarid landscape. *Conserv Biol* 17:420–432. <https://doi.org/10.1046/j.1523-1739.2003.01408.x>
- Hartig F (2022) DHARMA: residual diagnostics for hierarchical (multi-level/mixed) regression models\_R package version 0.4.6. <https://CRAN.R-project.org/package=DHARMA>
- Hobbs RJ, Huenneke LF (1992) Disturbance, diversity, and invasion: implications for conservation. *Conserv Biol* 6:324–337
- Hothorn T, Bretz F, Westfall P (2008) Simultaneous inference in general parametric models. *Biom J* 50:346–363
- Keeley JE, Lubin D, Fotheringham CJ (2003) Fire and grazing impacts on plant diversity and alien plant invasions in the southern Sierra Nevada. *Ecol Appl* 13:1355–1374. <https://doi.org/10.1890/02-5002>
- Keeley JE, McGinnis TW, Keeley JE, McGinnis TW (2007) Impact of prescribed fire and other factors on cheatgrass persistence in a Sierra Nevada ponderosa pine forest\*.

- Int J Wildland Fire 16:96–106. <https://doi.org/10.1071/WF06052>
- Keeley JE, Franklin J, D'Antonio C (2011) Fire and invasive plants on California landscapes. In: McKenzie D, Miller C, Falk DA (eds) The landscape ecology of fire. Springer, Dordrecht, pp 193–221
- Kerns BK, Tortorelli C, Day MA et al (2020) Invasive grasses: A new perfect storm for forested ecosystems? For Ecol Manag 463:117985. <https://doi.org/10.1016/j.foreco.2020.117985>
- Kleinhesselink AR, Kachergis EJ, McCord SE et al (2023) Long-term trends in vegetation on bureau of land management rangelands in the western United States. Rangel Ecol Manag 87:1–12. <https://doi.org/10.1016/j.rama.2022.11.004>
- Kulmatiski A (2006) Exotic plants establish persistent communities. Plant Ecol 187:261–275. <https://doi.org/10.1007/s11258-006-9140-5>
- LANDFIRE (2016a) LANDFIRE (LF) 2016 Remap (LF 2.0.0) reference database (LFRDB). <https://landfire.gov/lfrdb.php>
- LANDFIRE (2016b) LANDFIRE biophysical settings layer. <https://landfire.gov/bps.php>
- Lembrechts JJ, Pauchard A, Lenoir J et al (2016) Disturbance is the key to plant invasions in cold environments. Proc Natl Acad Sci 113:14061–14066. <https://doi.org/10.1073/pnas.1608980113>
- Liaw A, Wiener M (2002) Classification and Regression by random. Forest 2:5
- Mack RN, Simberloff D, Mark Lonsdale W et al (2000) Biotic invasions: causes, epidemiology, global consequences, and control. Ecol Appl 10:689–710. [https://doi.org/10.1890/1051-0761\(2000\)010\[0689:BICEGC\]2.0.CO;2](https://doi.org/10.1890/1051-0761(2000)010[0689:BICEGC]2.0.CO;2)
- Mahony CR, Wang T, Hamann A, Cannon AJ (2022) A global climate model ensemble for downscaled monthly climate normals over North America. Int J Climatol 42:5871–5891. <https://doi.org/10.1002/joc.7566>
- McGlone CM, Sieg CH, Kolb TE (2011) Invasion resistance and persistence: established plants win, even with disturbance and high propagule pressure. Biol Invasions 13(2):291–304. <https://doi.org/10.1007/s10530-010-9806-8>
- Melgoza G, Nowak RS, Tausch RJ (1990) Soil water exploitation after fire: competition between *Bromus tectorum* (cheatgrass) and two native species. Oecologia 83:7–13. <https://doi.org/10.1007/BF00324626>
- Merriam KE, Keeley JE, Beyers JL (2006) Fuel breaks affect nonnative species abundance in Californian plant communities. Ecol Appl 16:515–527. [https://doi.org/10.1890/1051-0761\(2006\)016\[0515:FBANSA\]2.0.CO;2](https://doi.org/10.1890/1051-0761(2006)016[0515:FBANSA]2.0.CO;2)
- Mi C, Huettmann F, Guo Y et al (2017) Why choose random forest to predict rare species distribution with few samples in large undersampled areas? Three Asian crane species models provide supporting evidence. PeerJ 5:e2849. <https://doi.org/10.7717/peerj.2849>
- Miller G, Friedel M, Adam P et al (2010) Ecological impacts of buffel grass (*Cenchrus ciliaris* L.) invasion in central Australia—Does field evidence support a fire-invasion feedback? Rangel J 32:353–365. <https://doi.org/10.1071/RJ09076>
- Montesinos D (2022) Fast invasives fastly become faster: invasive plants align largely with the fast side of the plant economics spectrum. J Ecol 110:1010–1014. <https://doi.org/10.1111/1365-2745.13616>
- Nagy RC, Fusco EJ, Balch JK et al (2021) A synthesis of the effects of cheatgrass invasion on US Great Basin carbon storage. J Appl Ecol 58:327–337. <https://doi.org/10.1111/1365-2664.13770>
- NALCMS (2015) <http://www.cec.org/north-american-environmental-atlas/land-cover-30m-2015-landsat-and-rapid-eye/>
- Omernik JM, Griffith GE (2014) Ecoregions of the conterminous United States: evolution of a hierarchical spatial framework. Environ Manag 54:1249–1266
- Pearson RG, Dawson TP (2003) Predicting the impacts of climate change on the distribution of species: Are bioclimate envelope models useful? Glob Ecol Biogeogr 12:361–371. <https://doi.org/10.1046/j.1466-822X.2003.00042.x>
- Perkins LB, Leger EA, Nowak RS (2011) Invasion triangle: an organizational framework for species invasion. Ecol Evol 1:610–625. <https://doi.org/10.1002/ece3.47>
- Petri L, Beaury EM, Corbin J, Peach K, Sofaer H, Pearse IS, Early R, Barnett DT, Ibáñez I, Peet RK, Schafale M (2022) SPCIS: standardized plant community with introduced status database. Ecology 104:e3947. <https://doi.org/10.1002/ecy.3947>
- Pilliod DS, Jeffries MA, Welty JL, Arkle RS (2021) Protecting restoration investments from the cheatgrass-fire cycle in sagebrush steppe. Conserv Sci Pract 3:e508. <https://doi.org/10.1111/csp2.508>
- Prevéy JS, Seastedt TR (2014) Seasonality of precipitation interacts with exotic species to alter composition and phenology of a semi-arid grassland. J Ecol 102:1549–1561. <https://doi.org/10.1111/1365-2745.12320>
- Ramcharan A et al (2018) Soil property and class maps of the conterminous United States at 100-meter spatial resolution. Soil Sci Soc Am J 82(1):186–201. <https://doi.org/10.2136/sssaj2017.04.0122>
- R Core Team (2022) R: a language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria
- Reisner MD, Grace JB, Pyke DA, Doescher PS (2013) Conditions favouring “*Bromus tectorum*” dominance of endangered sagebrush steppe ecosystems. J Appl Ecol 50:1039–1049
- Ridder LW, Perren JM, Morris LR et al (2021) Historical fire and *Ventenata dubia* invasion in a temperate grassland. Rangel Ecol Manag 75:35–40. <https://doi.org/10.1016/j.rama.2020.11.006>
- Roundy BA, Chambers JC, Pyke DA et al (2018) Resilience and resistance in sagebrush ecosystems are associated with seasonal soil temperature and water availability. Ecosphere 9:e02417. <https://doi.org/10.1002/ecs2.2417>
- Shea K, Chesson P (2002) Community ecology theory as a framework for biological invasions. Trends Ecol Evol 17:170–176. [https://doi.org/10.1016/S0169-5347\(02\)02495-3](https://doi.org/10.1016/S0169-5347(02)02495-3)
- Sherrill KR, Romme WH (2012) Spatial variation in post-fire cheatgrass: Dinosaur National Monument, USA.

- Fire Ecol 8:38–56. <https://doi.org/10.4996/fireecology.0802038>
- Smith JK, Zouhar K, Sutherland S, Brooks ML (2008) Fire and nonnative plants—summary and conclusions. USDA Forest Service Gen. Tech. Rep. RMRS-GTR-42-vol:6
- Smith JT, Allred BW, Boyd CS et al (2022) The elevational ascent and spread of exotic annual grass dominance in the Great Basin, USA. *Divers Distrib* 28:83–96. <https://doi.org/10.1111/ddi.13440>
- Sofaer HR, Jarnevich CS, Flather CH (2018) Misleading prioritizations from modelling range shifts under climate change. *Glob Ecol Biogeogr* 27:658–666. <https://doi.org/10.1111/geb.12726>
- Sofaer HR, Jarnevich CS, Buchholtz EK et al (2022) Potential cheatgrass abundance within lightly invaded areas of the Great Basin. *Landsc Ecol* 37:2607–2618. <https://doi.org/10.1007/s10980-022-01487-9>
- Stevens JT, Beckage B (2009) Fire feedbacks facilitate invasion of pine savannas by Brazilian pepper (*Schinus terebinthifolius*). *New Phytol* 184:365–375
- Storey EA, Lee West KR, Stow DA (2021) Utility and optimization of LANDSAT-derived burned area maps for southern California. *Int J Remote Sens* 42:486–505. <https://doi.org/10.1080/01431161.2020.1809741>
- Taylor K, Brummer T, Rew LJ et al (2014) *Bromus tectorum* response to fire varies with climate conditions. *Ecosystems* 17:960–973. <https://doi.org/10.1007/s10021-014-9771-7>
- Tortorelli CM, Krawchuk MA, Kerns BK (2020) Expanding the invasion footprint: *Ventennata dubia* and relationships to wildfire, environment, and plant communities in the Blue Mountains of the Inland Northwest, USA. *Appl Veg Sci* 23:562–574. <https://doi.org/10.1111/avsc.12511>
- Tortorelli CM, Kim JB, Vaillant NM et al (2023) Feeding the fire: annual grass invasion facilitates modeled fire spread across Inland Northwest forest-mosaic landscapes. *Ecosphere* 14:e4413. <https://doi.org/10.1002/ecs2.4413>
- US Census Bureau (2018) TIGER transportation data. <https://www.census.gov/geographies/mapping-files/time-series/geo/tiger-line-file.html>. Accessed June 2018
- USDA Forest Service (2022) Confronting the wildfire crisis, a 10 year implementation plan. FS-1187b
- USDA, NRCS (2022) The PLANTS database. National Plant Data Team, Greensboro, NC USA. <http://plants.usda.gov>. Accessed 12 July 2022
- USGS (2019) USGS 3D elevation program digital elevation model. <https://elevation.nationalmap.gov/arcgis/rest/services/3DEPElevation/ImageServer>. Accessed Jan 2020
- Wang T, Hamann A, Spittlehouse D, Carroll C (2016) Locally downscaled and spatially customizable climate data for historical and future periods for North America. *PLOS ONE* 11: e0156720. <https://doi.org/10.1371/journal.pone.0156720>
- Waller SS, Lewis JK (1979) Occurrence of C3 and C4 photosynthetic pathways in North American grasses. *J Range Manag* 32:12. <https://doi.org/10.2307/3897378>
- Welty JL, Jeffries MI (2021) Combined wildland fire datasets for the United States and certain territories, 1800s-Present: U.S. Geological Survey data release. <https://doi.org/10.5066/P9ZXGFY3>
- Wilder BT, Jarnevich CS, Baldwin E et al (2021) Grassification and fast-evolving fire connectivity and risk in the Sonoran desert, United States. *Front Ecol Evol* 9:655561
- Williams J (2013) Exploring the onset of high-impact megafires through a forest land management prism. *For Ecol Manag* 294:4–10. <https://doi.org/10.1016/j.foreco.2012.06.030>
- Williamson MA, Fleishman E, Mac Nally RC et al (2020) Fire, livestock grazing, topography, and precipitation affect occurrence and prevalence of cheatgrass (*Bromus tectorum*) in the central Great Basin, USA. *Biol Invasions* 22:663–680. <https://doi.org/10.1007/s10530-019-02120-8>
- Wolkovich EM, Cleland EE (2011) The phenology of plant invasions: a community ecology perspective. *Front Ecol Environ* 9:287–294. <https://doi.org/10.1890/100033>
- Woolman AM, Coop JD, Shaw JD, DeMarco J (2022) Extent of recent fire-induced losses of ponderosa pine forests of Arizona and New Mexico, USA. *For Ecol Manag* 520:120381. <https://doi.org/10.1016/j.foreco.2022.120381>
- Zefferman E, Stevens JT, Charles GK et al (2015) Plant communities in harsh sites are less invaded: a summary of observations and proposed explanations. *AoB Plants* 7:plv056. <https://doi.org/10.1093/aobpla/plv056>
- Zhu Z, Key C, Ohlen D, Benson N (2006) Evaluate sensitivities of burn-severity mapping algorithms for different ecosystems and fire histories in the United States. Final Report to the Joint Fire Science Program. JFSP 01–1–4–12

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