# Chapter 2 Exotic Annual *Bromus* Invasions: Comparisons Among Species and Ecoregions in the Western United States

Matthew L. Brooks, Cynthia S. Brown, Jeanne C. Chambers, Carla M. D'Antonio, Jon E. Keeley, and Jayne Belnap

**Abstract** Exotic annual *Bromus* species are widely recognized for their potential to invade, dominate, and alter the structure and function of ecosystems. In this chapter, we summarize the invasion potential, ecosystem threats, and management strategies for different *Bromus* species within each of five ecoregions of the western United States. We characterize invasion potential and threats in terms of ecosystem resistance to *Bromus* invasion and ecosystem resilience to disturbance with an emphasis on the importance of fire regimes. We also explain how soil temperature and moisture regimes can be linked to patterns of resistance and resilience and provide a

M.L. Brooks (⊠)

US Geological Survey, Western Ecological Research Center, Oakhurst, CA 93644, USA e-mail: matt\_brooks@usgs.gov

#### C.S. Brown

Department of Bioagricultural Sciences and Pest Management, Colorado State University, Fort Collins, CO 80523-1177, USA e-mail: cynthia.s.brown@colostate.edu

### J.C. Chambers

USDA Forest Service, Rocky Mountain Research Station, Reno, NV 89512, USA

e-mail: jchambers@fs.fed.us

#### C.M. D'Antonio

Environmental Studies Program, University of California, Santa Barbara, Santa Barbara, CA 93106, USA

e-mail: dantonio@es.ucsb.edu

#### J.E. Keelev

US Geological Survey, Western Ecological Research Center, Three Rivers, CA 93271, USA e-mail: jon\_keeley@usgs.gov

#### J. Belnap

US Geological Survey, Southwest Biological Science Center, Moab, UT 84532, USA e-mail: jayne\_belnap@usgs.gov

© Springer International Publishing Switzerland 2016 M.J. Germino et al. (eds.), *Exotic Brome-Grasses in Arid and Semiarid Ecosystems of the Western US*, Springer Series on Environmental Management, DOI 10.1007/978-3-319-24930-8\_2

12 M.L. Brooks et al.

conceptual framework that can be used to evaluate the relative potential for invasion and ecological impact of the dominant exotic annual *Bromus* species in the western United States.

**Keywords** Fire • Resilience • Resistance • Management • Moisture regime • Temperature regime

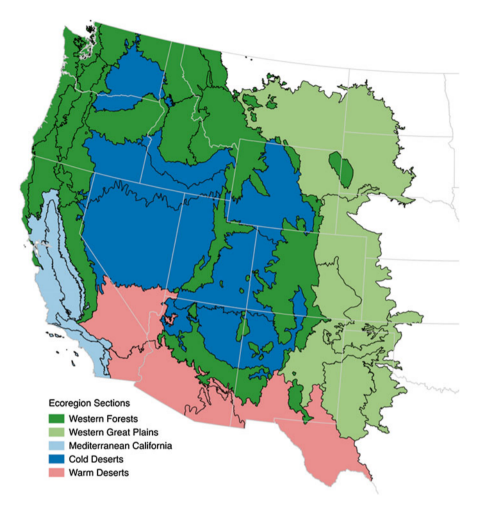
### 2.1 Introduction

Numerous *Bromus* species occupy a wide range of ecosystems around the globe (Atkinson and Brown 2015; Clayton et al. 2006), including 58 species within North America (Clayton et al. 2006), approximately half of which are exotic species that are nonnative to the continent (Pavlik 1995). Exotic annual *Bromus* (*Bromus* hereafter) are of significant concern to ecologists and land managers alike (Brooks and Pyke 2001). Much of what has been reported about the environmental and ecological controls on *Bromus* and their interactions with land uses and disturbance regimes have focused on the Cold Deserts ecoregion of the western United States (Fig. 2.1) and on *Bromus tectorum* L. (cheatgrass) in particular. This chapter reaches beyond that specific ecoregion and individual species to examine the invasion potential, ecosystem threats, and management strategies for the dominant *Bromus* species within each of five major ecoregions of the western United States (Table 2.1; Fig. 2.1). Other chapters within this book that are relevant to the topics in this chapter include Chambers et al. (2015), Germino et al. (2015), Monaco et al. (2015), and Pyke et al. (2015).

### 2.1.1 Resistance and Resilience to Bromus Invasions

The invasion potential, ecosystem threats, and management strategies for *Bromus* all vary based on ecosystem resistance to invasion and resilience to disturbance. Resistant ecosystems have the capacity to retain their fundamental structure, processes, and functioning (or remain largely unchanged) despite stressors, disturbances, or invasive species (Folke et al. 2004). Resilient ecosystems have the capacity to regain their fundamental structure, processes, and functioning following stessors like drought and disturbances like wildfire and then reorganize without crossing thresholds to alternative ecosystem states (Holling 1973; Walker et al. 2004). We use resistance and resilience as explanatory concepts throughout this chapter, along with temperature and precipitation gradients which are key factors affecting resistance and resilience (Brooks and Chambers 2011; Chambers et al. 2014a, b; Fig. 1.1).

Resistance to invasion is particularly important in the context of this chapter and is related to abiotic and biotic attributes and ecological processes that limit the population growth of invading species (D'Antonio and Thomsen 2004). Ecosystem resistance to invading species is a function of both the fundamental niche, which is largely related to climate suitability, and the realized niche, which is related



**Fig. 2.1** Ecoregions of the western United States that are characterized by distinct temperature and soil moisture regimes and influence resistance to *Bromus*. Each ecoregion is an amalgam of various level I, II, and III Ecoregions of North America (http://www.epa.gov/wed/pages/ecoregions/na\_eco.htm, accessed 17 Jun 2014; Wiken et al. 2011). Their specific delineations were based on breaks in soil temperature and moisture (amount and seasonality) regimes associated with ecological resistance to, and ecosystems threats posed by, *Bromus* invasions, and the availability of published information on these topics. The Western Forests ecoregion encompasses Level I ecoregions 6.0 and 7.0 (the Northwestern Forested Mountains and the Marine West Coast Forest). Mediterranean California corresponds to the Level I ecoregion 11.0 which bears the same name. The Cold Deserts and Warm Deserts correspond to Level II ecoregions 10.1 and 10.2, respectively. The Western Great Plains includes three level III ecoregions (Table 2.1) representing parts of two Level II ecoregions, the West-Central Semi-Arid Prairies and South-Central Semi-Arid Prairies (ecoregions 9.3 and 9.4)

to species interactions, both of which need to be understood to evaluate *Bromus* invasions (Chambers et al. 2014a). Temperature and precipitation amounts, and precipitation seasonality, are key attributes that influence ecosystem resistance

**Table 2.1** Major ecoregions of the western United States and their most widespread dominant and subdominant exotic annual *Bromus* species

	Dominant	Subdominant	Associated EPA III Ecoregions
	Bromus	Bromus	(mean and range of annual precipitation
Ecoregion	species	species	in mm) (Wiken et al. 2011)
Warm Deserts	B. rubens	B. tectorum	10.2.1, 2, 4: Mojave (167, 50–900), Sonoran (206, 75–560), Chihuahuan (340, 200–635) 12.1.1 Madrean Archipelago (421, 260–950)
Cold Deserts	B. tectorum	B. rubens	10.1.2–8: Columbia Plateau (334, 150–600), Northern Basin and Range (351, 150–1000), WY Basin (296, 130–500), Central Basin and Range (277, 4–1000), CO Plateaus (298, 130–800), AZ/NM Plateau (293, 125–380), Snake River Plain (316, 110–350)
Mediterranean California	B. rubens B. diandrus (= B. rigidus) B. hordeaceus	B. madritensis B. tectorum	11.1.1–3: CA coastal sage/chaparral/ woodlands (548, 200–1400), central valley (443, 125–760), southern California mountains (525, 220–1250)
Western Great Plains	B. arvensis (= B. japonicus) B. tectorum	None	9.3.3: NW great plains (393, 250–510) 9.4.1, 3: high plains (433, 305–530), southwestern tablelands (449, 255–710)
Western Forests	B. tectorum	B. diandrus	6.2.5, 7, 8, 11, 12, 13, 14: north cascades (1761, 300–6000), Cascades (824, 1150–3600), eastern cascades (649, 500–3500), Klamath mountains (1438, 500–3000), Sierra Nevada (1070, 150–2500), Wasatch/ Uinta mountains (602, 150–1400), southern rockies (588, 255–1750) 7.1.7–9: Puget lowland (1400, 300–2500), coast range (2149, 1000–5000), Willamette valley (1228, 900–1600) 13.1.1: AZ/NM mountains (477, 270–2000)

and define ecological niche space (Davis et al. 2000; Miller et al. 2013; Chambers et al. 2014a, b, 2015).

Site occurrence information for *Bromus* gleaned from the GrassPortal website (www.grassportal.org, Osborne et al. 2011) indicates that resistance to invasion should vary among species based on mean annual minimum and maximum temperature and precipitation (Fig. 2.2). *Bromus rubens* L. (red brome) is clearly found at the hottest and driest sites, whereas *Bromus arvensis* L. (field brome) and to some degree *B. tectorum* occur at the coldest and wettest sites in the database. Variability among species is low at the lower end of the precipitation gradient, but very high at the upper end of the gradient (Fig. 2.2c). The preponderance of outlier sites on the high end of the precipitation gradient may represent occurrences at early successional or otherwise disturbed sites embedded within high precipitation vegetation types, especially within the Western Forests ecoregions, that would otherwise have

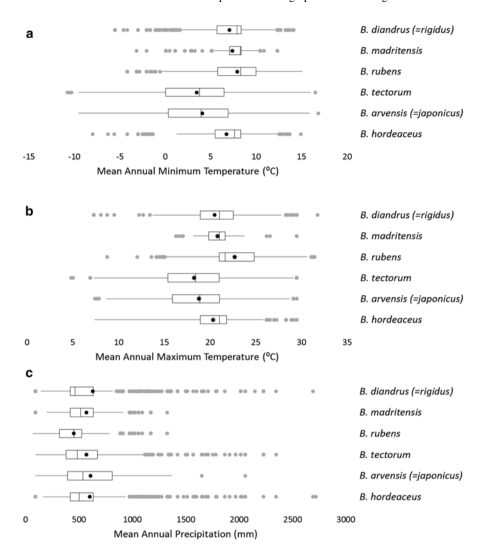


Fig. 2.2 (a) Annual minimum temperature, (b) annual maximum temperature, and (c) annual precipitation for site locations of *Bromus* in the western United States. Data were summarized for states intersecting the ecoregional range map (Fig. 2.1), and downloaded from www.grassportal. org (accessed 16 Feb 2015, Osborne et al. 2011). Species were only included if the database was represented by site locations in states throughout the species range (www.plants.usda.gov). Numbers of sites were as follows: *B. diandrus* (and *B. rigidus*), n=1241; *B. madritensis*, n=226; *B. rubens*, n=1568; *B. tectorum*, n=2004; *B. arvensis* (and *B. japonicus*), n=897; *B. hordeaceus*, n=1188. Climate data are 30-year mean annual values from 1961 to 1991; box plots show quartiles 1, 2, and 3, and upper and lower limits. Mean climate values are shown as *solid dots* and outliers as *gray dots*. Data sources and other database information are listed in Table 2.2

high resistance to *Bromus*. These data suggest that *Bromus* may be firmly established at the relatively lower precipitation sites that define one boundary of both their fundamental and realized niches. However, they additionally suggest there may be higher precipitation sites that are within their fundamental niche, but that are not currently within their realized niche due to competition from other species. The implications of this are that *Bromus* may be poised to expand and potentially dominate climatically suitable high precipitation sites in the event that land use or climate conditions change the competitive balance in their favor.

There is great interest in understanding the resistance of ecosystems to invasion by *Bromus* because of their negative ecological effects. Chief among these effects are altered fire regimes (Whisenant 1990; D'Antonio and Vitousek 1992; Brooks and Pyke 2001; Brooks et al. 2004; Balch et al. 2013; Germino et al. 2015). *Bromus* can alter fire regimes by changing fuel structure in ways that can lengthen the fire season, enhance ignition probabilities, increase fire extent, and alter fire severity (Brooks et al. 2004). The altered fire regime in turn creates conditions that promote dominance of the species that created the new fuels, thus creating a positive grass/fire feedback (D'Antonio and Vitousek 1992), the mechanisms of which are much better understood than perhaps any other potential ecosystem threat posed by *Bromus*. Important questions for evaluating the potential impacts of *Bromus* among ecoregions include the following (Brooks 2008). Has the invasion introduced a novel structure or property into an ecosystem? Has that novel feature in turn altered ecosystem processes or their controls (disturbance regimes)? Do the resultant conditions perpetuate a new, persistent ecosystem state (e.g., alternative state)?

Ecosystem resilience, and thus ecological risks posed by plant invasions, can be affected when disturbance regimes (perturbation type, timing, duration, size, and/or intensity) are altered from historical norms. Land uses are often at the root of these alterations, including inappropriate grazing of grasslands and shrublands by livestock and clear-cut harvests of forests. Plant invasions themselves may also facilitate changes in disturbance regimes in ways that increase ecological effects of the invading species, such as the grass/fire cycle.

# 2.1.2 Species, Ecoregions, and Chapter Structure

We focus on six widespread annual exotic *Bromus* taxa within the western United States, specifically *B. arvensis*, *Bromus diandrus* Roth (ripgut brome), *Bromus hordeaceus* L. (soft brome), *Bromus madritensis* L. (compact brome), *B. rubens*, and *B. tectorum* (Table 2.1). At times in the past, botanical taxonomic authorities have considered *B. arvensis* and *Bromus japonicus* Thunb. (Japanese brome) to be distinct species, but the taxonomic authority used in this book, namely the USDA, Natural Resources Conservation Service, Plants Database, currently considers them to be one in the same species within *B. arvensis* (http://plants.usgs.gov, accessed 21 Jul 2015). Similarly, *B. diandrus* and *Bromus rigidus* Roth (ripgut brome) have been considered by some as separate species, but are both currently subsumed within

B. diandrus in the Plants Database. Thus, where we reference B. arvensis or B. diandrus throughout this chapter we are also referencing their Plants Database synonyms.

Bromus madritensis and B. rubens are currently considered distinct species in the Plants Database (http://plants.usgs.gov, accessed 21 Jul 2015), although they intergrade so much in their taxonomic characteristics that they are often difficult to distinguish and may be misidentified in some publications. For example, an analysis of 28 field samples from central and southern California and southern Nevada (M. Brooks, unpublished data, 2015) indicated that every sample contained taxonomic characteristics of both species, although those of B. rubens were much more frequent (mean 9.1, median 9.0, of 11 characteristics).

We also identify five major ecoregions in the western United States (Fig. 2.1), each of which is characterized by a unique set of dominant and subdominant *Bromus* (Table 2.1). For each ecoregion, we recognize 3–5 major vegetation types which can be placed along hot-dry to cool-moist soil temperature and moisture gradients and which vary in their resistance to *Bromus* and resilience to disturbance. We use soil temperature and moisture regimes as indicators of relative ecosystem resistance and resilience to various *Bromus* species within each ecoregion (Chambers et al. 2014a, b, c).

Each of the five ecoregional sections in this chapter follows a common outline. For each ecoregion, we (1) describe the invasion potential of each widespread *Bromus* species; (2) explain the ecosystem threats posed by *Bromus* invasions; and (3) summarize the efficacy of typical management strategies for *Bromus*. The majority of citations focus on published studies conducted within each ecoregion. The relative resistance to invasion and resilience to fire are illustrated in a series of conceptual models for each ecoregion focusing on the dominant *Bromus* species in the ecoregion. We acknowledge that the amount of supporting science that underpins these figures varies greatly among ecoregions, and we refer to them as hypothetical resistance and resilience models.

### 2.2 Warm Deserts

Environmental conditions in the Warm Deserts ecoregion are the hottest and driest of anywhere in western United States (Fig. 2.3). Soil temperature regimes are either thermic (mean annual temperature 15–22 °C) or hyperthermic (≥22 °C). Soil moisture regimes are mostly aridic, meaning that the soil is dry for at least half of the growing season and moist for less than 90 consecutive days, which supports mostly desert shrubland vegetation types. Some areas are classified as ustic aridic, meaning they have slightly longer periods of moist soil during the growing season due to higher amounts of summer precipitation which is conducive to the growth of semi-arid desert perennial grasslands. These warm and dry conditions result in limited anthropogenic land uses compared to other ecoregions, although mining, livestock grazing, off-highway vehicle use, and most recently wind and solar energy developments can have broad landscape effects with implications for plant invasion (Lovich and Bainbridge 1999; Brooks and Pyke 2001; Brooks 2009).

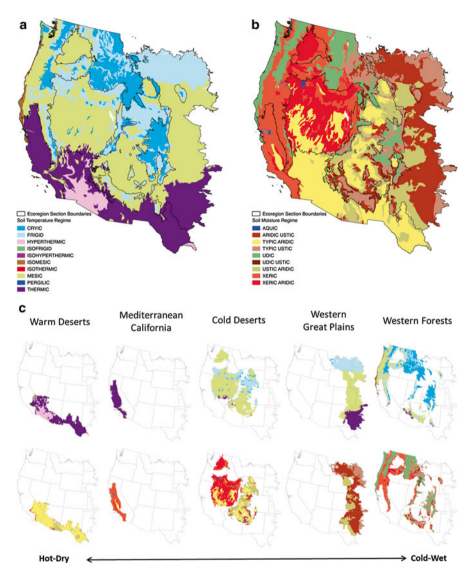
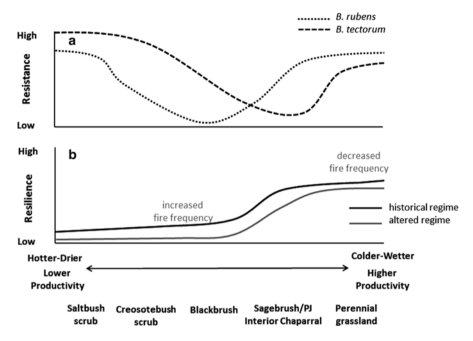


Fig. 2.3 (a) Soil temperature regimes, (b) soil moisture regimes, and (c) comparisons among the six major ecoregions of the western United States

Bromus rubens is the most ubiquitous Bromus species in the Warm Deserts ecoregion. It can occur in all but the most hyperthermic and xeric regions, but is most prevalent in middle elevations typified by creosote bush scrub and blackbrush shrubland which have the lowest resistance to its dominance (Fig. 2.4a) (Brooks and Berry 2006; Brooks 2009; Klinger et al. 2011a). Bromus tectorum can also be locally abundant, but is much more restricted in its geographic distribution in Warm Deserts than is B. rubens (www.calflora.org, accessed 8 Apr 2014). Although B. tectorum can



**Fig. 2.4** Hypothetical (a) resistance to common *Bromus* species and (b) resilience to historical and altered fire regimes for the Warm Deserts ecoregion (primary altered regime characteristic labeled in *gray*)

appear in lower elevation areas with locally higher soil moisture such as riparian zones and roadsides (Brooks 2009), it only occurs as a dominant *Bromus* species at higher elevations, such as those within the upper blackbrush ecotone, sagebrush, piñon–juniper, and interior chaparral (Fig. 2.4b) (Klinger et al. 2011a). *Bromus trinii* Desv. (Chilean chess) has also been reported from some locations, generally within the same ecological range as *B. rubens* (www.calflora.org, accessed 8 Apr 2014), but its relative dominance is typically very low (Brooks 2009; exception is Brooks 1999). *Bromus diandrus* can also be found in the Warm Deserts (www.calflora.org, accessed 8 Apr 2014), but almost always in localized areas of higher soil moisture such as along roadsides, in riparian areas, or associated with agricultural or urban developments (Dudley 2009; M. Brooks, pers. obs.).

# 2.2.1 Invasion Potential of Warm Deserts by Bromus

The lowest elevation ecosystems of the Warm Deserts are characterized by hyperthermic, typic aridic soils (Fig. 2.3c) and sparse cover of creosote bush scrub or saltbush scrub. Creosote bush scrub is dominated by the type species, *Larrea tridentata* (DC.) Coville (creosote bush), and is the most widespread vegetation

type. Saltbush scrub is dominated by one or more *Atriplex* species, including *Atriplex polycarpa* (Torr.) S. Watson (cattle saltbush), *Atriplex spinifera* J.F. Macbr. (spinescale saltbush), and *Atriplex canescens* (Pursh) Nutt. (fourwing saltbush). Productivity is strongly limited by low annual precipitation, and resistance to invasion by *Bromus* is generally high (Fig. 2.4a).

Blackbrush occurs on thermic and mesic soils at middle elevations above creosote bush scrub and is dominated by the type species, *Coleogyne ramosissima* Torr. (blackbrush), but may also contain *L. tridentata*, *Atriplex confertifolia* (Torr. & Frém.) S. Watson (shadscale saltbush), *Yucca brevifolia* Engelm. (Joshua tree), and *Juniperus* spp. (juniper). This vegetation type is the least resistant to invasion by *Bromus*, especially *B. rubens* (Fig. 2.4a). At higher elevations, sagebrush, piñon–juniper woodland, and interior chaparral occur on the more mesic soils of desert mountain ranges and at ecoregional ecotones along the western, northern, and north-eastern margins of the Warm Desert ecoregion. The specific ecosystem type that is present depends on local conditions associated with soils, topography, and disturbance history, but in most cases the dominant *Bromus* species at these higher elevations is *B. tectorum*.

Desert grasslands dominate the thermic soils in the monsoonal regions of the far eastern Mojave and the Sonoran and Chihuahuan deserts. Historically, these ecosystems were dominated by perennial grasses and a fire frequency of every 5–15 years (Humphrey 1958; McPherson 1995; Wright and Bailey 1982). Vigorous stands of perennial grasses with their extensive shallow roots very likely provided high ecosystem resistance to invasion by *Bromus* which share the same shallow rooting zone. During the twentieth century, overgrazing and fire suppression have allowed deeper rooting woody species to invade (McPherson and Weltzin 2000), reducing the vigor of perennial grasses and potentially the ecosystem resistance to *Bromus*. Although the exotic annuals *B. rubens* and *B. tectorum* occur in desert grasslands, they have not exhibited the widespread dominance observed for exotic perennial grasses such as *Eragrostis* von Wolf spp. (lovegrasses) and *Pennisetum ciliaris* (L.) Link (buffelgrass) which have replaced native grasses in many areas and maintain a fairly high level of resistance to *Bromus* invasion.

Bromus rubens was present at 44 % of 126 low elevation (149–1222 m) sites in the eastern Mojave Desert near Lake Mead National Recreation Area (Abella et al. 2012). It was twice as prevalent at middle elevation (800–1222 m) sites, which tended to have more precipitation, were more species rich, and were characterized by mixed perennial communities, compared to the lowest elevation sites (<800 m) which were drier, more species poor, and dominated by either Atriplex L. spp. or L. tridentata. Similar results were reported from the western Mojave Desert in lower to middle elevations ranging from 683 to 1402 m (Brooks and Berry 2006). Similar patterns were also found in post-fire landscapes of the eastern Mojave Desert (Klinger et al. 2011a), where B. rubens was the dominant Bromus species in lower elevation communities, with peak densities between 800 and 1200 m, whereas B. tectorum dominated higher elevation communities, with a peak density at 1800 m. Averaged over both low and high elevation sites in the eastern Mojave Desert, peak densities of

*B. rubens* were almost twice as high as those of *B. tectorum* (Klinger et al. 2011a). The prevalence of *B. rubens* at lower and middle elevation and *B. tectorum* at high elevations is likely a function of the lower minimum temperature tolerance of the latter (Fig. 2.2a), but may be modified by variation in the realized niches of these two species along this elevation gradient as a result of interspecific competition.

Livestock grazing may decrease resistance to *Bromus* by reducing vigor of perennial plants particularly grasses (Brooks and Pyke 2001; Brooks et al. 2007). Short-term, high-intensity grazing has been used to reduce fine fuel loads and, if applied in early spring or late summer, it can preferentially impact *B. tectorum* populations in Cold Deserts (Strand and Launchbaugh 2013). However, in most cases dominance of *Bromus* increases following cessation of grazing in both Warm Deserts and Cold Deserts, and the net long-term effects of livestock grazing are generally favorable for invasive plants (Brooks and Pyke 2001; Brooks et al. 2007).

Atmospheric nitrogen deposition may also reduce landscape resistance to *Bromus* species in Warm Deserts where deposition levels are high downwind from major urban or agricultural areas (Brooks 2003; Allen et al. 2009; Rao and Allen 2010). These studies explain that naturally low soil nitrogen levels in Warm Desert regions, along with low soil moisture availability, can hinder *Bromus* invasions, and elevated levels can reduce this resistance. Increased biomass of *Bromus* and other annual plants facilitated by nitrogen deposition may increase the potential for fire, which may further promote *Bromus* dominance (Rao et al. 2010, 2015).

### 2.2.2 Ecosystem Threats to Warm Deserts from Bromus

The role of *Bromus* in creating continuous biomass cover in the interspaces between perennial plants and increasing the size and frequency of fires represents its primary ecosystem threat in the Warm Deserts ecoregion (Brooks and Pyke 2001; Brooks and Esque 2002). This threat is most prevalent in areas with low resistance to invasion by Bromus and resilience of native vegetation to disturbance, typically at middle elevations dominated by mixed woody scrub and blackbrush communities, and to some degree in creosote bush scrub (Brooks and Minnich 2006; Brooks et al. 2013) (Fig. 2.4). Native perennial cover in blackbrush is already at the threshold of what can carry fire (Fig. 2.5a), and adding Bromus biomass can tip the balance to a more fire-prone landscape (Fig. 2.5b) (Brooks and Matchett 2006). Bromus dominance following fire can increase substantially after only a single fire in saltbush scrub and creosote bush communities (Fig. 2.5c), although subsequent fires can sequentially reduce cover and diversity of native perennials and promote recurrent fire (Fig. 2.5d) (Brooks 2012). This reduced resilience caused by frequent fires is operative primarily in saltbush, creosote bush scrub, and blackbrush, although it can also affect even perennial grasses in cases of extreme fire frequency (Fig. 2.4b).

This grass/fire cycle has been occurring in the Mojave Desert since at least the early 1900s, when post-fire blackbrush landscapes were recognized to be dominated



**Fig. 2.5** (a) Unburned blackbrush stand uninvaded by *Bromus*, (b) unburned blackbrush stand invaded by *B. rubens*, (c) once-burned blackbrush stand dominated by *B. rubens*, and (d) multiple-burned blackbrush stand dominated by *B. rubens* 

by B. rubens and land managers were concerned about the potential for recurrent fire (Fig. 2.6a) (Holmgren 1960; Brooks et al. 2007). Dominance by B. rubens in this region can be established after a single fire and may persist at least a half century even without subsequent burning (Fig. 2.6b). By some definitions this condition can be considered a type conversion from native shrubland to exotic annual grassland, even after a single fire. These fires are much more prevalent in the northeastern Mojave Desert (Brooks and Esque 2002), especially in areas of both high winter and high summer rainfall (Tagestad et al. in press). Modeled estimates of precipitation through the year 2100 forecast increased interannual variability including higher peak rainfall years (Tagestad et al. in press) which are conditions associated with increased probability of fire occurrence (Brooks and Matchett 2006). Higher elevation sagebrush, piñon-juniper, and interior chaparral ecosystems with mesic to warm frigid soils have relatively low resistance to invasion by B. tectorum (Fig. 2.4a), but they also have high resilience to disturbance and thus are at relatively low risk of conversion to a grass/fire cycle (Fig. 2.4b). Fire occurrences at these high elevation sites are also not associated with interannual rainfall amounts, providing further evidence that they are not significantly affected by altered fire regimes associated with exotic annual grasses (Brooks and Matchett 2006).



**Fig. 2.6** (a) 7-year post-fire scene dominated by a carpet of *B. rubens* following a stand-replacing fire in a blackbrush shrubland that burned once in 1939 (photo by A. Croft, 12 May 1946). (b) A 63-year post-fire scene showing *B. rubens* persisting with early successional shrubs following a half century of no additional fire (photo by D. Oldershaw, 9 May 2002). The view is looking southwest inside the mouth of Horse Spring Basin in the northeastern Mojave Desert, Lincoln County, NV

Bromus rubens has also been shown to effectively compete with native annual (Brooks 2000; DeFalco et al. 2003) and perennial plants in the Mojave Desert (DeFalco et al. 2007). These competitive effects can influence community dynamics and resilience following disturbance (Brooks and Chambers 2011). For example, during the first 3 years following the 2005 Southern Nevada Fire Complex woody plant seedlings were almost nonexistent except where densities of exotic annuals, primarily B. rubens and Erodium cicutarium (L.) L'Hér. ex Aiton (redstem stork's bill), were exceedingly low (Klinger et al. 2011a). Reduced resilience of native perennials is a key element of the process by which a grass/fire cycle becomes established (Brooks 2008). The reduced productivity and diversity of native annuals and loss of the habitat structure provided by perennial shrubs can negatively affect wildlife, such as the federally Threatened Gopherus agassizii Cooper (desert tortoise) (Brooks and Esque 2002).

# 2.2.3 Management Strategies for Bromus in Warm Deserts

Bromus are already so wide ranging in the Warm Deserts that early detection and eradication is not a relevant management strategy for these taxa. Even when direct control efforts such as herbicide use are implemented, they are typically economically feasible to implement for only a couple of years and in relatively small areas. In addition, even if dominance of one Bromus species is effectively reduced, other exotic species may fill the void if there is no plan for facilitating the growth of other more desirable species to compete with them. In one dramatic case from the riparian zone in Zion Canyon of SW Utah at the regional ecotone between the Warm Deserts, Cold Deserts, and Western Forests, an herbicide treatment effectively controlled B. diandrus, but inadvertently led to dominance of B. tectorum by the following year (Fig. 2.7). This is an important reminder that any effort to reduce abundance of



**Fig. 2.7** Effects of fall Imazapic treatment of *B. diandrus* on subsequent increase in dominance by *B. tectorum* the following spring in a riparian woodland in Zion Canyon, Utah (photo by M. Brooks, Spring 2006)

*Bromus* in any ecosystem should consider the likely net response of the plant community and include effective contingency plans.

Restoration guidelines in Warm Deserts generally focus on maximizing abundance and diversity of native species, diversity of functional types, and groups of species important for critical aspects of ecosystem function (Brooks et al. 2013). Thus, the most effective management approach for *Bromus* in the Warm Deserts is to maximize ecosystem resilience to disturbance. In doing so, landscapes will be less prone to ecosystem type conversions and can increase resistance to dominance by *Bromus* and other invasive plant species (Brooks and Chambers 2011; Chambers et al. 2014a). This can be done proactively by minimizing the extent and frequency of major disturbances such as wildfire (Brooks et al. 2007). Once these disturbances occur, management options become much more limited. Post-fire seeding is a standard management tool in other ecoregions to promote recovery of native species and minimize dominance of undesirable invasives such as Bromus. However, its long-term efficacy has recently been shown to vary depending on relative resistance and resilience in both the Cold Deserts (Beyers 2004; Shinneman and Baker 2009; Pyke et al. 2013; Knutson et al. 2014) and Warm Deserts (Klinger et al. 2011b). Thus, the most effective fire management strategy is to prevent fires from starting, and suppress fires once they start, especially at middle elevations characteristic of the upper creosote bush scrub ecotones and blackbrush shrublands (Brooks et al. 2013) where resistance and resilience are both low (Fig. 2.4).

### 2.3 Mediterranean California

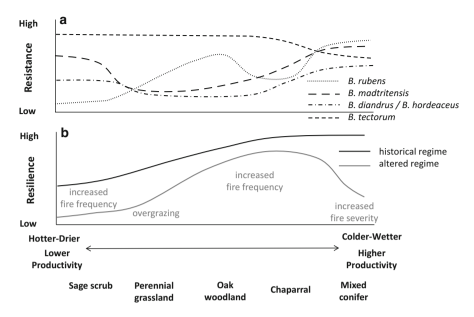
The central valley and surrounding foothills, coastal mountains, and coastal plains of California have a Mediterranean climate with a relatively cool and wet winter and spring and a hot and dry summer and autumn. Soil temperatures are mostly thermic and similar to much of the warm desert region, but moisture regimes are mostly classified as xeric, rather than drier typic aridic which characterizes the Warm Deserts (Fig. 2.3c).

The grasslands of Mediterranean California can be divided into various ecotypes dominated by different *Bromus* species, often with co-dominance by exotic annual grasses from other genera. The more arid southern and interior grasslands are defined by an abundance of *B. rubens*, although its taxonomic similarity with *B. madritensis* (Sect. 2.1.2) makes it difficult to attribute dominance to one or the other. By contrast, the coast range grassland is defined by dominance of *B. diandrus* and *B. hordeaceus* (Jackson and Bartolome 2002; Keeler-Wolf et al. 2007). Serpentine grasslands can be dominated by *B. hordeaceus*, especially following periods of high rainfall. The coastal prairie is more defined by native and invasive perennial grasses, but *B. hordeaceus* and *B. diandrus* are also common. Disturbed areas within the region's coastal sage scrub and chaparral shrublands tend to be dominated by *B. rubens*, often in association with *B. diandrus* and *B. hordeaceus*. In somewhat moister woodlands and riparian zones, these species are often replaced entirely by *B. diandrus*, and in the higher inland mountain forests with colder winters, they are replaced by *B. tectorum*.

# 2.3.1 Invasion Potential of Mediterranean California by Bromus

Ecosystems within the Mediterranean California ecoregion are diverse and vary in their resistance to *Bromus* invasion. Sage scrub is at the hottest and driest end of the productivity gradient in the Mediterranean California ecoregion and tends to have the lowest resistance to *Bromus* invasion particularly when it is disturbed or stressed by nitrogen addition (Fig. 2.8a). Perennial grasslands and oak woodlands tend to have moderate resistance to *Bromus* invasion (Fig. 2.8a), but only where disturbance regimes have not shifted outside of the range of historic variation due to grazing or increased fire frequency (Fig. 2.8b). Oak woodlands and oak savannas that have been grazed for many decades often have an understory of *B. diandrus* which increases in dominance after grazing ceases (Rice and Nagy 2000; Stahlheber and D'Antonio 2013). Chaparral and mixed conifer forests display the highest resistance to *Bromus* invasion due to their high canopy cover which shades the soil surface impeding annual plant growth.

*Bromus* species in Mediterranean California thrive best where perennial cover is slow to recover following disturbance, and their dominance is often transitory where



**Fig. 2.8** Hypothetical (**a**) resistance to common *Bromus* and (**b**) resilience to historical and altered fire regimes for the Mediterranean California ecoregion (primary altered regime characteristic labeled in *gray*)

perennial cover recovers more quickly. In a study of 250 chaparral sites (Keeley et al. 2008), *B. madritensis* cover was negatively correlated with pre-fire stand age and previous fire severity. In addition, 5 years post-fire cover of exotic species, of which *Bromus* was a major contributor, was best predicted by shrub cover and estimated exotic seed bank at the time of fire. In essence, the extent to which *Bromus* and other exotics can invade disturbed Mediterranean shrublands is a race between exotic propagules getting to the site (since most that were there before are killed by high-intensity fire) and shrub canopy reestablishment. Under historical disturbance regimes, productivity and resilience covaried (Fig. 2.8b). However, various anthropogenic factors have altered disturbance regimes and reduced ecosystem resilience, especially at the lowest and highest ends of the productivity gradient. Because chaparral tends to be more species rich with more post-fire resprouting species compared to sage scrub, it tends to be more resilient to increased fire frequency (Fig. 2.8b).

Exotic annual grasslands, most of which have a strong component of *Bromus*, owe their origins to either (1) disturbance that converts native woody vegetation or mixed native perennial grass/shrub stands or (2) direct invasion of native grasslands (Keeley 1990). These exotic grasslands are markedly unlike native perennial grasslands in that the former are lower in functional diversity than the latter, being dominated entirely by annual grasses and forbs with a different composition of native forbs (Molinari and D'Antonio 2014). Where remnants of native grasslands persist within larger exotic annual grasslands, *Bromus* are usually only co-dominants (e.g., Molinari and D'Antonio 2014).



**Fig. 2.9** Chaparral with three different fire histories in southern California. The entire scene burned in the 1970 Laguna Fire. The front and middle third burned in the 2001 Viejas Fire, and the front third burned a third time in the 2003 Cedar Fire. The background illustrates mature chaparral dominated by the native shrub *Adenostoma fasciculatum*, the middle third early seral chaparral dominated by the native shrub *Lotus scoparius*, and the front portion a 1-year post-fire scene dominated by the nonnative annual grass *B. madritensis* (photo by Richard Halsey, Spring 2004)

Fire history plays a critical role in the prevalence of exotic grasses within shrubland ecosystems of the region. Short intervals between fires can thin shrub density, which can both increase grass abundance and reduce native shrub seed banks and the vigor of potential resprouts (Zedler et al. 1983; Haidinger and Keeley 1993; Talluto and Suding 2008; Keeley and Brennan 2012). These thinned stands lead to reduced fire intensity which may further favor survivorship of exotic propagules (Keeley et al. 2008). A classic example of this dynamic is illustrated in Fig. 2.9 where a chaparral stand that burned three times over three decades was dominated by *B. madritensis* compared to areas burned only once or twice. However, in a region-wide remote sensing study only a few shrublands appeared to be undergoing type conversion via fires that occurred two times within 8 years (Meng et al. 2014), so other factors may interact with fire frequency to affect vegetation trajectories.

Atmospheric pollution in the form of nitrogen deposition has been implicated in the invasion of semiarid shrublands by annual grasses including *Bromus* (Westman 1979; Allen et al. 1998). In desert scrub, openings between widely spaced shrubs are readily invaded by invasive grasses and this is enhanced by nitrogen pollution (Brooks 2003). In contrast, within closed canopy shrublands there is no evidence

that pollution alone can promote type conversion from shrubs to these grasses, but there is evidence that following a disturbance that opens up the canopy (such as fire or human disturbance), *B. diandrus* and *B. rubens* can quickly establish and suppress native shrub seedlings (Eliason and Allen 1997). Recent work suggests that nitrogen enhances competitive suppression of native shrub seedlings in the presence of abundant *Bromus* at least in part due to interactions with mycorrhizae (E. Allen, pers. comm.). Also in coastal prairies, nitrogen addition such as by cyclical growth of native lupines enhances *B. diandrus* cover (Maron and Connors 1996; Maron and Jeffries 1999; Suttle et al. 2007) at the expense of native grassland species. In serpentine grassland, nitrogen addition enhances *B. hordeaceus* dominance at the expense of native species (Huenneke et al. 1990).

Sites dominated by native perennial grasses today appear to have some resistance to invasion by annual grass species including some *Bromus*. This has been demonstrated experimentally as well as by observation (Seabloom et al. 2003; Corbin and D'Antonio 2004; Lulow 2006). Invasive bromes in these settings include B. *diandrus*, B. hordeaceus, and B. madritensis. Resistance was disrupted historically by conversion of landscape to crop agriculture and by animal disturbance such as by livestock. The interaction of grazing with native versus exotic grasses in California is controversial and varies by region. However, a meta-analysis showed that exotic grass cover, including largely *Bromus*, decreases with livestock grazing (Stahlheber and D'Antonio 2013) whereas seasonal grazing can enhance native grass cover particularly in the more coastal regions of California. Exotic forb cover tends to increase with grazing while native forb richness (but not cover) increases with grazing. Despite the use of grazing as a tool to enhance native species and reduce exotic annual grasses such as B. diandrus, these grasses remain an important component of virtually all of the grazing lands in California (Jackson and Bartolome 2007).

Natural disturbance factors such as gopher excavation have also been shown to promote *Bromus* abundance in grassland settings (Hobbs and Mooney 1995) as do excavations by feral pigs (Cushman et al. 2004). These *Bromus* also invade ruderal environments created by various types of disturbance that remove native vegetation and disturb the soil. Invasion success is complicated and tied to soil nutrients and belowground root competition (Brown and Rice 2010). On severely disturbed sites, *Bromus* and other exotic annuals may persist for 70 years or more in what otherwise would have been sage scrub (Stylinski and Allen 1999).

In ecosystems such as oak woodlands and mixed conifer forests that are characterized mostly by frequent surface fire regimes, decreasing fire frequency can lead to understory fuel accumulations and increased potential for higher severity crown fire and lower post-fire resilience (Fig. 2.8b). The loss of forest canopy cover can change microclimates in ways that significantly reduce resilience and improve conditions for dominance by *Bromus*. In Oak savannas, the understory of trees is well known to promote dominance of *B. diandrus* over other annual species including other *Bromus* (Rice and Nagy 2000), and this is true over a wide latitudinal range of California oak savannas (Stahlheber 2013). Thus, loss of tree cover may change the relative dominance of specific *Bromus* in this vegetation type.

# 2.3.2 Ecosystem Threats to Mediterranean California from Bromus

Bromus invasions have had a multitude of impacts in Mediterranean California. Their success has largely been at the cost of losing native shrublands, primarily chaparral, sage scrub, and oak woodlands, although perennial grasslands have also been affected (Molinari and D'Antonio 2014). When this type conversion occurs it affects biodiversity, hydrology, fire regimes, and the global carbon balance. However, the direction of such effects can be variable. For example, Bromus invasion of coastal sage scrub in southern California has been linked to increased soil carbon storage if the shrublands remain unburned. If they burn and convert to grasslands, it is likely that soil carbon storage will decline (Wolkovich et al. 2009a).

Where type conversion has occurred, the dominant functional type has been changed from deep-rooted shrubs to shallow fibrous rooted annuals. This impacts soil water holding capacity and increases erosion and debris flows following heavy winter rains but also has the potential for enhancing stream flow (Meixner and Wohlgemuth 2003). Type conversion also impacts fire regimes in a couple of important ways. Length of the fire season is greatly expanded because fuel moisture in drying herbaceous biomass is lost rapidly and this can provide available fuel during all 12 months of the year. In addition, a substantial proportion of fires in the region ignite in herbaceous fuels (Syphard and Keeley 2015), which provides a wick that carries fires into woody vegetation, promoting repeat fires that further degrades mixed grass and shrub mosaics (Fig. 2.9) (Keeley et al. 2012; Monaco et al. 2015).

This grass/fire cycle where fire regimes are altered by grass invasion is well documented for several ecosystems (D'Antonio and Vitousek 1992). However, it is important to recognize that there are very diverse ways grass invasion impacts fire regimes depending on the natural fire regime in the region. For example, fire-prone perennial grass invasion into fire-sensitive tropical forests results from increased fire intensity, which acts to open up the forest to further invasion. California annual grass invasions by *Bromus* spp., *Avena* L. spp., and others are quite different because these annuals are invading a fire-prone shrubland ecosystem, one with a natural fire regime of high-intensity fires, and such fires are in some respects detrimental to grass invasion as they kill seeds of the invaders. However, these native shrublands require decades to recover and if grasses invade in the early post-fire stages they can produce flashy fuels that readily ignite and carry repeat fires when the native vegetation is still not mature. These short-interval low-intensity fires can result in stand thinning, further grass invasion, and increased probability of a repeat fire (Fig. 2.9). Sage scrub is especially susceptible to alteration due to repeat short-interval fire compared to chaparral, largely because productivity and levels of resilience are lower (Fig. 2.8b). In these stands, fire severity is lower and exotic seed survivorship is higher. As a general rule as stand age decreases, fire severity decreases and exotic seed survivorship and vegetative dominance increase after fire (Keeley et al. 2008).

One of the predominant concerns of reserve managers in California grassland and oak savanna habitats is the apparent decline of native wildflowers (Minnich 2008)

and native biodiversity in general. *Bromus*-dominated grasslands are depauperate in diversity of plant functional types and animal communities (Rochester et al. 2010; Molinari and D'Antonio 2014). An ongoing study suggests that it is the accumulated litter of *Bromus* that suppresses native forb germination and not direct competition for soil nutrients between *Bromus* and native forbs (Molinari 2014). Molinari and D'Antonio (2014) demonstrate that there are several native forbs that can coexist with *B. diandrus* and that although native forb richness is depressed where this species becomes dominant, several native forbs preferentially associate with this grass. The influence of *Bromus* invasion on native animals has been little studied. One study found that despite enhancing primary production, *Bromus* invasion into coastal shrublands caused either a decrease in or no effect on arthropod assemblages (Wolkovich et al. 2009b). Key native ant species declined as litter of *Bromus* increased (Wolkovich et al. 2009b).

# 2.3.3 Management Strategies for Bromus in Mediterranean California

Preventing further invasions of *Bromus* species is largely dependent on reducing the incidence of disturbance and maximizing ecosystem resilience (Fig. 2.8). In grasslands, this may involve manipulating livestock grazing patterns or removal of feral animals such as pigs. Where exotic grasses have invaded fire-tolerant or even fire-dependent ecosystems, it is important to appreciate that these shrublands require a particular fire regime that typically involves long fire-free intervals and high-intensity fire. Humans currently cause over 95 % of all fires in the foothills and coastal plain of California and have greatly increased the natural fire frequency (Safford and Van de Water 2014). Keeping unwanted ignitions out of these systems is a major challenge. A more tractable problem is reducing planned ignitions and resource managers in the region are beginning to appreciate that prescription burning, while possibly having some fire-hazard reduction benefit, is adding to the readily ignitable fuel load on these landscapes.

Bromus dominance may decline under passive management for some landscapes whereas other landscapes require active restoration. Sage scrub displaced by grasslands from various disturbances can return within a few decades if disturbance is reduced and native seed sources are in close proximity (Freudenberger et al. 1987; DeSimone and Zedler 1999; Callaway and Davis 1993), although some sites have proved resistant to passive restoration (Stylinski and Allen 1999). The type of disturbance associated with annual grassland appears important as soil disturbance that affects biological soil crusts (if present) can favor exotic annuals such as Bromus and inhibit native shrub seedlings (Hernandez and Sandquist 2011).

Native perennial grasslands invaded by exotic grasses show some promise of being restored to a greater native grass and forb composition through active management. In particular, winter/spring grazing has been shown to enhance dominance of

the bunch grass *Nassella pulchra* (Hitchc.) Barkworth (purple needlegrass) while reducing dominance of *Bromus* (Stahlheber and D'Antonio 2013). It has also been suggested that spring fires, which kill the exotic seed bank while still on the plant but do not diminish the resprouting capacity of native perennial bunch grasses and forbs, have potential (DiTomaso et al. 2006), although native grasses can still be highly susceptible to mortality from fire (Marty et al. 2005). One experiment showed that high-intensity fires in mixed grasslands can improve the ratio of native versus exotic (*Bromus* and *Avena* spp.) grasses for at least 2 years (Keeley and Brennan 2015). However, a key to the success of this effort was that these native-dominated grasslands possessed substantial fuel loads because it had been many years since the last fire or other major form of disturbance. Large fuel loads contributed to extraordinary flame lengths and presumably high soil temperatures that were sufficient to kill the exotic seed bank but allowed for the survival of native perennial grasses.

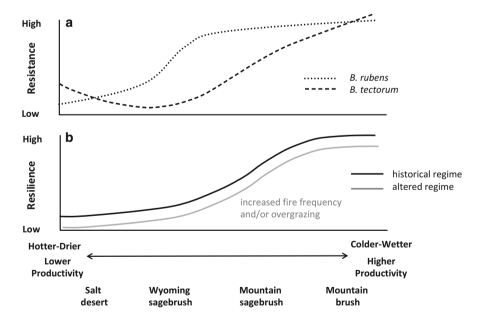
Studies from other vegetation types also illustrate the importance of the fuel load in creating fire behavior that can reduce *Bromus* dominance. For example, an investigation of spring burning on *B. tectorum* understory in a ponderosa pine forest failed to displace this exotic invader (Keeley and McGinnis 2007). In that case the low nutrient availability of the soils produced rather sparse grass populations and field measures of temperatures during the fire revealed that they failed to reach high enough levels to kill seed banks. The importance of high fuel loading to make this approach effective has been noted in studies of other *Bromus* (Sweet et al. 2008). Even when prescription burning to eliminate *Bromus* has been effective for the target species, they are replaced by other exotic annuals (Keeley et al. 2011). Other studies have also shown that effects of a single fire on annual grass reduction can be short-lived and repeated burning or burning followed by grazing may be necessary for sustained reductions in annual grasses (Corbin et al. 2004).

### 2.4 Cold Deserts

Environmental conditions in the Cold Deserts are on average moderately warm and dry in comparison to the range of soil temperature and moisture regimes across the western United States (Fig. 2.3). However, the topographic diversity of this ecoregion results in high temperature and moisture variability associated with local elevation gradients. The lower elevations dominated by salt desert and Wyoming big sagebrush ecosystems are relatively hot and dry whereas the higher elevations typified by mountain sagebrush and mountain brush are cold and moist. The most widespread invasive *Bromus* in the Cold Deserts ecoregion is *B. tectorum* which occurs over a broad range of temperature and precipitation conditions (Fig. 2.2). *Bromus rubens* also occurs in the Cold Deserts but appears less cold tolerant (Fig. 2.2a) and occurs at lower elevations and latitudes than *B. tectorum*. Colder winter conditions, in particular, severe and sudden freeze events, are lethal to *B. rubens* but are not lethal to *B. tectorum* (Bykova and Sage 2012).

# 2.4.1 Invasion Potential of Cold Deserts by Bromus

Salt desert vegetation types typically occur at relatively low elevations and have the warmest (mesic) and driest (aridic) soil temperature and moisture regimes in the Cold Deserts. Vegetation types that characterize the salt desert are typically dominated by members of the Chenopodiaceae such as Atriplex L. spp. and Sarcobatus Nees spp. (West 1983a, b). Resistance to B. tectorum and presumably B. rubens, which has received less study in this vegetation type, is low to moderate (Fig. 2.10a). Under the warmest and driest regimes, establishment of B. tectorum is limited by frequent low and sporadic precipitation years, and longer term persistence requires either the necessary environmental conditions to maintain viable populations or high propagule pressure (Meyer et al. 2001). Resilience to disturbance is generally low due to low productivity, infrequent plant establishment, and slow recovery (Fig. 2.10b, and see Haubensak et al. 2009). Recent expansion of B. tectorum and B. rubens into marginal salt desert environments has been attributed to favorable years for establishment during El Niño-Southern Oscillation (ENSO) events (Meyer et al. 2001; Salo 2005). Rising CO<sub>2</sub> levels (Smith et al. 1987) and high genetic variability (Ramakrishnan et al. 2006; Haubensak et al. 2014) also may help explain increased establishment and spread of *B. tectorum* and likely *B. rubens*. Invasion of B. tectorum has increased fine fuels and fuel continuity in these types



**Fig. 2.10** Hypothetical (a) resistance to common *Bromus* species and (b) resilience to historical and altered fire regimes for the Cold Deserts ecoregion (primary altered regime characteristic labeled in *gray*). Figure modified from Chambers et al. (2014a)

and wildfires are burning for the first time in recorded history (Haubensak et al. 2009; Balch et al. 2013).

Low to mid elevations have warm and dry (mesic/aridic) to warm and moist (mesic/xeric) soil temperature and moisture regimes and are characterized by Artemisia tridentata Nutt. ssp. wyomingensis Beetle and Young (Wyoming big sagebrush) and to a lesser degree A. tridentata Nutt. ssp. tridentata (basin big sagebrush) vegetation types (West 1983a, b; Miller et al. 2011). Resistance to B. tectorum is low under these regimes, largely due to high climatic suitability for establishment and persistence (Fig. 2.10a) (Chambers et al. 2007, 2014a; Davies et al. 2012). Bromus rubens can occur on warmer and drier sites, especially at the lowest elevations (Salo 2005), but its distribution and relative abundance have not been well quantified. Resilience to disturbance and management treatments is low to moderately low (Fig. 2.10b) (Miller et al. 2013; Davies et al. 2012; Chambers et al. 2014a, b). Because the Cold Deserts represent more moderate climates, they also have high levels of anthropogenic disturbance (agricultural, urban and energy development, livestock, wild horse and burro grazing, off-highway vehicle and recreation use, etc.) which has increased the spread and dominance of B. tectorum (Knick et al. 2011). Progressive expansion of B. tectorum has increased fire frequency and size (Balch et al. 2013), and threshold transitions to annual grass dominance are highest in these vegetation types (Chambers et al. 2014a).

Upper elevations typically have cool and moist (frigid/xeric) to cold and moist (cryic/xeric) regimes and are characterized by A. tridentata Nutt ssp. vaseyana (Rydb.) Beetle (mountain big sagebrush) and mountain brush (e.g., Symphoricarpos Duham. spp. [snowberry.], *Purshia tridentata* (Pursh) DC. [antelope bitterbrush]) vegetation types. Colder soil temperature regimes limit the amount of suitable niche space and increase resistance to *B. tectorum* invasion (Fig. 2.10a) (Chambers et al. 2007; Davies et al. 2012). Although B. tectorum can germinate at relatively low temperatures (Roundy et al. 2007), growth and reproduction under cool to cold (frigid to cryic) regimes are limited as indicated by low growth and reproduction (Chambers et al. 2007). At the warmer end of the gradient depletion of perennial herbaceous species due to either inappropriate livestock grazing or expansion of piñon and juniper trees can promote B. tectorum establishment following fire, but few of these systems become B. tectorum dominated (Chambers et al. 2014b). Favorable conditions for establishment and growth and relatively high levels of productivity result in moderate to high resilience on all but the coldest high elevation sites (Chambers et al. 2014a).

Slope, aspect, and soil characteristics modify soil temperature and water availability and influence occurrence of *B. tectorum* at landscape to plant community scales (Chambers et al. 2007; Condon et al. 2011; Reisner et al. 2013). At higher elevations with cooler soil temperatures, *B. tectorum* is found primarily on south facing slopes (Kulpa et al. 2012) with more solar radiation (Condon et al. 2011; Lovtang and Riegel 2012).

Seasonality of precipitation within the ecoregion also appears to influence establishment and persistence of *B. tectorum*. In the western portion of the ecoregion, most precipitation arrives in winter and spring months and climate suitability to

*B. tectorum* is high. Life history and physiological traits such as germination in early fall through winter (Mack and Pyke 1983) and high growth and nutrient uptake rates (James et al. 2011) make *B. tectorum* highly competitive with native plants. In the eastern portion of the ecoregion, summer precipitation is higher (ustic soil moisture regimes) and this competitive advantage appears to be greatly diminished (Bradley 2009), although establishment and persistence still can occur following disturbance (Bradford and Lauenroth 2006).

### 2.4.2 Ecosystem Threats to Cold Deserts from Bromus

The most widely cited effect of *Bromus* invasions in the Cold Deserts ecoregion is the alteration of fire regimes (Whisenant 1990; Brooks and Pyke 2001; Balch et al. 2013). The mechanisms of these changes in Cold Deserts are largely the same as those described for Warm Desert in Sect. 2.2 and the impacts are explained in more detail in Germino et al. (2015). Invasion of B. tectorum can also alter the size and dynamics of soil N pools, or the activity of the microbial community, with the type and magnitude of the effect depending on factors such as soil characteristics, plant community composition, and time since invasion (Rimer and Evans 2006; Hooker et al. 2008; Schaeffer et al. 2012). Bromus invasions can alter hydrologic conditions resulting in increased soil water recharge (Wilcox et al. 2012) which can decrease resistance to secondary invasion by deep-rooted forbs that mature later in the growing season (Kulmatiski et al. 2006). After wildfires, runoff and erosion can increase, particularly on steep slopes (>15 %) (Wilcox et al. 2012). Wind erosion also can occur after fire, resulting in redistribution or loss of nutrient-enriched soil fines and, under shorter fire return intervals caused by *Bromus*, homogenization of soil properties in Cold Deserts shrublands (Sankey et al. 2009).

Both resilience from disturbance and resistance to Bromus in the Cold Deserts are reduced by disturbances that decrease perennial species abundance and increase resource availability. Such disturbances include excessive livestock grazing and altered fire regimes, specifically increased fire frequency. Many of the dominant shrubs (e.g., Artemisia spp.) are not fire tolerant in the warmer and drier ecological types that are least resistant to Bromus. Plant establishment is often sporadic (Miller et al. 2013) even when these species are seeded (Knutson et al. 2014). More frequent fire coupled with increased competition due to invasion of Bromus and following livestock grazing too soon after reseeding can prevent native regeneration (Eiswerth and Shonkwiler 2006). Populations of B. tectorum often increase rapidly following fire or other disturbances that remove perennial native species and seed banks can grow to ≥20,000 seeds/m<sup>2</sup> within a few years (Humphrey and Schupp 2001; Meyer et al. 2007; Chambers et al. 2015). Typically small seed banks of most native species (Hassan and West 1986; Allen et al. 2008), coupled with low competitive ability of native seedlings with B. tectorum (James et al. 2011; Mazzola et al. 2011), can greatly decrease seedling establishment. Mature individuals of perennial herbaceous species, especially those with similar phenologies, are often strong competitors (Booth et al. 2003), but large individuals or populations of B. tectorum can reduce growth and seed production of these species. Decreases in perennial native species, especially grasses and forbs, biological soil crusts, and the distance between perennial herbaceous species (gaps) due to inappropriate livestock grazing or other disturbances are strongly associated with increases in B. tectorum across a range of ecological types (Chambers et al. 2007; Dettweiler-Robinson et al. 2013; Reisner et al. 2013; Pyke et al. 2015). Increasing plant community dominance of B. tectorum can alter species interactions, influence trophic interactions, and reduce species diversity (Germino et al. 2015). For example, increasing B. tectorum cover has a negative influence on nesting of *Centrocercus urophasianus* Bonaparte (greater sage-grouse) which has been considered for listing under the US Endangered Species Act of 1972 (Federal Register, 50 CFR Part 17), and on the abundance of Spermophilus townsendii Bachman (Townsend's ground squirrel) which is important prey for raptors and other predators (Yensen et al. 1992). Complete conversion to B. tectorum results in loss of shrub structure and shrubland-associated species like Amphispiza belli Cassin (sage sparrow) and an overall decrease in native species diversity (Earnst and Holmes 2012).

Ecosystem resilience to disturbances increases with increasing elevation, latitude, and associated primary productivity (Fig. 2.10). Because resistance to invasion follows a similar pattern, especially for *B. tectorum*, the net threat posed by this species differs with seasonality of precipitation, but is generally highest at low to mid productivity, and lowest in areas of higher productivity (Fig. 10.1, Chambers et al. 2015).

# 2.4.3 Management Strategies for Bromus in Cold Deserts

In the Cold Deserts, a suite of different vegetation management treatments are used to increase resilience to fire and other disturbances and to enhance resistance to invasive annuals (Chambers et al. 2014a). Primary objectives are to reduce woody fuel loads and thus fire severity and extent, decrease exotic annual species abundance and spread, and increase perennial herbaceous species dominance which both promote recovery after disturbance and compete with exotic annual plants. Treatments include various combinations of prescribed fire, mechanical treatment, and herbicide application to decrease sagebrush, piñon, and juniper abundance, herbicide application and grazing to control Bromus, and seeding with perennial herbaceous species and shrubs to restore native plant communities (Monsen et al. 2004; Pyke 2011). However, responses to these treatments often vary due to inherent differences in site resilience and resistance (Miller et al. 2013) and subsequent land use regimes (Eiswerth and Shonkwiler 2006). Knowledge of environmental factors, ecosystem attributes and processes, and disturbance and land use history effects that influence resilience and resistance (Fig. 1.1) can be used to determine appropriate management strategies at both site and landscape scales (Chambers et al. 2014a; Miller et al. 2013).



**Fig. 2.11** (a) A fire that started along a major highway and burned through a sagebrush ecosystem with *B. tectorum* in the understory and into a Jeffrey pine ecosystem, Washoe County, Nevada. (b) Landscape conversion of a sagebrush ecosystem to *B. tectorum* and other exotic annual dominance following repeated fire, Humboldt County, Nevada. Photos by Nolan E. Preece

The degree of uncertainty increases and management options become increasingly limited with decreases in resilience to fire and management treatments and resistance to invasion. In cool and dry to cool and moist regimes with moderate to high resilience and resistance, a variety of management treatments like carefully timed grazing, prescribed fire, mechanical treatments, and herbicide applications can be used to maintain or improve ecological conditions. However, in warm and dry to warm and moist regimes, widespread invasion and increasing dominance of *Bromus* and other invaders have altered vegetation dynamics and often limit options. Largely irreversible thresholds can occur following either fire or management treatments if sufficient perennial herbaceous species for recovery are lacking (Fig. 2.11).

### 2.5 Western Great Plains

The Western Great Plains comprises shortgrass steppe and northern and southern mixed-grass prairie. The foothills of the Rocky Mountains form the western border (Lauenroth and Milchunas 1992; Coupland 1992) and tallgrass prairie the eastern border, which is a vegetation transition zone (Sims and Risser 2000) at approximately the 100th meridian (Van Dyne 1975). Temperatures in the ecoregion are cooler in the north than the south (frigid, mesic, thermic from north to south; Fig. 2.3c). The mean annual temperature in shortgrass steppe is 8.6 °C (Lauenroth 2008) and ranges from 8.2 °C in northern Colorado to 17.7 °C in western Texas and southeastern New Mexico (Lauenroth and Milchunas 1992). Likewise, mean annual temperatures in the mixed-grass prairie increase from north to south (3.6 °C in Saskatchewan to 12.9 °C in central Kansas) (Coupland 1992). A precipitation gradient runs perpendicular to the temperature gradient; it is drier in the west than the east (ustic aridic, aridic ustic, typic ustic from west to east; Fig. 2.3c). Annual precipitation in the shortgrass steppe of northern Colorado ranges from 300 to 400 mm on the west side to 500–600 mm on the east side (Lauenroth and Milchunas 1992) with mean annual precipitation of 321 mm (Lauenroth et al. 2008). Mean annual precipitation in the mixed-grass prairie ranges from around 300 mm near the shortgrass steppe to around 600 mm in the eastern transition zone to tallgrass prairie (Coupland 1992; Moran et al. 2014). There is a smaller gradient of precipitation from south to north ranging from around 430 mm in southern mixed-grass prairie to 340 mm in the northern mixed-grass prairie (Van Dyne 1975).

B. tectorum and B. arvensis are the two most common invasive Bromus that occur in the shortgrass steppe, although they are largely restricted to roadsides and disturbed areas such as old fields because intact native vegetation is highly resistant to invasion (Kotanen et al. 1998; Milchunas et al. 1992) (Fig. 2.13a). The shortgrass steppe is dominated by the short-stature, warm-season species Bouteloua gracilis Willd. ex Kunth) Lag. ex Griffiths (blue grama) and Bouteloua dactyloides (Nutt.) J.T. Columbus (buffalograss). The mid-height, cool-season grass Pascopyrum smithii (Rydb.) Á. Löve (western wheatgrass) is commonly associated with B. gracilis

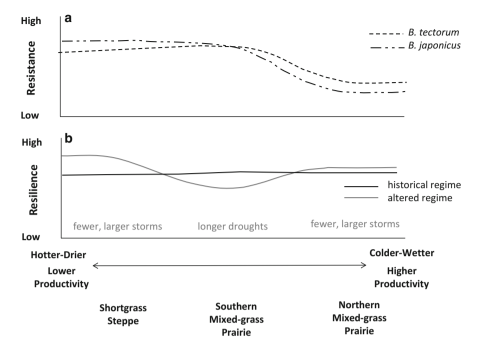
in the northern portion of the shortgrass steppe (Lauenroth and Milchunas 1992). This shifts to *B. gracilis* in association with *Sporobolus airoides* (Torr.) Torr. (alkali sacaton), *Pleuraphis jamesii* Torr. (James' galleta), and *B. dactyloides* toward the south (Lauenroth and Milchunas 1992).

Bromus tectorum and B. arvensis are also the most common invasive Bromus species in northern and southern mixed-grass prairie and are joined by Bromus catharticus Vahl (rescuegrass) and Bromus secalinus L (rye brome), but these species are of secondary dominance. Bromus gracilis remains a major component of the mixed-grass prairie, which comprises cool-season and warm-season species from tallgrass prairie and shortgrass steppe (Sims and Risser 2000). Pascopyrum smithii and Hesperostipa (Elias) Barkworth spp. (needlegrass) are codominant species with B. gracilis in the northern mixed-grass prairie (Van Dyne 1975). Other cool-season species such as Carex L. spp., Elymus lanceolatus (Scribn. & J.G. Sm.) Gould ssp. lanceolatus (thickspike wheatgrass), Koeleria macrantha (Ledeb.) Schult. (prairie Junegrass), and Nassella viridula (Trin.) Barkworth (green needlegrass) become more abundant in particular microsites. In the southern mixed-grass prairie, taller dominants are *P. smithii*, *Andropogon gerardii* Vitman (big bluestem), Schizachyrium scoparium (Michx.) Nash (little bluestem), and Bouteloua curtipendula (Michx.) Torr. (sideoats grama), with B. gracilis and B. dactyloides as a lower layer in the canopy (Van Dyne 1975).

# 2.5.1 Invasion Potential of Western Great Plains by Bromus

The relative resistance of Western Great Plains ecosystems to invasion by exotic annual *Bromus* species can be attributed to adaptation of the native plants to the climate and characteristic disturbances of drought, fire, and grazing. Of the three subregions of the Western Great Plains, northern mixed-grass prairie appears to be least resistant to exotic annual *Bromus* invasion (Fig. 2.12). *Bromus arvensis* (Heitschmidt et al. 1995; Karl et al. 1999) and *B. tectorum* (Heitschmidt et al. 1995) can be dominant species in northern mixed-grass prairie. In Wind Cave National Park in South Dakota, occurrence of exotic annual *Bromus* species was associated with cool-season grasses such as *P. smithii* and *N. viridula* and the shrub *Symphoricarpos occidentalis* Hook. (western snowberry) and negatively associated with warm-season grasses, especially *A. gerardii*, *B. curtipendula*, and *S. scoparium*, and the shrub *Rhus aromatica* Aiton (fragrant sumac) (Ogle and Reiners 2002).

Data from the U.S. National Park Service Northern Great Plains Network indicate that higher cover of *Bromus* can be associated with lower native species richness, but only in the South Prairie region of the network where *Bromus* cover is highest (Fig. 2.14). This is especially true at Scotts Bluff National Monument in northwestern Nebraska, Badlands National Park in southwestern South Dakota, and Fort Laramie National Historic Site in southeastern Wyoming where *Bromus* cover was repeatedly measured in excess of 30 % (Fig. 2.14) as illustrated in the foreground of Fig. 2.13d. This pattern was not exhibited at Agate Fossil Beds National



**Fig. 2.12** Hypothetical (a) resistance to common *Bromus* species and (b) resilience to historical and altered fire regimes for the Western Great Plains ecoregion (primary altered regime characteristic labeled in *gray*)

Monument (Fig. 2.13c), or elsewhere in the Northern Great Plains where *Bromus* cover rarely exceeded 10 % *Bromus* cover (Fig. 2.14).

Resistance to invasion by exotic annual *Bromus* species of southern mixed-grass prairie may be as high as in shortgrass steppe (Fig. 2.12). Although *B. catharticus*, *B. arvensis*, and *B. tectorum* occur in Oklahoma (Bidwell et al. 2004), none of the fact sheets for managing invasive species published by the Oklahoma Cooperative Extension Service are about exotic annual grasses or mention them as being problems (http://www.oces.okstate.edu/extension-fact-sheets, accessed 6 Jan 2015). The invasive species of concern are all shrubs and subshrubs (Cummings et al. 2007; Bidwell et al. 2009).

Drought has been a common feature of the Western Great Plains climate (Coupland 1992), where potential evaporation exceeds precipitation (Sims and Risser 2000). Many long-term studies found weather to be a more important influence on vegetation response than grazing or burning treatments (Teague and Dowhower 2003; Teague et al. 2004, 2010; Heitschmidt et al. 2005; White and Loftin 2000; Vermeire et al. 2008, 2014). Modeling efforts by Bradford and Lauenroth (2006) indicate that climate is a stronger influence on invasion of *B. tectorum* than soil characteristics. Bradley (2009) found that two of the strongest predictors of improved habitat suitability for *B. tectorum* in the Great Basin region

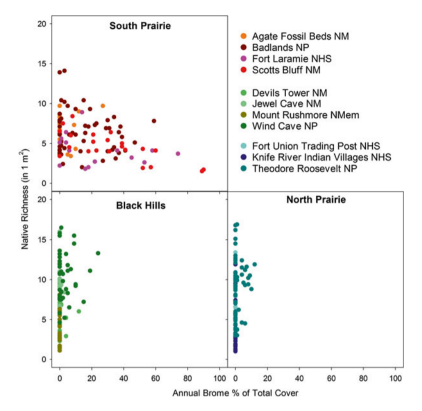


**Fig. 2.13** (a) Typical *B. tectorum* invasion of roadsides in the shortgrass steppe ecosystem, Weld County, Colorado. Photo by CS Brown. (b) Conservation Reserve Program land 3 years after seeding with native species. Increased beige vegetation on *left* side of fence is *B. tectorum*. Photo by M Vandever. (c) Little or no *Bromus* at Agate Fossil Beds National Monument. Photo by US Geological Survey. (d) Aerial cover at this site in Scotts Bluff National Monument is 60 % *B. tectorum* and 10 % *B. arvensis*. Photo by US National Park Service

were decreased annual and summer precipitation. This suggests that the normal pattern of summer precipitation in the Western Great Plains creates conditions unfavorable to *Bromus* and drought could improve their success. However, it does not appear that drought alone (see Van Dyne 1975; Munson and Lauenroth 2009; Moran et al. 2014) or in combination with grazing (Heitschmidt et al. 1999, 2005; Eneboe et al. 2002) has reduced resistance of these grasslands to invasion by *Bromus*.

Grazing of ungulates has always been important in the Great Plains, which was once inhabited by 2–4 million bison (*Bison bison*) and other grazing and browsing ungulates (Van Dyne 1975). As a result, this ecoregion is resilient to grazing (Vermeire et al. 2008). The abundances of *Bromus* and other weedy plants can be reduced (Fig. 2.13b) (Milchunas et al. 1992, 2011; Harmoney 2007; Vermeire et al. 2008) or remain unaffected by grazing, even in combination with drought (Heitschmidt et al. 1999, 2005).

Fire is a natural disturbance in the Western Great Plains and has either no effect on or reduces aboveground productivity (Scheintaub et al. 2009). Herbaceous productivity of shortgrass steppe (Ford and Johnson 2006; Augustine and Milchunas



**Fig. 2.14** In southern Northern Great Plains Network prairie parks, plots with high cover of *Bromus* have lower native plant species richness than plots with little *Bromus* cover. NM=National Monument; NP=National Park; NHS=National Historic Site; NMem=National Memorial. (Courtesy of the National Park Service, Northern Great Plains Network Inventory and Monitoring Program)

2009; Augustine 2010), northern mixed-grass prairie (Whisenant and Uresk 1990; Vermeire et al. 2011), and southern mixed-grass prairie (Whisenant et al. 1984; Ford and White 2007; Teague et al. 2008, 2010; Vermeire et al. 2014) may be unaffected by fire or can recover productivity and species composition within 6 months to 2 years of burning, depending on precipitation amounts during the recovery period (Teague et al. 2008; Augustine and Milchunas 2009). Grazing is often reported not to alter the effects of fire (Augustine et al. 2010; Vermeire et al. 2014). Not only do these ecosystems have high resilience to fire, but fire either reduces (Whisenant and Uresk 1990; Scheintaub et al. 2009; Teague et al. 2010; Vermeire et al. 2011, 2014) or does not affect (Teague and Dowhower 2003; Augustine and Milchunas 2009; Augustine et al. 2014) abundance of *Bromus*. However, in some cases when *Bromus* were initially reduced by spring burning, they increased in abundance the second year after the burn (Whisenant et al. 1984).

### 2.5.2 Ecosystem Threats to Western Great Plains from Bromus

Although several *Bromus* are considered invasive in the Western Great Plains, they are neither as widespread nor have they had the extensive, ecosystem altering effects documented in some other ecoregions. Even though the Western Great Plains is relatively resistant and resilient to invasion by *Bromus*, invasions can effect change in some circumstance. For example, *Bromus* can reduce the growth of native plants in northern mixed-grass prairie (Haferkamp et al. 1997, 1998; Ogle et al. 2003). Ogle et al. (2003) found *Bromus*-dominated northern mixed-grass prairie had lower above- and belowground biomass, slower decomposition, and sometimes greater litter than native vegetation. Modeling of *B. arvensis* and *B. tectorum* in northern mixed-grass prairie found non-significantly greater carbon storage with high *Bromus* cover, but the differences continued to diverge after 50 years and may become large in the future (Ogle et al. 2004).

Resilience of ecosystems to disturbance and resistance to invasion can be altered when historically normal patterns of disturbance change (Germino et al. 2015). The absence rather than presence of grazing, fire, or both represents disturbances for Western Great Plains grasslands (Milchunas et al. 1992; Haferkamp et al. 1993; Adler and Lauenroth 2000; Brockway et al. 2002; Ford and Johnson 2006; Vermeire et al. 2011) and may facilitate invasion by *Bromus* (Milchunas et al. 1992, 2011). Fire frequencies in the Western Great Plains may increase as a result of accumulation of fuels due to fire suppression (Brockway et al. 2002; Garfin et al. 2014) and increased lightning strikes due to climate warming (Romps et al. 2014).

As explained above, weather is an important determinant of productivity and composition of Western Great Plains grasslands. Thus, changes in climate may influence resistance of Western Great Plains ecosystems to *Bromus* invasion (Fig. 2.12). More frequent high temperatures and extreme heat are predicted for the region (Walsh et al. 2014). *Bromus tectorum* reportedly increases with temperature in northern mixed-grass prairie (Blumenthal et al., unpublished data). Longer periods between storms are predicted for much of Texas and Oklahoma (Walsh et al. 2014), which may decrease its resilience to disturbance and resistance to *Bromus* invasion (Fig. 2.12). Models that assume continued increases in heat trapping greenhouse gases predict that winter and spring precipitation and days with heavy rain will increase in the northern part of the Great Plains (Walsh et al. 2014). Changes such as these are not likely to reduce resilience to disturbance and may increase resilience in more water-limited shortgrass steppe (Fig. 2.12).

The drought in the early 2000s resulted in conditions similar to predicted future climates (Moran et al. 2014). Western Great Plains grasslands had linear responses to current year total precipitation and previous year productivity (Moran et al. 2014). No new species assemblages (e.g., dominance of *Bromus*) emerged in response to the drought (Moran et al. 2014). The system appears to be buffered against changes predicted in the next 50 years (Symstad et al. 2014), but it remains to be seen how far the climate system can be pushed before the biotic components of the ecosystem can no longer respond sufficiently to be sustained.

# 2.5.3 Management Strategies for Bromus in Western Great Plains

*Bromus* invasions can be prevented and minimized throughout the ecoregion by conserving intact grasslands and applying reasonable levels of grazing and burning to maintain their integrity and resistance to invasion. The Western Great Plains ecosystems are currently most threatened by large-scale disturbances such as conversion to cultivated agriculture, urban development, and oil and gas exploration and development, all of which may foster *Bromus* dominance.

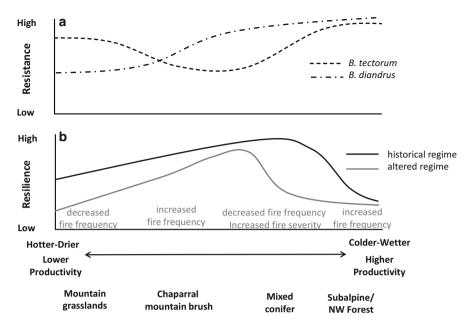
Management beyond good stewardship of the ecosystem can focus first on containing and then eliminating *Bromus* where it occurs. Herbicides that control *Bromus* with minimal negative effects on native species can be effectively used alone (Hewlett et al. 1981; Haferkamp et al. 2001) or in combination with burning (Masters et al. 1992; Calo et al. 2012). *Bromus* populations can be managed by reducing seed production through well-timed grazing, mowing, or burning (e.g., Fig. 2.13b) (Vermeire et al. 2008; Milchunas et al. 2011) or application of growth regulator herbicides used for broadleaf weed control (Rinella et al. 2010a, b, 2013).

### 2.6 Western Forests

The Western Forests range from the Marine West-Coast Forest to the Northwestern Forested Mountains Level I EPA ecoregions and are characterized by a very diverse range of environmental conditions. Collectively, they represent some of the coolest and wettest conditions in the western United States (Fig. 2.3). Soil temperature regimes are generally cryic or frigid in the east and north, but can also be mesic and even thermic to the west and south. These conditions combined with high precipitation are conducive to the growth of forests with relatively high canopy and surface cover of perennial vegetation and high accumulations of litter and duff. Shrublands or grasslands occur as early successional patches following disturbances that remove forest canopy cover, and where productivity levels are relatively low due to shallower soils or warmer and drier soil temperature and moisture regimes (Fig. 2.15).

Bromus species are generally not considered among the highest priority invasive plants in the Western Forests. In fact, one recent review of the impacts of exotic invasive species in US forests did not even mention Bromus (Moser et al. 2009). The exception are patches of chaparral and grasslands (Fig. 2.15) within ponderosa pine and Jeffrey pine mixed conifer forests (Pierson and Mack 1990; Keeley and McGinnis 2007). In these situations, soils are more frequently exposed to sunlight and warmer and drier conditions facilitate establishment, growth, and reproduction of Bromus. Bromus tectorum is the most widespread and abundant Bromus in these ecoregions, but B. diandrus can also occur within grassland and chaparral forest openings.

44 M.L. Brooks et al.



**Fig. 2.15** Hypothetical (a) resistance to common *Bromus* species and (b) resilience to historical and altered fire regimes for the Western Forests ecoregion (primary altered regime characteristic labeled in *gray*)

# 2.6.1 Invasion Potential of Western Forests by Bromus

Forests with cool to cold temperature regimes have low climatic suitability to *Bromus*. Also, forests with high amounts of canopy cover that accumulate large amounts of litter and duff in the understory and create low light conditions at the soil surface have limited establishment and growth of annual plant species. Drier and warmer mixed conifer, Jeffrey pine, and ponderosa pine forests that have less canopy cover and lower litter and duff accumulation are generally less resistant to *Bromus* than subalpine and northwestern forests due to lower levels of productivity and higher frequency of fire (Fig. 2.15).

Fire events that remove forest canopy, especially high severity fire, are often associated with decreased resistance to *Bromus* in climatically suitable forests (Crawford et al. 2001; Keeley and McGinnis 2007; McGlone et al. 2009; Fornwalt et al. 2010; Kerns et al. 2011). Decreased resistance is partly due to increased light availability at the soil surface, but also may be affected by increased availability of soil nutrients. Biomass of *B. tectorum* beneath *Pinus ponderosa* Lawson and C. Lawson (ponderosa pine) trees can be limited by low N availability. Fire can increase available nutrients due to deposition of ash onto the soil surface, release of available P and N from organic matter, and decomposition of belowground biomass

and result in increases in *B. tectorum* growth and reproduction (Covington and Sackett 1992; Monleon et al. 1997; Gundale et al. 2005, 2008). Also, charcoal generated by fire may adsorb phenolic compounds from pine litter inputs which can otherwise impede understory plant growth (Gundale and DeLuca 2006, 2007), and increase nitrification rates, further increasing the amount of plant available N in the soil following fire (DeLuca et al. 2006).

High severity human disturbances such as clear-cut logging can similarly decrease resistance of landscapes. Recruitment, survival, and reproduction of *B. tectorum* can all be higher in clear-cuts across a range of western forest types including those dominated by *P. ponderosa*, *Pseudotsuga menziesii* (Mirb.) Franco (Douglas-fir), *Abies grandis* (Douglas ex D. Don) Lindl. (grand fir), and *Thuja plicata* Donn ex D. Don (western redcedar) (Pierson and Mack 1990). Lower severity understory thinning can also decrease resistance to *B. tectorum* in these forest types, but only where significant understory cover of *B. tectorum* was present before thinning. Even lower severity human disturbances such as cattle grazing occurring at light intensity and short duration can lead to increased cover of *B. tectorum* in ponderosa pine forests under drought conditions (Sorensen and McGlone 2010), possibly due to competitive release of other vegetation such as *Elymus elymoides* (Raf.) Swezey (bottlebrush squirreltail) and *P. smithii* caused by preferential grazing by cattle.

Removal of litter and coarse woody debris can increase exposure of mineral soil and the probability that *Bromus* seeds can make contact with the soil and germinate (Pierson and Mack 1990; Keeley and McGinnis 2007; Gundale et al. 2008). This is partly due to the litter creating a physical barrier preventing seeds from coming into contact with mineral soil, but also may be due to phenolytic compounds in new pine litter that can inhibit germination.

Lower temperature limits may limit the upper elevation sites attainable by *Bromus* species (Fig. 2.2a), but climate change is increasingly recognized as a mechanism by which high elevation habitats such as Western Forests may become increasingly more invasible (Pauchard et al. 2009). There is evidence suggesting that *B. tectorum* already exhibits the ability to adapt to higher elevation sites (Rice and Mack 1991; Leger et al. 2009), and climatic warming may further reduce the resistance of these habitats to *Bromus* invasion (Compagnoni and Adler 2014). Rising CO<sub>2</sub> has also been shown to increase *B. tectorum* productivity through increased water use efficiency in Cold Deserts shrublands (Smith et al. 1987), and the operative mechanism could further promote growth at higher elevations as they warm.

# 2.6.2 Ecosystem Threats to Western Forests from Bromus

It is unclear how much of a threat *Bromus* pose to Western Forests compared to other ecoregions of the western United States. No doubt *Bromus* can have some of the same effects documented for other ecoregions, such as direct competition with

other plant species. They may even affect landscape susceptibility to fire by producing continuous fuelbeds at least where the forest canopy does not overshadow and inhibit their growth (e.g., Fig. 2.16). However, the potential for *Bromus* species to alter fire regimes (i.e., fire frequency, severity, seasonality, etc.) in Western Forests is more difficult to discern.

Many of the lower elevation forests that are most susceptible to invasion by *Bromus* (e.g., ponderosa pine) are also evolutionarily suited to fire return intervals as short as once every 2 years (reviewed by Hurteau et al. 2013). So if *Bromus* species, *B. tectorum* in particular, has the potential to increase landscape flammability and promote recurrent fire, is it reasonable to think that they might promote such short intervals that even these fire-adapted forests cannot persist? The general consensus is that they could, especially in areas where anthropogenic activities increase ignition sources and promote productivity of *Bromus*, thus enhancing their competitive suppression of regenerating forest species and their contributions to highly flammable fuelbeds (McGinnis et al. 2010; Hurteau et al. 2013). As explained above in Sect. 2.6.1, as the climate increasingly warms, resistance of Western Forests to invasion by *Bromus* will likely decrease. When this decreased resistance is coupled with increased frequency of large fires (Westerling and Bryant 2008) and length of the fire season (Westerling et al. 2006), the potential for *Bromus* to alter fire regimes in Western Forests may become more likely.



**Fig. 2.16** *Bromus tectorum* dominating a landscape 6 years after a high severity fire in a previous conifer forest stand (Star Fire, Tahoe National Forest, 2001). Note the conifer recruitment in the *upper right* side of this image despite the high cover of *B. tectorum* (photo by Jon Keeley, 22 June 2007)

## 2.6.3 Management Strategies for Bromus in Western Forests

Prevention and control of *Bromus* in the Western Forests have not be evaluated to the degree they have in other ecoregions, most notably in the Cold Deserts and Mediterranean California. Strategies developed in those regions should be generally applicable to the Western Forests, although there are some unique characteristics of Western Forests that lend themselves to approaches only possible in this ecoregion. For example, litter and duff can rapidly accumulate on the forest floor, which can suppress B. tectorum germination (Keeley and McGinnis 2007). However, some of the principles that have been applied to restore historical fire regimes where fire suppression and fuel accumulation have altered them may reduce litter and forest canopy cover, which could inadvertently improve conditions for *Bromus* species. Some of these include reducing surface and ladder fuels, decreasing crown density, and maintaining widely spaced large trees. Forest managers are challenged to find an effective balance between the restoration of historical fire regimes and the suppression of invasive plants such as Bromus, thus maximizing resilience to fire and resistance to invasion. Management of Bromus invasion should be linked with maintaining historical ecosystem structure and function, which may entail suppressing or facilitating fire, or reducing fire severity (e.g., by pre-fire fuels manipulations), as dictated by the historical fire regimes and current condition of local areas.

## 2.7 Management Implications

This chapter explained that invasion potential, ecosystem threats, and management strategies for *Bromus* vary both within and among ecoregions and vegetation types, and among *Bromus* species. In particular, it should not be assumed that just because a particular *Bromus* is documented to pose a significant threat in one ecoregion or vegetation type, that other *Bromus* will pose similar threats to other ecoregions or vegetation types. Variation among ecoregions and vegetation types in ecosystem resistance to *Bromus* invasion, and resilience to disturbance, can be used to explain the conditions associated with high probability of ecosystem impacts such as an altered fire regime caused by a grass/fire cycle.

A strategic, landscape-scale management approach that includes protection, prevention, and restoration (e.g., Brooks and Chambers 2011; Chambers et al. 2014a) can be used to better focus management activities across the environmental/productivity gradients in the western United States. Protection focuses on maintaining or increasing the resilience and resistance of ecosystems with high conservation value by eliminating or minimizing current and future stressors. Protection is a viable strategy for intact communities within ecosystem types that have inherently low resistance and resilience such as blackbrush ecosystems in the Warm Deserts and salt desert and warmer Wyoming big sagebrush ecosystems in the Cold Deserts. Protection may also be a viable strategy for critical habitat for threatened and endangered species that are sensitive to *Bromus* invasions and their associated impacts.

Prevention involves increasing both resilience and resistance of ecosystems that have not crossed thresholds, but that exhibit declining conditions and are at risk of high severity fire and invasion (Miller et al. 2013). The focus is on ecosystems with higher resilience, like cooler and moister *A. tridentata* ecosystems in the Cold Deserts (Fig. 2.10) and mixed conifer ecosystems of the Western Forests (Fig. 2.15). Management objectives are to improve or maintain ecosystem attributes and processes by reducing woody species dominance and promoting a functionally diverse community of shrubs, perennial grasses, and forbs.

Restoration involves increasing resilience and resistance of disturbed, degraded, or invaded areas by reestablishing functionally diverse plant communities. Integrated management strategies that include using herbicides to control/suppress *Bromus* and seeding perennial herbaceous and shrub species adapted to local conditions may be used to restore priority areas dominated by *Bromus* and other annual invaders (Monaco and Sheley 2012). Restoration activities that target relatively moister sites are likely to be more successful, and even then repeated intervention may be required. As the climate warms, it may be necessary to consider transformative restoration in areas with novel climates in which species are established that are climatically suitable but that have not occurred on a given landscape historically (Bradley et al. 2010).

### 2.8 Research Needs

This chapter indicates that patterns of ecological resistance to *Bromus* invasions have been well documented in the western United States, especially related to presence/absence of *B. tectorum* and *B. rubens* in the Cold Deserts, Mediterranean California, and Warm Desert Ecoregions. What is less understood are the patterns of relative abundance (e.g., cover or biomass) of these and other *Bromus*, and the ecological mechanisms associated with these patterns. Also, the range of conditions associated with the grass/fire cycle have only been studied in depth within these same three ecoregions, and even within them there is limited capacity to predict when and where they will establish. A framework exists for objectively evaluating the grass/fire cycle potential (Brooks 2008) which can be systematically applied to each major vegetation types within each ecoregion to better understand when and where *Bromus* invasions pose the threat of altering fire regimes. This information is needed to more reliably predict potential future conditions of spread, dominance, and ecological impact by *Bromus* species within each ecoregion of the western United States.

The relative resistance to invasion by the dominant *Bromus* species and resilience to fire of each of five major ecoregions in the western United States were presented in Figs. 2.4, 2.7, 2.9, 2.11, and 2.14. In the introduction to this chapter, we emphasized that the degree to which scientific studies supported these models varies widely among ecoregions, and we suggested that the conservative approach would be to consider them hypothetical models. We feel confident with the scientific

support for the Cold Desert and Warm Desert models, but encourage others to develop studies to test the predictions of all five ecoregional models. The refinement and validation of these models would improve their reliability and utility in evaluating the relative potential for invasion and ecological impact of *Bromus* in the western United States.

# **Appendix**

See Table 2.2.

**Table 2.2** GrassPortal (www.grassportal.org, accessed 16 Feb 2015, Osborne et al. 2011) site localities used to construct Fig. 2.2

	B. arvensis		B. diandrus	В.	В.	B.
	(= japonicus)	B. hordeaceus	(= rigidus)	madritensis	rubens	tectorum
Arizona	82	36	108	4	377	183
California	104	979	934	220	1109	664
Colorado	59	3	1	0	1	121
Idaho	31	16	2	0	10	48
Kansas	220	7	0	0	0	266
Montana	45	14	0	0	0	77
Nebraska	50	0	0	0	0	98
Nevada	8	7	9	0	4	25
New Mexico	45	1	9	0	11	61
North Dakota	0	0	0	0	0	34
Oklahoma	36	0	0	0	0	22
Oregon	40	55	71	2	35	141
South Dakota	7	1	0	0	0	36
Texas	89	8	6	0	4	30
Utah	31	10	9	0	13	45
Washington	27	12	92	0	4	113
Wyoming	23	39	0	0	0	40
Sum	897	1188	1241	226	1568	2004
Data Provider			Dataset			
USDA PLANTS			USDA PLANTS Database			
Consortium of California Herbaria			Consortium of California Herbaria			
University of Arizona Herbarium			UA Herbarium			
Arizona State University, International Institute for Species Exploration			Arizona State University Vascular Plant Herbarium			
University of Connecticut			CONN GBIF data			
Berkeley Natural History Museums			University and Jepson Herbaria DiGIR provider			
New Mexico Biodiversity Collections Consortium			New Mexico Biodiversity Collections Consortium database			
						(continue)

(continued)

Table 2.2 (continued)

Data Provider	Dataset		
Missouri Botanical Garden	Missouri Botanical Garden		
Utah Valley State College (UVSC)	Utah Valley State College Herbarium		
University of Kansas Biodiversity Research Center	Botany Vascular Plant Collection		
Oregon State University	Vascular Plant Collection		
University of Washington Burke Museum	Vascular Plant Collection – University of Washington Herbarium (WTU)		
Bernice Pauahi Bishop Museum	Bishop Museum Natural Sciences Data		
University of Alabama Biodiversity and Systematics	Herbarium (UNA)		
Colorado State University Herbarium (CSU)	Colorado State University Herbarium		
National Museum of Natural History	NMNH Botany Collections		
Canadian Museum of Nature	Canadian Museum of Nature Herbarium		
University of Colorado Museum of Natural History	Specimen Database of Colorado Vascular Plants		
The New York Botanical Garden	Herbarium of The New York Botanical Garden		

The 17 states represent the ecoregions of the western United States (Fig. 2.1). The seven *Bromus* species are represented by state site records spanning their current distributional range as indicated by the USDA Plants database (www.plants.usda.gov, accessed 18 Feb 2015). The 19 data providers represent the original sources of information for the database used in Fig. 2.2

#### References

- Abella SR, Embrey TM, Schmid SM et al (2012) Biophysical correlates with the distribution of the invasive annual red brome (*Bromus rubens*) on a Mojave Desert landscape. Invasive Plant Sci Manag 5:47–56
- Adler PB, Lauenroth WK (2000) Livestock exclusion increases the spatial heterogeneity of vegetation in Colorado shortgrass steppe. Appl Veg Sci 3:213–222
- Allen E, Chambers JC, Nowak RS (2008) Immediate and longer-term effects of a spring prescribedburn on the soil seed bank in an encroaching semi-arid woodland. West N Am Nat 68:265–277
- Allen EB, Padgett PE, Bytnerowicz A et al (1998) Nitrogen deposition effects on coastal sage vegetation of southern California. In: Proceedings of the international symposium on air pollution and climate change effects on forest ecosystems. Gen Tech Rep PSW-GTR-166. USDA, Forest Service, Pacific Southwest Research Station, Albany, CA, pp 131–140
- Allen EB, Rao LE, Steers RJ et al (2009) Impacts of atmospheric nitrogen deposition on vegetation and soils in Joshua Tree National Park. The Mojave Desert: Ecosystem processes and sustainability. University of Nevada Press, Las Vegas, NV
- Atkinson SY, Brown CS (2015) Attributes that confer invasiveness and impacts across the large genus *Bromus* lessons from the *Bromus* REEnet database. In: Germino MJ, Chambers JC, Brown CS (eds) Exotic brome-grasses in arid and semiarid ecosystems of the western USA: causes, consequences, and management implications. Springer, New York, NY (Chapter 6)
- Augustine DJ (2010) Spatial versus temporal variation in precipitation in a semiarid ecosystem. Landscape Ecol 25:913–925

- Augustine DJ, Brewer P, Blumenthal DM et al (2014) Prescribed fire, soil inorganic nitrogen dynamics, and plant responses in a semiarid grassland. J Arid Environ 104:59–66
- Augustine DJ, Derner JD, Milchunas DG (2010) Prescribed fire, grazing, and herbaceous plant production in shortgrass steppe. Rangel Ecol Manag 63:317–323
- Augustine DJ, Milchunas DG (2009) Vegetation responses to prescribed burning of grazed short-grass steppe. Rangel Ecol Manag 62:89–97
- Balch JK, Bradley BA, D'Antonio CM et al (2013) Introduced annual grass increases regional fire activity across the arid western USA (1980–2009). Glob Change Biol 19:173–183
- Beyers JL (2004) Postfire seeding for erosion control: Effectiveness and impacts on native plant communities. Conserv Biol 18:947–956
- Bidwell TG, Engle DM, Moseley ME et al (2009) Invasion of Oklahoma rangelands and forests by eastern redcedar and ashe juniper, vol E947. Oklahoma Cooperative Extension Service, Stillwater, OK
- Bidwell TG, Masters RE, Tyrl RJ (2004) A checklist of prairie, shrubland, and forest understory plants of Oklahoma: characteristics and value to deer, auail, turkey, and cattle. Fact sheet NREM-2872. Oklahoma Cooperative Extension Service, Stillwater, OK
- Booth MS, Caldwell MM, Stark JM (2003) Overlapping resource use in three Great Basin species: implications for community invasibility and vegetation dynamics. J Ecol 91:36–48
- Bradford JB, Lauenroth WK (2006) Controls over invasion of *Bromus tectorum*: the importance of climate, soil, disturbance and seed availability. J Veg Sci 17:693–704
- 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
- Bradley BA, Blumenthal DM, Wilcove DS et al (2010) Predicting plant invasions in an era of global change. Trends Ecol Evol 25:310–318
- Brockway DG, Gatewood RG, Paris RB (2002) Restoring fire as an ecological process in short-grass prairie ecosystems: initial effects of prescribed burning during the dormant and growing seasons. J Environ Manag 65:135–152
- Brooks ML (1999) Habitat invasibility and dominance by alien annual plants in the western Mojave Desert. Biol Inv 1:325–337
- Brooks ML (2000) Competition between alien annual grasses and native annual plants in the Mojave Desert. Am Midl Nat 144:92–108
- Brooks ML (2003) Effects of increased soil nitrogen on the dominance of alien annual plants in the Mojave Desert. J Appl Ecol 40:344–353
- Brooks ML (2008) Plant invasions and fire regimes. In: Zouhar K, Kapler-Smith J, Sutherland S et al (eds) Wildland fire in ecosystems: fire and nonnative invasive plants. Gen Tech Rep RMRS-GTR-42-volume 6. USDA, Forest Service, Rocky Mountain Research Station, Ogden, UT, pp 33–46
- Brooks ML (2009) Spatial and temporal distribution of non-native plants in upland areas of the Mojave Desert. In: Webb RH, Fenstermaker LF, Heaton JS et al (eds) The Mojave Desert: ecosystem processes and sustainability. University of Nevada Press, Reno, NV, pp 101–124
- Brooks ML (2012) Effects of high fire frequency in creosote bush scrub vegetation of the Mojave Desert. Int J Wildl Fire 21:61–68
- Brooks ML, Berry KH (2006) Dominance and environmental correlates of alien annual plants in the Mojave Desert. J Arid Environ 67:100–124
- Brooks M, Chambers J (2011) Invasive plants that alter fire regimes in the deserts of North America. Rangel Ecol Manag 64:431–438
- Brooks ML, Chambers JC, McKinley RA (2013) Fire history, effects, and management in southern Nevada. In: Chambers JC, Brooks ML, Pendleton BK et al (eds) The southern Nevada agency partnership science and research synthesis: science to support land management in southern Nevada. Gen Tech Rep RMRS-GTR-303. USDA, Forest Service, Rocky Mountain Research Station, Fort Collins, CO, pp 75–96
- Brooks ML, D'Antonio CM, Richardson DM et al (2004) Effects of invasive alien plants on fire regimes. Bioscience 54:677–688

- Brooks ML, Esque TC (2002) Alien annual plants and wildfire in desert tortoise habitat: status, ecological effects, and management. Chelonian Conserv Biol 4:330–340
- Brooks ML, Esque TC, Duck T (2007) Creosotebush, blackbrush, and interior chaparral shrublands. In: Hood S, Miller M (eds) Fire ecology and management of the major ecosystems of Southern Utah. Gen Tech Rep RMRS-GTR-202. USDA, Forest Service, Rocky Mountain Research Station. Fort Collins, CO, pp 97–110
- Brooks ML, Matchett JR (2006) Spatial and temporal patterns of wildfires in the Mojave Desert. 1980–2004. J Arid Environ 67:148–164
- Brooks ML, Minnich RA (2006) Southeastern Deserts Bioregion. In: Sugihara NG, van Wagtendonk JW, Shaffer KE et al (eds) Fire in California's Ecosystems. US Press, Berkeley, pp 391–414
- Brooks ML, Pyke D (2001) Invasive plants and fire in the deserts of North America. In: Galley K, Wilson T (eds) Proceedings of the invasive species workshop: the role of fire in the control and spread of invasive species. Fire conference 2000: the first national congress on fire ecology, prevention and management. Miscellaneous publications No. 11, Tall Timbers Research Station, Tallahassee, FL, pp 1–14
- Brown CS, Rice KJ (2010) Effects of belowground resource use complementarity on invasion of constructed grassland plant communities. Biol Invasions 12:1319–1334
- Bykova O, Sage RF (2012) Winter cold tolerance and the geographic range separation of *Bromus tectorum* and *Bromus rubens*, two severe invasive species in North America. Glob Change Biol 18(12):3654–3663
- Callaway RM, Davis FW (1993) Vegetation dynamics, fire, and the physical environment in coastal central California. Ecology 74:1567–1578
- Calo A, Brause S, Jones S (2012) Integrated treatment with a prescribed burn and post-emergent herbicide demonstrates initial success in managing cheatgrass in a northern Colorado natural area. Nat Area J 32:300–304
- Chambers JC, Germino MJ, Belnap J et al (2015) Plant community resistance to invasion by *Bromus s*pecies the roles of community attributes, *Bromus i*nteractions with plant communities, and *Bromus* traits. In: Germino MJ, Chambers JC, Brown CS (eds) Exotic brome-grasses in arid and semiarid ecosystems of the Western USA: causes, consequences, and management implications. Springer, New York, NY (Chapter 10)
- Chambers JC, Bradley BA, Brown CS et al (2014a) Resilience to stress and disturbance, and resistance to *Bromus tectorum* L. invasion in cold desert shrublands of western North America. Ecosystems 17:360–375
- Chambers JC, Miller RF, Board DI et al (2014b) Resilience and resistance of sagebrush ecosystems: implications for state and transition models and management treatments. Rangel Ecol Manag 67:440–454
- Chambers JC, Pyke DA, Maestas JD et al (2014c) Using resistance and resilience concepts to reduce impacts of invasive annual grasses and altered fire regimes on the sagebrush ecosystem and greater sage-grouse: a strategic multi-scale approach. Gen Tech Rep RMRS-GTR-326. USDA, Forest Service, Rocky Mountain Research Station, Fort Collins, CO, p 73
- Chambers JC, Roundy BA, Blank RR et al (2007) What makes Great Basin sagebrush ecosystems invasible by *Bromus tectorum*? Ecol Monogr 77:117–145
- Clayton W, Vorontsova MS, Harman KT, Williamson H (2006) GrassBase The Online World Grass Flora. http://www.kew.org/data/grasses-db.html. Accessed 20 Mar 2014
- Compagnoni A, Adler PB (2014) Warming, soil moisture, and loss of snow increase *Bromus tectorum*'s population growth. Elem Sci Anth 2:000020
- Condon L, Weisberg PJ, Chambers JC (2011) Abiotic and biotic influences on *Bromus tectorum* invasion and *Artemisia tridentata* recovery after fire. Int J Wildl Fire 20:597–604
- Corbin J, D'Antonio CM (2004) Competition between native perennial and exotic annual grasses: Implications for an historical invasion. Ecology 85:1273–1283

- Corbin JD, D'Antonio CM, Bainbridge SJ (2004) Tipping the balance in restoration of native plants. In: Gordon MS, Bartol SM (eds.) Experimental Approaches to conservation biology. UC Press, Berkeley, pp 154–179
- Coupland R (1992) Mixed prairie. In: Coupland R (ed) Ecosystems of the World. Elsevier, Netherlands, pp 151–182
- Covington WW, Sackett SS (1992) Soil mineral nitrogen changes following prescribed burning in ponderosa pine. Forest Ecol Manag 54:175–191
- Crawford JA, Wahren CHA, Kyle S et al (2001) Responses of exotic plant species to fires in Pinus ponderosa forests in northern Arizona. J Veg Sci 12:261–268
- Cummings DC, Bidwell TG, Medlin CR et al (2007) Ecology and management of Sericea lespedeza. Division of Agricultural Sciences and Natural Resources, Oklahoma State University
- Cushman JH, Tierney TA, Hinds JM (2004) Variable effects of feral pig disturbances on native and exotic plants in a California grassland. Ecol Appl 14:1746–1756
- D'Antonio CM, Thomsen M (2004) Ecological resistance in theory and practice. Weed Technol 18:1572–1577
- D'Antonio CM, Vitousek PM (1992) Biol invasions by exotic grasses, the grass fire cycle, and global change. Ann 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
- Davis MA, Grime JP, Thompson K (2000) Fluctuating resources in plant communities: a general theory of invasibility. J Ecol 88:528–534
- DeFalco LA, Bryla DR, Smith-Longozo V et al (2003) Are Mojave Desert annual species equal? Resource acquisition and allocation for the invasive grass *Bromus madritensis* subsp. *rubens* (Poaceae) and two native species. Am J Bot 90:1045–1053
- DeFalco LA, Fernandez GCJ, Nowak RS (2007) Variation in the establishment of a non-native annual grass influences competitive interactions with Mojave Desert perennials. Biol Invasions 9:293–307
- DeLuca TH, MacKenziea MD, Gundalea MJ et al (2006) Wildfire-produced charcoal directly influences nitrogen cycling in forest ecosystems. Soil Sci Soc Am J 70:448–453
- DeSimone SA, Zedler PH (1999) Shrub seedling recruitment in unburned Californian coastal sage scrub and adjacent grassland. Ecology 80:2018–2032
- Dettweiler-Robinson E, Bakker JD, Grace JB (2013) Controls of biological soil crust cover and composition shift with succession in sagebrush shrub-steppe. J Arid Environ 94:96–104
- DiTomaso JM, Brooks ML, Allen EB et al (2006) Control of invasive weeds with prescribed burning. Weed Technol 20:535–548
- Dudley TL (2009) Invasive plants in Mojave Desert riparian areas. In: Webb RH, Fenstermaker LF, Heaton JS et al (eds) The Mojave Desert: ecosystem processes and sustainability. University of Nevada Press, Reno, NV, pp 125–155
- Earnst SL, Holmes AL (2012) Bird-habitat relationships in interior Columbia Basin shrubsteppe. Condor 114:15–29
- Eiswerth ME, Shonkwiler JS (2006) Examining post-wildfire reseeding on arid rangeland: a multivariate tobit modelling approach. Ecol Model 192:286–298
- Eliason SA, Allen E (1997) Exotic grass competition in suppressing native shrubland reestablishment. Restor Ecol 5:245–255
- Eneboe EJ, Sowell BF, Heitschmidt RK et al (2002) Drought and grazing: IV. Blue grama and western wheatgrass. J Range Manag 55:197–203
- Folke C, Carpenter S, Walker B et al (2004) Regime shifts, resilience, and biodiversity in ecosystem management. Ann Rev Ecol Syst 35:557–581
- Ford P, White CS (2007) Effects of dormant-season fire at three different fire frequencies in short-grass steppe of the southern Great Plains. In: Masters R, Galley KEM (eds) 23rd Tall timbers fire ecology conference: fire in grassland and shrubland ecosystems. Tall Timbers Research Station, Tallahassee, FL, p 71

- Ford PL, Johnson GV (2006) Effects of dormant vs. growing-season fire in shortgrass steppe: biological soil crust and perennial grass responses. J Arid Environ 67:1–14
- 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 12:2683–2695
- Freudenberger DO, Fish BE, Keeley JE (1987) Distribution and stability of grasslands in the Los Angeles Basin. Bull South Calif Acad Sci 86:13–26
- Garfin G, Franco G, Blanco H et al (2014) Sounthwest. In: Melillo J, Richmond TC (eds) Climate change impacts in the United States: the third national climate assessment. US global change research program, pp 461–486
- Germino MJ, Belnap J, Stark JM et al (2015) Ecosystem impacts of exotic annual invaders in the genus *Bromus*. In: Germino MJ, Chambers JC, Brown CS (eds) Exotic brome-grasses in arid and semiarid ecosystems of the Western USA: causes, consequences, and management implications. Springer, New York, NY (Chapter 3)
- Gundale MJ, DeLuca TH (2006) Temperature and source material influence ecological attributes of ponderosa pine and Douglas-fir charcoal. Forest Ecol Manag 231:86–93
- Gundale MJ, DeLuca TH (2007) Charcoal effects on soil solution chemistry and growth of *Koeleria macrantha* in the ponderosa pine/Douglas-fir ecosystem. Biol Fertil Soils 43:303–311
- Gundale MJ, DeLuca TH, Fiedler CE et al (2005) Restoration management in a Montana ponderosa pine forest: effects on soil physical, chemical, and biological properties. Forest Ecol Manag 213:25–38
- Gundale MJ, Sutherland S, DeLuca TH (2008) Fire, native species, and soil resource interactions influence the spatio-temporal invasion pattern of *Bromus tectorum*. Ecography 31:201–210
- Haferkamp M, Volesky J, Borman M et al (1993) Effects of mechanical treatments and climatic factors on the productivity of northern great-plains rangeland. J Range Manag 46:346–350
- Haferkamp MR, Grings EE, Heitschmidt RK et al (2001) Suppression of annual bromes impacts rangeland: animal responses. J Range Manag 54:663–668
- Haferkamp MR, Heitschmidt RK, Karl MG (1997) Influence of Japanese brome on western wheat-grass yield. J Range Manag 50:44–50
- Haferkamp MR, Heitschmidt RK, Karl MG (1998) Clipping and Japanese brome reduce western wheatgrass standing crop. J Range Manag 51:692–698
- Haidinger TL, Keeley JE (1993) Role of high fire frequency in destruction of mixed chaparral. Madroño 40:141–147
- Harmoney KR (2007) Grazing and burning Japanese brome (*Bromus japonicus*) on mixed grass rangelands. Rangel Ecol Manag 60:479–486
- Hassan M, West N (1986) Dynamics of soil seed pools in burned and unburned sagebrush semideserts. Ecology 67:269–272
- Haubensak KA, D'Antonio CM, Saundra Embry S et al (2014) A comparison of *Bromus tectorum* growth and mycorrhizal colonization in salt desert vs. sagebrush habitats. Rangel Ecol Manag 67:275–284
- Haubensak K, D'Antonio C, Wixon D (2009) Effects of fire and environmental variables on plant structure and composition in grazed salt desert shrublands of the Great Basin (USA). J Arid Environ 73:643–650
- Heitschmidt R, Grings E, Haferkamp M et al (1995) Herbage dynamics on 2 Northern Great Plains range sites. J Range Manag 48:211–217
- Heitschmidt RK, Haferkamp MR, Karl MG et al (1999) Drought and grazing: I. Effects on quantity of forage produced. J Range Manag 52:440–446
- Heitschmidt RK, Klement KD, Haferkamp MR (2005) Interactive effects of drought and grazing on Northern Great Plains rangelands. Rangel Ecol Manag 58:11–19
- Hernandez RR, Sandquist DR (2011) Disturbance of biological soil crust increases emergence of exotic vascular plants in California sage scrub. Plant Ecol 212:1709–1721
- Hewlett D, Johnson J, Butterfield R et al (1981) Japanese brome response to atrazine in combination with nitrogen fertilizer in the mixed prairie. J Range Manag 34:22–25

- Hobbs RJ, Mooney HA (1995) Spatial and temporal variability in California annual grassland: results from a long term study. J Veg Sci 6:43–56
- Holling CS (1973) Resilience and stability of ecological systems. Ann Rev Ecol Syst 4:1-2
- Holmgren R (1960) Inspection tour of old blackbrush burns in BLM District N-5, southern Nevada. USDA, Forest Service, Intermountain Forest and Range Experiment Station, Reno Research Center, Reno, NV
- Hooker TD, Stark JM, Norton U et al (2008) Distribution of ecosystem C and N within contrasting vegetation types in a semiarid rangeland in the Great Basin, USA. Biogeochemistry 90:291–308
- Huenneke LF, Hamburg SP, Koide R et al (1990) Effects of soil resources on plant invasion and community structure in Californian serpentine grassland. Ecology 71:478–491
- Humphrey LD, Schupp EW (2001) Seed banks of *Bromus tectorum*-dominated communities in the Great Basin. West N Am Nat 61:85–92
- Humphrey RR (1958) The desert grassland. Bot Rev 24:193–253
- Hurteau MD, Bradford JB, Fulé PZ et al (2013) Climate change, fire management, and ecological services in the southwestern US. Forest Ecol Manag 327:280–289
- Jackson RD, Bartolome JW (2002) A state-transition approach to understanding non-equilibrium plant community dynamics in California grasslands. Plant Ecol 162:49–65
- Jackson RD, Bartolome JW (2007) Grazing ecology of California grasslands. In: Stromberg M, Corbin J, D'Antonio CM (eds) California grasslands: ecology and management. UC Press, Berkeley, CA, pp 197–206
- James JJ, Drenovsky RE, Monaco TA et al (2011) Managing soil nitrogen to restore annual grass-infested plant communities: effective strategy or incomplete framework? Ecol Appl 21:490–502
- Karl MG, Heitschmidt RK, Haferkamp MR (1999) Vegetation biomass dynamics and patterns of sexual reproduction in a northern mixed-grass prairie. Am Midl Nat 141:227–237
- Keeler-Wolf T, Evens JM, Solomeshsch AI et al (2007) Community classification and nomenclature. In: Stromberg M, Corbin J, D'Antonio CM (eds) California grasslands: ecology and management. UC Press, Berkeley, CA, pp 21–36
- Keeley JE (1990) The California valley grassland. In: Schoenherr AA (ed) Endangered plant communities of southern California. Southern California Botanists, Fullerton, CA, pp 2–23
- Keeley JE, Bond WJ, Bradstock RA et al (2012) Fire in Mediterranean ecosystems: ecology, evolution and management. Cambridge University Press, New York, NY, p 528
- Keeley JE, Brennan TJ (2012) Fire-driven alien invasion in a fire-adapted ecosystem. Oecologia 169:1043–1052
- Keeley JE, Brennan T (2015) Research on the effects of wildland fire and fire management on federally listed species and their habitats on San Clemente Island, California. Unpublished report submitted to the US Navy
- Keeley JE, Brennan T, Pfaff AH (2008) Fire severity and ecosystem responses following crown fires in California shrublands. Ecol Appl 18:1530–1546
- 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, Netherlands, pp 193–221
- Keeley JE, McGinnis TW (2007) Impact of prescribed fire and other factors on cheatgrass persistence in a Sierra Nevada ponderosa pine forest. Int J Wildl Fire 16:96–106
- Kerns BK, Buonopane M, Thies WG et al (2011) Reintroducing fire into a ponderosa pine forest with and without cattle grazing. Ecosphere 2:art59
- Klinger R, Brooks ML, Frakes N et al (2011a) Establishment of aerial seeding treatments in Blackbrush and Pinyon-Juniper sites following the 2005 Southern Nevada Complex. In: Derasaray L, Frakes N, Gentilcore D et al (eds) Southern Nevada complex emergency stabilization and rehabilitation final report. USDI, Bureau of Land Management, Ely, NV, pp 92–117 (Chapter 5)
- Klinger R, Brooks ML, Frakes N et al (2011b) Vegetation trends following the 2005 Southern Nevada Complex Fire. In: Derasaray L, Frakes N, Gentilcore D et al (eds) Southern Nevada

- complex emergency stabilization and rehabilitation final report. USDI, Bureau of Land Management, Ely, NV, pp 118–194 (Chapter 6)
- Knick ST, Connelly JW (2011) Greater sage-grouse: ecology and conservation of a landscape species and its habitats, vol 38, Studies in Avian Biology. UC Press, Berkeley, CA
- Knick ST, Hanser SE, Miller RF et al (2011) Ecological pathways of land use in sagebrush. In: Knick ST, Connelly JW (eds) Greater sage-grouse: ecology and conservation of a landscape species and its habitats, vol 38, Studies in Avian Biology. UC Press, Berkeley, CA, pp 203–251 (Chapter 12)
- Knutson KC, Pyke DA, Wirth TA et al (2014) Long-term effects of seeding after wildfire on vegetation in Great Basin shrubland ecosystems. J Appl Ecol 51:1414–1424
- Kotanen PM, Bergelson J, Hazlett DL (1998) Habitats of native and exotic plants in Colorado shortgrass steppe: a comparative approach. Can J Bot 76:664–672
- Kulmatiski A, Beard KH, Stark JM (2006) Exotic plant communities shift water-use timing in a shrub-steppe ecosystem. Plant Soil 288:271–284
- Kulpa SM, Leger EA, Espeland EK et al (2012) Postfire seeding and plant community recovery in the Great Basin. Rangel Ecol Manag 65(2):171–181
- Lauenroth WK (2008) Vegetation of the Shortgrass Steppe. In: Lauenroth WK, Burke IC (eds) Ecology of the shortgrass steppe: a longterm perspective. Oxford University Press, New York, NY, p 7083
- Lauenroth W, Milchunas DG (1992) Short-grass steppe. In: Coupland R (ed) Ecosystems of the world. Elsevier, Netherlands, pp 183–226
- Lauenroth W, Burke IC, Morgan JA (2008) The shortgrass steppe. In: Lauenroth W, Burke IC (eds) Ecology of the shortgrass steppe: a long-term perspective. Oxford University Press, New York, NY, pp 3–13
- Leger EA, Espeland EK, Merrill KR et al (2009) Genetic variation and local adaptation at a cheatgrass (*Bromus tectorum*) invasion edge in western Nevada. Mol Ecol 18:4366–4379
- Lovich JE, Bainbridge D (1999) Anthropogenic degradation of the southern California desert ecosystem and prospects for natural recovery and restoration. Environ Manag 24:309–326
- Lovtang SCP, Riegel GM (2012) Predicting the occurrence of downy brome (*Bromus tectorum*) in central Oregon. Invasive Plant Sci Manag 5:83–91
- Lulow ME (2006) Invasion by non-native annual grasses: the importance of species biomass, composition and time among California native grasses of the Central Valley. Restor Ecol 14:616–626
- Mack RN, Pyke DA (1983) The demography of *Bromus tectorum*: variation in time and space. J Ecol 71:69–93
- Maron JL, Connors P (1996) A native nitrogen-fixing shrub facilitates weed invasion. Oecologia 105:302–312
- Maron JL, Jeffries RL (1999) Bush lupine mortality, altered resource availability and alternative vegetation states. Ecology 80:443–454
- Marty JT, Collinge SK, Rice KJ (2005) Responses of a remnant native bunchgrass population to grazing, burning and climatic variation. Plant Ecol 181:101–112
- Masters RA, Vogel KP, Mitchell RB (1992) Response of central plains tallgrass to fire, fertilizer, and atrazine. J Range Manag 45(3):291–295
- Mazzola MB, Chambers JC, Blank RR et al (2011) Effects of resource availability and propagule supply on native species recruitment in sagebrush ecosystems invaded by *Bromus tectorum*. Biol Invasions 13:513–526
- McGinnis TW, Keeley JE, Stephens SL et al (2010) Fuel buildup and potential fire behavior after stand-replacing fires, logging fire-killed trees and herbicide shrub removal in Sierra Nevada forests. Forest Ecol Manag 260:22–35
- McGlone CM, Springer JD, Covington WW (2009) Cheatgrass encroachment on a ponderosa pine Ecol Restor project in northern Arizona. Ecol Restor 27:37–46
- McPherson GR (1995) The role of fire in the desert grasslands. In: McClaran MP, Van Devender TR, Thomas R (eds) The desert grassland. University of Arizona Press, Tucson, AZ, pp 130–151

- McPherson GR, Weltzin JF (2000) The role and importance of disturbance and climate change in US/Mexico borderlands: a state-of-the-knowledge review. Gen Tech Rep RMRS-GTR-50. USDA, Forest Service, Rocky Mountain Research Station, Fort Collins, CO, p 24
- Meixner T, Wohlgemuth PM (2003) Climate variability, fire, vegetation recovery, and watershed hydrology. First interagency conference on research in the watersheds, Benson, AZ, 27–30 October 2003, pp 651–656
- Meng R, Dennison PE, D'Antonio CM et al (2014) Remote sensing analysis of vegetation recovery following short-interval fires in southern California shrublands. PLoS One 9, e110637
- Meyer SE, Garvin SC, Beckstead J (2001) Factors mediating cheatgrass invasion of intact salt desert shrubland. In: McArthur D, Fairbanks DJ (eds) Shrubland ecosystem genetics and biodiversity. Gen Tech Rep RMRS-P-21. USDA Forest Service, Rocky Mountain Research Station, Fort Collins, CO, pp 224–232
- Meyer SE, Quinney D, Nelson DL et al (2007) Impact of the pathogen *Pyrenophora semeni*perda on *Bromus tectorum* seedbank dynamics in North American cold deserts. Weed Res 47:54–62
- Milchunas DG, Lauenroth WK, Chapman PL (1992) Plant competition, abiotic, and long-term and short-term effects of large herbivores on demography of opportunistic species in a semiarid grassland. Oecologia 92:520–531
- Milchunas DG, Vandever MW, Ball LO et al (2011) Allelopathic cover crop prior to seeding is more important than subsequent grazing/mowing in grassland establishment. Rangel Ecol Manag 64:291–300
- Miller R, Chambers JC, Pyke DA, et al (2013) A review of fire effects on vegetation and soils in the Great Basin Region: response and ecological site characteristics. Gen Tech Rep RMRS-GTR-308. USDA, Forest Service, Rocky Mountain Research Station, Fort Collins, CO
- Miller RF, Knick ST, Pyke DA et al (2011) Characteristics of sagebrush habitats and limitations to long-term conservation. Stud Avian Biol 38:145–184
- Minnich R (2008) California's fading wildflowers: lost legacy and biological invasions. UC Press, Berkeley, CA
- Molinari N (2014) Invasion, impact and persistence of an exotic annual grass. PhD dissertation, University of California, Santa Barbara, p 181
- Molinari N, D'Antonio CM (2014) Structural, compositional and trait differences between nativeand non-native-dominated grassland patches. Funct Ecol 28:745–754
- Monaco TA, Hardegree SP, Pellant M et al (2015) Assessing restoration and management needs for ecosystems invaded by exotic annual *Bromus s*pecies. In: Germino MJ, Chambers JC, Brown CS (eds) Exotic brome-grasses in arid and semiarid ecosystems of the Western USA: causes, consequences, and management implications. Springer, New York, NY (Chapter 12)
- Monaco TA, Sheley R (2012) Invasive plant ecology and management: linking processes to practice. CABI Invasive Series. CABI
- Monleon VJ, Cromack K, Landsberg JD (1997) Short- and long-term effects of prescribed underburning on nitrogen availability in ponderosa pine stands in central Oregon. Can J Forest Res 27:369–378
- Monsen SB, Stevens R, Shaw NL (2004) Restoring western ranges and wildlands. Gen Tech Rep RMRS-GTR-1136-vol 1, 2, 3. USDA, Forest Service, Rocky Mountain Research Station, Fort Collins, CO, p 294
- Moran MS, Ponce-Campos GE, Huete A et al (2014) Functional response of US grasslands to the early 21st-century drought. Ecology 95:2121–2133
- Moser WK, Barnard EL, Billings RF, Crocker SJ et al (2009) Impacts of nonnative invasive species on US forests and recommendations for policy and management. J Forestry 107:320–327
- Munson SM, Lauenroth WK (2009) Plant population and community responses to removal of dominant species in the shortgrass steppe. J Veg Sci 20:224–232
- Ogle SM, Ojima D, Reiners WA (2004) Modeling the impact of exotic annual brome grasses on soil organic carbon storage in a northern mixed-grass prairie. Biol Invasions 6:365–377
- Ogle SM, Reiners WA (2002) A phytosociological study of exotic annual brome grasses in a mixed grass prairie/ponderosa pine forest ecotone. Am Midl Nat 147:25–31

58

Ogle SM, Reiners WA, Gerow KG (2003) Impacts of exotic annual brome grasses (*Bromus* spp.) on ecosystem properties of northern mixed grass prairie. Am Midl Nat 149:46–58

- Osborne CP, Visser V, Chapman S et al (2011) GrassPortal: an online ecological and evolutionary data facility. www.grassportal.org. Accessed 16 Feb 2015
- Pauchard A, Kueffer C, Dietz H et al (2009) Ain't no mountain high enough: plant invasions reaching new elevations. Front Ecol Environ 7:479–486
- Pavlik LE (1995) Bromus L. of North America. Royal British Columbia Museum, Victoria, BC, p 160
- Pierson EA, Mack RN (1990) The population of biology of *Bromus tectorum* in forests: effect of disturbance, grazing, and litter on seedling establishment and reproduction. Oecologia 84:526–533
- Pyke D (2011) Restoring and rehabilitating sagebrush habitats. In: Knick S, Connelly JW (eds) Greater sage-grouse: ecology and conservation of a landscape species and its habitats, vol 38. University of California Press, Berkeley, CA, pp 531–548
- Pyke DA, Chambers JC, Beck JL et al (2015) Land uses, fire, and invasion exotic annual *Bromus* and human dimensions. In: Germino MJ, Chambers JC, Brown CS (eds) Exotic brome-grasses in arid and semiarid ecosystems of the Western USA: causes, consequences, and management implications. Springer, New York, NY (Chapter 11)
- Pyke DA, Wirth TA, Beyers JL (2013) Does seeding after wildfires in rangelands reduce erosion or invasive species? Restor Ecol 21:415–421
- Ramakrishnan AP, Meyer SE, Fairbanks DJ et al (2006) Ecological significance of microsatellite variation in western North American populations of *Bromus tectorum*. Plant Species Biol 21:61–73
- Rao LE, Allen EB (2010) Combined effects of precipitation and nitrogen deposition on native and invasive winter annual production in California deserts. Oecologia 162(4):1035–1046
- Rao LE, Allen EB, Meixner T (2010) Risk-based determination of critical nitrogen deposition loads for fire spread in southern California deserts. Ecol Appl 20:1320–1335
- Rao LE, Matchett JR, Brooks ML et al (2015) Relationships between annual plant productivity, nitrogen deposition and fire size in low-elevation California desert scrub. Int J Wildl Fire 24:48–58
- Reisner MD, Grace JB, Pyke DA et al (2013) Conditions favouring *Bromus tectorum* dominance of endangered sagebrush steppe ecosystems. J Appl Ecol 50:1039–1049
- Rice KJ, Mack RN (1991) Ecological genetics of *Bromus tectorum*. III. The demography of reciprocally sown populations. Oecologia 88:91–101
- Rice KJ, Nagy ES (2000) Oak canopy effects on the distribution patterns of two annual grasses: the role of competition and soil nutrients. Am J Bot 87:1699–1706
- Rimer RL, Evans RD (2006) Invasion of downy brome (*Bromus tectorum* L.) causes rapid changes in the nitrogen cycle. Am Midl Nat 156:252–258
- Rinella MJ, Haferkamp MR, Masters RA et al (2010a) Growth regulator herbicides prevent invasive annual grass seed production. Invasive Plant Sci Manag 3:12–16
- Rinella MJ, Masters RA, Bellows SE (2010b) Growth regulator herbicides prevent invasive annual grass seed production under field conditions. Rangel Ecol Manag 63:487–490
- Rinella MJ, Masters RA, Bellows SE (2013) Effects of growth regulator herbicide on Downy brome (*Bromus tectorum*) seed production. Invasive Plant Sci Manag 6:60–64
- Rochester CJ, Brehme CS, Clark DR et al (2010) Reptile and amphibian responses to large-scale wildfires in southern California. J Herpetol 44:333–351
- Romps DM, Seeley JT, Vollaro D et al (2014) Projected increase in lightning strikes in the United States due to global warming. Science 346:851–854
- Roundy BA, Hardegree SP, Chambers JC et al (2007) Prediction of cheatgrass field germination potential using wet thermal accumulation. Rangel Ecol Manag 60:613–623
- Safford HD, Van de Water KM (2014) Using fire return interval departure (FRID) analysis to map spatial and temporal changes in fire frequency on national forest lands in California. Gen Tech Rep PSW-RP-266. USDA, Forest Service, Pacific Southwest Research Station, p 59

- Salo LF (2005) Red brome (*Bromus rubens* subsp *madritensis*) in North America: possible modes for early introductions, subsequent spread. Biol Invasions 7:165–180
- Sankey JB, Germino MJ, Glenn NF (2009) Aeolian sediment transport following wildfire in sagebrush steppe. J Arid Environ 73:912–919
- Schaeffer SM, Ziegler SE, Belnap J et al (2012) Effects of *Bromus tectorum* invasion on microbial carbon and nitrogen cycling in two adjacent undisturbed arid grassland communities. Biogeochemistry 111:427–441
- Scheintaub MR, Derner JD, Kelly EF et al (2009) Response of the shortgrass steppe plant community to fire. J Arid Environ 73:1136–1143
- Seabloom EW, Harpole WS, Riechman OJ et al (2003) Invasion, competitive dominance and resource use by exotic and native California grassland species. Proc Natl Acad Sci U S A 100:13384–13389
- Shinneman DJ, Baker WL (2009) Environmental and climatic variables as potential drivers of post-fire cover of cheatgrass (*Bromus tectorum*) in seeded and unseeded semiarid ecosystems. Int J Wildl Fire 18:191–202
- Sims P, Risser PG (2000) Grasslands. In: Barbour M, Billings WD (eds) North American terrestrial vegetation, 2nd edn. Cambridge University Press, Cambridge, UK, pp 323–356
- Smith SD, Strain BR, Sharkey TD (1987) Effects of CO<sub>2</sub> enrichment on four Great Basin grasses. Funct Ecol 1:139–143
- Sorensen CD, McGlone CM (2010) Ponderosa pine understory response to short-term grazing exclusion (Arizona). Ecol Restor 28:124–126
- Stahlheber K (2013) The influence of savanna oaks on California grassland composition. PhD dissertation, University of California, Santa Barbara, p 275
- Stahlheber K, D'Antonio CM (2013) Using livestock to manage plant composition: a metaanalysis of grazing in California Mediterranean grassland. Biol Conserv 157:300–308
- Strand EK, Launchbaugh KL (2013) Livestock grazing effects on fuel loads for wildland fire in sagebrush dominated ecosystems. Great Basin fire science delivery report, p 21
- Stylinski CD, Allen EB (1999) Lack of native species recovery following severe exotic disturbance in southern Californian shrublands. J Appl Ecol 36:544–554
- Suttle B, Thomsen MA, Power ME (2007) Species interactions reverse grassland responses to changing climate. Science 315:640–642
- Sweet SB, Kyser GB, DiTomaso JM (2008) Susceptibility of exotic annual grass seeds to fire. Invasive Plant Sci Manag 1:158–167
- Symstad A, Long AJ, Stamm JF et al (2014) Two approaches for incorporating climate change into natural resource management planning at Wind Cave National Park. Gen Tech Rep NPS/ WICA/NRTR-2014.918. National Park Service, Fort Collins, CO
- Syphard AD, Keeley JE (2015) Location, timing and extent of wildfire vary by cause of ignition. Int J Wildl Fire 24:37–47
- Tagestad J, Brooks M, Cullinan V et al (in press) Precipitation regime classification for the Mojave Desert: implications for fire occurrence. J Arid Environ
- Talluto MV, Suding KN (2008) Historical change in coastal sage scrub in southern California, USA in relation to fire frequency and air pollution. Landscape Ecol 23:803–815
- Teague WR, Dowhower SL (2003) Patch dynamics under rotational and continuous grazing management in large, heterogeneous paddocks. J Arid Environ 53:211–229
- Teague WR, Dowhower SL, Baker SA et al (2010) Soil and herbaceous plant responses to summer patch burns under continuous and rotational grazing. Agr Ecosyst Environ 137:113–123
- Teague WR, Dowhower SL, Waggoner JA (2004) Drought and grazing patch dynamics under different grazing management. J Arid Environ 58:97–117
- Teague WR, Duke SE, Waggoner JA et al (2008) Rangeland vegetation and soil response to summer patch fires under continuous grazing. Arid Land Res Manag 22:228–241
- USDA NRCS (2015) The PLANTS Database. http://plants.usda.gov. Accessed 26 Feb 2015
- Van Dyne G (1975) An overview of the ecology of the Great Plains grasslands with special reference to climate and its impact. Grassland biome ecosystem analysis studies. Gen Tech Rep 290. Natural resources ecology laboratory, Colorado State University, Fort Collins, CO

60 M.L. Brooks et al.

Vermeire LT, Crowder JL, Wester DB (2011) Plant community and soil environment response to summer fire in the Northern Great Plains. Rangel Ecol Manag 64:37–46

- Vermeire LT, Crowder JL, Wester DB (2014) Semiarid rangeland is resilient to summer fire and postfire grazing utilization. Rangel Ecol Manag 67:52–60
- Vermeire LT, Heitschniidt RK, Haferkamp MR (2008) Vegetation response to seven grazing treatments in the Northern Great Plains. Agric Ecosyst Environ 125:111–119
- Walker BH, Holling CS, Carpenter SR et al (2004) Resilience, adaptability and transformability in social-ecological systems. Ecol Soc 9:5
- Walsh J, Wuebbles D, Hayhoe K et al (2014) Our changing climate. In: Melillo JM, Richmond TC, Yohe GW (eds) Climate change impacts in the United States: the third national climate assessment. US Global Change Research Program, pp 19–67
- West N (1983a) Great Basin-Colorado Plateau sagebrush semi-desert. Temperate Deserts and Semi-deserts. Elsevier, Amsterdam
- West N (1983b) Intermountain salt-desert shrubland. Temperate Deserts and Semi-deserts. Elsevier, Amsterdam, pp 375–378
- Westerling AL, Bryant BP (2008) Climate change and wildfire in California. Clim Change 87:S231–S249
- Westerling AL, Hidalgo HG, Cayan DR et al (2006) Warming and earlier spring increase western U.S. forest wildfire activity. Science 313:940–943
- Westman WE (1979) Oxidant effects on Californian coastal sage scrub. Science 205:1001-1003
- Whisenant SG (1990) Changing fire frequencies on Idaho's Snake River Plains: ecological and management implications. In: McArthur ED, Romney EM, Smith SD et al (eds) Symposium on cheatgrass invasion, shrub die-off and other aspects of shrub biology and management, 5–7 April 1989, Las Vegas, NV. Gen Tech Rep INT-276. USDA, Forest Service, Intermountain Research Station, Ogden, UT, pp 4–10
- Whisenant S, Ueckert D, Scifres C (1984) Effects of fire on Texas wintergrass communities. J Range Manag 37:387–391
- Whisenant SG, Uresk DW (1990) Spring burning Japanese brome in a western wheatgrass community. J Range Manag 43:205–208
- White CS, Loftin SR (2000) Response of two semiarid grasslands to cool-season prescribed fire. J Range Manag 53:52–61
- Wilcox BP, Turnbull L, Young MH et al (2012) Invasion of shrublands by exotic grasses: ecohydrological consequences in cold versus warm deserts. Ecohydrology 5:160–173
- Wiken E, Nava FJ, Griffith G (2011) North American Terrestrial Ecoregions—Level III. Commission for Environmental Cooperation, Montreal, Canada
- Wolkovich EM, Bolger DT, Holway DA et al (2009a) Invasive grass litter facilitates native shrubs through abiotic effects. J Veg Sci 20:1121–1132
- Wolkovich EM, Bolger DT, Holway DA (2009b) Complex responses to invasive grass litter by ground arthropods in a Mediterranean scrub ecosystem. Oecologia 262:697–708
- Wright HE, Bailey AW (1982) Fire ecology: United States and Canada. Wiley, New York, NY
- Yensen E, Quinney DL, Johnson K et al (1992) Fire, vegetation changes, and population fluctuations of Townsend ground-squirrels. Am Midl Nat 128:299–312
- Zedler PH, Gautier CR, McMaster GS (1983) Vegetation change in response to extreme events: the effect of a short interval between fires in California chaparral and coastal scrub. Ecology 64:809–818