Natural Range of Variation for Yellow Pine and Mixed-Conifer Forests in the Sierra Nevada, Southern Cascades, and Modoc and Inyo National Forests, California, USA

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Cover photos: Top left—Mixed-conifer stand in the Sierra de San Pedro Mártir (SSPM), Baja California, Mexico, elevation 2600 m. Tree species are Jeffrey pine, sugar pine, white fir, and canyon live oak. The SSPM National Park is an important reference site for eastern Sierra Nevada yellow pine and mixed-conifer restoration, as it was never logged and has only experienced a few decades of fire suppression.

Top right—Jeffrey pine-white fir stand on the Humboldt-Toiyabe National Forest, Walker River Canyon, elevation 1850 m. This site was burned 2 years previous to the photo by the Larson Fire.

Bottom—Moist mixed-conifer forest on the Tahoe National Forest, above the Middle Fork of the Yuba River, elevation 1000 m. Forest is dominated by Douglas-fir, with some incense cedar and ponderosa pine, and black oak. This area was logged multiple times between the mid-1800s and 1970s. The black oak patch in the middle of the photo occupies the site of an earlier clearcut.
Abstract


Yellow pine and mixed-conifer (YPMC) forests are the predominant montane forest type in the Sierra Nevada, southern Cascade Range, and neighboring forested areas on the Modoc and Inyo National Forests (the “assessment area”). YPMC forests occur above the oak woodland belt and below red fir forests, and are dominated by the yellow pines (ponderosa pine \textit{[Pinus ponderosa} Lawson & C. Lawson] and Jeffrey pine \textit{[Pinus jeffreyi} Balf.]); white fir \textit{(Abies concolor} (Gord. & Glend.) Lindl. ex Hildebr.)); sugar pine \textit{(P. lambertiana} Douglas); incense cedar \textit{(Calocedrus decurrens} (Torr.) Florin); and black oak \textit{(Quercus kelloggii} Newberry), along with a number of other hardwood and conifer species. We conducted an indepth assessment of the natural range of variation (NRV) of YPMC forests for the assessment area, focusing on ecosystem processes and forest structure from historical data sources from pre-Euro-American settlement times (16\textsuperscript{th} through mid-19\textsuperscript{th} centuries) and current reference forests (YPMC forests that have retained frequent fire and have suffered little human degradation), and comparing current conditions to the NRV. The Mediterranean climate of the assessment area, modified by strong latitudinal, topographic, and elevational gradients, plays an important role in shaping the structure and composition of YPMC forests. Fire was an historically important ecosystem process that occurred frequently, generally burned at low to moderate severity, created a heterogeneous forest structure at a fine spatial scale, and maintained pine dominance in many stands that would otherwise undergo succession to more shade-tolerant fir and cedar species. Forest structure at larger spatial scales was highly variable but was characterized mostly by relatively low tree densities, large tree sizes, high seedling mortality as a result of recurrent fire, and highly heterogeneous understory structure that could include locally abundant fire-stimulated shrub species. Following Euro-American settlement, wholesale changes occurred in YPMC forests in the assessment area, principally because of extensive logging followed by a century of highly effective fire suppression. Modern YPMC forests have departed from NRV conditions for a wide range of ecosystem processes and structural attributes. There is strong consensus among published studies that, on average, modern YPMC stands have much higher densities dominated by smaller trees (often of shade-tolerant species), much longer fire-return intervals, and less
area burned across the landscape compared to reference YPMC forests. In addition, fires that escape initial attack are much larger and higher severity on average than the average pre-Euro-American settlement fire. There is more moderate consensus among published studies that the average modern YPMC stand in the assessment supports greater fuels and deeper forest litter, higher canopy cover and fewer canopy gaps, more coarse woody debris, a higher density of snags, and experiences a longer fire season compared to reference YPMC forests. Among the variables assessed, only basal area, overall plant species richness, and percentage cover of grass/forbs and shrubs appear to be within or near the NRV.

Keywords: Yellow pine forests, mixed-conifer forest, ecosystem function, fire regime, forest structure, historical range of variation, HRV, natural range of variation, NRV, species diversity and composition, succession.
Preface

In 1976, President Gerald Ford signed the National Forest Management Act (NFMA), which—together with the Forest and Rangeland Renewable Resources Planning Act of 1974 (FRRPPA)—committed the Forest Service to developing and periodically updating land and resource management plans (LRMPs) at the national forest or national grassland level. The principal purpose of the LRMPs is to provide for “multiple use” and “sustained yield” of natural resources in the National Forest System.

The NFMA and FRRPPA required the development of regulations by the Forest Service to guide the LRMP revision process. These guidelines came to be known as the “Planning Rule,” and were first published in 1982. Various inadequacies of the original rule became apparent over time, and multiple abortive efforts were made to modify or “modernize” it. In 2012, a new rule was finally adopted (36 CFR 219) (USDA FS 2012), and new forest plans are following the revised process as of 2013.

According to 36 CFR 219.1(c), the purpose of the 2012 Planning Rule is “to guide the collaborative and science-based development, amendment, and revision of land management plans that promote the ecological integrity of national forests and grasslands...” The rule is focused on maintaining biological diversity on Forest Service units, and ensuring the “integrity of the compositional, structural, and functional components comprising... ecosystems.”

The 2012 Planning Rule places heavy emphasis on the concepts of “sustainability” and “ecological integrity.” In the rule, sustainability is defined as “the capability of ecosystems to maintain ecological integrity” (36 CFR 219.19 p 21272), and ecological integrity is defined as:

The quality or condition of an ecosystem when its dominant ecological characteristics (for example, composition, structure, function, connectivity, and species composition and diversity) occur within the natural range of variation and can withstand and recover from most perturbations imposed by natural environmental dynamics or human influence (36 CFR 219.19 p 21271).

The definition of ecological integrity in the 2012 Planning Rule thus inherently requires the determination of the “natural range of variation” (NRV) for a suite of important ecosystem variables, organized by their composition, structure, and function (connectivity belongs to structure, and species composition and diversity are nested in composition). Natural range of variation is defined by Forest Service Handbook 1909-12 Chapter 10 as:
Spatial and temporal variation in ecosystem characteristics under historic disturbance regimes during a reference period... The NRV can help identify key structural, functional, compositional, and connectivity characteristics, for which plan components may be important for either maintenance or restoration of such ecological conditions.

As mandated by NFMA, national forests in the “Sierra Nevada bioregional assessment area” will revise and update their forest plans over the course of the next 6 to 10 years. The assessment area (see the outline map in fig. 1) includes the Sierra Nevada proper, the southern Cascade Range in California, the Modoc Plateau, the Warner Mountains, and the White and Inyo Mountains, among other areas. It is essentially the same area that was analyzed by the Sierra Nevada Ecosystem Project (Erman and SNEP Team 1996) and the Sierra Nevada Forest Plan Amendments of 2001 and 2004 (USDA FS 2001, 2004). Between October 2012 and May 2013, the Ecology Program of the Forest Service’s Pacific Southwest Region carried out assessments of NRV for 11 major vegetation types across the Sierra Nevada bioregional assessment area. These NRV assessments fed the development of a more wide-ranging bioregional assessment that treats ecological, economic, social components and provides a broad-scale framework for the forest-level assessments that began in late 2013.

The intent is to publish the Ecology Program’s NRV assessments as Forest Service general technical reports over the course of the next few years. This document, the NRV assessment for yellow pine and mixed-conifer forests, is the first installment of these publications. An earlier draft of this document was available online between February 15, 2015, and June 6, 2016, at http://www.fs.usda.gov/detail/r5/plants-animals/?cid=stelprdb5434436.

Summary

Yellow pine and mixed-conifer (YPMC) forests are among the most widely distributed forest types in California. Within the assessment area, these forests are primarily mid-elevation montane forests, with the distributions of the dominant canopy species limited by cold conditions at upper elevations and by dry conditions at lower elevations. YPMC forests occupy a Mediterranean climate zone with prolonged summer drought and sufficient winter precipitation to support tree growth, with greater precipitation and productivity on the western slopes of the Sierra Nevada and southern Cascade Range. The yellow pines, ponderosa pine (Pinus ponderosa Lawson & C. Lawson) and Jeffrey pine (P. jeffreyi Balf.) are important canopy species within these forests, predominating at drier and frequently burned
sites while mixing with other more shade-tolerant conifer species (white fir \textit{Abies concolor} (Gord. & Glend.) Lindl. ex Hildebr.), incense cedar \textit{Calocedrus decurrens} (Torr.) Florin], and Douglas-fir \textit{Pseudotsuga menziesii} (Mirb.) Franco]) at sites with less moisture stress or less frequent fire; sugar pine \textit{(P. lambertiana} Douglas) and black oak \textit{(Quercus kelloggii} Newberry) are other important tree species in YPMC forests. Important ecological differences among these canopy species (e.g., tolerance of shade, fire, and drought) have influenced their distribution and abundance across climatic and topographic gradients, which in concert with frequent fires have led to high levels of heterogeneity within this general forest type. Despite climatic fluctuations during the Holocene Epoch, the assessment area has been at least partially forested for 10,000 years, with an increase in the mixed-conifer component over the past 4,000 years when fire also has been very frequent, at least partially because of burning by American Indians. The past 150 years have seen extensive changes to assessment area YPMC forests, which experienced large-scale logging and subsequent, nearly ubiquitous fire exclusion that have dramatically altered contemporary forest structure and ecological processes.

Fire has been a major force in YPMC forests for millennia owing to the productive yet seasonally dry climate of the assessment area. Prior to Euro-American settlement in the mid-1800s (“presettlement”), fires were generally frequent, with a mean fire-return interval of 11 to 16 years. Fire occurrence during the summer dry season was often fuel limited, but also increased during periods of drought. These frequent fires kept fuel loads low and heterogeneously distributed across the landscape, and therefore fire severity under presettlement conditions tended to be low to moderate. Most estimates of presettlement high-severity proportions ranged from 5 to 15 percent, based on historical observations and surveys, studies of contemporary reference forests, and landscape-scale modeling. Mean fire size (of all fires >10 ha) was relatively small (c. 200 to 400 ha), while the mean annual area burned was about 5 percent of total YPMC forest area.

Forest structure in presettlement YPMC forests was generally characterized by fine-grained (within-stand) heterogeneity controlled by fire interacting with geologic, topographic, and climatic features that influenced vegetation productivity and structure primarily via their effects on growing-season water availability (the primary limiting resource in YPMC forests). Models and present-day reference landscapes suggest that roughly 50 percent of the presettlement landscape would have been in an old-growth condition, but this old-growth condition included both open stands of yellow pine and dry mixed conifer, and more closed-canopy stands of moist mixed conifer. Thus, stand structure within YPMC stands was likely highly variable, but in general, stands were characterized by relatively low
The heterogeneity of presettlement landscapes in YPMC forests has been simplified over the past 150 years, primarily by logging and fire exclusion, which have led to an increase in small-tree density and a decrease in large-tree density relative to the natural range of variation. Contemporary mean tree density of YPMC forests is 397 trees/ha, with densities ranging from 238 to 755 trees/ha in stands for which presettlement reconstructions exist. Most of this increase is in trees <60 cm diameter at breast height (d.b.h.). Corresponding changes in forest and understory structure from the presettlement era include a decrease in the average tree size (d.b.h.) by 25 to 40 percent; an increase in canopy cover by ~25 percent; a near-complete loss of fine-scale canopy gaps; increases in snag density, coarse woody debris, litter, and duff depth; and surface fuel volume and continuity. These changes in stand structure have caused changes in other ecosystem processes. For instance, insect and pathogen activity has probably increased with stand density (although presettlement information on this is sparse), while litter and duff accumulation can increase nutrients leaching to surface and ground water, and, combined with increased small-tree density, increase vulnerability to high-severity fire.

Modern fire suppression practices have caused a strong departure from presettlement fire-return intervals in YPMC forests, with an average of 80 to 85 percent fewer fires per 100 years than during the presettlement era. Although the frequency of fires has been greatly reduced, the characteristics of fires that do burn have also changed. The mean proportion of high-severity fire in YPMC forests has increased to between 30 and 35 percent in the past 30 years, while the size of high-severity patches also has increased. The mean fire size (of all fires >10 ha) over the past 30 years was ~1400 ha. As future climates continue to warm, future fire size, severity, and frequency are likely to increase.

Regeneration and understory dynamics in YPMC forests are complex. Regeneration dynamics historically were greatly influenced by fire (along with other gap-forming processes and precipitation patterns), with pine regeneration particularly enhanced by low- and moderate-severity fires, which maintained low canopy cover and exposed bare microsites for improved germination success. Presettlement regeneration was patchy and likely determined by interactions between fire, climate, topography, surface litter depth, and canopy cover. Modern tree regeneration
is still highly variable, but survival of seedlings is promoted by fire exclusion, and
the seedling class is more dominated by shade-tolerant fir, Douglas-fir, and incense
cedar. Presettlement shrub cover was also quite variable. Areas of continuous
shrubfields may have dominated after one or more severe fires, as is the case with
modern severe fires, but shrubs could also be quite abundant in the understory of
open stands that experienced frequent lower severity fires, as is the case in many
modern reference stands. Contemporary YPMC forests may be within the NRV of
shrub cover at a landscape scale, but many stands have likely experienced increases
in shrub cover within large contiguous patches of high-severity fire, while others
have seen reductions in shrub cover in fire-suppressed stands owing to increased
shading. Presettlement herbaceous cover was likely low but diverse, and modern
forests may be within the NRV in terms of herbaceous cover, but contemporary
herbaceous diversity increases in stands that experience fire.
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Chapter 1: Introduction

Natural range of variation (NRV) assessments (essentially, equivalent to historical range of variation [HRV] assessments) provide baseline information on ecosystem conditions (composition, structure, and function) that can be compared to current conditions to develop an idea of trend over time and an idea of the level of departure of altered ecosystems from their “natural” state (Landres et al. 1999, Manley et al. 1995, Morgan et al. 1994, Wiens et al. 2012) (see chapter 2). These trend assessments form part of the basis for the assessment of ecological integrity that is required in the 2012 Forest Service Planning Rule. Natural range of variation assessments were carried out for 11 terrestrial ecosystems by the USDA Forest Service Pacific Southwest Region (Region 5) Ecology Program between October 2012 and May 2013 using historical information (primarily from the pre-Euro-American period, 16th century to the mid-19th century) as well as information from modern-day reference ecosystems and other sources (see “Methods” and the appendix). The assessments were later updated for more recent scientific findings, and can be considered generally current as of late 2015, with some sections updated to 2016 or even early 2017.

In Region 5, forest planning under the 2012 Planning Rule is moving forward in geographically contiguous groups of national forests. Each forest will carry out its own set of assessments, but Region 5 also elected to carry out a bioregional assessment covering the same area as the 1996 Sierra Nevada Ecosystem Project (SNEP) to update trends and conditions described in the SNEP report and the Sierra Nevada Forest Plan Amendment (USDA FS 2004), and to better provide consistency among national forest plans with respect to conditions and trends at spatial scales greater than the size of a single national forest (400 000 to 800 000 ha), and temporal scales greater than the typical lifespan of a forest plan (15 to 20 years). In this report, we provide (1) an estimate of the NRV for key elements of yellow pine and mixed-conifer forest ecosystems, including quantitative ranges for specific variables whenever possible; and (2) a comparison of our NRV analysis with current conditions in the bioregional assessment area. Our NRV assessments also go farther in time, and summarize the current science concerning projected future conditions for key ecosystem elements.
Physical Setting and Geographic Distribution

Yellow pine and mixed-conifer forests are found throughout the assessment area (fig. 1). In this contribution, we jointly consider the two forest types and refer to them collectively as “YPMC” forests (the reasoning behind this combination is given in “Ecological Setting” below). YPMC forests are the most widely distributed vegetation type in the assessment area. Based on the California Wildlife Habitat Relations vegetation types (Mayer and Laudenslayer 1988), forest types corresponding to YPMC forest cover about 3 million ha (7.5 million ac) of the assessment area (fig. 1).

YPMC forests are found on a variety of soils and bedrocks, and there is little evidence of strong soil chemistry-driven differences in plant species composition or forest structure in the assessment area, except in the case of ultramafic (“serpentine”) soils, which are extremely nutrient deficient but relatively rare in the YPMC forest belt, aside from parts of the northern Sierra Nevada (Alexander et al. 2007). On the other hand, variations in soil depth and texture, which are strongly related to topography (slope, aspect, elevation, slope shape, etc.) in the assessment area’s Mediterranean climate, are major drivers of variation in forest species composition, density, cover, and other related variables. A major role of soils in YMPC forests in the assessment area is their contribution to water availability (O’Geen et al. 2007).

Most of California, including the assessment area, falls within the North American Mediterranean climate zone. Five geographic areas on Earth share the unique Mediterranean climatic characteristics, which are typically expressed on the western boundaries of continents between about 30° and 45° north latitude. Under the Köppen (1931) classification, the Mediterranean climate (“Cs” in the Köppen classification) is described as a temperate rainy climate with warm to hot, dry summers. YPMC forests in the assessment area fall primarily within the Köppen subcategory “Csb,” in which the average temperature of the warmest month is <22 °C and at least 2 months average above 10 °C (oak woodland, foothill pine, and chaparral vegetation types fall mostly in the warmer “Csa” type). As Bailey (2009) noted, the combination of wet winters with dry summers is rare along world climate types and leads to a distinctive vegetation dominated by evergreen trees and shrubs. The major environmental stress is the severe summer drought, which can last from 3 to 6 months, and leads to high evaporative demand during much of the growing season. In a Mediterranean climate, the demand for water and its supply are exactly out of phase (Major 1988). Within the Cs climate zone, there is a gradient in intensity of the dry summer season from areas with relatively mild summer temperatures (northern assessment area and higher elevations) to areas with hot summer temperatures (lower elevations, especially in the southern and eastern assessment areas) (SSP 2010).
Figure 1—Distribution of yellow pine and mixed-conifer forests in the assessment area, with climate station locations indicated.
Figure 2 shows Walther-type climate diagrams for six National Weather Service weather stations in or near YPMC forests in the assessment area; data are from the Western Regional Climate Center (2013). The diagrams are organized to approximately match their locations in figure 1. As per Walter and Lieth (1967), the temperature and precipitation axes are scaled such that 20 mm precipitation = 10 °C temperature. The dry season is found approximately where the precipitation line undercut the temperature line; the wet season occurs where precipitation supersedes temperature. We say approximately because moisture stored in the soil and snowpack is available for plant use after the end of the wet season and can reduce true dry season length by a month or more depending on annual precipitation (see Major 1988).

The climate diagrams underline the major climatic gradients in the assessment area (fig. 2). Dry season length is shortest along the west slope, especially in the north (3 to 4 months), and longest in the extreme south and on the east slope (5 to 6 months). Precipitation is higher in the northwest than the rest of the assessment area. Stations east of the hydrologic divide (which is either the Sierra Nevada or Cascade Range crest in most of the assessment area) are subject to a rain shadow effect, as is the Modoc Plateau, represented by Alturas. All stations receive some monsoon-derived precipitation in the summer months, with the magnitude and importance of that precipitation increasing to the east and southeast in concert with the influence of the Great Basin climate. Of the stations shown, Alturas and Lee Vining are the most influenced by periodic summer precipitation. Temperatures are warmest in the south, and coolest in the north and at higher elevations (fig. 2).

Topography drives major differences in ecosystem distribution across the assessment area. Elevation increases from north to south, and local relief is generally much higher in the central and southern Sierra Nevada than the rest of the assessment area. The Sierra Nevada is also narrower in the south, and the combination of less land area, higher mountains, and steeper slopes means that the YPMC forest belt is much more compressed south of the latitude of Sacramento (fig. 1). The southern Sierra Nevada also rises to a high plateau relatively quickly from the floor of the Central Valley. As a result, the area on the west slope dominated by ponderosa pine (*Pinus ponderosa* Lawson & C. Lawson) in the southern Sierra Nevada is relatively narrow, and higher elevation mixed-conifer forests with an important white fir (*Abies concolor* Gord. & Glend.) component are more widespread. Steeper terrain, a drier climate, and the predominance of granitic bedrock in the central and south Sierra Nevada also lead to much greater prevalence of bare rock substrates and cliffs. Overall, the natural lay of the land means that physical barriers to ecological processes like animal migration, plant seed dispersal, and fire spread are greater in the southern and central Sierra Nevada than elsewhere, but land ownership and land use follow the opposite pattern, with much more fragmentation of habitat in the northern subregion.
Figure 2—Walther-type climate diagrams for six weather stations in or near yellow pine and mixed-conifer forest in the assessment area; elevations are provided. Locations shown in figure 1. Dry season length is approximately the period during which the precipitation curve undercuts the temperature curve.
Because of dry growing season conditions in the assessment area, water availability is a major driver of ecosystem distribution and condition (Barbour et al. 2007, Loik et al. 2004, Major 1988). Topography exerts strong influence on water availability, both directly through its influence on temperature, solar insolation, and evaporation, and indirectly through its influence on soil depth and texture. Forest conditions in the YPMC forest belt differ substantially along topographic gradients (Barbour et al. 2007, North et al. 2012b, Sugihara et al. 2006). It can be generalized that, all other environmental factors being equal, YPMC forest cover and density tend to be higher on north-facing (“cool”) aspects (where water availability is higher), and more open and less dense on south- and west-facing (“warm”) aspects. Because higher elevations receive more precipitation because of orographic effects and also lose less water to heat evaporation, they also tend to support denser stands of forest than lower elevations. The upper portions of mountain or canyon slopes lose substantial water to gravity-driven flow and tend to support more open forest conditions, whereas lower slopes are net importers of water from higher slopes, and they support denser forest conditions. The most open forest conditions tend to be on south- or west-facing, upper, convex canyon slopes at lower elevations; the densest forest conditions tend to be on north-facing, concave, lower slopes. Overlaying on these topographic effects are precipitation and temperature gradients across the broader assessment area. In general, forests are more open (and more composed of drought-tolerant species) in the southern and eastern portions of the assessment area than in the western and, especially, the northern portions.

Ecological Setting

Yellow pine forests in the assessment area are those that are dominated by one or both of the “yellow pine” species present in California: ponderosa pine and Jeffrey pine (P. jeffreyi Balf); Washoe pine is now considered a variety of ponderosa pine (Pinus ponderosa Lawson & C. Lawson var. washoensis (H. Mason & Stockw.) J.R. Haller & Vivrette) (Baldwin et al. 2012). Ponderosa and Jeffrey pine are closely related (they are both in the subgenus Pinus, section Pinus, subsection Ponderosae), and they occasionally hybridize. Ponderosa pine, the most widely distributed pine species in North America, is found throughout the mountainous regions of California, whereas Jeffrey pine is primarily a California tree, with some occurrences in westernmost Nevada, southwestern Oregon, and northern Baja California. Of the two species, Jeffrey pine is more stress tolerant, and replaces ponderosa pine at higher elevations, on poorer soils, and in colder or drier climates (Barbour and Minnich 2000, Haller 1959). Ponderosa pine-dominant forests can occur from about 300 m elevation to about 1800 m in the northern subregion of the assessment area,
and from about 1200 to 2100 m in the southern subregion (Fites-Kaufman et al. 2007). Jeffrey pine-dominant forests occur mostly between 1500 and 2400 m in the northern subregion and from 1700 to 2800 m in the southern subregion, with the highest elevations usually being on the east side of the Sierra Nevada (Barbour and Minnich 2000, Fites-Kaufman et al. 2007). Both yellow pine species can also occur in other forest types. A large area of the Lassen, Plumas, and Tahoe National Forests east of the Sierra Nevada crest supports a mixed-yellow pine forest with codominance by ponderosa and Jeffrey pine; this forest type is often called “east-side pine.”

Many conifer species with overlapping geographic distributions are found in the assessment area, and difficulties in differentiating obvious forest types have led to the general recognition of a “mixed-conifer” belt in the lower montane zone, usually intermixed with yellow pine-dominant stands at its lower edge and up to 2000 to 2200 m elevation at its highest, depending on latitude. Major tree species include the yellow pine species, sugar pine (Pinus lambertiana Douglas), white fir, incense cedar (Calocedrus decurrens (Torr.) Florin), Douglas-fir (Pseudotsuga menziesii (Mirb.) Franco), and black oak (Quercus kelloggii Newberry). Red fir (Abies magnifica A. Murray bis), lodgepole pine (Pinus contorta Douglas ex Loudon), and western white pine (Pinus monticola Douglas ex D. Don) are upper montane species that also make appearances in higher mixed-conifer stands; a variety of hardwood species also occur in lower elevation stands, including canyon live oak (Quercus chrysolepis Liebm.), interior live oak (Q. wislizenii A. DC.), and tanoak (Notholithocarpus densiflorus (Hook. & Arn.) P.S. Manos, C.H. Cannon, & S.H. Oh). As Barbour and Minnich (2000) noted, the yellow pine species (especially ponderosa pine) are “the biological thread that holds the [mixed-conifer] forest together,” but the very fine-grained pattern of local dominance makes it difficult to clearly discriminate the various “phases” for classification or mapping purposes. It should be noted that the topography of the southern Sierra Nevada (see “Physical Setting”) leads to a relatively narrow belt of ponderosa pine-dominant forest, and many YPMC stands in this part of the assessment area support a notable component of white fir.

Although Sawyer et al. (2009) described nearly 20 different forest alliances that make up the montane mixed-conifer forest, we stick with common practice and treat the mixed-conifer forest as a single, if geographically mutable, entity. In some cases in this assessment we refer to “dry mixed-conifer” or “moist mixed-conifer” forests. These terms are used only in a general sense, and differentiate mixed conifer based on the yellow pine and fir components (yellow pine dominance in dry mixed conifer; greater fir presence in moist mixed conifer) and the
annual precipitation (mostly <1000 mm in dry mixed conifer, mostly >1000 mm in moist mixed conifer); moist mixed-conifer stands are also more common at higher elevations. We also follow Barbour and Minnich (2000) in combining yellow pine forest and mixed conifer in this chapter. This is because yellow pine forests have become rare on the west side of the Sierra Nevada, owing to logging of the ponderosa pine or fire-exclusion-driven succession to mixed stands dominated by more shade-tolerant and fire-intolerant species like white fir and incense cedar (the same can be said of sugar pine, which was even more targeted by logging). Indeed, in drier sites throughout, and up to 2000–2500 m elevation at its highest Yellow pine-dominant forests remain common on the east side of the Sierra Nevada, however.

We do not explicitly treat giant sequoia (Sequoiadendron giganteum (Lindl.) J. Buchholz) in this NRV assessment. Giant sequoia occurs in scattered groves in the southern assessment area, with a few occurrences in the central assessment area. Giant sequoia is a locally dominant member of moist mixed-conifer stands with particular topographic and soils conditions (Barbour et al. 2007, Sugihara et al. 2006). Our treatment of moist mixed-conifer forest, which is otherwise dominated by species like white fir and sugar pine, is generally applicable to giant sequoia stands, but we direct the reader to the forest-level assessments for more detail. Stephenson (1999) is an excellent treatment of ecological, management, and restoration issues in giant sequoia-dominant mixed-conifer forest.

An understanding of past, current, and possible future conditions in YPMC forests in the assessment area requires consideration of the ecological differences between the major tree species. Forestry experience and scientific investigation have provided us with a wealth of information vis-à-vis the relative ecological tolerances of major tree species in the YPMC forests of the assessment area. Table 1 offers some summary information about geographical and elevational distributions of the seven major tree species in assessment area YPMC forests. Four of the seven species are broadly restricted to the North American Mediterranean zone (Jeffrey pine, sugar pine, incense cedar, black oak), and three are more widely distributed (ponderosa pine, Douglas-fir, white fir), with the two former species among the most widely distributed tree taxa in North America. Species restricted primarily to the YPMC forest zone include ponderosa pine, sugar pine, white fir, incense cedar, and black oak. Douglas-fir is also found at lower elevations in the northern and central parts of the assessment area, usually in moist, mixed-evergreen forests with a number of hardwood/broadleaf associates. Jeffrey pine ranges into higher elevation forests and is a common member of red fir forests and even some warm sites in the subalpine zone (table 1).
We have compiled and condensed important information related to comparative ecological traits and tolerances for the important tree species in assessment area YPMC forests. Table 1 provides information on seed weights, which are inversely related to dispersal distances (and therefore scale with the relative rapidity by which these species can recolonize or migrate to distant habitat), and leaf flammability, which is a trait strongly associated with the importance of fire to species ecology (Fonda et al. 1998, Keeley and Zedler 1998, Keeley et al. 2012, Schwilk and Caprio 2011). Table 2 ranks tree species in assessment area YPMC forests by their relative tolerances to shade, frost, temperature, drought, and fire. Figure 3 compares growth rates of young shade-tolerant and -intolerant tree species, and figure 4 compares bark thickness of young trees of the six major conifer species from table 1.

Table 1—Dominant tree species of yellow pine and mixed-conifer forests in the assessment area, their geographic distribution, approximate elevational range within the assessment area, and comparisons of leaf flammability and seed weight

<table>
<thead>
<tr>
<th>Species</th>
<th>Scientific name</th>
<th>Geographic distribution</th>
<th>Elevationa Meters</th>
<th>Leaf flammabilityb</th>
<th>Seed weightc Grams</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ponderosa pine</td>
<td><em>Pinus ponderosa</em></td>
<td>Western United States; southwest Canada; northern Mexico</td>
<td>300–1800 N; 1200–2100 S</td>
<td>Very high</td>
<td>0.02–0.07</td>
</tr>
<tr>
<td>Jeffrey pine</td>
<td><em>Pinus jeffreyi</em></td>
<td>Eastern edge of North American Mediterranean zone; southern Oregon to Baja California</td>
<td>1500–2400 N; 1700–2800 S</td>
<td>Very high</td>
<td>0.08–0.2</td>
</tr>
<tr>
<td>Sugar pine</td>
<td><em>Pinus lambertiana</em></td>
<td>Oregon to Baja California</td>
<td>1000–2000 N; 1400–2700 S</td>
<td>High</td>
<td>0.15–0.3</td>
</tr>
<tr>
<td>Incense cedar</td>
<td><em>Calocedrus decurrens</em></td>
<td>Oregon to Baja California</td>
<td>600–2100</td>
<td>Moderate</td>
<td>0.015–0.07</td>
</tr>
<tr>
<td>Douglas-fir</td>
<td><em>Pseudotsuga menziesii</em></td>
<td>Western United States; southwest Canada; northern Mexico</td>
<td>300–2100 N; 600–2100 S</td>
<td>Low</td>
<td>0.01–0.02</td>
</tr>
<tr>
<td>White fir</td>
<td><em>Abies concolor</em></td>
<td>Southern Rockies; Southwest United States; southern Oregon to Baja California</td>
<td>800–2300 N; 1500–2500 S</td>
<td>Moderate</td>
<td>0.015–0.055</td>
</tr>
<tr>
<td>Black oak</td>
<td><em>Quercus kelloggii</em></td>
<td>Southern Oregon to Baja California</td>
<td>900–1500 N; 1400–2100 S</td>
<td>Very high</td>
<td>3–9</td>
</tr>
</tbody>
</table>

Note: elevations are rounded to the nearest 100 m; N = north, S = south.

McCune (1988) defined groups of North American pines that could be reliably differentiated based on their ecological characteristics. Ponderosa, Jeffrey, and sugar pine all formed part of his “fire resisters” species group, which was characterized by traits and tolerances that promoted the ability to survive low- to moderate-intensity surface fire. This group was generally marked by long needles, thick twigs (which protect growing tissues), thick bark, thick cone scales (which may protect seeds from heat), relatively slow growth, and an extended adolescent period (the mean time to seed-producing age was 16.3 years). McCune (1988) noted that this group of pines was tolerant of frequent surface fire “and reacts to fire more as a stress than as a disturbance.”

Bark thickness of young YPMC tree species as a function of tree diameter is shown in figure 4. Among YPMC tree species, the two yellow pine species have the thickest bark at young ages, but are equaled and even passed by incense cedar at about 10 cm diameter at breast height (d.b.h.) and other species between 25 and 70 cm d.b.h.

### Table 2—Comparative ecological tolerances of common tree species in assessment area yellow pine and mixed-conifer forests.

<table>
<thead>
<tr>
<th>Shade</th>
<th>Frost</th>
<th>Temperature</th>
<th>Drought</th>
<th>Fire</th>
</tr>
</thead>
<tbody>
<tr>
<td>Black oak/western juniper</td>
<td>Madrone</td>
<td>Lodgepole</td>
<td>Red fir</td>
<td>Lodgepole pine</td>
</tr>
<tr>
<td>Ponderosa pine</td>
<td>Douglas-fir</td>
<td>Red fir</td>
<td>White fir</td>
<td>Sugar pine/white fir</td>
</tr>
<tr>
<td>Lodgepole pine</td>
<td>White fir</td>
<td>Jeffrey pine</td>
<td>Western white pine</td>
<td>Incense cedar</td>
</tr>
<tr>
<td>Sugar pine</td>
<td>Sugar pine</td>
<td>White fir</td>
<td>Sugar pine</td>
<td>Jeffrey pine/ponderosa pine/Douglas-fir</td>
</tr>
<tr>
<td>Incense cedar/western white pine</td>
<td>Incense cedar</td>
<td>Douglas-fir/sugar pine/</td>
<td>Douglas-fir incense cedar</td>
<td></td>
</tr>
<tr>
<td>Douglas-fir</td>
<td>Ponderosa pine/Jeffrey pine/red fir</td>
<td>Lodgepole pine/incense cedar/madrone</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Red fir</td>
<td>Lodgepole pine, western white pine</td>
<td>Ponderosa pine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>White fir</td>
<td></td>
<td></td>
<td>Jeffrey pine</td>
<td>Black oak</td>
</tr>
</tbody>
</table>

*a Species arranged from low tolerance (top) to high tolerance (bottom).

*b Least heat tolerant/most cold tolerant on top.

*c Fire tolerance of mature trees. Fir and Douglas-fir seedlings and saplings are less tolerant of fire than yellow pine and sugar pine.

Source: Burns and Honkala 1990, Minore 1979, USDA FS 2013b.

Figure 3—Comparison of predicted small tree (between 3.8 and 7.6 cm diameter at breast height) growth rates of yellow pines, sugar pine, and true firs as a function of the basal area of trees larger than the subject tree. Curves shown are for a site of moderate productivity (site index of 80) and crown ratio of 50 percent.
McCune (1988) defined groups of North American pines that could be reliably differentiated based on their ecological characteristics. Ponderosa, Jeffrey, and sugar pine all formed part of his “fire resisters” species group, which was characterized by traits and tolerances that promoted the ability to survive low- to moderate-intensity surface fire. This group was generally marked by long needles, thick twigs (which protect growing tissues), thick bark, thick cone scales (which may protect seeds from heat), relatively slow growth, and an extended adolescent period (the mean time to seed-producing age was 16.3 years). McCune (1988) noted that this group of pines was tolerant of frequent surface fire “and reacts to fire more as a stress than as a disturbance.”

Bark thickness of young YPMC tree species as a function of tree diameter is shown in figure 4. Among YPMC tree species, the two yellow pine species have the thickest bark at young ages, but are equaled and even passed by incense cedar at about 10 cm diameter at breast height (d.b.h.) and other species between 25 and
50 cm d.b.h. (Dolph 1984, also see van Mantgem and Schwartz 2003). Note that its thick bark notwithstanding, mature incense cedar is somewhat more susceptible to cambial injury than its bark thickness might indicate owing to the bark’s dry, stringy, and highly furrowed nature (Lachmund 1923). Yellow pines are thus well protected from fire during the sapling stage, which is critical in an ecosystem characterized by fires every 5 to 20 years on average (Van de Water and Safford 2011). Other fire-related traits of ponderosa, Jeffrey, and sugar pine include self-pruning of lower branches (Keeley and Zedler 1998) and highly flammable needles and cones, which promote fires that kill competitors that are less fire tolerant (Fonda and Varner 2004, Fonda et al. 1998, van Wagendonk et al. 1998). Black oak, which is a common member of drier YPMC forests in the assessment area, also produces extremely flammable litter (it is the most flammable Western U.S. oak) (Engber and Varner 2012). Douglas-fir is an interesting case. Young Douglas-firs are quite susceptible to fire mortality, but mature individuals can be extremely fire resistant (Agee 1993, Skinner et al. 2006). The species develops very thick bark when mature, and the bark is less prone to sloughing and collecting at the base of the tree than pine bark. Douglas-fir needles are also short and relatively inflammable (Fonda et al. 1998; see below). It has been hypothesized that Douglas-fir’s presence in frequent-fire forests was favored by a variable fire frequency over time, one that periodically included fire-free intervals sufficiently long to allow recruitment of Douglas-fir seedlings into age classes that are more fire resistant (Agee 1993).

Fonda et al. (1998) tested flammability and burning characteristics of the needles of 13 common conifer species from the Western United States. Based on six burning characteristics, ponderosa and Jeffrey pine were ranked first and second, and sugar pine was seventh (giant sequoia was sixth). Douglas-fir and the two tested fir (Abies) species were three of the four lowest ranked species (Fonda et al. 1998). Fonda et al. (1998) noted that the highest ranked species were characterized by fire regimes supported by nonwoody fuels (e.g., needle litter, herbaceous fuels), whereas the lower ranked species typically required woody fuel accumulation over decades to support fires. In another flammability study, de Magalhaes and Schwilk (2012) found that black oak, ponderosa pine, and Jeffrey pine had the highest flammability across most parameters tested. White fir and red fir were less flammable, but they were both more flammable than the fir species tested by Fonda et al. (1998), which is not surprising given their presence in the highly fire-prone Sierra Nevada. Also, de Magalhães and Schwilk (2012) found that flammability of species mixtures of needles was mostly driven by the most flammable species in the mixture, which were Jeffrey pine, ponderosa pine, sugar pine, and black oak. See table 2 for a fire tolerance ranking of assessment area species.
Major tree species in YPMC forests differ notably with respect to the size of their seeds (table 1), the distances these seeds are dispersed, and the number of seeds they produce. In general, heavier seeded species disperse shorter distances, although animal vectors play an important role in the dispersal of some species. Fowells and Schubert (1956) measured seed rain in a YPMC forest during a 9-year period and found that, on average, white fir produced about 2.5 times as much seed as either ponderosa or sugar pine, and incense cedar produced about four times as much seed. Fowells and Schubert (1956) found very high interannual variability, and in some years the pines outseeded the two shade-tolerant species. Working in Yosemite and Sequoia & Kings Canyon National Parks, van Mantgem et al. (2006) concluded from a 5-year study that, standardized by basal area, fir species produced between seven and eight times as much seed as pines (yellow and sugar), and incense cedar produced about three times as much seed. Zald et al. (2008) found the seed rain of shade-tolerant white fir and incense cedar to be 5 to 26 times greater than Jeffrey and sugar pine in southern Sierra Nevada mixed conifer. Stark (1965), in a study of natural tree regeneration after logging, reported that, over a 13-year period, more than 80 percent of surviving seedlings were incense cedar or white fir, 10 percent were sugar pine, and 4.3 percent were yellow pine. Fowells and Schubert (1956) found that white fir and ponderosa pine had similar dispersal distances, which were 50 to 75 percent farther than sugar pine. Vander Wall (2003) found that rodents and birds were important dispersers of pine seeds in YPMC forests. Rodents moved seeds of ponderosa, Jeffrey, and sugar pine about 25 m on average from the parent plants, which approximately doubled the dispersal distances for the latter two species.

Growth rates of YPMC forest trees differ by species and environmental conditions. Yeh and Wensel (2000) found that diameter growth of assessment area YPMC species typically occurs between late March/early April and mid-September, but growth ends earlier at lower elevations (owing to summer drought) and starts later at high elevations (e.g., growth at Blacks Mountain Experimental Forest in the northern assessment area, which ranges from 1700 to 2100 m elevation, generally begins in May). Yeh and Wensel (2000) found that the pines were less disadvantaged by water stress than other tree species. Overall, the most important factors to growth were tied to water availability and included the previous winter’s precipitation (related to water storage) and the current summer temperature (related to water loss).

Light availability is an important environmental driver of growth for plants, and YPMC tree species differ notably in the effects that canopy shading has on

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1 Taylor, A.H. 2013. Personal communication. Professor, Department of Geography, Pennsylvania State University, University Park, PA 16802.
their growth rates (table 2). Keyser (2010) provided equations for growth of major tree species in the assessment area. Small tree (d.b.h. between 3.8 and 7.6 cm [1.5 and 3 in]) growth is modeled as a function of site productivity (site index, or the mean potential height in feet of a free-grown tree at 100 years); crown ratio (ratio of crown length to tree height); and competition/shading by larger trees (basal area of trees larger than the subject tree) (Keyser 2010). Figure 3 shows how both the yellow pine species and sugar pine grow more rapidly than white fir and red fir in open stands but are outgrown by the fir species once the basal area of larger trees exceed about 30 m²/ha (130 ft²/ac). Bigelow et al. (2011) examined the crossover-point irradiance (CPI), the light at which the height-growth rank of pairs of species changes, and found that an understory light environment with 41 percent or more of full sunlight favored ponderosa pine regeneration over white fir. Moghaddas et al. (2008) showed that black oak and ponderosa pine seedling survival was strongly connected to high light environments. Oliver and Dolph (1992) found that ponderosa pine grew more slowly than the other YPMC tree species at even moderate shade levels, while sugar pine showed quite strong growth at 50 to 60 percent of full sunlight. They noted that sugar pine appeared to be adapted to exploit small forest gaps and showed some characteristics of the more shade-tolerant species (e.g., fir).

YPMC tree species also differ with respect to their tolerances to frost, heat and cold, and drought. In general, shade-tolerant species from low- and middle-elevation forests (e.g., Douglas-fir and white fir) are more susceptible to frost damage than the pine species and red fir (table 2). Not surprisingly, the most cold-tolerant species are those from higher elevations, and the most heat tolerant are those from lower elevations and warmer microsites (table 2). Drought tolerance is a very important trait, as precipitation in the assessment area is highly variable between years, and models of future climate in the assessment area project increased occurrence of drought (Dettinger 2011, Liu et al. 2010). The true fir species are the least drought-tolerant members of YPMC forests in the assessment area, whereas sugar pine, Douglas-fir, and incense cedar are intermediate. The yellow pines and black oak are the most drought-tolerant species in YPMC forests (table 2).

In summary, the major YPMC tree species are differentially adapted to the physical and biotic environment in the assessment area, and the different tolerances of these tree species play a major role in determining forest composition, structure, and function. Considering the overwhelming importance of fire and water availability to YPMC forest ecology and management in the assessment area, perhaps the most important distinction to be made is between those species that are highly tolerant of fire and drought but intolerant of shade (black oak and the yellow pines), versus those that are less tolerant of fire and drought but grow relatively well in
low-light conditions (white fir, incense cedar, and Douglas-fir; recall, however, the high-fire tolerance of mature individuals of the latter species). These ecological differences are at the root of general topographic tendencies in forest composition across the assessment area. Locations with higher water availability (north aspects, lower slopes, concave topography, higher elevations) tend to support higher densities of the shade-tolerant, fire-intolerant species, whereas sites with lower water availability (south aspects, upper slopes and ridgetops, convex topography, lower elevations) are more likely to be dominated by shade-intolerant, fire-tolerant species.

These ecological differences among species translate into successional differences as well. In regions that can support forest cover, trees that specialize in high-light environments will be generally restricted to early-successional stages, and locations where ecological factors slow successional processes. The very high frequency of fire before Euro-American settlement (see below) played such a role, and essentially kept YPMC forest in the assessment area in a state of arrested development; where early-successional species such as the yellow pines and black oak were able to maintain canopy dominance, even in places of relatively high water availability. In the general absence of fire disturbance over the past century, successional processes have become “unblocked” and more competitive; later-successional species like the firs and incense cedar have come to dominate most of the YPMC belt in the assessment area. Early-successional species continue to dominate where fire regimes have not been much perturbed, and where local soil productivity acts to slow succession. We will make reference to these differences in fire, shade, and drought tolerance and successional relationships throughout this report.

**Temporal Variability in the Ecological Setting**

The mutable nature of the climate has never been more apparent than today, with human inputs to the atmosphere rapidly increasing greenhouse gas levels and global temperatures. Although the concept of “climate” is one that suggests long-term stability, climates are constantly changing, and climatic variability throughout the Holocene Epoch has had major effects on YPMC forests in the assessment area. It is also important to note that temporal changes in climate have not been uniform across the assessment-area landscape, and the timing of changes in temperature and precipitation, as well as biotic responses, differ from one area to another. Nonetheless, some useful generalizations can be made.

The Holocene Epoch is now considered to have begun about 12,000 YBP (years before present). The entire epoch falls within a broadly defined “interglacial” period. The Earth’s climate has been in “glacial” periods for about 90 percent of the past 850,000 years, and interglacial periods as warm as the Holocene are
relatively rare (Tausch et al. 2004). Overall, mean annual temperatures have fluctuated by 3 to 6 °C through the Holocene, and precipitation has also risen and fallen. Researchers generally divide the Holocene into three periods that are defined by broad (and somewhat ill-defined) changes in temperature and precipitation. The Early Holocene stretches from the beginning of the epoch to 8,000 or 7,000 YBP, and was characterized by post-glacial warming, but generally cool and moist conditions compared to today; however, data from the northern Great Basin portion of the assessment area (e.g., Warner Mountains) suggest that post-glacial conditions were somewhat warmer and drier than today (Minckley et al. 2007). Many closed basins east of the Sierra Nevada crest supported large rain- and snowmelt-fed lakes at the beginning of the Holocene, but most of these dried or decreased greatly in size over the ensuing couple of millennia (Minnich 2007). At the beginning of the Holocene, elevations that currently support YPMC forests were largely vegetated by high-elevation sagebrush and grass species, with a minor presence of pine and juniper (Anderson 1990, Woolfenden 1996). By 9,000 to 10,000 YBP, however, conifer forests had established themselves in most of these areas (Minnich 2007), except in the Great Basin portions of the assessment area (Minckley et al. 2007).

The Middle Holocene, about 8,000 to 4,000 YBP, is also often referred to as the “Xerothermic,” “Hypsithermal,” or “Altithermal” Period. Climates became much warmer and drier, with the driest and warmest conditions occurring around 6,000 YBP. Glaciers completely disappeared from the Sierra Nevada (Clark and Gillespie 1997), and lake levels dropped precipitously. For example, Lake Tahoe was apparently permanently below the Truckee River outlet elevation during the period between about 6,300 and 4,800 YBP (Lindström 1990). Fire frequency also increased during the Middle Holocene, at least in places that were able to maintain plant cover (Beaty and Taylor 2009, Woolfenden 1996). Paleoecological data suggest that forests of fir and pine were replaced by oak, sagebrush, and juniper in many areas, and forest structure was likely very open, with abundant understory shrubs. Conifers invaded formerly moist areas of meadow, and desert plant and animal taxa migrated upslope (Anderson 1990, Minnich 2007).

The Late Holocene (4,000 YBP to present) has been generally characterized by cooling, with some warmer periods. Precipitation increased, and small glaciers began to form again in the Sierra Nevada. Millar and Woolfenden (1999) suggested that the basic spatial and compositional outlines of modern Sierra Nevada ecosystems developed by the beginning of the Late Holocene (note that Minckley et al. [2007] suggested that ecological conditions similar to today were already in place in the northwestern Great Basin by the Middle Holocene). As temperatures cooled, available moisture rose, and fir and incense cedar abundance increased relative to pine and oak; giant sequoias began to colonize their current groves (Millar and Woolfenden 1999).
White fir was a mostly minor component of assessment area YPMC forests before the general cooling and increase in precipitation seen over the past 4,000 years. Oaks have declined in importance since the end of the Middle Holocene. Earlier in the Holocene, sagebrush was a major component of low-elevation west-side landscapes, but now it is primarily restricted to higher elevations on the east side of the assessment area (Anderson 1990). Fire has been present as an important ecosystem process ever since deglaciation at the beginning of the epoch, but the presettlement period, characterized by frequent fire and large areas of fire-adapted vegetation, began during the Late Holocene for most of the assessment area (Millar and Woolfenden 1999).

The past 1,000 years of the Holocene have been marked by short-term changes in temperature and precipitation that have had major impacts on assessment-area ecosystems (Millar and Woolfenden 1999, Minnich 2007, Woolfenden 1996). Between about 900 and 1100 CE, and from 1200 to 1350, two long drought periods (the “medieval droughts,” or collectively, the Medieval Warm Period) led to very low levels in lakes and streams (Stine 1994) and increased fire frequencies. This was followed by a shift to cooler temperatures known as the Little Ice Age, initiated apparently by a series of massive volcanic eruptions that caused atmospheric reflection of solar radiation (Miller et al. 2012a). The Little Ice Age lasted from about 1400 to 1880, and the period between 1650 and 1850 was the coolest since the Early Holocene (Stine 1996). Glaciers expanded in the Sierra Nevada, tree line dropped, and fire frequencies moderated. Minnich (2007) noted, however, that there is little evidence for major changes in vegetation composition during the Little Ice Age. It is important to underline that the period of the Little Ice Age is also the period most commonly used as an historical reference period for restoration planning in the Western United States.

Most recently, human emissions to the atmosphere have resulted in renewed warming, even though Earth orbital cycles should be resulting in a cooling trend (Ruddiman 2005). Current temperature trends include increased temperatures especially at nighttime, a decrease in the number of days with below-freezing temperatures, and an increase in the number of extreme-heat days. The 20th century was one of the wettest centuries in the Late Holocene (Stine 1996), and recent precipitation trends in most of the assessment area have been steady or positive (Safford et al. 2012b). Interannual variability in precipitation is up at many stations in the assessment area, and the proportion of precipitation falling as rain versus snow is increasing; as a result, the depth of the winter snowpack is decreasing, except in the southern assessment area, where mountain elevations are very high (Safford et al. 2012b). Recent trends in fire activity are positive, with burned area, fire size, and fire frequency all rising in assessment area YPMC forests, accompanied by an increase in fire severity on lands where fuels have accumulated as a result of long-term fire exclusion policies (Mallek et al. 2013, Miller and Safford 2012, Miller et al. 2009b, Steel et al. 2015).
Cultural Setting

Humans have actively managed yellow pine and mixed-conifer forests for resource benefits for thousands of years. YPMC forests are generally productive and accessible to denser human settlements at lower elevations, which has made them attractive to humans for a variety of uses. The period of human settlement and forest management in California may be divided into four time periods—the American Indian era, the Euro-American settlement era, the fire-exclusion era, and the ecosystem management era (Sugihara and Barbour 2006). Each of these periods was characterized by distinct cultural values and practices that shaped the management of YPMC forests. Of particular importance to the management of YPMC forests throughout these eras are the role of fire and use of timber resources. The suite of human uses in YPMC forests has been fairly consistent among the different parts of the assessment area. The southern Cascade Range and Modoc Plateau have generally had lower human population densities than the Sierra Nevada region and may have been less affected by some human activities (e.g., urbanization, agriculture, mining), but the high accessibility of much of the forest land in the southern Cascades and Modoc Plateau led to high levels of logging and grazing use (Dasmann 1965, Riegel et al. 2006, Skinner and Taylor 2006, van Wagtendonk and Fites-Kaufman 2006).

American Indians have lived in present-day California for at least 11,000 years (Anderson 2005, Sugihara and Barbour 2006). During this period, human use of YPMC forests was common, although permanent settlements were often located at lower elevations (Anderson 2006). Forest uses were generally related to providing food and materials for building, and often incorporated fire. American Indians set fires in YPMC forests to promote oak growth for acorn harvesting, encourage sprouting of hardwoods for basketry materials, increase understory vegetation for wildlife forage and hunting purposes, and reduce woody fuels that could otherwise lead to severe fires (Anderson 1993). Impacts to YPMC forests were likely not uniform, but rather targeted at specific areas, including those near water sources, hunting grounds, and settlements (Vale 1998). This variable pattern of management likely led to increased heterogeneity of forest structure across the landscape (Anderson 2006, Vale 2002).

European settlement in California began with settlement by Spanish explorers and missionaries during the late 1700s, but impacts in YPMC forests were minimal until settlement by Euro-Americans from the Eastern United States, beginning with the 1850s gold rush that initiated a period of intense resource use and extraction (Beesley 1996, Stephens and Sugihara 2006). The primary impacts of this wave of settlement were felt in the Sierra Nevada portion of the YPMC forest belt,
where mining operations were established, and extensive logging was conducted to support these mining operations (Stephens and Sugihara 2006, van Wagtendonk and Fites-Kaufman 2006). Initial logging operations were conducted with animal teams, while railroad logging opened previously inaccessible areas to extraction after the 1890s (Beesley 1996, Conners 1997, McKelvey and Johnston 1992, TCHS 2016). Yellow pine and mixed-conifer forests were the most intensively logged of all forest types in California, and logging often focused on removing the largest trees from a stand (Leiberg 1902, Stephens and Sugihara 2006). In some more accessible locations, such as the Lake Tahoe basin, entire watersheds were clearcut during this period. In addition to direct impacts to YPMC forests from mining and logging, grazing from domesticated livestock increased dramatically during this period (Sugihara and Barbour 2006), although there is uncertainty about the extent to which grazing reduced grass cover in YPMC forests, which is presently quite low (North et al. 2015).

During the early 20th century, with the professionalization of America’s forestry industry and the creation of protected forest reserves for conservation, resource managers began to argue that YPMC forests were understocked because of wildfires and excessive historical use of fire (Show and Kotok 1924, Stephens and Sugihara 2006). This perspective became more widespread after 1910 when large fires burned through the northern Rocky Mountains, and extensive resources for managing federal lands were dedicated to fire suppression, including within the YPMC belt (Sugihara and Barbour 2006). This fire-suppression era coincided with a large increase in logging demand following World War II, when timber yields in Sierra Nevada YPMC forests increased tenfold (Beesley 1996). Much of this demand was met by high-grade logging on national forest lands, as most of the private land in the YPMC belt had already been harvested (Ruth 1996). The combination of timber harvesting and fire suppression led to structural changes throughout YPMC forests, with losses in the large tree component, general increases in stand density, shifts toward even-aged structure, loss of structural heterogeneity, and increases in fir dominance within areas that had previously been primarily yellow pine stands (Benedict 1930, McKelvey et al. 1996, North et al. 2015, Skinner and Taylor 2006, Thorne et al. 2008).

The 1960s marked the beginning of the transition to the ecosystem management era, although fire suppression and timber harvesting remained dominant practices through much of this era (Sugihara and Barbour 2006). With much of the YPMC belt situated on Forest Service land, the multiple-use mandate of that agency began to be cited in legal challenges to the large-scale timber harvesting operations. Other cultural values placed on YPMC forests began to gain traction among the public,
including ecosystem services such as water and carbon storage, wildlife habitat, and recreation (Ruth 1996). The Wilderness Act of 1964 and subsequent wilderness designations protected many alpine and subalpine regions from extractive activities, but most designated wilderness in California did not cover YPMC forests, with the exception of some national parks. Most YPMC forests on federal lands therefore fall under the purview of both the National Environmental Policy Act of 1969 and the National Forest Management Act of 1976. Environmental organizations have used these pieces of legislation to prevent timber harvests in YPMC forests on the grounds of their negative impacts to habitat for wildlife species including the California fisher and the California spotted owl (Ruth 1996, Truex and Zielinski 2013). During the 1970s and early 1980s, approximately 69 percent of growth on timbered national forest lands in the Sierra Nevada (primarily from YPMC forests) was harvested; however, recent rates of harvesting have slowed dramatically (Ruth 1996). Associated with this downward trend in timber harvesting has been a decline in the processing capacity for timber and other wood products in the bioregion: California-wide, the number of forest product mills dropped by approximately 85 percent between 1968 and 2000, and the number has declined further since then (Laaksonen-Craig et al. 2003). Today, many communities that were once supported by the timber industry have suffered economic downturns or have turned to other sources of jobs such as tourism and forest recreation.

Despite growing awareness of the importance of frequent fire to YPMC forest health, fire suppression remains the dominant federal policy in these forests (Stephens and Ruth 2005). A number of obstacles prevent the increased use of fire as a tool for ecosystem management in these forests. Foremost among these are the public health consequences of smoke in communities, the rapidly increasing extent of wildland-urban interface within YPMC forests, and opposition to landscape-level fuel-reduction treatments the Forest Service asserts are needed to safely reintroduce low- to moderate-severity fire (North et al. 2012a, North et al. 2015). Human population densities in some areas are greater than 100 times what they were prior to Euro-American settlement, leading to considerable restrictions on fire use (Sugihara and Barbour 2006). The Sierra Nevada foothills region is one of the most rapidly growing areas of California, with much of the development occurring directly below the YPMC forest belt and susceptible to risks associated with fire spread and smoke production (Bryant and Westerling 2014).
Chapter 2: Methods

The natural range of variability (NRV), or “range of natural variation,” was defined by Landres et al. (1999) as:

…the ecological conditions, and the spatial and temporal variation in these conditions, that are relatively unaffected by people, within a period of time and geographical area appropriate to an expressed goal.

“Historical range of variation” (HRV) is a related concept, and was defined by Wiens et al. (2012) as:

…the variation of ecological characteristics and processes over scales of time and space that are appropriate for a given management application.

HRV was developed to allow the explicit incorporation of human influences on ecosystems into the analysis, because in most places on Earth humans have been major ecological players for millennia. We evaluate human influences on assessment area ecosystems in our analyses, thus they may have been more properly called HRV assessments than NRV assessments, but Forest Service guidance for implementation of the 2012 Planning Rule adopts the term “natural range of variability,” so we have stuck with this term.

The Forest Service Handbook 1909-12 chapter 10 (USDA FS 2013c), which guides the development of the resource assessments that must precede forest plan development, defines NRV as:

Spatial and temporal variation in ecosystem characteristics under historic disturbance regimes during a reference period. The reference period considered should be sufficiently long to include the full range of variation produced by dominant natural disturbance regimes, often several centuries, for such disturbances as fire and flooding and should also include short-term variation and cycles in climate. “Natural range of variation” (NRV) is a term used synonymously with historic range of variation or range of natural variation. The NRV is a tool for assessing ecological integrity, and does not necessarily constitute a management target or desired condition. The NRV can help identify key structural, functional, compositional, and connectivity characteristics, for which plan components may be important for either maintenance or restoration of such ecological conditions.

NRV/HRV assessments are tools used by managers to bring insights from historical ecology to resource management (Hayward et al. 2012). NRV/HRV characterizes variations in ecosystem function, structure, and composition over scales of time and space. The basic purpose of NRV/HRV is to define the bounds of ecosystem behavior or trends in those bounds. As Morgan et al. (1994) put it:
“The concept of HRV (NRV) provides a window for understanding the set of conditions and processes that sustained ecosystems prior to their recent alterations by humans.” Morgan et al. (1994), Manley et al. (1995), Landres et al. (1999), and Wiens et al. (2012) all listed the purposes of conducting HRV/NRV assessments and the issues that must be considered in the assessment. These include the ecosystems of interest, the spatial and temporal scales of analysis, the ecological indicators to be assessed, whether to include human influences, whether to use only historical information or to use contemporary reference conditions and modeling as well, and so on.

According to Manley et al. (1995), HRV/NRV assessments should include the following steps:

1. Determine key ecosystem elements (e.g., functions/processes; structures/patterns; composition),
2. Identify measurable indicators for those ecosystem elements (e.g., fire frequency, tree density, species diversity), and
3. Estimate values for the indicators over the selected HRV reference period.

The appendix contains details, in outline form, of the process by which the Forest Service Pacific Southwest Region Ecology Program carried out these steps for the 11 ecosystem types assessed.

We are greatly indebted to the thorough HRV assessments funded by the Forest Service’s Rocky Mountain Region (Region 2) in the early 2000s (e.g., Dillon et al. 2005, Meyer et al. 2005, Veblen and Donnegan 2005). These valuable documents provided a sort of template for our efforts and set a high bar against which to gauge our own efforts.

**Historical Reference Period**

Morgan et al. (1994) and Wiens et al. (2012) noted that the temporal scale of analysis will always be constrained by our ability to look clearly back through time. Certain data types permit insight into ecological patterns and processes thousands or millions of years in the past, but most reasonably decipherable data sources extend back only decades and, in some cases, a few centuries. In addition, climate and other environmental conditions tend to diverge more from current conditions the farther one goes back into the past. As a result, most NRV/HRV assessments use the past 100 to 400 years as their baseline or “reference” period. In this NRV assessment, our principal reference period was the three to four centuries before significant Euro-American settlement of the assessment area, i.e., the 16th century to the late 19th century. It is important to underline that this reference period is coincident with the Little Ice Age, and current warming trends are making future use of NRV/HRV reference conditions
as management targets gradually more tenuous (this, however, does not diminish the value of NRV/HRV assessments, just the ways in which they are used) (Millar et al. 2007; Safford et al. 2012a, 2012c). Therefore, as recommended by Manley et al. (1995) and Jackson (2012), we also collected and interpreted information as far back as the beginning of the Holocene Epoch (12,000 years before present [YBP]) when it was available. We were especially interested in patterns and processes from warmer, drier periods in the past (Xerothermic Period, medieval droughts), because most future climate projections for the assessment area project much warmer and somewhat drier conditions (at least during the growing season) by the end of the current century.

**Spatial Scale**

NRV/HRV analyses are focused on change over time, but variation in space must also be considered. Our historical and contemporary reference data sources are more often than not derived from specific locations or landscapes in the assessment area, but our analysis is intended to apply to the bioregion as a whole. Wherever possible, we sought data that represented the variety of different geographic regions and environmental situations that are found in the assessment area. Usually, though, we simply had to accept the limitations of those data we could find, and use inference and our understanding of environmental variation across the bioregion to extend those data points to the larger assessment area. We report the geographic locations of our data sources throughout the report.

**Information Sources**

Since the Sierra Nevada Ecosystem Project report was published in 1996, there has been a veritable explosion of scientific information on the ecology and current and past status of yellow pine mixed-conifer forests in the assessment area. Although we would like more data on every indicator we assessed, our principal problem was trying to assemble and filter all of this information into a coherent whole.

This NRV assessment is based on both historical and contemporary reference sites and information sources. Historical data are especially useful when they precede the onset of major anthropogenic disturbances and alterations that have degraded assessment area ecosystems. These sorts of data are few, however, and, in most cases, we were forced to resort to evaluation of information sources that postdated the settlement of Euro-Americans in the assessment area. We used modern-day data from reference ecosystems whenever possible. By “contemporary reference ecosystems,” we mean current-day ecosystems that have suffered relatively little degradation and may serve as a more natural reference against which degraded ecosystems may be compared. There are many advantages to using
contemporary reference sites in NRV/HRV analyses, including the availability of modern data on ecosystem condition, and the fact that climatic and atmospheric conditions in current reference sites and current degraded sites are more or less equivalent. Safford et al. (2012a: 57) noted that:

We use historical data principally to understand ecological events and processes that we cannot observe firsthand, but directional changes in the baseline state (climate, air, water, soil, etc.) mean that historical conditions may make poor templates for the future. To compensate, contemporary reference ecosystems that are functioning as we desire should form part of the package of information that underlies restoration and resource management.

The problem is that human alteration and degradation of assessment area ecosystems is so pervasive that identification of appropriate reference ecosystems is difficult in all instances, and impossible in some.

In our assessment, we used direct data analysis and interpretation whenever possible, and we resorted to inference where necessary and justifiable. This NRV assessment includes comparisons to current conditions, as well as a summary of the literature regarding possible future trends, whenever that literature existed. Our focus was on peer-reviewed publications, including papers in press or soon to be in press; government publications; Forest Service and other federal and state agency data; and, in some cases, academic theses or dissertations. Because information on the historical state of some ecosystems and ecological processes and patterns is scarce, we also refer to published anecdotal information from the mid-19th to early 20th centuries in some cases. We do not refer to anecdotal information from more recent times.

The Forest Service’s Forest Inventory and Analysis (FIA) program is the U.S. national forest inventory. Plots are found across the United States and are located randomly within a grid defined by latitude and longitude. We used a compilation of the most recent FIA data in the assessment area provided by the Pacific Southwest Region Remote Sensing Laboratory to provide current-day data on many forest structure and composition variables. It is important to stress that FIA is a statistically robust sample of all stand conditions across the assessment area, including areas with reduced tree density and cover owing to natural disturbances or harvest.

**Determination of Deviation From NRV**

Like the Rocky Mountain Region HRV analyses (e.g., Meyer et al. 2005), our NRV assessments are based primarily on a “range of means” approach. Variation in data can be characterized in a number of ways. For example, the entire range of variation in a dataset is captured by reporting the extremes (minimum and maximum), but
these extreme values more often than not represent rare cases that do not provide a clear picture of central tendencies. Use of extreme values to bound ecosystem variation makes NRV/HRV assessment more or less impossible, as nearly all values for a variable are theoretically possible at some time and at some place on the landscape. Generation of a standard deviation or standard error (the latter is standard deviation scaled to the sample size) is the usual statistical method for reporting variation, but in NRV/HRV analyses, the availability of data and sample sizes are often inadequate. A middle path is to base the assessment on an estimate of the range of means from multiple sources for a given variable. This produces a narrower, more discernible, and probably more meaningful range of variation that can be quantitatively or qualitatively compared to modern data. See figure 1 in Meyer et al. (2005) for an excellent depiction of how NRV/HRV varies with the type of variation used and the spatial scale of analysis.

Determination of deviation from NRV was accomplished by comparing the modern range of variation for some indicator variable (ideally represented by a mean, median, and standard deviation) with the range of means for the same variable from the NRV period or contemporary reference sites. In practice, direct statistical comparison was rarely possible, owing to small sample sizes in the reference sources, the lack of measures of statistical variation, orders-of-magnitude differences in sample sizes between current and historical data when multiple historical data points did exist, or not uncommonly, the lack of concrete quantitative measures in the historical dataset. Our assessment of current deviations from NRV was necessarily deductive in nature, and we came to conclusions about the status of specific variables based to a great extent on our general knowledge about the ecosystems in question. Table 11 on page 178 summarizes our conclusions about current deviation (or “departure”) from the NRV for key ecosystem elements in YPMC forests in the assessment area.

Note that measuring departure from historic or reference conditions and incentivizing and operationalizing realistic action are not the same thing, and this NRV assessment attempts only to measure departure. See Moritz et al. (2013), Thompson et al. (2009), and Wiens et al. (2012) for discussions on how socioeconomic considerations are really at the heart of the matter, and how they might be integrated with NRV/HRV types of information.
Chapter 3: Natural Range of Variability Descriptions

Function (Including Disturbance)

Extreme Climatic Events

Drought—

NRV—California’s Mediterranean climate is characterized by an annual drought of 3 to 6 months at most weather stations in the assessment area (Major 1988). This is a longer drought than most stations in the northern Rocky Mountains experience in a 50-year period. Aside from the predictable annual warm season drought, California is also characterized by extremely wide variation in annual precipitation and a high dependence of annual precipitation on a small number of very wet winter storms (note that this variability in precipitation decreases with latitude) (Dettinger et al. 2011). Most California plant taxa are therefore adapted to significant periods of time without access to atmospheric water. Nonetheless, periods of multiple, consecutive dry years can have major impacts on yellow pine and mixed-conifer (YPMC) forests (e.g., Guarín and Taylor 2005), as evidenced by the massive die-off of conifer trees in the San Bernardino Mountains after the drought of the late 1990s and early 2000s, and the even more massive tree mortality event that was beginning in the southern Sierra Nevada as we completed this assessment. Drought itself is usually not the proximal cause of tree mortality, however, as drought-induced stress also leads to greater susceptibility to insects and disease, and dry years tend to support more fire as well (Allen et al. 2010, Fettig et al. 2007, Logan et al. 2003, Savage 1994). Forest density can also intensify the effects of drought on tree mortality, through stress brought about by increased competition for water (Dolph et al. 1995, Innes 1992, Young et al. 2017).

The Holocene Epoch, which began about 12,000 years ago, has been characterized by a fairly stable climate, but that is only in geological terms. Over the past 7,000 to 8,000 years, dry climatic periods have occurred on average every 80 to 260 years, with durations of droughts lasting 20 to 100 years on many occasions. Between 8,000 and 6,300 years before present (YBP), drought conditions dominated in the northern and eastern assessment area (during the Xerothermic Period), and many large lakes in and near the assessment area either dried completely (e.g., Owens Lake) or dropped significantly in depth (e.g., Lake Tahoe). Aridity is thought to have been less pronounced in the southwestern assessment area (Yosemite National Park and points south) and to have eased earlier, perhaps as early as ±7,500 to 7,000 YBP. Over the ensuing 2,000 to 3,000 years, aridity lessened and distinct wet and dry climatic phases occurred. During the past 2,500 years, decades-long droughts have occurred on multiple occasions, with especially severe events occurring around 800 and 650 YBP (Benson et al. 2002, Mensing et al. 2004, Minnich 2007, Stine 1994).

Most California plant taxa are therefore adapted to significant periods of time without access to atmospheric water. Nonetheless, periods of multiple, consecutive dry years can have major impacts on yellow pine and mixed-conifer forests.
Biogeographically, forest vegetation in the assessment area responds to warmer and drier climatic periods by retreating to moister, cooler locations. For example, the alpine tree line rises during warmer periods and retreats with cooling; under extended drought, trees become more restricted to areas with permanent water. Lake Tahoe dropped below its sill during a number of Holocene droughts, and tree stumps can now be found under the surface of the lake. The Walker River between Bridgeport and Topaz Valleys was dry during these droughts, and trees grew directly in the current river channel (Stine 1994). Species composition also changes in response to drought. During dry periods in the Holocene, pollen data show that oaks, pines, junipers (*Juniperus*), and dryland shrubs (e.g., *Artemisia*, *Atriplex*) dominated much of the assessment area; less drought-tolerant species like the firs and incense cedar were reduced, but then increased when conditions became cooler or wetter (Minnich 2007).

One of the main drivers of vegetation structure and composition is fire, and long-term shifts to drier conditions can greatly enhance fire activity (frequency). Note, however, that climatic drying can also act to decrease the size of fires, as such conditions may lead to less accumulation of live fuel (Swetnam 1993). Beaty and Taylor (2009) studied Holocene fire in the Lake Tahoe basin and showed that maximum fire activity occurred at around 6,500 YBP, at the height of the Xerothermic Period. Fire became gradually less prevalent thereafter, with notable peaks occurring in later drought periods at about 3,000 and 1,000 to 800 YBP (Beaty and Taylor 2009). Other authors have shown similar drought-fire interactions in and around the assessment area (e.g., Enfield et al. 2001, Swetnam and Baisan 2003, Trouet et al. 2010, Whitlock et al. 2003). Because the yellow pine species and black oak are more fire and drought tolerant (table 2), drying climates and increasing fire frequency would be expected to benefit them over less tolerant tree species like the firs. **Comparison to current**—By many accounts, the late 19<sup>th</sup> and 20<sup>th</sup> centuries were anomalously wet and have experienced few drought events compared to earlier periods of the Holocene (Haston and Michaelsen 1997, Hughes and Brown 1992). Numerous authors have argued that the climate of the recent past is likely not a reliable roadmap for the future of the Western United States (Millar et al. 2007, Saxon et al. 2005, Williams and Jackson 2007). Nonetheless, Allen et al. (2010) noted that scientific documentation of climate-related forest mortality has been rising over the past few decades, and some recent drought episodes have been among the most severe of the past few centuries (e.g., the recent 4-year drought in California may have been the worst in 1,200 years or more) (Griffin and Anchukaitis 2014). The increasing abundance of drought-intolerant species, and the very high stem densities and structural homogeneity that characterize much of the contemporary YPMC forest in the assessment area, have primed these forests for more severe impacts of drought.
and related disturbance factors than would have likely been the case under reference conditions. A major concern is that larger and older trees are often more prone to drought-induced mortality (Allen et al. 2010). In many places, this may be exacerbated by higher densities of younger trees in the surrounding forest (Dolph et al. 1995, Ritchie et al. 2008). Higher levels of drought stress have recently been linked to higher probabilities of mortality resulting from fire (van Mantgem et al. 2013).

**Future**—Forest-landscape change driven by drought tends not to be gradual, but rather episodic and rapid, because trees grow relatively slowly but die quickly (Franklin et al. 1987). Increased drought and heat have been amply documented to increase tree mortality around the globe, principally owing to secondary factors like insects, disease, and fire (Adams et al. 2009, Allen et al. 2010, van Mantgem et al. 2013). McDowell et al. (2008) outlined three interacting mechanisms by which drought can lead to broad-scale forest mortality:

1. Extreme drought and heat kill trees through cavitation of the xylem water column.
2. Long-term water stress results in carbon deficits and metabolic limitations that result in carbon starvation and reduced capacity to defend against attack by biotic agents like insects and disease.
3. Extended warm spells during droughts can facilitate rapid population growth in these biotic agents, which then overwhelm the already stressed tree hosts.

Under warmer future climates, both drought-avoiding and drought-tolerating tree species may thus be negatively affected; the former by carbon starvation, the latter by hydraulic failure (McDowell et al. 2008).

Bachelet et al. (2001) modeled future vegetation cover and biomass under a number of different global circulation models (GCMs). One of their modeled response variables was a drought area index related to the Palmer Drought Severity Index. The authors found that the area of the United States subjected to drought stress rose approximately linearly with increasing temperatures. The assessment area was not identified as especially sensitive to increasing drought under their scenarios, but most of their scenarios assumed increasing precipitation in California during the 21st century, which runs counter to most of the more recent GCMs (Dettinger 2005).

Liu et al. (2010) modeled the Keetch-Byram Drought Index for the 2070 to 2100 period under different GCMs. They projected increasing drought potential across most of the contiguous United States, including eastern and southern California and the assessment area. In the Western United States, most of the summer drying trend was predicted to depend on warming temperatures more than reductions in precipitation. They concluded that much of their study area will experience greatly increased fire potential during the course of the coming century.
Extreme precipitation events—

*NRV and comparison to current*—On a year-to-year basis, California experiences the most variable precipitation regime in the United States. In addition, no other state depends as much on a few large storms to generate precipitation: in the Mediterranean-climate part of California, a third to half of annual precipitation falls in only 5 to 10 wet days per year. So-called atmospheric rivers (ARs) generate 20 to 50 percent of the state’s precipitation totals (Dettinger et al. 2011). Such ARs are narrow bands (less than 200 km wide typically) of concentrated water vapor that develop over the oceans and direct large amounts of moisture toward continental areas. Owing to the influence of periodic ARs developing over the tropical Pacific, California experiences more extreme precipitation events than any other part of the United States, including the hurricane-affected Gulf Coast (Dettinger et al. 2011).

The dependence of annual precipitation totals on extreme precipitation events leads to highly variable streamflows, as well as the propensity for landslides and other earth movement on unstable geologic substrates (Kerr 2006). Because they are more maritime, more tectonically active, and constituted of less consolidated bedrock, the California Coast Ranges are more heavily affected by flooding and earth movement associated with ARs, but the Sierra Nevada also experiences massive precipitation events that cause widespread flooding and landsliding. An example is the New Year’s storm of January 1997, which forced evacuations of 120,000 people, flooded Yosemite Valley, and closed Highways 140, 50, and 395 for months. Dettinger (2011) noted that ARs accounted for all major historical floods in some California river systems.

There is little direct evidence to compare the occurrence of ARs and other extreme precipitation events in the assessment area between current and past time periods. Paleoecological studies have documented the enhanced occurrence of extended droughts at different times of the Holocene Epoch (see “Drought” above), and because a large proportion of California precipitation stems from AR events, we can hazard the inference that major changes in precipitation must involve changes in the frequency of ARs.

*Future*—Dettinger (2011) modeled the occurrence and intensity of ARs under a variety of future climate change scenarios. He found that, under the most realistic emissions scenarios, average AR statistics did not change much, but the extremes changed markedly. For example, the frequency of larger than average ARs increased, the number of years with many AR episodes rose, and the temperatures of AR storms rose, which could feed into greater occurrence and magnitude of winter flooding and associated earth movement (Dettinger 2011).
Fire

Background: fire regime—
Ecological disturbances can be classified according to their characteristics, including frequency, size, season, intensity, severity, pattern, and so on. A “fire regime” describes the manner in which fires tend to occur in a given ecosystem, in a generalized sense and averaged over many fires over a long period of time. Fire regimes necessarily simplify a very complex phenomenon, but they offer a convenient and useful way to better understand and manage wildland fire (Sugihara et al. 2006).

Under presettlement conditions, yellow pine and mixed-conifer forests in the Sierra Nevada supported fire regimes characterized by frequent, low- to moderate-severity fires (Agee 1993; Arno 2000; Barbour et al. 1993, 2007; Skinner and Taylor 2006; van Wagtendonk and Fites-Kaufman 2006). These characteristics placed presettlement YPMC forests in Fire Regime I (fire-return intervals [FRI] 0 to 35 years, low to moderate severity), using the Schmidt et al. (2002) classification, which, although it uses only two fire regime attributes, has become somewhat of a national standard. The YPMC forests in the assessment area supported fire regimes limited principally by the amount of available fuels (rather than by fuel moisture), because a 3- to 6-month drought is typical in California’s Mediterranean climate, and temperatures at low and moderate elevations are very warm in the summer (Steel et al. 2015). As elevation increases, the role of fuel moisture becomes gradually more important, until it becomes a more important driver of fire regime than fuel load (Agee 1993, Miller and Urban 1999b). Various authors have identified the boundary between high-elevation mixed-conifer and red fir forests (2000 to 2300 m or more, depending on latitude), which is the elevation of the average freezing limit in winter storms and the zone of maximum snowfall, as a zone of major transition in fire regimes (Mallek et al. 2013, Miller and Urban 1999b, Safford and Van de Water 2014, Sugihara et al. 2006). Temporal and spatial variability in regional climate also plays a major role in driving fire regimes, and ocean-atmosphere phenomena like ENSO (El Niño-Southern Oscillation), the PDO (Pacific Decadal Oscillation), and the PNA (Pacific-North America Pattern) are important determinants of burning patterns across the Western United States (Taylor and Scholl 2012; Trouet et al. 2006, 2010).

Today, because of human influences since Euro-American settlement, including timber harvest, grazing, and fire suppression, many YPMC forests in the assessment area now support conditions that are more characteristic of Fire Regimes III (FRIs of 35 to 200 years, moderate severity) and IV (FRIs of 35 to 200 years, high severity) (Schmidt et al. 2002). These are often referred to as “climate-limited” fire regimes, as fuel moisture, rather than fuel load, tends to drive fire occurrence and behavior (Agee 1993, Schoennagel et al. 2004, Steel et al. 2015). Evidence for this
shift can be seen in the growing importance of climate in driving fire characteristics in the assessment area over the past century (Heyerdahl et al. 2008, Miller and Urban 1999a, Miller et al. 2009b, Running 2006).

In this section, we summarize information available on the different components of the fire regime for YPMC forests before Euro-American settlement and compare this to current conditions. At the end of the section, we summarize the results of studies that have generated qualitative or quantitative projections for future fire conditions in YPMC forests in the assessment area.

Background: the role of ignitions by humans prior to the 20th century—
The presettlement fire record is mostly derived from fire-caused injury lesions in tree stems or charcoal in layers of sediment or peat, and we are mostly unable to discern lightning-ignited fires from anthropogenic fires. Although lightning occurrence varies temporally (van Wagendonk and Cayan 2008), lightning strike densities (LSDs) may provide a rough idea as to how the ratio of lightning to anthropogenic ignitions might have varied across the landscape (with the important caveat that LSDs will only correlate reasonably well with lightning ignition densities if fuels are in sufficient quantity and sufficiently dry). California is one of the least lightning-prone states in the United States, with most of the Mediterranean part of the state averaging 0 to 0.25 strikes/km²/yr (compare to the Gulf Coast, with more than 6 strikes/km²/yr, or the southeastern and Midwestern United States, with more than 3 strikes/km²/yr) (Orville 2008). The highest LSDs in California are in the deserts of southeastern California and the eastern and higher western slopes of the Sierra Nevada, where average annual LSDs range from 0.3 to 0.55 strikes/km²/yr.

Given very low lightning-strike densities in westernmost California, pre-Euro-American settlement fire frequencies in the California Coast Ranges clearly resulted primarily from human use of fire. The relative magnitude of human inputs to the fire regime in the Sierra Nevada before Euro-American settlement is much less certain, although anthropogenic fire was certainly a significant factor within some radius of many American Indian cultural sites (Anderson 2005, Kilgore and Taylor 1979, Sugihara et al. 2006, Vale 2002). Rightly or wrongly, some observers in the late 19th century were convinced that much of the very frequent fire that characterized presettlement YPMC forests had been set by American Indians (e.g., Greeley 1907, Manson 1906). Pyne (1982) opined that American Indian management of vegetation through fire was widespread in the Western United States for thousands of years before Euro-American arrival. Kilgore and Taylor (1979 (see also Taylor et al. 2016)) noted a sharp decline in fire frequency after the 1870s in their southern Sierra Nevada study area, and correlated this with the replacement of American Indian populations by Euro-American settlers.
Fire frequencies did not follow the same late 19th century pattern everywhere, and some researchers have come to different conclusions. For example, Burcham (1960) suggested that fires set by Euro-Americans between 1850 and 1900 created an artificially high frequency of fire that observers mistook as representative of American Indian influences (but Burcham came from a school that discounted natural fire ignitions and did not base his opinion on actual data). Wagener (1961) summarized fire scar sites in the central and northern Sierra Nevada and determined that fire frequencies did not change until after 1900, when the national forests were established. Because the arrival of Euro-Americans after 1850 did not change fire frequencies in the records he studied, Wagener (1961) concluded that most presettlement fires were probably caused by lightning, not humans. Swetnam and Baisan (2003) noted that recent data on lightning-strike densities had documented much higher lightning incidence than assumed by earlier researchers. Based on their research in the Southwestern United States and the Sierra Nevada, they concluded that, “fire regimes in large portions of these regions would probably have had similar characteristics… if people had never entered the Americas.” Parker (2002) maintained that vegetation patterns in assessment area forests when Euro-Americans arrived were largely explainable based on topography, lightning ignitions, and climate. Probably the truth lies somewhere between the extreme views. Lewis (1973) noted that, “Despite a precontact population that is estimated to be among the highest in North America, (Indians) probably lacked sufficient numbers to burn all or even most of the vegetation on any regular and consistent basis, even had they so wanted.”

In the last decades of the 1800s, there was a general decrease in overall fire frequency, but an increase in large destructive fires in many parts of the Sierra Nevada, ignited by shepherds, miners, loggers, and other forest users (Barbour et al. 1993, Cermak 2005, Erman and SNEP Team 1996, Jackson et al. 1982, Kilgore and Taylor 1979, Leiberg 1902, Miller and Safford 2017, Sudworth 1900, Vankat and Major 1978). Shepherds were especially singled out for blame in the literature (although some authors contend that burning by shepherds was simply an extension of American Indian practices in their absence) (Vankat 1977). Forest managers decried the loss of forest cover and timber that was accompanying Euro-American settlement and the role that indiscriminate fire was playing (Greeley 1907, Jackson et al. 1982, Show and Kotok 1924). Cermak (2005) noted the plethora of photos from the period between 1880 and 1910 showing huge brush fields in and around

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settled or logged watersheds in the Sierra Nevada, areas that are now forest. By the end of the 19th century, much of the Sierra Nevada was being policed for fire, and by the second or third decade of the 20th century (and in some places earlier), fire frequencies began to noticeably drop as federal and state fire suppression policies and tactics were successfully implemented.

Components of the fire regime: fire frequency—

NRV—Fire frequencies are usually measured as FRIs, which measure the number of years between fire events. The most commonly referenced measures of FRIs in conifer-dominant vegetation are derived from small-scale (<4 ha) “composite” dendrochronological fire histories (which measure fire scars in the annual growth rings of trees), which include records from multiple trees in a defined area. Composite FRIs tend to represent the fire history of a given area better than point FRIs (derived from a single tree) because some fire events fail to scar every recording tree within the fire perimeter, and most trees are not recording trees, especially in regimes characterized by frequent low-intensity fire (Collins and Stephens 2007, Falk et al. 2011, Stephens et al. 2010). Composite FRIs are also more sensitive and better suited to analyzing changes in fire occurrence than point FRIs (Dieterich 1980, Swetnam and Baisan 2003). Although there is some variability introduced by using composite FRIs from different size areas, they are less likely to underestimate presettlement FRI values than point (single tree) FRIs (Farris et al. 2010, van Horne and Fulé 2006).

Van de Water and Safford (2011) conducted an exhaustive review of the published and unpublished literature pertaining to mean, median, minimum, and maximum FRIs observed prior to significant Euro-American settlement (i.e., the middle of the 19th century). Sources included fire histories derived from dendrochronological and charcoal deposition records, modeling studies, and expert quantitative estimates; a total of 298 sources were accessed. Van de Water and Safford’s (2011) results for yellow pine and mixed-conifer forests are given in table 3. These values represent the NRV reference period, i.e., the 16th century to about 1850. Averaged across the state, they found that mean FRIs ranged from 11 to 16 years in YPMC forests, and median FRIs ranged from 7 to 12 years. Mean minimum FRIs were around 5 years for both forest types, and mean maximum FRIs ranged from 40 to 80 years (table 3) (Van de Water and Safford 2011).

As table 3 makes clear, presettlement fire frequencies were highest in the drier, lower elevation forest types (YPMC) and lower in moister and higher elevation stands (Caprio and Swetnam 1995, Fites-Kaufman et al. 2007, Gill and Taylor
Natural Range of Variation for Yellow Pine and Mixed-Conifer Forests in National Forests of California

Fire frequencies are also driven directly and indirectly by local topographic variables, and various researchers have documented differences in FRIs between cool (mostly north-facing) and warm (mostly south-facing) slopes, and lower and higher slope positions and elevations. Overall, fire frequencies tended to be higher on warm slopes and upper slope positions, and lower on cool slopes and lower slope positions (Beaty and Taylor 2001, Fites-Kaufman 1997, Kilgore and Taylor 1979, Taylor 2000), but some site-to-site variability occurred, and obvious topographic patterns in fire frequencies were not always apparent (Beaty and Taylor 2007, Scholl and Taylor 2010).

Through the Holocene, fire activity has changed significantly with major changes in climate (Millar and Woolfenden 1999). Working at Lake Tahoe, Beaty and Taylor (2009) found that fire frequencies were low in the Early Holocene, but then reached their highpoint at the height of the Middle Holocene (Xerothermic Period). Frequencies dropped gradually through the Late Holocene, with upward trends during the Medieval Warm Period. Beaty and Taylor (2009) noted that current fire frequencies (under fire suppression) are at their lowest in probably the last 14,000 years. Other researchers have found similar broad patterns (see citations in Beaty and Taylor 2009).

Comparison to current—Safford and Van de Water (2014) compared pre-Euro-American settlement FRIs to FRIs from the last century of fire records in California, using a set of fire return interval departure (FRID) metrics. Figure 5 shows one of these metrics, mean PFRID, generalized to the ecological subregions of California mapped by Miles and Goudey (1997), using only the Forest Service and National Park Service lands within each subregion. “Mean PFRID” represents the current percentage departure from the mean presettlement fire return interval (see Safford and Van de Water [2013] for details). Clearly most of the assessment area is highly positively departed, which means that FRIs are much longer than under

<table>
<thead>
<tr>
<th>Forest type</th>
<th>Mean</th>
<th>Median</th>
<th>Mean minimum</th>
<th>Mean maximum</th>
<th>Number of sources</th>
</tr>
</thead>
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<tr>
<td>Yellow pine</td>
<td>11</td>
<td>7</td>
<td>5</td>
<td>40</td>
<td>24</td>
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<tr>
<td>Dry mixed conifer</td>
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<td>9</td>
<td>5</td>
<td>50</td>
<td>37</td>
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<tr>
<td>Moist mixed conifer</td>
<td>16</td>
<td>12</td>
<td>5</td>
<td>80</td>
<td>53</td>
</tr>
</tbody>
</table>

a Mean minimum and mean maximum are rounded to the nearest multiple of 5
Source: Van de Water and Safford 2011.
presettlement conditions. Areas in figure 5 that are greater than +33 percent departed have current FRIs that are at least 1.5 times longer than under presettlement conditions; areas greater than +67 percent departed have current FRIs that are at least 3 times longer than in presettlement times. To put this into perspective, yellow pine and dry mixed-conifer forests supported mean presettlement FRIs of about 11 years according to Van de Water and Safford (2011), which means that an average of 9.1 fires would occur over any given period of 100 years. Areas in figure

Figure 5—Mean percentage of fire return interval departure (PFRID), generalized to the ecological subregion (Miles and Goudey 1997) from U.S. Forest Service and National Park Service lands found within each subregion. Warm colors are experiencing more fire than under pre-Euro-American conditions; cool colors are experiencing less fire. Illustration from Safford and Van de Water (2014).
5 that are greater than 67 percent departed from this presettlement FRI have experienced three fires or fewer over the past century. The map shows that the southern and eastern regions of the assessment area are less departed than the central and northern regions. Most YPMC forest types in the latter two regions are more than 85 percent departed; i.e., most of the landscape has seen zero to one fire over the past century. Safford and Van de Water (2014) also calculated mean PFRID by vegetation type (see their table 4). They found that yellow pine and dry mixed-conifer were the most departed of 28 vegetation types analyzed (averaging 84 percent and 85 percent departure across California); moist mixed-conifer was tied with aspen for third (average of 80 percent departure). Steel et al. (2015) showed that about three-fourths of YPMC forest in California had not experienced a fire since 1908, after experiencing an average of 5 to 10 fires per century for many centuries before 1908.

Table 4—Reference fire rotations (in years) from the centuries preceding Euro-American settlement for yellow pine and mixed-conifer forests in California.

<table>
<thead>
<tr>
<th>Forest type</th>
<th>Mean</th>
<th>Low</th>
<th>High</th>
<th>Number of sources</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yellow pine</td>
<td>22</td>
<td>11</td>
<td>34</td>
<td>9</td>
</tr>
<tr>
<td>Dry mixed conifer</td>
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<td>8</td>
</tr>
<tr>
<td>Moist mixed conifer</td>
<td>31</td>
<td>15</td>
<td>70</td>
<td>12</td>
</tr>
</tbody>
</table>

* Reference area is the assessment area

Components of the fire regime: fire rotation—

NRV—Fire rotation is the time in years required for fire to burn an area that is equivalent to the study area’s total extent (Agee 1993, Heinselman 1973). Fire rotation can be thought of as a landscape-level FRI, and avoids the difficulties of extrapolating point or composite FRI data to landscapes (indeed, it is roughly equivalent to the grand mean of all point FRIs) (Mallek et al. 2013). Mallek et al. (2013) summarized information from the published literature on presettlement fire rotations for seven major forest types in the bioregional assessment area. Results for YPMC are given in table 4. Mean fire rotations in YPMC forests in the assessment area ranged from 22 to 31 years. As with the FRIs, presettlement fire rotations were longer in the higher and moister mixed-conifer forests.
Comparison to current—Miller et al. (2012b) calculated fire rotations for the period 1984-2009 for YPMC forests in the assessment area, and compared rotations between Yosemite National Park, where fire during the study period was mostly managed rather than suppressed, and Forest Service lands in the assessment area, where most fires are suppressed. From their table 3, the area-weighted average fire rotation for Forest Service-managed mixed-conifer forest over the study period was about 258 years (range 95 to 355), for Forest Service-managed yellow pine forest, it was about 280 years (range 115 to 516); the longest rotations were in the southern Cascades and Modoc Plateau, the shortest in the eastern Sierra Nevada. This compared to about 55-years fire rotation for both forest types in Yosemite National Park. In summary, YPMC fire rotations were about 10 times longer than presettlement on Forest Service lands in the assessment area, but only about 2 times longer in Yosemite (Miller et al. 2012b).

Components of the fire regime: fire severity—Fire severity is a measure of the magnitude of the effect that fire has on an ecosystem (Sugihara et al. 2006). The severity of fire can be assessed from the standpoint of any ecosystem characteristic; here we focus on the severity of fire on the vegetation. Fire “severity” is not synonymous with fire “intensity,” although it is usually correlated. Fire intensity is a measure of the rate of energy released by fire, whereas severity results from the interactions among intensity, fire residence time, fuel and air moistures, the plant species involved, other sources of plant stress, and other factors that cause change in ecosystem condition (Keeley 2009, Sugihara et al. 2006).

In yellow pine mixed-conifer and related forest types, fires in the presettlement period were dominated by areas of low- and moderate-severity effects in which low flame lengths consumed shrubs, herbs, and tree regeneration but only infrequently killed large groups of adult trees. Large, severe fires could occur, but before Euro-American settlement they were comparatively rare. Show and Kotok (1924, 1925, 1929; Sudworth 1900). Large, severe fires could occur, but before Euro-American settlement they were comparatively rare. Show and Kotok (1925: 2–3) concisely summarized the observations of many authors:

Fires run through these pine forests, but comparatively lightly. Only occasionally and in the younger stands do they flare up into the tops of trees and become crown fires. For the most part the fires are confined to the underbrush and forest litter, burning fallen dead trees, and killing by heat rather than actual flame the crowns of live, full grown trees.
In another report, the same authors (Show and Kotok 1924) wrote:

The virgin forest is uneven-aged, or at best even-aged by small groups, and is patchy and broken; hence it is fairly immune from extensive, devastating crown fires. Extensive crown fires, though common in the forests of the western white pine region, are almost unknown in the California pine region.

Even today, with a greatly increased component of high-severity fire in YPMC forests, the proportion of fire area experiencing low- and moderate-severity effects is still generally larger than the proportion experiencing high-severity fire (Miller and Safford 2008, 2012; Miller et al. 2009b), however, the proportional balance between the three fire severity classes has changed considerably (fig. 6).

Aside from historical accounts, estimates of past fire severity patterns in
conifer forests are usually developed from studies of forest stand structure. “High-severity” fires kill most trees on the landscape and leave their footprint through the mostly single-aged stands of regeneration that replace the original forest. Classic examples of vegetation types characterized by high-severity fires include lodgepole pine (*Pinus contorta* Douglas ex Loudon) in the northern Rocky Mountains, chaparral in southern California, and closed cone conifers like knobcone pine (*Pinus attenuata* Lemmon), whose reproduction is closely linked to the passage of stand-replacing fires.

The problem with estimating presettlement fire severity patterns in YPMC forests in the assessment area is that stand structures are and were extremely diverse, with stands typically composed of a mixture of young and old trees, often from a number of different species. Even where groups of even-aged trees are found, other factors than fire could be the cause (windfall, insect outbreak, disease, auspicious climatic conditions, etc.). Very few studies have found evidence of abundant large canopy gaps (>100 ha) in presettlement YPMC forests that after fire were repopulated by single-aged stands of trees (such large gaps, although still comparatively rare, are more commonly found in northern assessment area YPMC forests that supported higher densities of fir species). The prevalence of large canopy gaps caused by high-severity fire is partly related to precipitation and forest productivity, as—assuming everything else equal—wetter forest types grow fuels more rapidly and support denser forest stands, and moister fuels require more heat input to combust, which results in longer fire-free intervals but higher severity fires when fire does occur. These moister forests also tend to support higher densities of shade-tolerant, relatively fire-intolerant tree species like fir, whose canopy structure is much more conducive to rapid movement of surface flames into the forest canopy. Higher elevation forests in the Sierra Nevada generally receive more precipitation and support more fir and other relatively fire-intolerant species. A similar gradient occurs from south to north, with precipitation and the importance of fire-intolerant species increasing with latitude (along the west side of the assessment area; the eastern regions are drier in general and more dominated by drought-tolerant species of pine, etc.) The elevation- and precipitation-driven increase in the proportion of high-severity fire moving from lower and drier vegetation types to higher and wetter vegetation types can be seen in the reference conditions in figure 6, moving from left to right.

The relatively fine-grained heterogeneity that characterized forest structure in YPMC forests (see “Structure” on page 87) was the product of a largely low- and moderate-severity fire regime (Fire Regime I), with occasional occurrences of high-severity fire in larger patches under the right weather and fuels conditions.
It is important to remember that “low-severity” fire does not imply the absence of tree mortality. Most classification systems define low severity as occurring below 25 to 30 percent canopy mortality. Taking the midpoint of that class, an average of around 12.5 to 15 percent of canopy trees in low-severity areas would be expected to be killed by fire. “High-severity” fire is usually defined as encompassing those areas where fire-caused mortality exceeds 75 to 80 percent, although recent satellite-imagery derived severity assessments (relative differenced Normalized Burn Ratio [RdNBR] assessments) usually map high-severity areas in polygons where mortality is more than 90 or 95 percent (Miller et al. 2009a, 2009b, 2012b). Thus, these RdNBR-based studies are mapping “stand-replacing” fire, where tree mortality is almost complete. In this assessment, we use >95 percent mortality as our definition of high-severity fire. “Mixed-severity” or “moderate-severity” fire is simply the area between low and high (we use the term “moderate” in this assessment). Moderate-severity areas are really areas where the intermix of low- and high-severity effects is at a fine enough grain to not be discernible with the analysis tools at hand. For most classifications, the midpoint of the moderate-severity class falls between 50 and 60 percent mortality, so canopy tree loss in such areas may actually be substantial.

There are many ways to quantitatively estimate fire severity in presettlement YPMC forests. One is through direct evaluations of fire severity patterns in contemporary reference forests, in which fire suppression does not occur or has been relaxed, and where extensive logging did not occur. Such forests occur in Yosemite National Park and in the Sierra de San Pedro Mártir of northern Baja California, Mexico. Yosemite was mostly unlogged, but fire management in the park practiced strict fire suppression until 1972 (van Wagtenonk et al. 2002). Since then, extensive areas of the park have been managed under a policy allowing most naturally ignited fires to burn. As such, Yosemite is not a surrogate for presettlement conditions, but it is the closest large landscape we have today in the assessment area. Collins et al. (2009) used RdNBR data to study fire severity in Yosemite’s Illilouette Valley, which was relaxed from strict fire suppression in the early 1970s. The watershed is dominated by moist mixed conifer, red fir, and lodgepole pine, so it is most useful as a reference for high-elevation mixed-conifer stands. Collins et al. (2009) did not differentiate their fire severity results by forest type, but overall they found that over a 31-year period, a total of 13 percent of the fire areas they assessed had burned at high (stand-replacing) severity. Van Wagtenonk et al. (2012) studied the same area and found that the proportion of first-entry fire area that burned at high severity was 1 percent in red fir, 3 percent in white fir, 8 percent in Jeffrey pine/western white
pine, 12 percent in Jeffrey pine/shrub, 4 percent in lodgepole, and 19 percent in montane chaparral. Among subsequent fires that burned through the same vegetation types, the proportion of high severity was 13 percent in red fir, 16 percent in white fir, 13 percent in Jeffrey pine/western white pine, 14 percent in Jeffrey pine/shrub, 23 percent in lodgepole, and 59 percent in montane chaparral.

Miller et al. (2012b) analyzed RdNBR fire severity data from the entirety of Yosemite National Park, for the period 1984–2009. During their study period, about 72 percent of the fires they analyzed were managed for resource benefit (although five were later suppressed after they exceeded their prescriptions), while the remaining 28 percent were suppressed. They found that YPMC forests in the park burned at about 14 percent high severity (13.7 percent mixed conifer, 14.6 percent yellow pine).

Another contemporary source of reference fire regime information for YPMC forests in the assessment area is the Sierra de San Pedro Mártir National Park in northern Baja California, Mexico. This area is in the southernmost part of the North American Mediterranean climate zone and supports yellow pine (in this case Jeffrey pine) and dry mixed-conifer forests that are very similar to those of drier portions of the Sierra Nevada, for example, on the eastern slope (Dunbar-Irwin and Safford 2016, Stephens and Fulé 2005). Unlike the Sierra Nevada, most of the Sierra de San Pedro Mártir was not logged, and fire suppression has only been in effect for the past few decades, so the forests are much closer to pre-Euro-American settlement conditions. Minnich et al. (2000) reported results from aerial photointerpretation of two fires that burned in the Sierra San Pedro Mártir in 1989. Using photos from 1991, they estimated that 16 percent of the analyzed fire area had experienced >90 percent mortality, thus perhaps 8 percent or so experienced stand-replacing fire effects (>95 percent mortality). Aerial photo analysis is known to underestimate the area of low-severity fire, as fire extent is primarily mapped based on fire effects to canopy trees, so surface fires are difficult to pick out, especially when a number of years have passed since the fire event. Minnich’s analysis also took place 2 years after fire, whereas most RdNBR assessments occur 1 year after fire. Consequently, we view Minnich et al.’s (2000) numbers as an upper estimate of fire severity.

Stephens et al. (2008) used field plots to measure severity in a fire area in the Sierra San Pedro Mártir. Only 1 of their 27 plots (4 percent of their sample area) experienced high-severity effects (>95 percent mortality). Rivera-Huerta et al. (2016) carried out an RdNBR-based assessment of 25 years of fire severity patterns in the Sierra San Pedro Mártir, and found that the proportion of YPMC forests burning at high severity was about 3 percent, although most of the assessed fires had entered YPMC stands from chaparral under severe-fire weather conditions.

Sudworth (1900) inventoried forestlands in the central Sierra Nevada (Lake
Tahoe basin to the latitude of Yosemite National Park) at the end of the 19th century. Concerning the nature of fire in his study region, he wrote (Sudworth 1900: 557–558):

The fires of the present time are peculiarly of a surface nature, and with rare exception there is no reason to believe that any other type of fire has occurred here… The instances in this region where large timber has been killed outright by surface fires are comparatively rare. Two cases only were found… One of these burns involved less than an acre, and the other included several hundred acres. They are exceptional cases, and the killing of the trees is accounted for by the fact that long protection from fire and from all but cattle grazing had resulted in the accumulation of much fallen timber, considerable humus in depressions and on benches, and a dense undergrowth of brush and seedlings. The fires burned deep enough to badly injure the surface roots, which resulted in the subsequent death of the timber.

Leiberg (1902) carried out a field inventory of forestlands in the northern Sierra Nevada (American River watersheds to the Feather River watersheds) at the beginning of the 20th century and made estimates of the amount and severity of burning that had occurred in the region over the previous century. Euro-American presence in the Sierra Nevada was minimal until after 1850, and exclusion of fire from most Sierra Nevada forests is not noticed in the fire scar record until at least the 1870s or 1880s (Sugihara et al. 2006), so Leiberg’s (1902) results at least partly reflect pre-settlement conditions. That said, northern Sierra Nevada forests suffered heavier Euro-American degradation between 1850 and the early 20th century than the southern Sierra Nevada (Miller and Safford 2017). Leiberg tallied burned area by watershed for the northern Sierra Nevada and estimated that 8 percent of the 19th century fire area had experienced “total destruction,” i.e., stand replacement. Leiberg’s (1902) assessment did not quantitatively discern among forest types, although most of the fires he visited had taken place in upper elevation mixed-conifer and red fir forests.

Cooper (1906) noted: “Forest fires in California are usually ground fires, and as the mature trees of all species are more or less fire resisting, these fires seldom kill the timber outright, nor does a single fire greatly affect the virgin forest as a whole.” In the same year, Hodge (1906) wrote: “In virgin timber ground fires are the rule, and it is seldom the flames reach up into the foliage of large trees, even in stands of fir.”

Show and Kotok (1925) stated that fires in the “California pine region,” which equates to YPMC forest, rarely burned the forest canopy, but killed canopy trees through heat from surface fires and successive scarring and hollowing out of the
trunk, which resulted in typical fire-caused losses of about 5 percent of the “merchantable forest” (mature trees). In another publication, the same authors noted that the typical fire type for yellow pine and drier mixed-conifer forests was surface fire, while moister forests with a high sugar pine and white fir component would tend to experience creeping fires that also consumed the duff layer (Show and Kotok 1929).

Beaty and Taylor (2001) used tree-age data from forest plots and aerial photos from 1941 to reconstruct spatial variation in fire severity in the Cub Creek Research Natural Area on the Lassen National Forest. Vegetation in the area is dominated by moist mixed conifer. Severity classes were assigned based on a categorization of the numbers of emergent trees seen in the aerial photos. The authors found that forests in lower slope positions experienced mostly low-severity fire, middle slope positions experienced mostly moderate- and low-severity fire, and upper slope positions experienced mostly high-severity fire (Beaty and Taylor 2001). Assuming that the landscape was evenly divided among lower, middle, and higher slopes, the areal proportions amounted to about 32:30:38 (low:moderate:high). Also working on the Lassen National Forest and using the same methodology, Bekker and Taylor (2001) mapped fire severity using 1939 and 1943 aerial photos of the Thousand Lakes Wilderness on the Lassen National Forest. For mixed-conifer and white fir-Jeffrey pine stands, fire severity was calculated to have been “high” on 52 to 63 percent of the landscape. In both studies, the earliest fires assessed for severity occurred in the 1880s, so results may or may not be representative of pre-Euro-American conditions, as many destructive fires were set by settlers during the mining, logging, and grazing boom that occurred between 1850 and the early 1900s (Cermak 2005, Jackson et al. 1982, Leiberg 1902, Miller and Safrord 2017, Sudworth 1900). Also, because severity classes were based on the numbers of emergent trees seen in aerial photos, they are not easily correlated with measures of biomass loss like RdNBR, and they could have just as easily resulted from a series of moderate/mixed-severity fires over time rather than from one fire event (see footnote 2). It would be useful to determine (1) how to translate the “number of emergent trees” measure to RdNBR, and (2) to what extent these late 19th and early 20th century fires were representative of the presettlement fire regime.

Stephens et al. (2015) used historical timber survey data from 1911 in the Greenhorn Mountains on the Sequoia National Forest to estimate the occurrence of high-severity fire across an 11 500-ha landscape. Forty-one percent of the landscape was dominated by ponderosa pine, while 59 percent was classified as mixed conifer. High-severity fire was estimated using surveyor notes as well as surveyed areas that were dominated by montane chaparral and stands of dense immature timber, which are common vegetation conditions following stand-replacing fire. The percentage of surveyor transects that included evidence of high-severity fire ranged from 2.5 to
7.7 percent in mixed-conifer forests and 7.1 to 13.5 percent in ponderosa pine forests (depending on whether the estimate was based strictly on surveyor notes or also included vegetation condition). Accounting for the fact that only a proportion of each surveyor transect usually included evidence of high-severity fire, Stephens et al. (2015) calculated that high-severity fire occurred across 1 to 3 percent of all mixed-conifer forests and 4 to 6 percent of all ponderosa pine forests in the study area.

Fire severity patterns in presettlement forests can also be modeled. The national, interagency LANDFIRE project collaborated with The Nature Conservancy to convene dozens of workshops nationwide, where state-and-transition models were built of hypothesized presettlement fire and forest successional dynamics by forest and fire ecology professionals (Long et al. 2006, Rollins and Frame 2006). Different models were developed for major vegetation types, which were known as biophysical settings (BpS) (http://www.landfire.gov/NationalProductDescriptions24.php). The models were peer reviewed and extensively refined and calibrated, and then delivered to the LANDFIRE modeling group, which used the BpS models to develop national fuels, vegetation, and Fire Regime Condition Class spatial data layers. In contrast to maps depicting current vegetation distributions, the LANDFIRE BpS layer is a modeled output of potential vegetation representing the distributions of vegetation types as they are hypothesized to have existed prior to Euro-American settlement, based on topography, climate, soils, and the presettlement disturbance regime (Rollins 2009). The advantage of the LANDFIRE BpS layer, as opposed to maps of current vegetation, is that fire regimes both influence and are influenced by vegetation, so maps representing current vegetation may confound the ability to detect shifts in fire regime over time. The disadvantages of the BpS layer include the fact that the models are hypotheses and are difficult to validate empirically. Local inaccuracies in the mapped product can also make it difficult to use the map at fine scales, but at regional and statewide scales, the BpS map conforms closely to the broad biophysical gradients (e.g., Miller and Safford 2012, Safford and Van de Water 2014). In addition, the LANDFIRE BpS output is the only map of potential vegetation in California that is based on a transparent and peer-reviewed modeling process, incorporates the effects of fire, includes lands of all management jurisdictions, and extends across the entire assessment area.

Each BpS model includes documentation that summarizes, among other things, the fire severity outputs from 100 runs of 500 simulation years under presumed presettlement conditions. Mallek et al. (2013) summarized the percentage of high-severity outputs from these models and computed area-weighted averages (since multiple BpS models correspond to each of the major vegetation types analyzed in the study) for the assessment area: the yellow pine models averaged 5.9 percent high
severity; dry mixed conifer, 7.1 percent high severity; and moist mixed conifer, 11.6 percent high severity (see also fig. 6). Note that the LANDFIRE project based their severity categories on the Fire Regime Condition Class (FRCC), which defines high severity as >75 percent mortality. Using the 95-percent cutoff used by the RdNBR assessments and our assessment, the percentage of high-severity values from these BpS models would be somewhat lower than those reported.

Stephens et al. (2007) provided literature- and field observation-based estimates of pre-Euro-American fire severity in 12 forest types in California. Their measure of severity was “percent(age of) crown burned,” which correlates closely with the RdNBR-derived severity measures, as RdNBR is driven primarily by canopy loss from fire (Miller et al. 2009a). Stephens et al. (2007) used vegetation types described by Barbour and Major (1988) and estimated that an average of 5 percent of the tree canopy suffered burning (torching) in mixed-conifer and ponderosa/shrub forest types, and 2.5 percent in Great Basin pine (Jeffrey pine) forests.

Quigley et al. (1996) summarized scientific work that was done to support forest planning in the interior Columbia River basin, just north of the assessment area. According to Quigley et al. (1996), presettlement burning in eastern Oregon and Washington, which support large areas of semiarid ponderosa pine-dominant forest averaged between 10 and 15 percent “lethal” (high-severity) fire, and 55 to 60 percent “nonlethal” (low-severity) fire. Contemporary fires were very different, with annual burning averaging 30 to 35 percent lethal fire and 25 to 30 percent nonlethal fire.

Comparison to current—Miller et al. (2009b, 2012b), Miller and Safford (2008, 2012), and Mallek et al. (2013) studied contemporary fire severity patterns on Forest Service lands in the assessment area, using RdNBR data. The time periods, the minimum sizes of fires considered, and the vegetation type classification used differed somewhat among the studies. Miller and Safford (2008) and Miller et al. (2009) based their analysis on the existing forest types used in the Sierra Nevada Forest Plan Amendment, and considered severity trends between 1984 and 2004. Based on an area-weighting of their severity results, fires in yellow pine forests (ponderosa pine plus east-side pine) burned about 33 percent of their total area at high (stand-replacing) severity, while mixed-conifer forests (mixed conifer plus white fir) burned at about 29 percent high severity. Miller et al. (2012b), Miller and Safford (2012), and Mallek et al. (2013) based their analyses on potential forest types (biophysical settings) from the LANDFIRE national mapping project, and considered severity trends between 1984 and 2009 (2010 in Miller and Safford 2012). After area-weighting their results, they found that yellow pine types burned at about 42 percent high severity during the study period,
and mixed-conifer types at 23 to 30 percent, depending on whether mixed conifer was divided into dry mixed-conifer and moist mixed-conifer types (Mallek et al. 2013, Miller et al. 2012b). Miller and Safford (2012) joined yellow pine and mixed conifer, and found the combination burned at 33 percent high severity during the study period. By contrast, upper elevation mixed-conifer forests in Yosemite National Park during the same period burned at about 14 percent high severity (Miller et al. 2012b).

Using potential vegetation types (which are modeled from environmental and vegetation data) in comparisons between historical and current conditions avoids the complication introduced when vegetation type has changed over time, e.g., owing to human management. However, it is difficult to assess the accuracy of modeled vegetation typing. Miller and Safford (2012) combined yellow pine and mixed conifer precisely because of inaccuracies they found in the vegetation classification. When we consider the combined YPMC forest type, the results using the existing vegetation mapping (~30 percent high severity) (Miller and Safford 2008, Miller et al. 2009b) are very close to those derived using the potential vegetation mapping (33 to 35 percent) (Mallek et al. 2013, Miller and Safford 2012, Miller et al. 2012b). Either way, current fires in YPMC forests managed by the Forest Service in the assessment area are burning at much higher severity (30 to 35 percent high severity as an assessment area average) than was generally the case under presettlement conditions (long-term assessment area-wide average probably <10 percent, ranging from 3 to 15 percent; see fig. 6). Furthermore, since 1984, fire severity in Forest Service YPMC forests has risen substantially, from an average in the low 20 percent in the mid 1980s to more than 30 percent in 2010 (fig. 7). Severity in related forest types, such as black oak and white fir, has also risen over the past few decades (Miller et al. 2009b). In comparison, fire severity since the 1980s has not risen appreciably in upper elevation mixed-conifer forests in Yosemite National Park (Collins et al. 2009, van Wagendonk and Lutz 2007), where five decades of wildland fire use (also referred to as “fires managed for resource benefit”) have reduced fuels and restored forest structure.

Although we have generally been discussing proportional fire severity as applied across all YPMC forests in the assessment area, note that fire severity is not expected to be uniform within this forest type, either in presettlement or in modern times. Fire severity at multiple scales is known to be influenced by a range of factors, including gradients in precipitation, elevation, slope position, species composition, and previous fire history. Within the assessment area, fuel accumulation at the landscape scale is driven by both climate and land management practices (among other things); so higher proportions of high-severity fire
would be expected in areas with higher precipitation, lower evapotranspiration, higher elevation, and/or a longer legacy of fire suppression (Agee 1993, Noss et al. 2006, Schoennagel et al. 2004, Sugihara et al. 2006). Severity also varies at the stand scale, with greater severity expected in stands on north-facing slopes, in riparian areas, and with higher concentrations of fir (North et al. 2012b, Perry et al. 2011). Geographically, the higher precipitation and higher fir component found in the northwestern assessment area probably led to somewhat more severe fires on average than in the rest of the area.

An Alternative Viewpoint of Fire Severity in YPMC Forests

A small school of researchers and environmentalists maintains that the standard model of mostly low and moderate severity in presettlement YPMC forests is mistaken, and that modern efforts to reduce fuels in such forests are misguided. It is important to acknowledge the existence of this school of thought, although the vast bulk of evidence (summarized throughout this assessment) suggests otherwise. Science produced by this group can be characterized fairly as controversial. For example, Odion and Hanson (2006) used soil-burn-severity mapping of three fires

Figure 7—Temporal trends in the percentage of high-severity fire for yellow pine–mixed-conifer forests in the assessment area between 1984 and 2010. The figure shows annual percentages of high-severity fire, the time series (ARIMA) model for the 1984–2010 period, and linear trend lines for seven time series beginning in 1984 and ending in 2004 to 2010. Illustration adapted from Miller and Safford (2012); the definition of high severity is as described in this document.
to generalize about fire effects on vegetation across the entire Sierra Nevada, suggesting that modern levels of fire severity and sizes of high-severity patches were within NRV. Safford et al. (2008) responded to Odion and Hanson, and showed that the soil-burn-severity mapping Odion and Hanson used had little correlation to vegetation-burn severity, and that their analyses of fire patchiness and relationship between fire-return interval and fire severity were flawed.

In another example, Williams and Baker (2012) and Baker (2012, 2014) used General Land Office (GLO) data from the mid- and late-1800s (see “Forest structure,” “Tree size,” and “TreeSize-Class Distribution” below) to generalize about tree densities in YPMC forests in a number of locations across the Western United States (Baker [2014] specifically refers to the Sierra Nevada), and the role of fire in driving these densities. These studies similarly claim that current fire patterns—i.e., high levels of high-severity fire—are within the NRV. Aside from the obviously tenuous use of GLO data to compute landscape tree densities, which involves extrapolating from ≤8 trees per square mile, Baker and colleagues’ work also requires the belief that any area of denser-than-average small- to medium-size trees is necessarily the result of high-severity fire, even where there is no evidence of fire and even where the relationship between tree size and tree age is not known (Fulé et al. 2014). Furthermore, in the absence of fire, simple differences in site productivity and antecedent forest structure are sufficient to explain major variation in tree density in YPMC forests (Maxwell et al. 2014). Comparison of direct forest inventories from the early 20th century demonstrates that the Baker (2012, 2014) studies greatly overestimated forest stand density. Such inventories, which sampled orders of magnitude more trees than the GLO effort used by Baker, showed his stand density estimates to be from 2.1 to 5.9 times too high (Collins et al. 2011; Hagmann et al. 2013, 2014; Stephens et al. 2015). Similarly, Levine et al. (2017) used Baker’s algorithm to predict tree densities on a series of 6-ha mapped forest plots in YPMB forests in the Sierra Nevada and NW Mexico and found that Baker methodology notably overpredicted densities in every case (1.2 to 4.8 times higher than the true density). Fulé et al. (2014), Stephens et al. (2015), and Collins et al. (2015) also showed that the Baker methods greatly overestimated historical fire severity.

Hanson and Odion (2013) claimed to refute the trends of increasing high-severity fire in the Sierra Nevada national forests found by Miller et al. (2009b) and Miller and Safford (2012). However, Safford et al. (2015) uncovered a high number of compounded errors in Hanson and Odion’s analysis, including issues with their data layers, errors in cartographic analysis, unorthodox forest groupings that
conjoined different fire regimes, and the inclusion in their analysis of prescribed fires. In their dataset, Hanson and Odion (2013) also included large areas of fire on private lands (where high-severity fire areas are generally salvage logged before the postfire imagery used to determine the level of severity is acquired, which makes a credible severity assessment on such lands impossible) and wildland fire use areas in the national parks, which have no place in a test of fire patterns on Forest Service lands.

Odion et al. (2014) suggested that Forest Inventory and Analysis (FIA) “stand-age” data proved that YPMC forests across the Western United States were largely younger forests that had arisen after high-severity fire events. Odion et al. (2014) used data only from wilderness areas and national parks, which are at higher elevations in the assessment area and are not very representative of typical YPMC forest. The major problem with Odion et al. (2014), however, is that FIA stand-age data, which are a very rough estimate of the canopy-area-weighted mean age in a plot, are uninterpretable and essentially meaningless in the highly heterogeneous age structure of YPMC forests. Stevens et al. (2016) showed that most of the plots analyzed by Odion et al. (2014) included numerous trees that were older, in some cases hundreds of years older, than the “stand-age” computed in their study, which makes Odion et al.’s (2014) claim that high-severity fire had reset succession in all of these plots an impossibility, given that high-severity fires kill all or nearly all trees in the affected stand (Miller and Quayle 2015).

A standard claim made by this group is that their findings challenge the reigning model for presettlement YPMC forests, which they define as permitting only low-severity fire and open, park-like stands of large trees. This definition does not represent the scientific consensus. In 1996, the SNEP report (Erman and SNEP Team 1996) summarized the consensus regarding fire severity in the Sierra Nevada thusly: “In most lower-elevation oak woodland and conifer forest types of the Sierra Nevada, presettlement fires were frequent, collectively covered large areas, burned for months at a time, and, although primarily low to moderate in intensity, exhibited complex patterns of severity.”

This was the consensus 20 years ago, and it remains the consensus today. As noted throughout this assessment, YPMC forests in their reference state were extremely heterogeneous, and were characterized by areas of low-, moderate-, and high-severity fire (but with the balance more shifted to low severity than is currently the case).

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3 Fried, J. 2013. Personal communication. Research forester, USDA Forest Service, Pacific Northwest Research Station, Forestry Sciences Laboratory, 620 SW Main St., Ste. 400, Portland, OR 97205.
Components of the fire regime: high-severity patch size—

**NRV and comparison to current**—In low- and middle-elevation forests, high-severity patch size has also increased, with a dominance of small, scattered patches in presettlement and reference estimates, versus more contiguous coarser grained patchiness in modern fire-suppressed forests. High-severity patches more than a few hectares in size were relatively unusual (although not unknown) in fires in Sierra Nevada YPMC forests before Euro-American settlement (Agee 1993, Kilgore 1973a, Show and Kotok 1924, Skinner 1995, Skinner and Chang 1996, Stephenson 1990, Sudworth 1900, Weatherspoon and Skinner 1996), but in recent years high-severity patches >500 ha have become a regular occurrence (Miller and Safford 2008, Miller et al. 2012). Between 1984 and 2006, mean high-severity patch size in Forest Service fires in the assessment area almost doubled (Miller et al. 2009b). Comparisons between current reference YPMC forests (in which full fire suppression is not practiced) and Forest Service forests (managed primarily under full fire suppression) further reflect these changes. For example, Minnich et al. (2000) reported that patches of high-severity fire (>90 percent mortality in their study) in the Sierra de San Pedro Mártir were almost entirely <16 ha in size; mean patch size was 1.7 ha. Scholl and Taylor (2010) used a forest reconstruction of an 1899 mixed-conifer stand in Yosemite National Park to show that even-aged patches indicative of high-severity fire were generally <0.2 ha. In contrast, Miller et al. (2012b) found that the average size of high-severity patches in modern Forest Service-managed fires in the assessment area was 12.2 ha (1.6 SE); high-severity patches during the same time period in Yosemite, where many fires have not been suppressed for the past four decades, averaged 4.2 ha (0.6 SE). Van Wagtendonk and Lutz (2007) found that mean high-severity patch size for Yosemite from 1984 to 2005 was approximately 1.6 ha for prescribed fires, 2.5 ha for “fire-use” fires (natural ignitions that were not suppressed), and 6.7 ha for wildfires (any ignition that was managed for suppression). Collins and Stephens (2010) analyzed fire-severity patchiness in Yosemite’s Illilouette watershed and found that 48 percent of the total high-severity area was in patches >60 ha, which comprised only about 5 percent of the total number of patches. Data from Miller and Safford (2008) showed that patches >60 ha comprise 70 percent of high-severity fire area in recent Forest Service fires.

To summarize, the NRV of high-severity fire patch size in assessment-area YPMC forests was strongly dominated by a “salt-and-pepper” pattern of small areas mostly (much) less than a few hectares in size. Patches larger than a few hectares did occur, but they were rarely more than 100 ha. Nonetheless, such larger patches comprised perhaps half of the total high-severity area.
Components of the fire regime: fire size—

NRV—When we consider only fires >4 ha (the minimum fire size recorded in the California Fire Perimeter Database), mean fire size in California YPMC forests before Euro-American settlement was much smaller than under current conditions (fig. 8). We were able to find eight published estimates (Beaty and Taylor 2001, Bekker and Taylor 2001, Collins and Stephens 2007 [which contains estimates for two separate national park areas; raw data provided to authors], Scholl and Taylor 2010, Taylor 2000, Taylor and Skinner 1998, and Taylor and Solem 2001) and two unpublished estimates of presettlement fire size statistics in YPMC-type forests in northern California. Restricting our consideration to fires ≥10 ha to facilitate comparison, mean

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Figure 8—Comparison of mean fire sizes from yellow pine and mixed-conifer reference sites with intact fire regimes and the assessment area (forest fires only) since the institution of fire suppression in the early 20th century (the latter are divided into four equal time periods [1908–1934, 1935–1960, 1961–1986, 1987–2011] to show temporal trends). Only fires ≥10 ha are included. Ten sources were used for historical reference numbers; see text. SSPM = Sierra de San Pedro Mártir; data are from Minnich et al. (2000). Illilouette watershed wildland fire uses fires from B. Collins (see footnote 6); 20th- and 21st-century fires are from the California Fire Perimeter Database (see footnote 4).

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5 Taylor, A.H. 2012. Personal communication. Professor, Department of Geography, Pennsylvania State University, University Park, PA 16802.
presettlement fire size averaged from these 10 sources is 210 ha (±33.6 SE, range of means = 85 ha to 457 ha). Finally, there are two published fire size estimates from contemporary reference areas (again, we only include fires ≥10 ha in size to permit comparison among studies): mean fire size in the SSPM, Baja California, Mexico was measured at 221 ha (167 ha if all measured fires are included) between 1925 and 1991 (Minnich et al. 2000); mean fire size in the Illilouette Creek watershed in Yosemite National Park, where most naturally ignited fires are allowed to burn, was 456 ha (61.4 ha if fires of all sizes are included) between 1973 and 2006 \(^6\) (see van Wagendonk et al. 2012 for info on fires >40 ha in the same watershed). The overall average of these 12 sources is 296 ha. In addition, Show and Kotok (1923) reported fire sizes in the assessment area and northwestern California forests (then, as now, most fires occurred in the YPMC belt) for the 10-year period between 1911 and 1920, just as the federal government was beginning to engage in fire suppression. They found the average size of all recorded fires to be 48.3 ha (± 12.8 SE), and the average of all fires ≥4 ha was 177.8 ha (± 32.7 SE).

Another estimate of mean presettlement fire size in assessment-area YPMC can be made by dividing the total area extent of YPMC forest by the presettlement fire rotation (area-weighted mean for yellow pine, dry mixed conifer, and moist mixed conifer from Mallek et al. [2013] to generate an annual area burned, then dividing the area burned by the mean annual number of lightning ignitions in YPMC forests, also area-weighted by forest type), under the assumption that modern fire ignition data are representative of presettlement patterns (Short 2013). This approach yields a mean prehistorical fire size of 435 ha for YPMC forests (391 ha in yellow pine, 692 ha in dry mixed conifer, and 434 ha in moist mixed conifer), \(^7\) but ignores ignitions by humans and therefore overestimates mean fire size, especially in the lower elevation forest types (yellow pine and dry mixed conifer).

Note that there is evidence that fires in areas of simple topography may have often burned areas in excess of this ±300- to 400-ha mean. Norman and Taylor (2003) reported evidence of recurrent landscape-scale fires in an area of subdued topography in the southern Cascade Range; data from other parts of the southern Cascades show similar patterns.\(^8\) Their interpretation was that these larger fires

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\(^6\) Collins, B.M. 2013. Personal communication. Fire ecologist, Center for Fire Research and Outreach, University of California, Berkeley, CA 94720.

\(^7\) Miller, J.D. 2013. Personal communication. Remote sensing specialist, USDA, Forest Service, Pacific Southwest Region, 3237 Peacekeeper Way, Suite 101, McClellan, CA 95652.

\(^8\) Skinner, C.N. 2012. Personal communication. Research fire geographer, USDA, Forest Service, Pacific Southwest Research Station, Redding Silviculture Laboratory, 3644 Avtech Parkway, Redding, CA 96002.
were driven by herbaceous fuels (mostly in and around meadow complexes, but not exclusively there), where fuel loadings increased in wet years and then burned in subsequent dry years (Norman and Taylor 2003). Clearly topography and landscape conditions (presence of previously burned areas, rock outcrops, large streams, etc.) played a significant role in limiting fire size in the assessment area.

Comparison to current—Data from the California Fire Perimeter Database (see footnote 4) shows that average fire size (all fires >10 ha) over the 25-year period up to 2011 for fires in YPMC forests in the assessment area (all jurisdictions) was 1400 ha (±55 SE) (fig. 8), about five times larger than either the assessment area historical data sources or the modern SSPM data, and about three times larger than the average in the Illilouette Creek watershed in Yosemite National Park (not an entirely clean comparison, as Illilouette is included in the assessment area data). Note that modern YPMC fire areas (since 1908) in figure 8 were calculated by removing those portions of fires that burned in other vegetation types.

Figure 9 provides a graphic comparison of fire size frequency distributions in the assessment area (1910 to 2006) (see footnote 4) versus the SSPM (1925 to 1991) (data from Minnich et al. 2000). The climates and vegetation are similar in the two

![Graph showing comparison of 20th-century fire size frequency distributions, assessment area versus Sierra San Pedro Mártir. Proportionally speaking, the assessment area experiences many more large fires and many fewer small fires. Only fires >10 ha in size are included.](attachment:figure9.png)

\[ \chi^2 = 263.97, \text{ df} = 7, \ P < 0.0001 \]
areas (the latter being drier than the western and northern portions of the former). The former has experienced significant timber harvest and a century of fire suppression; the latter was not logged, and at the time of the last fire had experienced fewer than 15 years of fire management. The assessment area has experienced proportionally more than twice as many large (>400 ha) fires and proportionally less than half as many small (<50 ha) fires compared to the SSPM over the study period (fig. 9).

Comparing the same two datasets, fires greater than and smaller than 1600 ha contributed equally to the total area burned in the Sierra de San Pedro Mártir, but large fires (>1600 ha) contributed 70 percent of all burned area in the assessment area.

It is important to finish with a caveat: we have focused our comparison on fires >4 ha, as there are no presettlement data on small fires, and the California Fire Perimeter Database includes only fires >4 ha. However, a recent unpublished analysis by Miller and Safford (manuscript in preparation) of all fire sizes compares a post-1992 dataset for the assessment area against a 1911–1920 dataset from Show and Kotok (1923) and a 1980–2012 dataset from managed lightning fires in Yosemite National Park. Miller and Safford (in prep) found that, because of fire suppression, the proportion of fires that remain below 4 ha in size is much higher in the general assessment area (mostly Forest Service and private lands under fire suppression policies) today than at the beginning of the 20th century or in managed fires in Yosemite National Park. In summary: when considering all fire sizes, the average fire size before fire suppression was actually larger than today, but when considering fires >4 ha, fires today are larger on average. This is due to the strong statistical “mass” provided by the growing set of very large modern fires.

Components of the fire regime: annual area burned—

NRV—A number of published studies have estimated probable presettlement annual burned area, but only a few have split their estimates by vegetation type. Stephens et al. (2007) estimated burning rates for mixed-conifer and yellow pine forests across California, and found that 5 to 15 percent of the total area of YPMC forests burned per year before Euro-American settlement, depending on the fire frequency estimate used. Using the LANDFIRE BpS estimates of forest area before Euro-American settlement (3.65 million ha for YPMC) (Mallek et al. 2013), this would result in a range of about 180 000 to 550 000 ha burned per year in YPMC forest types in the assessment area. North et al. (2012a) estimated fire areas for the Sierra Nevada sensu stricto, which does not include lands north of the North Fork Feather River or east of the eastern Sierra Nevada foothills. Their “active-management” forest types, which essentially conform to YPMC plus some hardwood forests, were estimated to experience between 77 000 and 203 000 ha of burning
per year before Euro-American settlement, or 5 to 18 percent of the total land area in the YPMC forest types (North et al. 2012a). Mallek et al. (2013) carried out an assessment of fire area for the same study area as the bioregional assessment, basing their calculations on fire rotation data found in the scientific literature. Mallek et al. (2013) found that presettlement burning probably affected about 4 percent of the land area in the YPMC forest types in an average year, or around 145 000 ha. Stephens et al. (2007) suggested that their low-end estimates of burning were probably more trustworthy than their high-end estimates, so it seems reasonable to suggest that somewhere around 5 percent of the area of YPMC forest types probably burned in an average year (and there was likely very high variability, year to year), or somewhere between 150 000 and 175 000 ha across the assessment area.

Comparison to current—Mallek et al. (2013) used the California Fire Perimeter Geodatabase (see footnote 4) to calculate the total area of fire within yellow pine, dry mixed-conifer and moist mixed-conifer forests in the assessment area for the period 1984 to 2009. During the period, 489 000 ha of fire are recorded in the database for these forest types, which comes to an annual average of about 18 800 ha in YPMC forests. This is between about 10 and 13 percent of our presettlement estimate above. Using data up to 2010, North et al. (2012a) suggested that current wildfire in the Sierra Nevada proper (about 60 percent of the assessment area) accounts for 10 to 30 percent of presettlement fire area. Note that since these publications, the area burned by wildfire in the Sierra Nevada has increased dramatically. Between 2013 and 2015, the three largest fires—Rim, King, and Rough—burned over 200 000 ha on their own, principally in YPMC forests. According to the California Fire Perimeter Database, the average annual burned area in YPMC forests for the period 2010-2017 was more than twice the average annual burned area between 1984 and 2009 (~39 000 ha vs. 18 800 ha). Even so, this accounts for less than 1/3 of the average annual burned area before Euro-American settlement.

Components of the fire regime: fire season—

NRV and comparison to current—The concept of “fire season” refers qualitatively to the average period of the year during which wildfire activity can be expected. Qualitatively or quantitatively, there are a variety of ways to define the fire season. For example, Westerling et al. (2006) defined fire season as the number of days between the first wildfire occurrence of the season and the final wildfire control date. Thonicke et al. (2001) defined the annual length of the fire season by summing the probability of at least one fire in a day across the calendar year.

The assessment area experiences one of the longest fire seasons in North America because of the combination of fuels fed by winter and spring moisture and
the 3- to 6-month period of drought that coincides with the warm season (Keeley et al. 2012, Sugihara et al. 2006). Years of widespread burning in the assessment area are driven strongly by drier-than-average years, and this pattern stretches back to the beginning of the fire scar record (Trouet et al. 2010).

The fire season is driven to a great extent by temporal patterns in fuel moisture, which are in turn largely dependent on winter and spring precipitation. YPMC forests in the assessment area receive from about 25 to 50 percent of their precipitation as snow (Minnich 2007), depending on elevation and latitude, among other things. Fire season in YPMC forests north of the Tehachapi Mountains (i.e., north of approximately latitude 35° N) is generally considered to run from late June to October (Fried et al. 2004, Sugihara et al. 2006), but the average length varies with latitude, elevation, topography, and of course annual precipitation and temperature. Fire season tracks and slightly lags the climatic dry season. An approximation of the fire season length and timing can be derived from the Walter climate diagrams in figure 2. Because of fuel and soil moisture carryover from the wet season, coarse forest fuels are typically too moist to burn for some time after the beginning of the climatic dry season, and they remain dry enough to burn for some time after the beginning of the wet season (see fig. 2); fine fuels respond much more rapidly to meteorological wetting and drying. Thus, the fire season may be reasonably approximated with a 2- to 4-week lag on either end of the climatic dry season. The fire scar record shows that fires historically tended to occur late in the fire season (after cessation of summer tree growth) in the northern assessment area, but more in the middle of the fire season in the southern assessment area; areas farther south continue the earlier trend, with fires in the Baja California YPMC forests tending to occur in the early to middle fire season (Skinner 2002, Skinner et al. 2008, Stephens and Collins 2004).

In a study of wildfire in the Western United States, Westerling et al. (2006) found that fire season length had increased by more than 2 months during the period between 1970 and 2003. They attributed the earlier start of the fire season to earlier snowmelt from higher spring and summer temperatures.

Future fire regimes—
Flannigan et al. (2009) summarized the results of dozens of future fire activity projections under different climate change scenarios and concluded that most evidence pointed to increased fire occurrence and area burned. Referring to North American in general, Gedalof (2011) similarly noted that “nearly all… (modeling) efforts predict a substantial increase in wildfire activity over the next century…” Bachelet et al. (2007) concluded that “despite imprecise knowledge of future climate and human behavior, it is reasonable to conclude that fires will likely increase in the West.” Safford et al. (2012b) stated that “modeling studies specific to California
expect increased fire activity to persist and possibly accelerate under most future climate scenarios.” Miller and Urban (1999a) noted that the altered water balance that results from climate change can influence fire regimes both directly, through its influences on fuel moisture, and indirectly, through its influences on forest structure, composition, and fuel loads. Miller and Urban (1999a) concluded by stating, “If GCM predictions of future climate prove true, fires could be both more frequent and of greater spatial extent at certain sites.” Many modeling efforts have been undertaken that either focus on or include California, and we summarize these efforts below.

**Fire frequency and fire rotation**—Miller and Urban (1999a) simulated the effects of climate change on forest biomass, composition, and fire regimes across an elevational gradient in Sequoia & Kings Canyon National Parks. For their lowest two sites, which occur within YPMC forests, fire frequencies rose markedly during the first century of their climate change simulations, then dropped over time as woody biomass was increasingly lost. By the end of their 400-year climate change simulations, woody biomass had decreased at the 1800-m site from 0 to 4 m²/ha overall, leaving little fuel to burn; under their most extreme scenario, forest fuels at this site were completely replaced by fine (grassy) fuels, and fire frequency rose. At the 2200-m site, biomass loss was also high but not as extreme as at the 1800-m site, and fire frequencies remained similar to the baseline conditions, although fire area decreased with the decrease in biomass (Miller and Urban 1999a). The 2600-m site, which is currently in red fir forest but was predicted to transition to a mixed-conifer composition, experienced very large increases in fire frequency.

Loudermilk et al. (2013) used LANDIS to model carbon and fire dynamics under future climates for the Lake Tahoe Basin. Under their more severe future climate scenario (+4.6 °C and –22 percent precipitation compared to today), the projected fire rotation period dropped by 18.6 percent compared to today. Under a more benign future climate scenario, the rotation period dropped by about 6 percent. Both of these projections assumed similar fire behavior to today, however, as well as continued success of fire suppression efforts, such that fire rotations continued to be quite long even under the much warmer future scenario (around 200 years).

Yang et al. (2015) used a averaged spatial point process modelling approach to project future “fire occurrence density” for the Lake Tahoe basin. Using the same future climate scenarios as Loudermilk et al. (2013), they found that—compared to today—the density of fires per unit area increased by 210 percent under the more extreme future climate and 70 percent under the more benign scenario.
Fire severity and intensity—Flannigan et al. (2000) modeled the seasonal severity rating (SSR) across North America under two global circulation model (GCM) scenarios for the year 2060. SSR measures the difficulty of fire control. Flannigan et al. (2000) found that SSR increased by an average of 10 percent under both GCM scenarios for California. Lenihan et al. (2003a, 2003b) modeled vegetation and fire response to different GCM-based future climate scenarios for California, using the MC1 dynamic vegetation model. One of their mid-stream outputs was fireline intensity, measured as Btu/ft/sec (see Lenihan et al. 2003b). Lawler et al. (2012) summarized the fireline intensity outputs for the range of the California fisher in the southern Sierra Nevada (most of the west slope of the Sierra Nevada in the assessment area southern region), under a moderately warmer, slightly drier future climate scenario. They found that Lenihan et al.’s (2003a, 2003b) model projected higher fire intensity across about 35 percent of the fisher range, lower intensity on 15 percent of the area, and similar fireline intensity on 50 percent of the area.

Fried et al. (2004) modeled fire behavior while accounting for suppression response for lands under state of California fire management. Under a conservative future climate scenario, Fried et al. (2004) found that decreased fuel moistures and increased seasonal windspeeds under climate warming led to faster burning and more intense fires. Most of their increased fire activity was due to an increase in the number of fires burning under extreme conditions. Flannigan et al. (2013) linked the Canadian Forest Fire Weather Index to three GCMs and predicted the cumulative severity rating (CSR), a fire danger metric based on weather conditions, for the Northern and Southern Hemispheres for the periods 2041–2050 and 2091–2100. They projected that severity as measured by CSR would increase by 10 percent to ~30 percent by the later period over most of the assessment area. Yue et al. (2013) modeled trends in biomass consumption, which scales with severity, for the Western United States. They projected mean increases of 80 to 90 percent in biomass loss by the middle of the 21st century for their Pacific Northwest region, with the largest changes projected for the Sierra Nevada and coastal forests in northwestern California and western Oregon. Van Mantgem et al. (2013) showed that high prefire climatic water deficit increases the probability of postfire tree mortality, thus—aside from their well-known effects on fuel moisture—climate warming and increasing growing season drought can enhance fire severity independently of fire intensity. This suggests that future fire severities could be even higher than predicted by most climate modeling studies.
**Burned area**—In Miller and Urban’s (1999a) model, area burned at their 1800-m and 2200-m sites rose strongly during the first century of their climate change simulations, then decreased over time as woody biomass was gradually lost. At the lowest site, little woody biomass remained at the end of their simulation, and the abundance of grassy fuels led to a large increase in area burned. At the 2200-m site, fire area decreased as biomass was lost over time (Miller and Urban 1999a). Their red fir-forest site at 2600 m experienced very large increases in area burned. Fried et al. (2004), using a relatively conservative future climate scenario, compared 6-year simulations for current and future conditions. Their future simulation burned 43 percent more area in mixed-conifer forest than the current scenario.

Lenihan et al. (2008) simulated fire and vegetation dynamics in California under three future climate change scenarios. In all three of their scenarios, the assessment area experienced some of the greatest increases in simulated area burned (20 to 50 percent increases), especially on the Modoc Plateau, at higher areas of the southern Cascade Range, and in middle- to high-elevation forests along the Sierra Nevada axis. Spracklen et al. (2009) built regression models linking observed climate with observed wildfire area burned and used a GCM to project burned area out to 2050. They projected an increase of about 180 percent in annual area burned for their Pacific Northwest region, which included most of the assessment area. National Research Council (NRC 2011) modeling projected that compared to the average for the 1950–2003 period, median annual area burned would increase by more than 300 percent for the assessment area with a 1 °C increase in average temperature. The report also noted that, over time, extensive warming and wildfire could ultimately exhaust the fuel for fire in some regions, as forests were completely burned (NRC 2011).

Westerling et al. (2011) modeled burned area across California under a range of future climate and development scenarios. They found that, under the most realistic future climate and emissions scenarios and compared to the average for the period from 1960 to 1990, area burned by wildfire would increase by more than 100 percent by 2085 for most of the forested area of northern California. Middle and higher elevation forests in the assessment area would be among the most severely affected, with some future climate scenarios producing increases in burned area of more than 300 percent. Yue et al. (2013) used regression and parameterization approaches to project burned area under 15 GCMs for the Western United States for the middle of the 21st century. Their Pacific Northwest region, which included most of the assessment area, would experience projected median increases of 42 and 154 percent in area burned by wildfire, depending on the approach. Loudermilk et al. (2013) used LANDIS to model carbon and fire dynamics under future climates for the Lake
Tahoe basin. They found that under their more severe future climate scenario, the mean annual area burned would climb by 43 percent by the year 2100.

McKenzie et al. (2004) calculated correlations between mean summer temperature and precipitation and annual burned area for 11 Western States between 1916 and 2002, then employed regression models to project burned area into the future under two emissions/climate scenarios. They found strong relationships between their summer climate variables and fire area for all states but California and Nevada, and concluded that most of the Western United States was likely to experience large increases in annual area burned by wildfire in the 21st century. However, they concluded that “fire in California and Nevada appears to be relatively insensitive to summer climate, and area burned in these states may not respond strongly to changed climate.” In their study, McKenzie et al. (2004) made two errors with respect to their analysis in California. First, they neglected to account for California’s Mediterranean climate, which features a summer drought of 3 to 6 months. Second, McKenzie et al.’s (2004) analysis combined southern and northern California, which each contribute about half of California’s total burned area in an average year but which are extremely different in their fire-climate relationships, and often do not share the same extreme years. Their analysis thus buried the relatively strong relationship that exists between fire and summer climate variables (in this case, temperature) in the assessment area and other parts of central and northern California (Miller et al. 2009b, Trouet et al. 2006, Westerling et al. 2006) under the southern California fire-climate relationship, which is essentially independent of summertime temperature or precipitation (Keeley 2004). In summary, changes in summer temperature and precipitation may not have strong effects on southern California fire area, but McKenzie et al.’s (2004) predictions for the Western United States in general are likely to have validity for most of the assessment area.

Fire ignitions—Because the human population of California is expected to increase to nearly 50 million by 2050—and a large proportion of that increase is forecast for the Central Valley and Sierra Nevada foothills (http://www.dof.ca.gov/Forecasting/Demographics/projections/documents/P1_County.xlsx)—it is reasonable to conclude that the density of human fire ignitions will also increase over time. Educational efforts can help to reduce fire ignitions and improve public safety, but more people usually means more fire (Syphard et al. 2009).

Ignitions by natural causes may also increase through the 21st century. Price and Rind (1994) simulated lightning distributions and frequencies under a GCM-based future climate scenario incorporating 4.2 °C global warming by 2100. They projected that lightning incidence could increase by up to 30 percent globally. Romps et al. (2014) found similar results, based on the linear relationship between lightning
flash rate and the product of precipitation (per hour) times convective available potential energy (which measures atmospheric convective instability). Romps et al. (2014) projected a 12 percent average increase in lightning per degree Celsius of temperature rise. This could translate into 50 percent more lightning across much of the United States by the end of the 21st century. Although neither study highlighted the assessment area as a global “hotspot” for increased lightning occurrence, the combination of greater lightning incidence, warmer climates, and drier fuels inexorably leads one to conclude that fire activity will likely rise in most semiarid areas currently supporting forest. Yang et al. (2015) found that lightning ignitions could increase in the future even without an increase in lightning occurrence, as the lengthening of the fire season, loss of snowpack, and drying of fuels combine to expand the conditions under which lightning strikes led to fuel combustion.

**Fire effects on vegetation**—Fire is a major driver of vegetation change in both space and time. Fire activity and behavior are tied strongly to fuel amount, fuel moisture, and weather conditions at the time of burning (Agee 1993, Sugihara et al. 2006). All these factors are influenced by both temperature and precipitation. There is near-universal agreement that the assessment area climate will become notably warmer through this century, but projections of future precipitation patterns are much less certain (Dettinger 2005). The effects of fire on vegetation in the assessment area will depend greatly on precipitation trends, but Bachelet et al. (2007) noted that in either wetter or drier conditions, forest could be notably reduced in much of the Western United States in a warmer future. Under drier conditions, enhanced fire frequency could favor drought-tolerant grasses, which would further enhance ecosystem flammability and reduce woody cover. Under wetter conditions, expansion of woody plants might promote more intense fires and high mortality when drought conditions occur, ultimately reducing tree biomass.

Bachelet et al. (2007) projected that most of the assessment area would see an increase in biomass consumption by fire during the 21st century, whether warming was extreme or moderate, and whether carbon emissions were relatively high or low. Most pixels in the assessment area experienced increases in biomass loss of 25 to 67 percent when compared to the 1961–1990 period.

Using the same deterministic vegetation dynamics model as Bachelet et al. (2007), Lenihan et al. (2008) simulated the future distribution of terrestrial ecosystems in California under three GCM-based future climate scenarios (see fig. 10). The “MC1” model used by Lenihan et al. (2003a, 2003b) explicitly models fire and its effects on vegetation; modeling results were driven to a notable extent by fire. Total annual area burned in California increased under all three scenarios, ranging from 9 to 15 percent above the 1961–1990 mean by the end of the 21st century.
Figure 10—Lenihan et al. (2008) modeling results for the Sierra Nevada (A) and Sierra Nevada Foothills (B) ecological sections, comparing current conditions to future projections of vegetation extent. These ecological sections include most of the Sierra Nevada western slope. The GFDL-B1 scenario = moderately drier than today, with a moderate temperature increase (<3.1 °C); PCM-A2 = similar precipitation to today, with a <3.1 °C temperature increase; GFDL-A2 = much drier than today and much warmer (>4 °C higher). All scenarios project significant loss of subalpine and alpine vegetation. Most scenarios project lower cover of shrubland (including west-side chaparral and east-side sagebrush), owing principally to increasing frequencies and extent of fire. Large increases in the hardwood component of forests are projected in all scenarios except for the hot-dry scenario in the Foothills section. Large increases in grassland cover are projected for the Sierra Nevada section. The drier scenarios project moderate expansion of arid lands. In the Sierra Nevada section, coniferous forest decreases in cover under all scenarios. Illustration is adapted from Safford et al. (2012b).
Fire drove grassland expansion into former shrublands and woodlands, even under the coolest and wettest future scenario; by 2099, under the warmest and driest scenario, grassland almost completely replaced shrublands on the Sierra Nevada west slope and also expanded greatly in the California portion of the Great Basin. Broadleaf woodland and forest replaced large areas of evergreen conifer forest under all three scenarios, with fire playing an important role in the transition, especially in the relatively warmer and drier scenarios (Lenihan et al. 2008). Under the more moderate climate scenario, annual consumption of biomass by fire by the end of the 21st century across California was about 18 percent greater than the 1961–1990 mean. Under the two drier and warmer scenarios, simulated biomass consumption by fire was greater at first, but as woody biomass was burned, vegetation type conversions led to large reductions in overall biomass available for burning by the end of the century.

Fire season—Although overall precipitation in much of the assessment area has been steady to rising slightly over the past three-fourths of a century, the snow:rain ratio and snowpack depth and persistence have all been decreasing (Safford et al. 2012b). Combined with warming temperatures, these negative trends in snow amount and storage result in fuels drying earlier. Current trends and projections of future patterns in the snow:rain ratio and snowpack persistence thus portend longer fire seasons (Mote 2006, Mote et al. 2005, Safford et al. 2012b, Westerling et al. 2006).

Liu et al. (2010) projected future changes in “wildfire potential” by modeling the Keeetch-Byram Drought Index under a number of different future GCM-based scenarios. Under their projections, fire season will be a couple of months longer for much of the contiguous United States, including the assessment area, by the end of the 21st century. Flannigan et al. (2013) projected that fire season length would increase by more than 20 days for most of the assessment area by 2050, and for all of the assessment area by 2100. Basing their work on 15 GCMs, Yue et al. (2013) projected a median increase of more than 3 weeks in the fire season for the Pacific Northwest, which included most of the assessment area, by the middle of the 21st century.

Fire Suppression

The evolution of many plant species has been greatly influenced by their long-term relationships with fire (Bond and van Wilgen 1996, Keeley et al. 2012, Sugihara et al. 2006). Human-caused changes to natural fire regimes can have significant impacts on the diversity and composition of native plant and animal communities (Hobbs and Huenneke 1992, Mutch 1970). Because of the high ecological and evolutionary importance of frequent fire to assessment area YPMC forests, it has
been argued that the **exclusion of fire** from most of the assessment area for the last century is one of the most significant human-caused ecological disturbances currently in play (Barbour et al. 1993, Fites-Kaufman et al. 2007, Sugihara et al. 2006). The scientific and management literature is overflowing with assessments of, and references to, the deleterious effects of fire exclusion on assessment area ecosystems (summaries in Agee 1993; Barbour et al. 1993, 2007; Erman and SNEP Team 1996, Keeley et al. 2012; Sugihara et al. 2006, and others.). These include altered species composition and dominance patterns, increased fuels and forest density, impacts to soils and hydrological cycles and carbon sequestration, loss of important wildlife habitat, increased fire intensity and severity, decreased human safety, threats to infrastructure, and so on.

Cermak (2005) provided a detailed consideration of the development of the fire control organization and policies in California. The desire to control fire came largely as a response to the destructive burning practices of early settlers, and also the belief that frequent fires were destroying timber and reducing the capacity of the forest to regenerate. Controversies surrounding the use of “light burning” to reduce forest fuels and protect old growth developed in the 1910s and 1920s, and again in the 1950s, but in both cases proponents of fire exclusion prevailed. In 1910, Region 5 (Pacific Southwest Region) Regional Forester Coert DuBois directed his forest supervisors that fire control was the top management priority in the Forest Service’s Pacific Southwest Region. He followed this with the 1914 publication of *Systematic Fire Protection in the California Forests*, which Cermak (2005) called the “most influential single document in U.S. fire control history.” It set fire control standards (forest fires were to be controlled before they reached 10 ac [4 ha]), and it described the outlines for a formal fire control organization and the processes for coordinated fire planning. In 1919, Region 5 directed forest supervisors to suppress all fires, even on neighboring private land. In 1924, the California Board of Forestry endorsed “fire exclusion” from forest lands as state policy. A policy of overnight fire control was discussed at a national Forest Service meeting in 1935, and emerged as the famous “10 a.m.” rule in May of that year (whereby Forest Service units were expected to have fire starts controlled by 10 a.m. the day after discovery).

Federal land managers were already actively working to extinguish fires when the first forest reserves were established in California at the end of the 19th century, but the lack of training, coordination, planning, and technology meant that their ability to stop large fires was very limited (Cermak 2005). The first trained fire crews were established in the late 1920s, and the adoption of more modern techniques and technologies gradually led to increasing success in fire suppression.

Consultation of the California Fire Perimeters database (see footnote 4) shows a
strong drop in fire frequency and annual burned area in the 1930s and 1940s. This was helped by a series of wet years in the late 1930s, but by the end of the 1940s a number of innovations had markedly improved firefighting success, including the deployment of tanker trucks and bulldozers, the institution of “hotshot” fire crews and “smokejumpers,” and the expanded use of planes and helicopters in patrols and aerial water drops (Cermak 2005).

Patterns in fire frequency show remarkable success in fire control through most of the rest of the 20th century (the Forest Service succeeds in extinguishing 98 percent of all ignitions before they reach 300 ac [120 ha]; [Calkin et al. 2005]), but beginning in the 1980s, the area of forest burned began to climb. By the 1990s, 10-year running averages for annual burned area and average fire size were at their highest points since formal recordkeeping began in 1908, and the upward trend continues (Calkin et al. 2005; see figures in Miller et al. 2009b). Since 1910, 15 fires have exceeded ~20,000 ha (~50,000 ac) in size in the assessment area, but 14 of these have occurred since the late 1970s, and 12 since 1995.

Part of the trend in area burned and fire size is due to changed federal fire management policies. In the face of research and management reviews showing the detrimental ecological effects of fire exclusion on Western forest ecosystems (e.g., Biswell 1961, Leopold et al. 1963), the National Park Service began permitting prescribed fires in California in the late 1960s and early 1970s, and allowed some lightning ignitions to burn under prescribed conditions (van Wagtendonk et al. 2002). At the same time, Forest Service wilderness areas experimented with management, rather than suppression, of naturally ignited fires (Stephens and Ruth 2005). The Forest Service changed its policy from strict fire control to fire management in 1974, and formally abandoned the 10 a.m. rule in 1978 (Pyne 1982). By limiting direct attack on difficult fires, and taking greater advantage of topography, natural barriers, and weather to “indirectly” control fires, fire management agencies themselves have played a role in the growth of large fires since the late 1970s.

Nonetheless, the evidence is overwhelming that accumulated fuels and changes in forest structure resulting from a century of fire exclusion have led to major ecosystem changes in forest types that experienced frequent, primarily low-severity fires before Euro-American settlement (e.g., Agee 1993; Barbour et al. 1993, 2007; Erman and SNEP Team 1996; Leopold et al. 1963; Parsons and DeBenedetti 1979; Steel et al. 2015; Sugihara et al. 2006; etc). In interaction with climate warming, these forest changes are now resulting in larger and more severe fires throughout the YPMC forest belt, not only in the assessment area but across the southwestern United States (Dillon et al. 2011, Mallek et al. 2013, Miller and Safford 2012, Miller et al. 2009b, Skinner and Chang 1996). In summary, fire suppression is a major
Natural Range of Variation for Yellow Pine and Mixed-Conifer Forests in National Forests of California

disturbance factor in assessment area YPMC forests, both in its direct modification of ecosystem composition, structure, and function, and in its contribution to increased forest fuels amounts and continuity, which are leading to deleterious effects when forest fires escape control. Nearly every other section in this chapter contains additional information pertaining to the negative ecological effects of fire suppression on YPMC forests.

Grazing

American Indian inhabitants of the assessment area did not herd animals, and livestock grazing occurred only after Euro-American settlement. A short summary of the grazing history of YPMC forests is offered here to provide context to current conditions and to the early observations that Euro-Americans made of the assessment area. For additional grazing-related information, see “Grass and forbs” on page 153.

Appreciable livestock grazing began in assessment area YPMC forests after the arrival of Euro-American settlers after 1849. By the 1860s, valley and foothill ranchers were using public lands in the Sierra Nevada on a seasonal basis to graze their herds of cattle (Dasmann 1965, Jackson et al. 1982, Pease 1965). Sheep grazing was also practiced in much of the Sierra Nevada after about 1860 (McKelvey and Johnston 1992). The herding habits, huge numbers, and more general diet of sheep caused major effects on Sierra Nevada ecosystems, especially riparian areas and meadows, and probably affected fire regimes as well by reducing fine fuels. Leiberg (1902) viewed grazing, especially by sheep, as a “destructive agent to the forest by preventing reforestation.” Muir (1894) referred to sheep as “hoofed locusts.” Sudworth (1900) militated for stricter control of sheep grazing. Conflicts developed between cattle ranchers and shepherds; and public concern with the effects of grazing, particularly by sheep, was one of the factors leading to the designation of the forest reserves in the 1890s and early 1900s. Shepherds and cattlemen also often set fire to the forest in the late summer or fall to clear the forest understory and ostensibly to improve forage; in some cases these fires caused major damage to YPMC and red fir forests, mostly where previous logging had increased surface fuel loads (Cermak 2005, Greeley 1907, Jackson et al. 1982, Leiberg 1902, Sudworth 1900).

Vankat and Major (1978) noted that livestock grazing, especially by sheep, had affected most of Sequoia National Park. However, their references to specific records of overgrazing refer almost exclusively to montane meadows and high-elevation forests, and they do not list grazing as a major change agent for YPMC forests. Sheep grazing in the park ceased in the early 20th century, so there has been nearly a century for park ecosystems to recover.
Brewer’s memoirs from 1861 to 1864 (Brewer 1930) referred to the scarcity of good grass cover in the Sierra Nevada, and lush areas of grass were highlighted where they occurred. Brewer’s team traveled on horseback, so they were reliant on the availability of forage. Brewer’s time in the Sierra Nevada predated heavy cattle or sheep grazing, although he mentioned grazing in his memoirs. Sudworth’s team also relied on pasturage, and Sudworth (1900) noted that unfenced forest land supported very “short forage” and expressed the opinion that sheep grazing had decimated herbaceous and grass cover in much of the central and southern Sierra Nevada, basing his statement on the “study of long-protected forest land in the same region” and conversations with older settlers. Most of Sudworth’s unpublished notes refer to higher elevation locations, however, not mixed conifer. Leiberg (1902) also primarily referred to higher elevations (red fir, principally) when discussing the deleterious effects of grazing.

The period between 1894 and 1904 was extremely dry across southern California (but not as catastrophically dry in northern California, except between 1897 and 1899) (USDI 1951); most of the oft-cited observations of deleterious impacts of heavy grazing on Sierra Nevada ecosystems are from this period (e.g., those cited in McKelvey and Johnston 1992). Note also that the years in which Sudworth (1900) and Leiberg (1902) conducted their field studies coincided with the third longest recorded period of profound drought in California (as measured by the Palmer Drought Severity Index—PDSI), which included the 2nd driest year on record (1898), and the 3rd driest 2-year span (1898–1899) (NOAA National Climate Center data for 1895 to 2015: https://www.ncdc.noaa.gov/cdo-web/search?datasetid=GHCND). The extreme dryness of the soil and depleted herbaceous cover noted by observers during this period was ascribed by many of them entirely to sheep grazing, but the extreme climatic conditions certainly played a major role. Old settler’s memories were of times before significant sheep grazing but also of much more abundant rainfall.

Whatever the case, heavy grazing in much of the assessment area clearly reduced understory cover and affected soil in parts of the assessment area for many decades. It probably also reduced fire frequency in some parts of assessment area YPMC forests by reducing the amount of fine fuel. Swetnam and Baisan (2003) noted that many Sierra Nevada fire histories show a virtual absence of fire after the 1850s, which they attributed to the introduction of large herds of sheep into the range after the 1859–1860 drought. A soon to be published fire history study from the southern Modoc Plateau (Adin Pass area) shows a very early cessation of fires that coincides closely with the introduction of cattle to the study area, and other studies showing very low local FRIs (<8 years) also suggest that herbaceous fine fuels would have been necessary to support such high fire frequencies (see footnote 8). These studies are in northern and northeastern assessment area forests dominated by ponderosa pine, and may follow the model suggested for the Southwestern United
States (Arizona and New Mexico) by Swetnam and Betancourt (1998), where fire regimes in ponderosa pine forest were hypothesized to respond strongly to herbaceous fuel production, while fire regimes in mixed-conifer forests were suggested to be driven primarily by woody fuels. Note, however, that the general lack of summer precipitation in the assessment area results in much less grass cover on average than in yellow pine forests in the Southwestern United States, which receives much summertime rainfall; see “Forest understory and nonforest vegetation” on page 146.

Given that most heavy (especially sheep) grazing ceased before World War I, one question is to what extent the effects of this disturbance have lasted over the ensuing century. Grasses are extremely resilient to disturbance, and their seeds are very easily dispersed. A further issue is that the institution of fire suppression and the cessation of heavy grazing happened at about the same time. This complicates our ability to discern the independent effects of the two disturbances. It also means that forests were densifying (and reducing understory light availability) just as understory plant communities were being freed from decades of heavy pasturage. This probably stalled understory recovery and possibly led to different patterns of succession than would have occurred had fire not been suppressed.

Insects and Disease

Background information—

Table 5 lists major insects and diseases found in tree species of assessment-area YPMC forests. A more complete listing and description of injurious insects and diseases can be found at http://www.fs.usda.gov/Internet/FSE_DOCUMENTS/fsbdev3_046410.pdf.

Noteworthy increases in ponderosa pine, Jeffrey pine, and sugar pine mortality in the Sierra Nevada range can usually be attributed to moisture stress, high tree density, and elevated bark beetle activity (Young et al. 2017). The primary bark beetles associated with ponderosa pine mortality are western pine beetle (*Dendroctonus brevicomis*) and mountain pine beetle (*Dendroctonus ponderosae*). Mountain pine beetle also kills sugar pine. Jeffrey pine beetle is the primary killer of Jeffrey pine. Dwarf mistletoe (*Arceuthobium* M. Bieb.) and *Heterobasidion* root disease cause additional stress on host trees; the spread of *Heterobasidion* is abetted by logging when cut stump faces are not treated with borax (Slaughter and Rizzo 1999). Black stain root disease is scattered throughout the northern Sierra Nevada range and can be found in ponderosa and Jeffrey pine. White pine blister rust has been devastating to sugar pine since the disease entered northern California around 1930.

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9 “Insects and Disease” text primarily by Sheri Smith, USDA, Forest Service, Pacific Southwest Region regional entomologist, Lassen National Forest, 2550 Riverside Drive, Susanville, CA 96130.
White fir mortality throughout the Sierra Nevada is typically attributed to fir engraver beetle, moisture stress, and *Heterobasidion* root disease. High tree density and true fir dwarf mistletoe (*Arceuthobium abietinum* (Engelm.) Hawksw. & Wiens) also contribute toward decline in some areas. Douglas-fir tussock moth readily defoliates white fir in the Sierra Nevada. Population cycles trend upward every 7 to 10 years, and significant levels of tree mortality have been recorded during past outbreaks.

Insects rarely kill incense cedar. Bark beetles that attack incense cedar are not considered aggressive tree killers; however, when combined with drought stress, they can cause mortality. During drought periods in some areas of the Sierra Nevada range, small incense cedars are the first trees to decline and die. *Heterobasidion* root disease and true mistletoe also weaken incense cedar.

Douglas-fir in the Sierra Nevada can be heavily affected by insects or diseases typical of more northerly latitudes, but their incidence in the Sierra Nevada is reduced. However, Douglas-fir beetle, flatheaded fir borer, and black stain root disease can be found in some Douglas-fir stands. Both insects are capable of killing trees, particularly drought-stressed ones. The detected incidence of black stain root disease in Douglas-fir in the Sierra Nevada is low.

Several insects and diseases can be found on native oaks. Typically the extent or severity of their effects are not widespread or protracted. Foliar injury can result

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**Table 5**—Major insects and diseases found in tree species of assessment-area yellow pine and mixed-conifer forests

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<td>Heterobasidion root disease, <em>Heterobasidion</em> spp.</td>
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<td>Black stain root disease, <em>Leptographium wageneri</em></td>
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<td>Armillaria root disease, <em>Armillaria</em> spp.</td>
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<td>Dwarf mistletoe, <em>Arceuthobium</em> spp.</td>
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<td>White pine blister rust, <em>Cronartium ribicola</em></td>
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<td>Western pine beetle, <em>Dendroctonus brevicomis</em></td>
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<td>Jeffrey pine beetle, <em>Dendroctonus jeffreyi</em></td>
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<td>Mountain pine beetle, <em>Dendroctonus ponderosae</em></td>
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<td>Fir engraver beetle, <em>Scolytus ventralis</em></td>
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<td>Douglas-fir beetle, <em>Dendroctonus pseudotsugae</em></td>
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<td>Douglas-fir tussock moth, <em>Orgyia pseudotsugata</em></td>
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<td>Flatheaded fir borer, <em>Melanophila drummondi</em></td>
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*a* Table courtesy of Sherri Smith, USDA Forest Service Pacific Southwest Region regional entomologist.

from a variety of diseases, insects, and mites. Wood-boring beetles are usually restricted to dead or dying branches, although the recent emergence of gold spotted oak borer in southern California is a worrying sign, as it readily kills adult black oak, canyon live oak, and coast live oak. Damage by these agents is normally secondary in nature, rather than the primary cause of branch or tree decline. *Armillaria* root disease and true mistletoe can commonly be found on oaks.

Uprooting and stem breakage of giant sequoia is not uncommon and can be a problem along roads and in recreation areas. *Heterobasidion* root disease is sometimes found infecting the roots of fallen trees. Tree killing of giant sequoias by insects or diseases is rare.

**NRV and comparison to current**—We have little information on insect or disease occurrence in presettlement YPMC forests in the assessment area. Based on insect and forest ecology, however, some inferences can be made about probable changes over time. Fettig (2012) provided a list of the bark beetle species that cause “significant” mortality in the assessment area. Within YPMC forests, most research has been done on the beetles affecting yellow pine species, especially beetles from the genus *Dendroctonus*, as they can have major impacts on mortality rates in commercially important stands of trees.

It has been understood for some time that tree stand densities have a strong relationship to bark beetle-induced mortality. Higher density stands increase competition for resources (especially water and light) and reduce tree vigor, which makes individual trees less able to withstand insect attack. Various studies demonstrate that lower density YPMC stands are much less susceptible to bark beetle attack and subsequent mortality (Fettig et al. 2007, Young et al. 2017).

In the current absence of frequent understory fire, bark beetles have become one of the principal agents of tree mortality in the assessment area (Fettig 2012, Manley et al. 2000). Under reference conditions, frequent fire would have interacted with insects and disease, as well as abiotic and biotic site conditions, to drive stand structure (Bonnicksen and Stone 1982, North et al. 2012b). Much more open and heterogeneous forest structure resulted, and—based on the strongly inverse stand density versus bark beetle relationship—we can infer that bark beetle-caused mortality was probably lower than under current conditions.

Evidence from comparisons between contemporary forests in the assessment area and reference sites in Baja California support this inference. The Lake Tahoe Watershed Assessment compared modern disease and insect incidence in old-growth forest versus mid-seral forests in the Lake Tahoe basin and also versus old-growth forests in the SSPM (Manley et al. 2000). Mid-seral forests in the Lake Tahoe basin generally supported more mortality (29 percent mortality), more pest incidence, and more dwarf
mistletoe infestation than old-growth stands (21 percent mortality; note that these values are overall standing mortality and not annual mortality rates). Assessment was made after a 5-year drought (1987 to 1992), and compared to conditions in the SSPM, which experienced similar levels of drought. Overall mortality in the SSPM, which was not logged and has seen frequent fire throughout most of the 20th century, was only 12 percent. Savage (1997) assessed mortality in the SSPM versus mortality in YPMC forests in the Transverse Ranges of southern California. In her stands, she found only 4 percent mortality in the SSPM forests versus 14 percent in the southern California sites. Her interpretation was that low densities of trees in the Mexican sites (kept low by recurrent fire) reduced soil moisture competition and thus tree stress. Stephens and Gill (2005) measured mortality before and after a major drought, and found that cumulative mortality in Jeffrey pine/mixed-conifer forests in the SSPM ranged from 2.7 to 3.6 percent. It is unclear why there is such a discrepancy between the Savage (1997) and Stephens and Gill (2005) mortality measures and the Manley et al. (2000) number. Manley et al. (2000) concluded that fire continues to be responsible for most stand thinning in the SSPM, but insects have replaced fire as the major thinning agent in the Lake Tahoe basin.

One assessment area insect species for which presettlement outbreak dynamics have been documented is the pandora moth (*Coloradia pandora*), which is a defoliator of yellow pines in and around areas with very loose, usually pumice-based soils (which are required for pupation); such soils occur frequently in the northern and eastern portions of the assessment area. The pandora moth is generally not a mortality agent, but serious defoliation can result in significantly slowed growth among affected trees. Data are not from the assessment area itself, but rather from central Oregon in ponderosa pine forests. Pohl et al. (2006) identified six major outbreaks between 1800 and 2000 (mostly at 40- to 50-year intervals) and demonstrated a strong correlation between outbreaks and the occurrence of drought, which weaken trees’ resistance and increases susceptibility to herbivory. It seems likely that similar drought-driven dynamics also characterize pandora moth outbreaks in the east side of the assessment area.

Most recently, as a result of the interactions of high stand densities with 4 years of below-average precipitation and the warming climate, YPMC stands in the southern Sierra Nevada have begun to exhibit very high levels of beetle mortality, both in terms of local intensity as well as the geographic extent. Ponderosa pine, sugar pine, and white fir are being the hardest hit, but direct drought-driven mortality is beginning to be evident in species like incense cedar as well. Some smaller watersheds in the southern Sierra Nevada have seen >75 percent of their ponderosa pine die over the past few years. As this mortality wave moves northward, large areas of YPMC forest are being left in conditions that have no precedent in the historical (i.e., since 1850) record.
Future—Fettig (2012) noted that climate warming may benefit bark beetles in multiple ways. For example, warming may allow extra generations to complete their life cycles each year, and adult beetle emergence and flight could occur in the season and continue further into the fall. Under warming temperatures, cold-induced mortality during the winter may also decrease. Bentz et al. (2010) maintained that future thermal regimes, assuming continued warming, will be very favorable for many bark beetle species. Mountain pine beetles are likely to become especially damaging to higher elevation conifer forests. Huge warming and drought-driven beetle outbreaks have recently occurred in the United States and Canadian Rockies, and such events may become more common in the future. Bark beetles currently restricted to the southwestern-most United States or Mexico will also likely move northward as climates warm.

Kliejunas (2011) carried out a risk assessment of the likelihood that current and projected future climate change trends would lead to increased adverse effects by eight pathogens on Western U.S. forests. Kliejunas (2011) estimated risk potential as the likelihood of increased disease damage multiplied by the potential consequences (which included both ecological and economic components) of increased disease damage. High to very high risk potentials were identified for dwarf mistletoe and *Armillaria* root disease under both drier and wetter future climate projections; both pathogens affect numerous conifer species in the assessment area. A moderate risk potential was identified for white pine blister rust (which affects sugar pine and western white pine) under both precipitation scenarios. Swiss needle cast (which affects Douglas-fir) and *Dothistroma* needle blight (which affects pines and Douglas-fir) were assigned moderate risk potentials under wetter future conditions, but low potential under drier future conditions.

Logging

Although American Indians used trees for a variety of purposes, large-scale timber harvest did not begin in the assessment area until after Euro-American settlement. Like grazing, a short summary of the history of Euro-American logging in the assessment area is provided here in order to provide some context to current conditions, and to the observations of forest conditions that were made by Euro-Americans in the late 19th and early 20th centuries.

McKelvey and Johnston (1992) summarized the logging history of the Sierra Nevada up to about 1900. Most logging before the turn of the 19th century was done to support mining and the communities that arose to support mining. Timber was cut to build homes and commercial buildings, tunnels, mine and ore processing infrastructure, and railroad lines, and it was the fuel for heating, railroad engines
and other machines, and the various types of mills used for processing ore; in some areas, a very valuable market in sugar pine shakes (for roofing or siding) also arose (McKelvey and Johnston 1992). The great majority of timber harvest both before and after 1900 occurred in yellow pine and mixed-conifer forest.

According to McKelvey and Johnston (1992), most 19th-century logging occurred on private lands near the mines and communities, at elevations below the eventual national forests. Exceptions to this rule included the canyon of the South Fork American River, the Yuba River watershed, the Tahoe National Forest (which supported a high density of mines even at higher elevations and was crossed by the transcontinental rail line) (Jackson et al. 1982), and the basin of the Truckee River including Lake Tahoe. Where cutting did extend into what would become national forest lands, it was often selective rather than general, and focused on the most valuable trees, usually sugar pine and yellow pine (Leiberg 1902, Sudworth 1900), although less valuable species might be cut for fuel for steam engines or smelting (see footnote 3).

Between the 1890s and 1920s, railroad lines were extended throughout the lower and middle elevation Sierra Nevada to allow access to timber resources that were beyond reach of animal-drawn transport. More than 80 private railroad logging companies existed at one time (Beesley 1996). Use of railroads for transport allowed for much more generalized and intensive logging to take place, and the northern assessment area was particularly affected, as well as areas of the Truckee River basin and Lake Tahoe, the South Yuba River, and the Eldorado and Stanislaus National Forests (Conner 1997, Jackson et al. 1982, Laudenslayer and Darr 1990, TCHS 2016). According to Beesley (1996), by 1934 more than half of the mixed-conifer forest land in the “north-central” Sierra Nevada had experienced some level of harvest, primarily of the more valuable pine species. Private lands were the focus of most of the harvesting, and by the 1940s, almost all the remaining uncut forest land in the Sierra Nevada was found on national forests or national parks.

After World War II, demand for timber from federal lands increased dramatically, and the Forest Service greatly expanded its involvement in the timber industry. As an example, harvest on the Eldorado National Forest expanded from about 3.8 million board feet per year between 1902 and 1940, to 35.1 million board feet during the war, to more than 56 million board feet per year between the end of the war and 1959 (Beesley 1996). Before the war, Forest Service lands provided about 5 percent of the nation’s supply of lumber. By the end of the war the total had risen to 10 percent, and by 1970 Forest Service lumber supplied almost one-third of the U.S. need (Beesley 1996). In the Sierra Nevada, the period of the heaviest timber
harvest occurred in the 1950s and 1960s. Harvest techniques were more generalized than before the war, and large areas of forest were clearcut. Since the 1960s, national legislation, regulations, changing economics, and environmental concerns have acted in concert to greatly reduce the amount of logging on public lands in the Sierra Nevada and elsewhere, although private lands have made up some of the difference. In the end, Barbour et al. (1993) estimated that, “half the original acreage of the mixed-conifer forest has been… cut at least once in the last 150 years.”

The effects of logging on forest stand structure have been sufficiently described in the literature and will only be summarized here. Historically, most cutting in the assessment area has been at least partly selective, with the largest and most valuable trees from a handful of species (mostly sugar pine and yellow pine) being targeted. Areas of clearcutting also occurred, mostly between the 1950s and 1970s. In both cases the tree canopy is opened up, increasing light incidence in the understory and often resulting in a pulse of herbaceous and then shrub growth, followed by ingrowth of mostly single-aged tree regeneration (either planted or not). In selectively cut areas, smaller and less valuable trees were retained, usually from fir species or incense cedar, which greatly increased their relative presence in the forest stand. Without further management and in the absence of fire, harvested stands, which may originally have been dominated by fire-tolerant species like sugar pine or yellow pine, mature to much denser stands of more evenly aged individuals of shade-tolerant/fire-intolerant species, sometimes with scattered remnant overstory pines. Most of the shade-intolerant/fire-intolerant tree species are competitively dominant in the absence of recurrent disturbance (Burns and Honkala 1990, USDA FS 2013b) (see “Introduction”). Compared to the original, preharvest forests, these forests tend to be much denser, structurally much more homogeneous, more shaded, and less biodiverse (Agee 1993, Barbour et al. 1993, Fites-Kaufman et al. 2007). Fuel structure in these secondary forests is also highly altered. Fuel continuity is often greatly increased over the preharvest condition in both horizontal and vertical directions. Surface fuels are often increased owing to accumulation of timber harvest “slash,” in addition to the accumulation of fuels resulting from fire exclusion. Fir species support low canopy-base heights and often grow under the canopy of overstory dominant trees; should fire occur, fir species are easily ignited from surface fuels and act as “fire ladders” into the canopy of large trees that are otherwise protected by their thick bark or lack of lower branches. Modern timber harvest practices are designed to avoid or even to mitigate many of these negative effects, but many forest stands in the assessment area harvested before the last few decades are highly departed from structural and compositional conditions that characterized presettlement forests.
Nutrient Cycling

NRV and comparison to current—
Like most of the semiarid Western United States, tree growth in assessment area ecosystems is primarily limited by water availability rather than by nutrients (Major 1988, Stephenson 1990). However, in some situations and on some substrates significant nutrient limitation may occur, such as on ultramafic (“serpentine”) rock types, or where water availability is sufficient to permit dense vegetation growth. Where nutrient limitation does occur, it is usually for nitrogen (N) or, much less commonly, phosphorus (P). We know of no studies of nutrient dynamics in YPMC forests that have not experienced long-term fire exclusion. Because fire is such an important contributor to decomposition and nutrient flux (Johnson et al. 2009, Wohlgemuth et al. 2006), we focus on studies that (1) have investigated the nutrient dynamics of YPMC forests that have experienced fire, and (2) provide some insight into how fire suppression may have changed nutrient dynamics.

Compared to other temperate and boreal forest types, assessment area YPMC forests support very low rates of litter decomposition (Hart et al. 1992, Stohlgren 1988). This is due to the Mediterranean-type climate of the assessment area and general lack of precipitation during the growing season. Most decomposition occurs in the spring, often under snow, as temperatures begin to warm but soil moisture is still high (Johnson et al. 2009, St. John and Rundel 1976, Stohlgren 1988). Nitrogen and phosphorus are immobilized in needle litter and slowly released. Because of the near absence of precipitation in the growing season, trees in upland YPMC forests do not tend to generate extensive roots in the soil O horizon, and the slow decomposition rates of litter in most of the assessment area mean that nutrients mineralized from the O horizon and not taken up by plants are often leached into runoff waters (Johnson et al. 2009). Spatiotemporal variability in nutrient pools and fluxes is very high in YPMC forests in the assessment area. Spatial and temporal variation in moisture availability (mostly from seasonal snowmelt) is a major driver of nutrient dynamics, but fire is an even greater source of variability in some types of nutrient flux. Johnson et al. (2009) noted that fire exclusion has allowed large increases in the depth of the O horizon and subsequent buildups in N and P, which are released to stream waters by overland flow during rain events and may be significant contributors to the deterioration of Lake Tahoe water quality.

St. John and Rundel (1976) studied the effects of fire on nutrient dynamics in a giant sequoia/mixed-conifer forest. Phosphorus, cations (calcium [Ca], potassium, and magnesium), and pH levels were all higher in burned plots, while nitrogen and carbon (C), as well as cation exchange capacity (CEC), all decreased in burned plots. St. John and Rundel (1976) noted that although the cations were made avail-
able for plant uptake by fire, they were also susceptible to loss by runoff unless they were taken up by plants or held in exchange sites, which are in low supply after fire owing to the loss of much of the prefire organic layer and needle litter. The loss of N to volatilization could potentially also be balanced over time by any increase in N fixation promoted by fire, such as the postfire succession of N-fixing shrubs (e.g., *Ceanothus* spp.) or herbs (legumes) (Wohlgemuth et al. 2006).

Other studies have found similar results. For example, Chorover et al. (1994) studied soil solution chemistry before and after prescribed fire in YPMC forests in Sequoia National Park and documented very large increases in cation concentrations in soil water, and a drop in ammonium (a source of N) to below prefire levels within the year after fire; however, nitrate, another important source of N, rose and remained elevated for 3 years (Chorover et al. 1994). Stephens et al. (2004) found that prescribed fires in YPMC forests in the Lake Tahoe basin released Ca and raised pH of both soil and nearby stream water. Oliver et al. (2012) studied the effects of a severe wildfire on stream chemistry in a YPMC forest and found that P, nitrate, and cations (measured with electrical conductivity) were all elevated after fire.

Sobota et al. (2013) recently reported on geographic patterns of anthropogenic N input in the United States. Western portions of the assessment area are affected by atmospheric deposition of N from automobile exhaust and Central Valley use of synthetic fertilizers, especially in the central and southern sections. Central California is highlighted as one of the most heavily affected areas for N input in the entire United States. Maps in Sobota et al. (2013) indicated that the western portions of the southern assessment area are receiving from 5 to 10 times (or more) the annual input of N than they probably received before Euro-American settlement. In terrestrial ecosystems, high levels of N are known to increase invasion and survival of fast-growing annual weeds (Porter et al. 2013), which compete strongly for water, light, and nutrients with native plants, and in some cases have major effects on fire regimes (e.g., cheatgrass [*Bromus tectorum* L.]). High levels of N input can also have major effects on tree ecology, including stimulation of growth, changed root:shoot ratios, induced nutrient deficiencies or imbalances, and reduced drought tolerance (Bytnerowicz and Fenn 1996). Gaseous N can also act as a pollutant, and various forms of toxic and acidifying N compounds are formed with oxygen. Effects of N deposition on YPMC forests in the assessment area are not as extreme as in southern California. In both places, most N arrives as dry deposition, and the summer drought and limited dry season stomatal conductance mean that plant utilization of gaseous N is more limited than in more humid regions (Bytnerowicz and Fenn 1996).
Future—
Current trends in the assessment area are for continued suppression of most fires, with escaped fires becoming larger and more severe over time. This is a fundamentally different situation than what characterized presettlement YPMC forests, where fires were frequent and largely of lower severity and intensity. It is very difficult to generalize about how these trends may affect nutrient cycling in assessment area YPMC forests. One of the most significant outcomes of continued fire suppression is likely to be the enhancement of the nutrient pool in the forest litter layer and its episodic release by severe fires. Further enhancement of nutrients will continue to be carried by wind in the form of nutrient deposition (mostly N) from human sources to the west. Higher levels of soil N will likely abet further invasion by annual invasive weeds and subsequent changes in forest understories and fire regimes.

Miesel et al. (2009) conducted an interesting study in which they compared belowground soil and microbial variables in forest stands mechanically treated to favor ponderosa pine versus stands treated to favor large trees regardless of species. They found notable differences between the two treatment strategies, especially in soil organic C content and N availability, and pH. This work suggests that future soil nutrient status may depend to some degree on the tree species favored in forest management.

Successional Processes
Surprisingly little empirical and quantitative documentation of successional patterns in YPMC forests in the assessment area has been published. Nonetheless, early observers of YPMC forests in the assessment area were already well acquainted with the different ecological tolerances and successional tendencies of the major tree species. For example, Sudworth (1900), Leiberg (1902), and Greeley (1907) all referred to the strong potential within YPMC forest for dense seedling recruitment of the shade-tolerant/fire-intolerant species in the absence of fire (see chapter 1). The species differences referred to in tables 1 and 2, and figures 3 and 4 interact with the environment and ecological disturbances to drive successional processes in YPMC forests.

Leiberg (1902) stated that the relative proportion of tree species in assessment area YPMC forests was changing because of timber harvest and fire. In general, he noted that the relative proportions of sugar pine and yellow pine were decreasing, as recruitment of young trees was not keeping up with their removal from the overstory by logging (one exception was the northern part of his central survey area). At the same time, he described “a uniform increase” in the proportions of incense cedar and white fir across the survey area. Overall, the YPMC forests that Leiberg surveyed had low densities of tree seedlings and saplings, owing to the effects of
frequent fire. However, he noted that stands of YPMC forest that had escaped fire for 12 to 15 years were often filled with stands of saplings “so dense that a man can with difficulty force his way through” (Leiberg 1902: 43). Sudworth (1900) also noted that, “The frequent open spaces in yellow pine forests are sooner or later covered with dense patches of young trees, but these thickets may in turn be swept off by fire.” Show and Kotok (1924) made the same point, namely that fire protection in the pine belt in the Sierra Nevada had resulted in “an enormous number of young forest trees that have appeared as individuals and in groups, or, in the more open virgin stand, as a veritable blanket under the mature timber.”

The rate of forest infilling in the absence of fire varies along environmental gradients. For example, studies in assessment area YPMC forests have found that seedling recruitment, survival, and growth are inversely related to elevation (Hunter and Van Doren 1982, van Mantgem et al. 2006), and topographic exposure and insolation are also major drivers of seedling survival and growth rates (Kolb and Robberecht 1996, Maguire 1955). Local soil conditions and topographic- and vegetation-defined (e.g., nurse plants) microhabitats can also play a major role in seedling survival, young tree growth, and rates of forest succession and densification in the absence of frequent disturbance (Gomez et al. 2002, Tappeiner and Helms 1971). In the assessment area, dense shrub cover can have a major effect on future forest composition as well, as shade-tolerant trees (e.g., white fir) are more likely to survive the decades it may take to overtop the shrub canopy (assuming that fire can be kept out of the stand, in which case succession will be reset) (Stark 1965). Another major driver of seedling density and forest infilling is temporal coincidence between large seed crops and years with favorable climate (high precipitation, occurrence of summer thundershowers, low summer temperatures, etc.) (Burns and Honkala 1990).

Bonnicksen and Stone (1982) provided a summary of successional dynamics in moist mixed-conifer forest (including giant sequoia). Bonnicksen and Stone (1982) popularized the notion of the “shifting mosaic” of successional stages on the landscape, where neighboring sites of the same ecological “potential” could be in dissimilar vegetation states owing to different spatiotemporal processes and their rates. They stressed that the nature and rate of different successional pathways depended on abiotic and biotic conditions of the site in question. That said, a generally recognized truism is that white fir is the competitive dominant in most YPMC forests in the assessment area, and the long-term absence of fire will ultimately lead to white fir forests (or Douglas-fir at lower elevations in the north assessment area). Fires in the presettlement period were frequent and mostly of low severity, but some aggregations of mature trees would nonetheless be periodically killed by fire, while others were left untouched, and in yet others the
understory vegetation and tree regeneration would be consumed by the passage of
fire. Such patches of high-severity fire would often be succeeded by dense areas of
fire-promoted shrubs, and return to forest in such areas might take many decades
(Show and Kotok 1924). Although fire was frequent, there were always tree clumps
that had escaped fire for several FRIs, and thickets of shade-tolerant species (white
fir, incense cedar) would often develop in these places. Regeneration of species
like giant sequoia and the yellow pine requires mineral soil and canopy gaps large
enough to bring sunlight to the forest floor (Burns and Honkala 1990, Meyer and
Safford 2011). Recruitment of these species thus required fire intense enough to kill
clumps of canopy trees; sugar pine tends to favor smaller gaps (Burns and Honkala
1990, Oliver and Dolph 1992). Note that Bonnicksen and Stone’s (1982) ideas
were applicable to patchiness at a very fine scale (they explicitly state that they are
referring to patches between 0.0135 and 0.16 ha). As is noted in multiple locations
throughout this document, coarse-grained patchiness on the order of thousands,
hundreds, or even tens of hectares was uncommon in YPMC forests before Euro-
American settlement.

Overall, the general picture is one of very high potential for forest recruitment,
especially by shade-tolerant species, with frequent fire or soil conditions maintain-
ing the dominance of the pine (and in some places, giant sequoia) overstory and a
more open forest condition (Bonnicksen and Stone 1982; Kilgore and Taylor 1979;
Leiberg 1902; North et al. 2002, 2005; Show and Kotok 1924, Sudworth 1900).

Models of YPMC forest successional dynamics—
Kercher and Axelrod (1984) developed a Monte Carlo-based model of YPMC
forest succession (SILVA) at the stand level to better understand the effects of
fire on forest dynamics in the Sierra Nevada. The SILVA model is complex, and
includes more than 30 subroutines that model such phenomena as species-specific
demographic rates (recruitment, growth, death, injury, etc.), stand structures, fire,
and brush and litter dynamics. Fire effects on trees were estimated as a function of
scorch height and tree diameter, but weather inputs were mostly held constant, so
the simulated fire regime was relatively crude (Agee 2000). Kercher and Axelrod
(1984) used SILVA to compare forest succession after a simulated clearcut for 500
years at two different elevations, 1520 m (5,000 ft) and 1830 m (6,000 ft). The lower
elevation site is at the upper reaches of YPMC forests historically dominated by
ponderosa pine; the upper site is nearer the upper limits of YPMC forests in the
Sierra Nevada and historically included a significant component of fir species.

The time-averaged results of Kercher and Axelrod’s (1984) lower elevation
simulation are shown in figure 11. Figure 12 shows the successional progression of
the lower elevation YPMC stand through the 500 years of the SILVA simulations.
Fire was modeled as a stochastic process with a mean return interval of 7 years. Ponderosa pine and black oak dominate the stand immediately after the initial stand-replacing fire, but black oak becomes subordinate to incense cedar and then white fir by 70 to 100 years and almost completely drops out of the stand by 300 years (fig. 12). After 200 years, overall basal area varies between 50 and 54 m²/ha, and the relative dominance of species changes but the proportions of shade-intolerant to shade-tolerant species fluctuate around 70:30 until after 400 years, when the proportion of ponderosa drops. Simulations without fire supported much higher basal area of shade-tolerant/fire-intolerant species like white fir and incense cedar. The higher elevation simulation is in the “fir zone” and supported much more white fir than ponderosa pine, even under frequent fire (Kercher and Axelrod 1984).

![Figure 11](image-url)
Van Wagtendonk (1984) carried out simulation modeling of YPMC forest succession using an improved version of the FYRCYCL stand-level forest dynamics model. The model included subroutines on vegetation, fuels, fire, weather, and lightning ignitions. Historical fire weather was used to drive the fire regime, and fires could be of any intensity. Van Wagtendonk (1984) carried out 200-year simulations under three management conditions: no fire, natural lightning ignition regime, and fire suppression (which permitted fires to burn under certain extreme conditions). Starting conditions were a seedling patch of 40 percent (by basal area) ponderosa pine, 25 percent sugar pine, 20 percent white fir, and 15 percent incense cedar. Under the no-fire scenario, ponderosa pine increased to >55 percent of basal area by 90 years (taking advantage of originally open stand conditions), then began to drop as white fir accrued individuals and basal area; white fir dominated the stand after 150 years. Under the lightning-ignition scenario, fires occurred on average every 8.9 years (the first fire occurred at 34 years) and ponderosa pine comprised more than 90 percent of stand basal area by the end of the simulation. Sugar pine was the most important codominant species in this scenario, but by the end of the simulation it was less than 8 percent of the
stand as reckoned by either basal area or density. Stand densities varied widely in the lightning-ignition scenario, depending on fire frequency and intensity (van Wagtendonk 1984). Under the fire-suppression scenario (basically a modern business-as-usual scenario), two moderate- to high-severity fires occurred, reducing white fir density much more than basal area (as some white fir had reached sufficient size to survive intense fire), and resulting in a system dominated by a fluctuating balance of ponderosa pine and white fir (but with ponderosa pine always dominant in terms of basal area). Sugar pine and incense cedar were only minor players in all three scenarios.

Keane et al. (1990) developed the FIRESUM successional process model as an upgrade to SILVA and applied it to understanding successional dynamics in ponderosa pine/Douglas-fir forests in the inland Northwest under different fire regimes. Keane et al.’s (1990) study area includes the northern tip of the assessment area, and four of the forest types (“fire groups”) modeled by FIRESUM either occur in assessment-area YPMC forests or are similar (warm, dry ponderosa pine; grand fir [Abies grandis is ecologically similar to white fir]; warm, dry Douglas-fir; moist Douglas-fir). In Keane et al. (1990), FIRESUM was used to carry out a 200-year model of successional dynamics in a semiarid ponderosa pine-dominant stand beginning in 1900. The major findings were that Douglas-fir was able to establish in the stand only when FRIs reached 50 years, but ponderosa pine still dominated the site under these conditions, with about 50 percent of the basal area at year 200 (Larix occidentalis and Douglas-fir comprised the remainder). Under no fire, Douglas-fir comprised one-third of the total basal area by year 100, and dominated the stand by 130 years; at the end of the simulation, Douglas-fir was about 65 percent of the total basal area (65 m²/ha). Under the frequent fire scenarios (<50-year FRIs), most basal area was contributed by large trees, but at FRIs of 50 years and above, fuels accumulated and fires were intense, which resulted in stands of small to intermediate trees at high densities.

Miller and Urban (1999a, 1999b) and Urban et al. (2000) described an adaptation of the ZELIG forest gap model for forests along an elevational gradient in Sequoia & Kings Canyon National Parks in the southern Sierra Nevada. In various publications, Miller and Urban and colleagues employed the modified ZELIG model to study climate change scenarios, carbon dynamics, the effects of fire on stand parameters, the importance of the physical habitat and moisture availability, and so on, but they did not publish results on the actual successional dynamics between species within their simulations. We refer to these studies in various other places in this assessment.
All the simulation models referred to above make clear the initial advantage that the yellow pine species have in frequent fire scenarios. Seedlings and saplings of ponderosa pine and Jeffrey pine (and sugar pine) grow rapidly in high light environments (fig. 3), and as young trees they support thicker bark than their competitors (fig. 4). Both adaptations provide for higher survival under recurrent fire. Where fire is not frequent, or overstory cover is high, the yellow pines are ultimately outgrown by shade-tolerant species.

**Future**—It is unknown how future climates and conditions may affect basic successional processes in assessment-area YPMC forests. If future environmental conditions differentially affect key species in YPMC forest, then successional relationships among species may change. An example is the effect of white pine blister rust on the five-needle pines, which in YPMC forests are represented by sugar pine, and, to a lesser extent, western white pine. Aside from these sorts of effects, it appears likely that warming temperatures and increasing fire activity on some of the landscape, but continued fire exclusion on most of the landscape, will simply accelerate the sorts of successional changes we have already witnessed for the past half century.

**Tree Mortality**

**NRV and comparison to current**—

We have little historical information on tree mortality processes or patterns in assessment-area YPMC forests. Peter Decker, an observant miner whose journal from his 1849 to 1851 trip to California has been published (Giffen 1966), noted after 3 years in the Sierra Nevada that “the woods in California seem like the original or first growth. How seldom to be seen a dead tree unless fired.” Greeley (1907) came to a different conclusion, but his viewpoint was certainly colored by his formal silvicultural training. Greeley decried what he saw as “high densities” of dead, broken-topped, and fire-hollowed trees. Greeley’s (1907) estimate was that 1 to 5 percent of the stems in sugar and yellow pine were dead. Because this is a cumulative mortality estimate and snags remain standing for years, the actual annual mortality rates in the forests Greeley visited were probably very low.

Although periodic drought and high interannual variability in precipitation are characteristic of the California climate, assessment-area YPMC forests are generally much denser today than under presettlement conditions. Higher stand densities lead to higher competition for light, nutrients, and water. In the standard scenario, “self-thinning” ensues, in which younger, smaller, or weaker individuals succumb to competition and die, while larger, healthier trees benefit from the release of competition (Barbour et al. 1987, Westoby 1984). Although the mortality rates of smaller trees tend to be higher, recruitment in the small size classes is also higher, and many
recent studies documented surprisingly high mortality rates of large trees in YPMC and other forests in and around the assessment area, even before the 4-year drought and subsequent mortality wave (e.g., Dolanc et al. 2013, 2014b; Dolph et al. 1995; Fellows and Goulden 2012; Lutz et al. 2009; Smith et al. 2005; van Mantgem et al. 2009; Walker et al. 2006). Most researchers have concluded that some combination of water stress, stand density, and insect-driven mortality are to blame.

The few data available suggest that background mortality rates (averaged over multiple years) in assessment area forests are between about 0.25 and 1.4 percent for fire-excluded forests and less than 0.5 percent for contemporary reference forests with a largely intact fire regime (Ansley and Battles 1998, Maloney and Rizzo 2002, Stephens and Gill 2005). Modern plantation studies broadly corroborate these numbers, and show large increases in annual mortality between low- and high-density stands. Data in Zhang et al. (2006), for example, give annual mortality rates of between 0 and 0.8 percent in thinned stands of fewer than 332 trees/ha, versus rates of 0.6 to 2.3 percent in stands of more than 2,450 trees/ha.

Van Mantgem et al. (2009) showed that mortality rates in Western U.S. forests have strongly increased over the last four to five decades. Of the three large regions compared—Pacific Northwest, western interior United States, and California (in California all of van Mantgem et al.’s [2009] study sites are located within the bioregional assessment area)—the assessment area experienced the highest mortality rates, and the greatest rate of increase in mortality rates. Averaged over many sites, the rate changed from less than 1 percent in the 1980s to about 1.6 percent in the early 2000s. Furthermore, the greatest rates of increase in mortality were found in species of pine, in (formerly) frequent-fire forests, within the elevation belt occupied by YPMC forests; overall, the highest annual rate of mortality was found in the studied species of pine, which died at an annual rate of about 2 percent during the early 2000s (van Mantgem et al. 2009). Of course, mortality rates, especially among the pines, have skyrocketed since 2014, with tens of millions of dead trees mapped by the Forest Service as a result of the 4-year drought and subsequent beetle outbreak.

**Future**

Adams et al. (2009) showed that increases in temperature increased mortality in two-needle pinyon (*Pinus edulis* Engelm.) in the Southwestern United States via carbon starvation. When coupled with drought and other factors, such as insect outbreaks, Adams et al. (2009) concluded that future warming will increase background mortality rates, and also greatly increase the risk of regional die-off events. They called for similar experimental assessments of other tree species. Carnicer et al. (2011) studied tree species responses to increased temperatures in the Iberian Peninsula, which supports a similar Mediterranean climate to much of California. They assessed
16 species, mostly pines and oaks. They found that interactions between drought, temperature, forest density, insects, and disease were resulting in increased defoliation and mortality among the studied species, and they noted that climate change-driven drought pressures on Mediterranean tree species were likely to lead toward increased damage and mortality in the future. Allen et al. (2010) summarized the results of about 150 worldwide studies of tree mortality and concluded that the data were consistent with an increase in mortality resulting from climate warming and drought.

Wind

**NRV and comparison to current**—
There is little information on the effects of windthrow on YPMC species in the assessment area. YPMC stands were historically relatively open, and species adapted to these conditions must have some resistance to high winds. Larsen and Woodbury (1916) noted that wind was not generally an important disturbance factor in sugar pine stands. According to maps in Peterson (2000), California and neighboring states are subject to fewer major wind events like tornados and convective wind events (“downbursts”) than any other part of the contiguous United States. Very high winds are common when winter storms arrive at the Sierra Nevada crest, but these elevations support red fir and subalpine forests for the most part. It seems unlikely that wind as a disturbance agent has changed significantly between presettlement and current times.

The most famous wind event to hit California was the 1962 Columbus Day storm, which was precipitated by a strong tropical storm entering northern waters and interacting with the jetstream (Lynott and Cramer 1966). Many windspeed records at northern and northwestern California meteorological stations were set by this storm, and gusts above 90 kph (56 mph) were common in the northwestern part of the assessment area (southern Cascades). Huge areas of forest were blown down, with the most damage done outside the assessment area (around 11 billion board feet were blown down in Oregon and Washington) (Lynott and Cramer 1966).

Hillmire et al. (2012) reported on what they termed the most extensive windthrow event on record in the Sierra Nevada sensu stricto. An event with winds exceeding 145 kph (90 mph) occurred in the central Sierra Nevada in the upper San Joaquin River basin in the fall of 2011. Thousands of mature trees were downed, mostly red fir, white fir, and lodgepole pine. In some areas, more than 70 percent of live trees were downed. Large trees were more susceptible to uprooting than small trees, and the effects were relatively evenly distributed across species. Larger snags were also more likely to uproot than small ones (Hillmire et al. 2012). This size-dependent effect has a very different impact on forest structure than fire, which preferentially kills smaller trees.
Future—
Changing weather patterns resulting from climate change may increase windthrow as large-scale pressure systems shift at the continental scale (Peterson 2000).
However, in the assessment area, it seems likely that windier conditions would have greater effects on fire effects to forest than directly on the forest itself. For example, Fried et al.’s (2004) model of future fire activity under fire exclusion showed that increased windspeeds during the fire season were an important factor behind the increased number of modeled fire escapes. High winds during the wet season can also influence the outcome of future fires. The 1999 Megram Fire in northwest California burned primarily through a landscape full of wind-thrown trees and broken tops that resulted from a strong wind event a few years earlier. This led to more severe fire effects to the soil in some places and hampered fire control efforts (see footnote 3).

Structure
Forest Landscape Structure
NRV—
Analyzing landscape structure in assessment-area YPMC forests is difficult. Historically YPMC forests were not characterized by large, stand-replacing disturbance events, but rather by highly frequent, low- and moderate-severity events that did not generally leave a coarse-grained pattern on the landscape that could be easily mapped. Even after a century of fire exclusion, an increase in forest density, and the vestiges of logging and increased occurrence of high-severity fire, it can be challenging to conduct the sorts of classic landscape structure assessments that are routinely done in forests of the Eastern United States or northern Europe. Because of that, we urge some caution in interpreting “patchiness” in assessment-area YPMC forests from standard vegetation maps or model outputs that track successional/seral stages.

Although it may have been difficult to find much coarse-grained patchiness in presettlement YPMC forests, gradients of soil depth, moisture, temperature, and insolation combined to drive differences in forest structure between cool and warm aspects, high and low elevations, and upper, middle, and lower slopes (species changes across these gradients are treated in the “Forest composition and species diversity” section under “Composition” below). Such landscape differences in forest structure are still noticeable today (Barbour et al. 2007; Fites 1993; Lydersen and North 2012; North et al. 2002, 2012b). The major driver of this variation is water availability, but it was greatly modified by fire before fire exclusion. See chapter 1 for a summary of the effects of topography on forest structure in the assessment area. In general, areas with higher water availability (north aspects, higher
elevations, lower slopes) tend to support higher forest density and cover (and other related variables) than areas with lower water availability (south and west aspects, lower elevations, upper slopes). When exposed to fire, more densely forested areas tend to exhibit a more coarsely grained landscape structure than open forest areas, because fire effects are more severe where fuels are more continuous and fuel loadings higher (Agee 1993, Sugihara et al. 2006).

Soil depth and texture are the long-term products of interactions between topographically driven water availability and vegetation, and both of these variables also play a major role in driving vegetation conditions. Aside from the few truly unproductive soil types (serpentine soils, for example), the major effect of soil on vegetation in assessment-area YPMC forests is its contribution to water availability (see chapter 1). Meyer et al. (2007) found that soil depth had a strong influence on the basal area and canopy cover of YPMC forest at a site in the southern Sierra Nevada, and North et al. (2012b) described soil depth as a major driver of forest structural and compositional heterogeneity at the landscape scale.

Fire is also an important driver of forest landscape structure in assessment-area YPMC forests. When high-severity fire occurs, it normally results in a successional process dominated in its initial stages by montane chaparral, which includes shrub species that germinate in response to fire (van Wagendonk et al. 2012). The boundaries between forest and chaparral stands are quite abrupt in the first decades after fire, but in the absence of further burning, trees slowly reoccupy these sites. This process can take many decades. If the sites burn again, the process may take centuries, or the chaparral may become a semipermanent feature in the landscape (Nagel and Taylor 2005, Show and Kotok 1924, Skinner and Taylor 2006). The denser forests typically found in areas of higher water availability are more likely to burn at high severity owing to high fuel loadings and fuel continuity (unless fire has been very frequent in the stand over sufficient time to reduce fuels), and owing to the enhanced presence of fire-intolerant species like white fir (Agee 1993, North et al. 2012b, Sugihara et al. 2006). Such environments are therefore more likely to support coarse-grained landscape structures than drier forests supporting higher densities of fire-tolerant species.

The proportion of presettlement landscapes occupied by defined patches of chaparral was certainly variable, but there are few sources of information to base an estimate on. Bonnicksen and Stone (1982) carried out a reconstruction study in a YPMC-giant sequoia forest in Kings Canyon National Park, and estimated that about 19 percent of the study site in the 1890s was occupied by shrub “aggregations” (shrubfields); by 1977, the proportion had dropped to 11 percent. Show and Kotok (1924) reported on the area of national forests in northern California that
supported “brushfields” in the early 1920s, which were seral chaparral stands that
had resulted from (often human-caused) fires in previously forested areas. Their
estimate of 11.1 percent of the landscape on six national forests in the assessment
area is slightly higher than the current area of montane and mixed chaparral that
occurs on productive forest land on the same national forests (8.6 percent). See table
9 and “Shrubs” in “Forest understory and nonforest vegetation” for more detail.

Franklin and Fites-Kaufman (1996) used contemporary conditions in assessment
area national parks (which were not logged) to estimate that a maximum of
55 percent of assessment-area conifer forests would have been in “old-growth”
status before Euro-American arrival. It is important to qualify that Franklin and
Fites-Kaufman’s (1996) concept of “old growth” was not restricted to areas of high
 canopy cover, and they explicitly noted that much assessment-area old growth
would have been in areas of open canopy.

Models of landscape structure—The interagency LANDFIRE Program developed
a nationwide map of potential vegetation representing the distributions of vegetation
types as they are hypothesized to have existed prior to Euro-American settlement.
The vegetation types are referred to as biophysical settings (BpS), and are linked
to state and transition models (see “Fire severity” above for more detail). State and
transition models are nonequilibrium, nonlinear models linking defined states (in
this case, vegetation seral stages defined by tree size and canopy cover) via path-
ways that are driven by forces that can transition a site between states (e.g., succes-

Using Van de Water and Safford’s (2011) crosswalk between their presettlement
fire regime types and the LANDFIRE BpS types, we determined that BpS models
610270 (Mediterranean California Dry-Mesic Mixed Conifer Forest and Wood-
land), 610280 (Mediterranean California Mesic Mixed Conifer Forest and Wood-
land), and 610310 (California Montane Jeffrey Pine–Ponderosa Pine Woodland)
best represented dry mixed conifer, moist mixed conifer, and yellow pine, respec-
tively (see http://www.landfire.gov for BpS descriptions). One of the outputs of the
BpS state and transition models is an average end state for many runs of each
model over a 500-year period. The end states are assumed to represent a snapshot
of the average landscape condition during presettlement times, with the caveat that
the landscape is greater than about 5000 ha.10 LANDFIRE uses these reference
values to compare to current conditions in order to develop a measure of departure
(“fire regime condition class,” FRCC). Figure 13 provides the distribution of

10Keane, B. 2009. Personal communication. Research ecologist, USDA, Forest Service,
Rocky Mountain Research Station, 5775 W. Broadway Street, Missoula, MT 59808.

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reference values for the three models above, portrayed as percentages of the landscape occupied by each of five seral stages: early successional (after severe fire), mid successional (closed canopy and open canopy), and late successional (open and closed canopy). The early-successional stage represents anything not dominated by trees >10 cm diameter at breast height (d.b.h.) (>4 in), and therefore includes areas dominated by herbaceous plants, shrubs, and seedling and saplings. The “open” mid- and late-successional stages include areas of medium and large trees with canopy cover <50 percent (<40 percent in the yellow pine type), the “closed” mid- and late-successional stages include areas of medium and large trees with canopy cover >50 percent (>40 percent in yellow pine).

The LANDFIRE BpS models predict that, under the presettlement fire regime, 15 to 20 percent of the average YPMC landscape would have been in early seral stages (herbs, shrubs, seedlings/saplings) and young forest, about 35 percent in areas dominated by trees between 12.5 and 53 cm d.b.h. (5 to 21 in), and 45 to 50 percent in areas dominated by trees >53 cm d.b.h. (fig. 13). Furthermore, the BpS models predict that most of the landscape was under open forests of less than 50 percent canopy cover (“open” stages), especially in the yellow pine and dry mixed-conifer types (fig. 13). Dense, older stands (“late closed”) are predicted to have occupied around 5 percent of the landscape in the yellow pine and dry mixed-conifer types, but around 20 percent of the moist mixed-conifer type.

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Figure 13—Average landscape conditions for presettlement yellow pine–mixed-conifer forests in the assessment area, as predicted by LANDFIRE BpS state and transition models for LANDFIRE modeling region 6. These are applicable only on landscapes greater than about 5000 ha in area. LANDFIRE BpS reference conditions do not include error bars and are rounded off to the nearest 5 percent by LANDFIRE for ease of use by managers. See text for definitions of successional classes.
The LANDFIRE BpS models are only models, but the BpS outputs are surprisingly close to other, independent estimates of presettlement forest landscape structure in assessment-area YPMC forests. Compare, for example, Franklin and Fites-Kaufman’s (1996) estimate of around 55 percent of assessment area conifer forests in “old-growth” status before Euro-American arrival with the 45 to 50 percent of landscape in the BpS late-successional classes (fig. 13). Also, Bonnicksen and Stone’s (1982) estimate of about 19 percent chaparral cover in their 1890 landscape is very close to the landscape proportion for early-seral vegetation projected by the BpS model, as is the 15 to 25 percent range in shrub cover found in studies of unlogged (and in some cases periodically burned) YPMC forests in and near the assessment area (see “Shrubs” on page 148). Overall, we urge some caution in applying the LANDFIRE outputs, as they assume that distinct seral stages can be mapped and followed through time. As noted above, this is a tenuous assumption in YPMC forests, which under reference conditions were characterized by fine-grained heterogeneity that does not lend itself well to landscape-level mapping of structure (forest densification and increased occurrence of high-severity fire in modern stands makes seral stage mapping and modeling somewhat more tractable).

**Comparison to current**—Overall, it can be safely generalized that the landscape structure of current assessment-area YPMC forests is more coarsely grained (characterized by larger, more defined patches) than under presettlement conditions. This is largely due to the interacting effects of timber harvest and fire suppression. Early selective cutting removed the large pine trees from many areas of YPMC forest, and with the long-term absence of fire, natural succession has led to infilling by higher densities of young, mostly fire-intolerant species. More recent clearcutting, mostly on private lands, has increased the representation of coarse-grained and “hard-edged” patchiness in assessment-area YPMC forests. Fire exclusion has greatly increased forest fuel loadings and continuity, and driven a species dominance shift to fire-intolerant trees, which together have led to an increase in the occurrence of large, stand-replacing fires. Like clearcutting, these types of fires also lead to a much coarser-grained, more hard-edged pattern in assessment area forests than was typical before Euro-American settlement. Fry et al. (2014) documented a decrease in spatial complexity in fire-suppressed forests of the Sierra Nevada, compared with forests in the Sierra de San Pedro Mártir in Mexico with a relatively intact frequent fire regime. As noted in the “Fire regime” section above, because of human management over the past 150 years, assessment-area YPMC forests have largely
transitioned from Fire Regime I, characterized by frequent, low-severity fires and fine-grained heterogeneity in forest and fuels structure, to Fire Regimes III and IV, which are characterized by infrequent fires with a much greater high-severity component and a much coarser grained forest structure. More information on these fire regime and forest structure transformations is found in “Fire regime” above (see especially “High-severity patch size”), and “Forest structure” below (e.g., “Forest gaps and tree clumps”).

Many studies and reviews refer to the high structural homogeneity of contemporary YPMC forests (Agee 1993, Barbour et al. 1993, 2007; Erman and SNEP Team 1996, Sugihara et al. 2006), and the return of more heterogeneous landscape structure is currently a major management focus (North 2012, North et al. 2012a, North et al. 2012b). The high homogeneity of fire suppressed YPMC forest landscapes in the assessment area is being increasingly broken up by large, severe wildfires, but most of the assessment area has not experienced fire in the past century (Safford and Van de Water 2014, Steel et al. 2015) (see fig. 13). The coarse-grained landscape structure that results from these fires is also very different from the fine-grained landscape structure that characterized presettlement forests. It is important to underscore that the forest “heterogeneity” referred to by North and colleagues (North et al. 2012a, 2012b; North 2012) is a fine-grained heterogeneity driven by the interactions between high frequencies of mostly low-severity fire and topographically driven variations in water availability.

**Future**—The few models that have been run not only suggest increased transition of forest to chaparral, but increased transition of chaparral to grassland as well, both trends being driven by increased fire activity (Lenihan et al. 2003a, 2003b, 2008). Cole (2010) studied paleoecological data from earlier periods of rapid climate warming in the Pleistocene and suggested that current and projected future warming trends could be expected to greatly increase the amount of early-seral vegetation on the landscape. McKenzie et al. (2004) noted that, given current and projected trends in climate and fire, the long-term persistence of late-seral forest in much of the Western United States was questionable. Based on projections as well as trends already in play in southern California, it seems likely that—especially at lower elevations—some proportion of the YPMC forest belt will transition to shrubland and grassland over the next century. It also seems likely that forest landscape structure will become gradually more coarse-grained as fire frequency and severity continue to increase, and fire suppression efforts continue to lead to forest densification in the rest of the landscape.
Natural Range of Variation for Yellow Pine and Mixed-Conifer Forests in National Forests of California

Forest Structure

General forest structure—

NRV—Early Euro-American visitors to assessment-area YPMC forests with interests in vegetation tended to focus on the large size of individual trees, and the generally open nature of the forest canopy. William Brewer was part of the California State Geological Survey of 1860–1864, and traveled across much of the state during that period. Brewer’s memoirs (Brewer 1930) contain much reference to vegetation conditions. In general, Brewer was impressed by the large size of the trees in the Sierra Nevada and he noted in multiple places how they dwarfed anything he had been familiar with in the Eastern United States. Where Brewer (1930) refers to forest densities in the conifer belt, he often describes “open” conditions, or conditions of “scattered” trees. For example, his description of the Crane Flat area in current-day Yosemite National Park is of “open forests of enormous trees.” Brewer also describes some areas of “dense” forest, however, and the general impression is one of a very heterogeneous forest landscape.

John Fremont’s memoirs of his 1843–1844 expedition to California (Fremont and Smucker 1856) refers often to the immense size of conifer trees his group encountered in the Sierra Nevada. He also makes mention of open groves of pines in a number of locations, but does not once refer to forests he encountered as being dense or closed. Indeed, as his group left the San Joaquin Valley southwest of Bakersfield on their return journey to Colorado, he remarks, as they reenter the pine belt, that “we found ourselves again traveling among the old orchard-like places.”

Miners’ journals occasionally contain interesting and useful information about forest structure in the mid-19th century. Peter Decker’s journals from 1849 to 1851 (Giffen 1966) refer periodically to forest conditions. He wrote both of thickets of trees and areas of open or regular spacing. J.G. Bruff’s diary entries similarly contain short descriptions of both open and dense forest along the path of his travels, although—unlike other early observers—he described the latter condition more often.

Clarence King worked for Brewer’s team on the geological survey, and his memoirs also contain much reference to vegetation conditions (King 1871). Referring to the Sierra-Cascade axis, which stretches from California to British Columbia, King noted the transition from more open and “grove-like” forests in California, where individual trees tended to be larger, to denser forests in Oregon and Washington, where pines ceased to be a major component. King noted that the transition was almost imperceptible from close range, but very clear at the broad scale. King (1871: 28–29) described the YPMC forest above Visalia thusly:
Passing from the glare of the open country into the dusky forest, one seems to enter a door, and ride into a vast covered hall... You are never tired of gazing down long vistas, where, in stately groups, stand tall shafts of pine... Here and there are wide open spaces around which the trees group themselves in majestic ranks.

John Muir (1894; chapter 8) wrote that nowhere in the Sierra Nevada would one find, either “on the rocky heights (or) down in the leafiest hollows,” anything approaching the dense forests found in the Amazon, or the Himalaya, the Black Forest, or the Douglas-fir forests of Oregon. In a classic passage reproduced innumerable times, he noted that:

The inviting openness of the Sierra woods is one of their most distinguishing characteristics. The trees of all the species stand more or less apart in groves, or in small, irregular groups, enabling one to find a way nearly everywhere, along sunny colonnades and through openings that have a smooth, park-like surface, strewn with brown needles and burs.

In general, both Leiberg (1902) and Sudworth (1900) (see also Stephens 2000, Stephens and Elliott-Fisk 1998) described highly heterogeneous forest structure in the Sierra Nevada. Both surveyors referred qualitatively and quantitatively to the large size (height and diameter) of adult trees. Referring to the “middle timber belt,” which occupied elevations between 2,000 and 6,000 ft (600 to 1800 m) and therefore corresponds approximately to the YPMC forest type, Sudworth (1900: 515) wrote: “As a rule the growth is continuous but rather open... there are, however, areas of considerable extent on broad benches where the forest is dense... The trees are usually of large dimensions.”

Leiberg (1902: 32), made similar statements about the YPMC forests of the northern Sierra Nevada:

In the eastern and trans-Sierran districts... the old-growth forests... are generally open on all slopes except the northern and on tracts with much seepage... In the central district, outside the canyon areas, the forest is of moderate density and is rarely what might be called open, except in stands of very old growth. Elsewhere large quantities of white fir and Douglas-fir with oak combine to form thickset stands. On the rocky slopes of canyons and in the great gorges of the rivers the forest is always very open and scattered.

Concerning yellow pine-dominated forests, Sudworth (1900) noted that they were rarely if ever dense, and single big trees or groups of three to six trees often stood far apart, forming a clumped but open stand structure. He observed that
younger forests (up to 60 years old), which would establish in the frequent open spaces in the forest, were often very dense, but that successional processes and fire would thin them greatly over time. Incense cedar was a regular associate of ponderosa pine, and was often also in such open stands, although it was also a common riparian tree.

Manson (1906) wrote that the frequent burning in YPMC forests (he expressed the opinion that the American Indians annually burned everything) suppressed seedlings, and as a result the forests were “mainly composed of old trees, many badly burned at the butt.” From his wide travels in the Sierra Nevada, he attested that lower and middle-elevation conifer forests of all types were “devoid of middle-aged and young trees… The light fires gave open forests through which one could readily see for great distances.”

Greeley (1907) observed that the characteristic structure of YPMC forest was composed of smaller, often even-aged groups, usually with a core of pine. Weaver (1943) gave a very similar description, noting that when considered together these different-aged groups scaled up to an overall uneven-aged forest. Greeley (1907) wrote that:

…in deep, moist soils… and on the north slopes, one sees bunch after bunch of six or eight mature sugar pines of nearly the same size. Where... seedlings and saplings occur at all they are usually in groups under broken cover or in narrow openings in the stand. On the same sites fir and cedar crowd the sugar pine closely and bunches or large patches of these species occur in among the groups of sugar pine. Yellow pine… seeks the drier and warmer sites… Here it is also commonly found in large even-aged groups, from open bunches of mature trees to dense thickets of saplings and seedlings.

In the tone of a forester, Greeley (1907) expanded on some of the unfortunate aspects of YPMC forests for silvicultural management. One of the chief unfavorable features was the “widespread over-maturity of the timber” (i.e., the general lack of regeneration and small trees), and the “high density” of dead, broken-topped, and fire-hollowed trees. He estimated that that 1 to 5 percent of the stems in sugar and yellow pine were dead, 10 to 25 percent were “decadent” (needed to be cut immediately to realize any timber value), 30 to 40 percent were “mature” (should be cut in 10 to 15 years), and only the remainder was composed of “thrifty” trees that were still vigorously growing and could be the basis for a 30- to 40-year cutting rotation. In fir and incense cedar, the percentage of dead and decadent trees was higher yet, and in many stands, 30 percent of the firs over 61 cm (24 in) d.b.h. were unmerchantable from decay.
According to Greeley (1907), other unfavorable features of the YPMC forest from the standpoint of a forester were the very open and “irregular” nature of the stand and the fundamental role of fire in reducing stand densities, and the very strong successional pressure of the shade-tolerant species, principally white fir, on YPMC forest composition. Regarding the former, Greeley decried the effects of intermittent fires that were common in YPMC forests, especially at lower elevations. These fires removed leaf litter and humus and killed young trees, while “simply scorching the butts of larger trees.” The whole effect reminded him of the pineries of the Southeastern United States. The consequent open canopy resulted in much wasted growing space in YPMC forests. Greeley (1907), like Sudworth (1900) and Leiberg (1902), also described how dense stands of chaparral would arise where fires burned the tree canopy, and how such areas would be lost to tree production without human intervention.

Areas of shrubs were an important component of YPMC forests, as fire stimulates germination of species of the most important shrub genera (especially *Arctostaphylos* and *Ceanothus*), and the open canopy meant that much light reached the forest floor. Even stands of sugar pine, which we tend to equate today with denser, more productive mixed-conifer stands, were often very open and with considerable underbrush (Knapp et al. 2013, Larsen and Woodbury 1916).

In summary, in comparison to today, early observers described YPMC forests that were generally more open, more heterogeneous, and more dominated by (clumps of) large trees (Agee 1993; Barbour et al. 1993, 2007; Brewer 1930; Greeley 1907; Jepson 1923; King 1871; Laudenslayer and Darr 1990; Leiberg 1902; Muir 1894; North et al. 2012b; Sudworth 1900; Sugihara et al. 2006; Vankat and Major 1978; Weaver 1943).

Comparison to current—The Sierra Nevada Ecosystem Project executive summary (Erman and SNEP Team 1996) included the following statement about post-settlement human impacts on assessment area forest structure:

**Forest Simplification.** The primary impact of 150 years of forestry on middle-elevation conifer forests has been to simplify structure (including large trees, snags, woody debris of large diameter, canopies of multiple heights and closures, and complex spatial mosaics of vegetation), and presumably function, of these forests. By reducing the structural complexity of forests, by homogenizing landscape mosaics of woody debris, snags, canopy layers, tree age and size diversity, and forest gaps, species diversity has also been reduced and simplified. At low elevations along the western boundary, ponderosa pine was preferentially removed, and throughout its range, sugar pine has decreased in abundance first through selection and
later by blister rust disease. Although the situation in the Sierra differs from that in forests in the Pacific Northwest, where fragmentation leaves remnant old-growth patches surrounded by large openings, functionally the Sierran forests have been fragmented to a lesser degree by simplification.

In the sections below, we refer more specifically to components of forest structure, including tree density, tree size and size-class distribution, tree basal area and volume, canopy cover, forest gaps and tree clumps, snags and coarse woody debris, and forest understory and nonforest vegetation.

**Tree density—**

**NRV and comparison to current**—In the assessment area, the average YPMC forest stand today is much denser than during presettlement times. Figure 14 shows comparisons between historical reconstructions of stand conditions in the late 19th and early 20th centuries with current conditions at the same sites. In all cases, modern densities are much higher than the earlier reconstructed densities; differences range from 80 to 600 percent (fig. 14). These differences may be slightly inflated by the inability of reconstruction studies to accurately account for very small trees in the historical period, especially from species that rapidly decay such as firs (Barth 2014).

![Figure 14](image-url)

Figure 14—Comparisons of current and historical tree densities at eight yellow pine–mixed-conifer sites in the assessment area. 1 = Taylor 2004 (≥10 cm diameter at breast height [d.b.h.]); 2 = North et al. (2007) (trees ≥5 cm d.b.h.); 3 = Lydersen et al. (2013) (≥10 cm d.b.h.); 4 = Scholl and Taylor (2010) (≥10 cm d.b.h.); 5 = USDA (1911) vs. current from Scholl and Taylor (2010) (≥15.2 cm d.b.h.); 6 = Collins et al. (2011) (≥15.2 cm d.b.h.); 7 and 8 = Parsons and Debenedetti (1979) (≥12 cm d.b.h.).
Numerous other empirical studies in the assessment area have also documented similar patterns to those portrayed in figure 14. These include Ansley and Battles (1998), Beaty and Taylor (2007, 2008), Taylor et al. (2014), Knapp et al. (2013), Dolanc et al. (2014a, 2014b), and Barth (2014). Dolanc et al. (2014a) compared the 1930s Forest Service vegetation type mapping (VTM) inventory of the central Sierra Nevada (Wieslander 1935) with the modern FIA inventory, and found that mixed-conifer forests had experienced increases over the 70-year period of about 69 percent, and ponderosa pine had seen density increase by 40 percent. The VTM inventory ignored trees below 10 cm (4 in) d.b.h., so Dolanc et al. (2014a) also removed small trees from the FIA data to allow comparison. Including the smaller trees, we hypothesize that the overall increase in density would probably be much greater, because the major response to climate and management trends through the period has been in the smallest size classes.

Stephens and Gill (2005) sampled 49 0.1-ha plots in the Sierra de San Pedro Mártir (SSPM), in Baja California and found an average of 145.3 trees/ha (±10.4 SE, range 30 to 320). Dunbar-Irwin and Safford (2016) measured an average of 188 trees/ha (±15.1 SE) in a more widespread sample of the SSPM, and Rivera and Safford (unpublished data) measured 216 trees/ha in a similar assessment of Jeffrey pine forests in the Sierra Juarez, another Baja California site. These values fall well within the reconstructed values in figure 14.

Baker (2014) used a complicated algorithm to suggest that GLO section-corner tree data (see “Tree size and size-class distribution” for more information) from four large areas in the Sierra Nevada showed mean densities of 293 trees/ha (±477 standard deviations [SD]; the median was 206). This is very much on the high end of prefire suppression estimates. Baker’s results run counter to early 20th century forest inventories that were conducted in the same areas. For example, Hagmann et al. (2014) summarized data from a 1922–1925 inventory in the eastern Oregon Cascade Range and found tree densities that were about four times lower than Baker’s estimates for the same area (in Baker 2012). The dataset Hagmann et al. (2014) analyzed had sampled >16,000 trees/mi² (6,265 trees/km²), whereas the GLO sampling densities are at most 8 trees/mi² (a 0.004 percent sampling effort based on Baker’s density estimates). Hagmann et al. (2013) also found that Baker’s (2012, 2014) methodology notably overestimated density in YPMC-type forests in south-central Oregon. Collins et al. (2011) resampled a 1911 inventory in YPMC forest near Yosemite National Park and found an average of <70 trees/ha (only trees >15 cm d.b.h. included) over

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a 4000-ha area. Baker’s (2014) GLO-based estimates for the same area suggested densities between 150 trees/ha and 450 trees/ha (>10 cm d.b.h.). Stephens et al. (2015) summarized a 1911 Forest Service inventory from the Greenhorn Mountains, west of Kernville, and found densities of 45 to 132 trees/ha (yellow pine and mixed-conifer stands [trees >15 cm d.b.h.]) in an area that Baker’s (2014) GLO algorithm calculated 210 to 548 trees/ha. Stephens et al. (2015) summarized discrepancies between direct inventories in Oregon and California and Baker’s estimates, and noted that (1) Baker’s sampling effort is multiple orders of magnitude lower than the forest inventory studies from the same areas, and (2) Baker’s methodology consistently overestimated forest densities by an average of more than 400 percent. Most recently, Levine et al. (2017) deconstructed Baker’s algorithm and found that his methodology overestimated true tree densities from an array of 6-ha mapped forest plots by 1.2 to 4.8 times.

The compiled FIA data from YPMC forest plots (USDA FS 2013) show an average modern density (all trees greater than 10 cm d.b.h.) of 396.5 trees/ha (±292 SD). This is 2.75 times more dense than the reference density obtained by averaging the eight studies in figure 14, Taylor (2004), Stephens and Gill (2005), Minnich et al. (1995), and Dunbar-Irwin and Safford (2016): 143.8 trees/ha (±80.9 SD). Recall that the FIA plots include logged and severely burned areas, so the difference between FIA and reference site studies is less than it would be if we restricted the comparison only to undisturbed FIA plots.

Further information on density by species is found in “Forest composition and species diversity” on page 161. Information on tree densities by size class (diameter) is in “Tree size and size-class distribution” below.

**Future**—Continuation of current trends will lead to further increases in forest density, primarily among small- and medium-sized trees. Trends in fire and in forest mortality (see below) may counteract these tendencies to some extent.

**Tree size and size-class distribution**—

**NRV and comparison to current**—Average and maximum tree sizes in YPMC forest stands appear to have been much larger in presettlement times. Taylor et al. (2014), working in Jeffrey pine-dominated forests in the Lake Tahoe basin, found the average tree size (d.b.h.) in the modern forest to be only about 60 percent of the average tree size in the forest in 1873. Lydersen et al. (2013) compared stand conditions on the Stanislaus National Forest in 1929 with conditions in 2008 and found that mean tree diameter had decreased by about 26 percent.

Miners like Peter Decker and Joseph Bruff referred often to the large trees they encountered. In one place on the present-day Lassen National Forest, Bruff described a stand of pines “ten feet diameter, 200 ft high, straight as arrows” (Read and Gaines 1949: 204). For ponderosa pine, Muir (1894) suggested that the average
size of “full-grown” trees on the western slope was over 65 m in height and from 1.5 to 1.8 m in diameter. Sudworth (1900) was somewhat less generous, estimating the averages to be 45 to 55 m in height and 0.9 to 1.2 m d.b.h. Sugar pine grew much larger; Muir (1894) listed a height of 67 m and 1.8 to 2.4 m d.b.h. as common measurements for full-grown trees, but Sudworth (1900) suggested that only the largest of all sugar pines reached those sizes. Table 6 lists the average sizes of “full-grown” adult trees provided by Sudworth (1900) for uncut stands in the central Sierra Nevada. These sizes are so large as to convince various investigators that Sudworth’s numbers can only refer to the best growing sites in his survey area (Bouldin 1999, Stephens 2000, Stephens and Elliott-Fisk 1998).

Early GLO land surveys also provide an idea of the sizes of trees on the late 19th century landscape. At section corners and halfway between section corners (1 mi apart), the original land survey teams took formal notes on “bearing” trees, which were the nearest trees >10 cm d.b.h. in each compass quadrant that were likely to survive over the long term. Trees between 25 to 36 cm d.b.h. (10 to 14 in) were apparently preferred if they were available (Hyde 2002). The survey teams identified the species, marked the tree, and measured the diameter, bearing, and distance from the section corner to each tree. At quarter-section corners along the section lines, two trees were marked and measured (Hawes 1882).

Hyde (2002) summarized GLO data from the Stanislaus, Sierra, and Sequoia National Forests. The average witness/bearing tree sampled by the GLO teams was <61 cm d.b.h., with the average sampled oak being between 38 and 50 cm d.b.h., the average sampled pine between 56 and 76 cm d.b.h., and the average sampled fir between 51 and 71 cm d.b.h. (values estimated from table 4.5 in Hyde 2002). On the Stanislaus National Forest, the average sampled pine was more than 78 cm d.b.h.; half of all pines sampled there were above 86 cm d.b.h., and more than one-third were

<table>
<thead>
<tr>
<th>Species</th>
<th>Height</th>
<th>D.b.h.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yellow pine</td>
<td>150–180</td>
<td>45–55</td>
</tr>
<tr>
<td>Sugar pine</td>
<td>180–200</td>
<td>55–61</td>
</tr>
<tr>
<td>Incense cedar</td>
<td>80–100</td>
<td>24–31</td>
</tr>
<tr>
<td>White fir</td>
<td>175–190</td>
<td>53–58</td>
</tr>
<tr>
<td>Jeffrey pine</td>
<td>125–160</td>
<td>38–49</td>
</tr>
<tr>
<td>Douglas-fir</td>
<td>150–175</td>
<td>46–53</td>
</tr>
</tbody>
</table>

Table 6—Mean “full-grown” tree sizes (height and diameter at breast height [d.b.h.]) for major tree species in uncut yellow pine and mixed-conifer forests in the central Sierra Nevada at the turn of 19th century, according to Sudworth (1900)
greater than 102 cm d.b.h.; fir on the Stanislaus were nearly as big on average, but the
median was between 61 and 76 cm d.b.h. Trees on the Sierra and Sequoia National
Forests were 15 to 30 percent smaller in diameter on average (Hyde 2002). General
Land Office data from the Lake Tahoe basin give an average tree diameter of 61 cm
in YPMC forests (Manley et al. 2000), and unpublished GLO data from the Eldorado
National Forest show average yellow pine and sugar pine diameters between 76 and
83 cm d.b.h. Baker’s (2014) GLO-based study of forest structure in the northern and
southern Sierra Nevada found that quadratic mean diameters (QMD) ranged from 50
to 59 cm (north and south), but Baker did not provide species-specific values.

Modern FIA data from the assessment area (USDA FS 2013a) for trees >10 cm
d.b.h. give an average diameter of approximately 26 cm and a QMD of 32 cm.

The drop in the average size of trees in YPMC forests over the past century
is the result of two trends: a great and general increase in the density of small
trees, combined with a decrease in the number of large trees. Many studies have
documented the former pattern (referenced throughout this assessment), but the
latter may be ecologically just as significant. Figure 15 is reproduced from Dolanc
et al. (2014b) and shows changes in density between the 1930s and early 2000s,
as documented by the Forest Service VTM and FIA inventories. The pattern of
increasing small tree density as well as the pattern of decreasing large tree density
are seen clearly in the figure. Although overall forest density is up at all elevations,
by far the greatest changes are in the YPMC belt (500 to 2000 m in figure 15),
where fire suppression has played a major role. Other studies documenting the loss
of larger trees in assessment-area YPMC forests include Dolph et al. (1995), Smith
et al. (2005), Ritchie et al. (2008), Lutz et al. (2009), and van Mantgem et al. (2009);
note that Collins et al. (2011) did not find this pattern. Although timber harvest
certainly explains some of this trend across the assessment area (e.g., Lydersen et al.
2013, Taylor 2004), the patterns also occur in unlogged forests. Other factors might
include insect outbreaks, pathogens, and drought stress, probably exacerbated by
the much higher stand densities that characterize modern YPMC forests.

Figure 16 shows reconstructed size class distributions combined from four study
sites in the assessment area. Figure 16 is given as the percentage of measured trees, to
better allow comparison of the shape of the reference distributions among sites. Figure
17 reproduces size class distributions from three modern reference sites that were not
logged and have not experienced total fire exclusion. In both figures, the roughly flat
or even hump-shaped distribution of tree sizes in the historical reconstructions can

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12 Fites-Kaufman, J. 2013. Personal communication. Regional planning ecologist, USDA,
Forest Service, Pacific Southwest Region, 1323 Club Drive, Vallejo, CA 94592.
Figure 15—Change in density (trees/ha) between vegetation type mapping (VTM) plots (from the 1930s) and Forest Inventory and Analysis (FIA) plots (from the early 2000s) for all species combined, for six elevation bands (x-axis), and by size class: (A) 10.2 to 30.4 cm diameter at breast height (d.b.h.); (B) 30.5 to 60.9 cm d.b.h.; (C) >61.0 cm d.b.h.; (D) percentage change in density from VTM to FIA plots across all size classes. Statistically significant differences are indicated by * = 0.01 < \( p \leq 0.05; ** = 0.001 < p \leq 0.01; \) and *** = \( p < 0.001. \) Yellow pine–mixed-conifer forests are found principally between 500 and 2000 m. Illustration adapted from Dolanc et al. (2014b).
Figure 16—Historical tree-size distributions from four studies of yellow pine–mixed-conifer forest in the assessment area, standardized by percentage of trees measured. (A) Sierra National Forest, 1865 (North et al. 2007); Lake Tahoe west shore, 1873 (Taylor et al. 2014). (B) Lake Tahoe east shore, 1870–1900 (Taylor 2004); Yosemite National Park, 1899 (Scholl and Taylor 2010).
be appreciated. This is very different from the classic “reverse J-shaped” age or size distribution that is typical of stable, self-replacing climax communities (Barbour et al. 1987, Oliver 2001). Assuming that size is roughly correlated with age, the $x$-axes of figures 16 through 19 represent the spectrum from juveniles through immature, mature, and then senescent individuals. Given the relatively high probability of mortality for juveniles, a high number of young trees is usually necessary to replace the relatively few adults that die in any given period. This leads to the J-shaped (or “hockey stick”) size distribution in the contemporary forest stands in figure 18. Such a distribution is typical of a forest community relatively free of disturbance (Smith et al. 2009). If a population is composed mostly of mature and senescent individuals, the population may be in decline (which is how early foresters interpreted the situation they encountered) (e.g., Greeley 1907, Sudworth 1900), or it may be one that is replaced only by episodically successful recruitment (Barbour et al. 1987). This was largely the case with presettlement YPMC forests in the assessment area, where recurrent fire killed most juvenile trees and successful recruitment was somewhat of a stochastic event, when seed production happened to coincide with a period of sufficient precipitation and little or no fire. Roughly flat or hump-shaped age or size distributions are characteristic of old-growth forests in areas that experience frequent, low-severity fire (Lydersen and North 2012, North et al. 2007, Oliver 2001). As a result, they are also found in stand reconstructions of presettlement ponderosa pine.
Figure 18—Comparisons of reconstructed historical tree size class distributions and contemporary distributions from three sites in the assessment area. (A) North et al. (2007); (B) Taylor (2004); (C) Scholl and Taylor (2010).
forest in the U.S. Southwest (Mast et al. 1999), and in forests of longleaf pine (*Pinus palustris* Mill.) in the Southeastern United States (Heyward 1939). The distribution of size classes in the GLO data from the Lake Tahoe basin show a similar humped shape (Manley et al. 2000). Oliver (2001: fig. 2) graphed the size-class distribution from the Beaver Creek Pinery, a well-known reference ponderosa pine-black oak site in the northern assessment area that was not logged and experienced a number of fires in the 20th century; it is strongly hump-shaped. Using a project at Blacks Mountain Experimental Forest in the same general area, Oliver (2001) showed that prescribed fire could convert a reverse J-shaped distribution to a hump-shaped distribution. Taylor (2010) showed the same effect of fire in his study of the Beaver Creek Pinery and the nearby Devil’s Pinery.

Not all reference YPMC forests show the hump-shaped distribution of tree size classes. In their array of plots in the contemporary Sierra de San Pedro Mártir, Stephens and Gill (2005) found a J-shaped distribution of size classes.

Figure 19 shows the average size-class distribution in 20-cm classes for the FIA plot data compilation (USDA FS 2013a). The obvious reverse J-shaped curve is very different from the more flat or hump-shaped size distribution found in presettlement forests.

**Future**—Continued high levels of recruitment among shade-tolerant species, combined with increasing mortality among larger trees (see “Tree mortality” above), will likely intensify the tendency toward a reverse J-shaped size-class distribution.
Tree basal area/volume (forest biomass)—

NRV—A number of stand reconstruction studies report late 19th century basal areas. Scholl and Taylor (2010) found that overall reconstructed basal area in 1899 was between 29 and 30 m²/ha, depending on the smallest tree size considered. A U.S. Forest Service (USDA FS 1911) report referenced by Scholl and Taylor (2010) found basal areas in western Yosemite Park of about 21 m²/ha. Taylor et al. (2014) reported an overall basal area of about 27 m²/ha in YPMC forests in the Lake Tahoe basin (reconstructed date = 1893). Taylor (2004) reconstructed Jeffrey pine stands on the east shore of Lake Tahoe and found average basal areas of 25.5 m²/ha for the period between 1870 and 1900. Parsons and DeBenedetti (1979) provided data that produced basal areas (in 1875) of 38 to 89 m²/ha in yellow pine and productive mixed-conifer forests in the southern Sierra Nevada. North et al. (2007) reported an 1865 basal area of about 51 m²/ha for a moist mixed-conifer forest on the Sierra National Forest, and Knapp et al. (2013) found a mean basal area of 53.9 m²/ha for a productive site on the Stanislaus National Forest. Using 1911 forest inventory data, Collins et al. (2011) reported basal areas of about 14 to 18 m²/ha from the east side of Yosemite National Park. Collins et al. (2015) reported basal areas of 11 to 30 m²/ha for the west side of Yosemite National Park, and Stephens et al. (2015) reported basal areas of 11 to 30 m²/ha for the Greenhorn Mountains. Baker’s (2014) GLO-based estimates were between 32.5 and 35.5 m²/ha (north and south Sierra Nevada sites, respectively).

Outside of the assessment area but in similar forest, Avery et al. (1976) used long-term stand records in ponderosa pine forest in Arizona to determine that basal area in 1920, just as federal fire suppression was gaining traction, averaged about 14.2 m²/ha across 16 forest inventory plots.

Data from contemporary reference sites give similar numbers. Lydersen and North (2012) found that modern old-growth stands with recent recurrent fire averaged 54 m²/ha in a sample of 48 sites across much of the assessment area. Stephens and Gill (2005) measured stand characteristics in a limited area of Jeffrey pine/mixed-conifer forest in the Sierra de San Pedro Mártir in Baja California and found that basal area averaged 19.9 m²/ha, but with an enormous range (5.7 to 50.7 m²/ha). Dunbar-Irwin and Safford (2016) found an average basal area of 22.5 m²/ha for Jeffrey pine and mixed-conifer stands across the plateau of the Sierra de San Pedro Mártir, and in an unpublished study they measured an average of 15.5 m²/ha for Jeffrey pine stands in the Sierra Juarez,13 which is at the lowest altitudinal limit of Jeffrey pine in Baja California. Taylor (2010) measured basal areas of 27.1 m²/ha

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in the Beaver Creek Pinery on the Lassen National Forest, which is an old-growth ponderosa pine-black oak forest with multiple 20th century fire entries.

Outlying values are given by Sudworth (1900, and in his unpublished notes), who sampled plots in highly productive sites in the central and southern Sierra Nevada. Mean basal areas from plots in mixed-conifer stands ranged from 130 to 270 $m^2$/ha. The representativeness of Sudworth’s (1900) plot data has been questioned (Baker 2014, Bouldin 1999, Stephens 2000, Stephens and Elliott-Fisk 1998).

**Comparison to current**—A compilation of the most recent FIA data in the assessment area (USDA FS 2013a) suggests that mean basal area in YPMC forests has not changed dramatically over the past 100+ years. The average of the reference values given above (not including Sudworth and Avery) is 31.7 $m^2$/ha ($\pm$18.5 SD), while the average of the 2,508 FIA plots in YPMC forests included in the compilation is 32.9 $m^2$/ha ($\pm$20.4 SD; all trees $>5$ cm d.b.h.). Mean basal area in YPMC FIA plots on the west side of the assessment area (north, central, and south subregions) is 34.6 $m^2$/ha; on the east side (northeast and southeast subregions), it is 26.5 $m^2$/ha.

Obviously, it is statistically somewhat tenuous to compare the results of 14 local studies with an average derived from 2,508 plots regularly spaced across the entire assessment area, especially when the latter comprise all stand conditions, including logged and severely burned areas. Studies that have directly reconstructed early stand conditions on a site and then compared them to the current forest stand have mostly found that basal area has increased, although not at the rate that tree density has increased (see above). Figure 20 summarizes temporal changes in basal area by species for four sites in assessment-area YPMC forests; the increase in basal area is nearly entirely in shade-tolerant/fire-intolerant species. Overall increases in basal area from these studies and others referenced above range from 10 to 150 percent, averaging about 60 percent across studies.

Ansley and Battles (1998) compared modern YPMC forest structure to structure in a stand first sampled 40 years before. They found that basal area rose by 15 percent over the 40 years, and it increased for all species except for sugar pine (which remained the same) and ponderosa pine (which dropped). Lydersen et al. (2013) compared forest stand structure conditions in 1889 (estimated through reconstruction), 1929, and 2008 on the Stanislaus-Tuolumne Experimental Forest and found that basal area had increased from 52.8 to 56.6 to 70.5 $m^2$/ha, respectively. Increases in basal area were primarily in shade-tolerant species; the percentage of pine in the sampled stands dropped from 25 to 11 percent over the same period. Ritchie et al. (2008) also found that the contribution of pine to stand basal area dropped by about 12 percent over the 20th century in unlogged stands at Blacks Mountain on the Lassen National Forest.
Because presettlement YPMC forests in the assessment area generally supported more large trees than current forests, a rise in basal area over time would be driven mostly by increases in small and medium tree densities. In places where those young tree densities have not risen as rapidly, or where large tree densities have not dropped (e.g., unlogged areas or areas where large tree mortality has not been otherwise high), basal areas of current forest may be similar to or even lower than presettlement forests (North et al. 2007).

Interestingly, comparisons between current FIA data and the Forest Service VTM dataset from the 1930s have mostly found that basal area and volume have decreased over time (e.g., Bouldin 1999, Fellows and Goulden 2008, McIntyre et al. 2015). It is difficult to determine why this is. There is some concern that VTM plots may have been preferentially sited in areas of better growing conditions or larger
trees (Bouldin 2009), whereas the FIA data are a statistically unbiased sample. Little note is made in the VTM plot notes of previous logging, so it may be that they tended to avoid previously harvested stands. This would perhaps provide a better idea of natural forest conditions in the early 20th century, but it would complicate comparisons with modern inventories, which include all stand conditions to permit a statistically robust estimate of landscape-level conditions. Another issue with the VTM plots is that they did not sample any trees <10 cm d.b.h. The general understanding is that presettlement YPMC forests generally supported very low densities of saplings and small trees because of frequent fire. The VTM plots were sampled a few decades after the institution of fire suppression, so it seems likely that the absence of the smallest tree sizes in the VTM data does not constitute a major omission when calculating biomass. However, modern YPMC forests are highly enriched in the smallest size classes, and high densities of <10 cm d.b.h. trees can contribute measurable amounts of biomass to a forest stand. In the modern FIA data from assessment-area YPMC forests (USDA FS 2013a), the size class between 5 and 10 cm d.b.h. contributes an average of about 0.7 m²/ha in basal area. Combining this with a rough estimate of the biomass in the 0 to 5 cm d.b.h. class (seedlings and small saplings), which is not measured in size by FIA but only enumerated, omission of trees <10 cm d.b.h. can reduce measured forest biomass from 1 to 2 m²/ha or more. As a result, removing trees <10 cm d.b.h. from FIA plots to permit an unbiased comparison with VTM (a standard practice in FIA to VTM comparisons) may result in modern plots appearing to support less biomass than the early plots when they actually do not.

In summary, FIA data and local forest reconstruction studies suggest that modern basal areas are broadly similar to or slightly higher than basal areas in presettlement times. Local trends will depend on the relative effects of fire exclusion (which would increase basal area on its own) versus logging and other forest mortality agents (which decrease basal area). Remember that most (but not all) reconstruction studies are done in areas that did not experience logging, whereas the FIA dataset samples the forest without bias.

Future—Miller and Urban (1999a) simulated the effects of climate change on forest biomass and composition along an elevational gradient in Sequoia & Kings Canyon National Parks (see “Forest species composition and species diversity” on page 161 for more detail). At their two lowest elevation sites, which fall in YPMC forests, woody biomass was completely or almost completely lost by the end of the simulations. Under their less extreme warming and drying scenarios, basal area at 1800 m declined from 29 to 4 m²/ha, and was completely lost under the more extreme scenarios. At 2200 m, basal area declined from 52 m²/ha to 25–32 m²/ha under the less extreme future scenarios, and to 8 m²/ha under the most extreme scenario. Miller and Urban
Plates

The following 24 photographs provide an overview of the variety of stand conditions (except recently logged stands) found in yellow pine and mixed-conifer forests in the assessment area and a reference site in northwest Mexico. The approximate locations, forest types, and management agencies are identified, along with brief notes on site histories and conditions.

Plate title: Open mixed-conifer stand.

Plate Notes: Near the Wawona Hotel (Big Trees Lodge), Wawona, Yosemite National Park, Mariposa County, 1310 m (4300 ft). This stand is dominated by ponderosa pine, with sugar pine, incense cedar, and black oak. The understory is dominated by bear clover (Chamaebatia foliolosa), which is a common ground cover in open stands characterized by frequent fire. This area has experienced up to three prescribed fires over the last 35 years. Some ponderosa pine in the stand were being killed by western pine beetle when the photo was taken in 2015.
Plate title: Large patch of high-severity fire in moist mixed-conifer forest

Plate Notes: The Rich Fire (2008) burned across this south-facing slope above North Fork Feather River, Plumas National Forest, Plumas County. Elevation in the photo ranges from 1130 to 1800 m (3700 ft to 5900 ft). Prefire species composition was moist mixed conifer, with Douglas-fir, white fir, incense cedar, ponderosa pine, sugar pine, Jeffrey pine, black oak, and canyon live oak. This is an example of the very large patches of high-severity fire that are becoming common in yellow pine and mixed-conifer forests in the Sierra Nevada. The distance from the photo point to the top of the peak is about 2.7 km, the patch of nearly complete mortality is >600 ha. Before the Rich Fire, there was no fire here in at least 100 years, and much of the area was heavily logged in the late 19th and early 20th centuries.
Plate title: Smaller patch of high-severity fire in moist mixed-conifer forest

Plate Notes: The Meadows Fire (2004) burned within the Illilouette Creek Basin, Yosemite National Park, Mariposa County, 1980 m (6,500 ft). The predominant species are white fir and Jeffrey pine. This is an example of one of the larger high-severity patches seen in this wildland-fire-use area. Distance across patch is 140 m, patch size is about 2 ha. This site also burned previously in 1981, after at least 80 years of fire suppression.
Plate title: Large legacy trees

Plate Notes: Rubicon River drainage, Eldorado National Forest, Eldorado County, 1200 m (3,950 ft). This is one of the few remnant stands of older trees in the Rubicon River drainage; very light levels of cutting occurred here. Most of the forest in this area was heavily logged at least once since the mid-19th century. This stand is dominated by ponderosa pine, sugar pine, white fir, incense cedar, Douglas-fir and black oak, with buck brush (Ceanothus cuneatus) in the understory. Trees in the foreground are 80 to 140 cm diameter at breast height (32 to 55 in). No fire has burned here in at least 100 years, but stand conditions are still somewhat open owing to rocky, excessively well-drained soils. This stand escaped the very severe King Fire that burned much of this drainage in 2014.
Plate title: Jeffrey pine stand following fuel treatment

Plate Notes: Lake Tahoe Basin, just outside of Angora Fire perimeter, Eldorado County, 2010 m (6,600 ft). Stand dominated by Jeffrey pine, with some white fir, red fir, and lodgepole pine. This site on the left (panel A) experienced three forest fuel treatments between 1996 and 2005, beginning with a mechanical thin, followed by a hand thin and then a pile burn/prescribed fire. The shrub resprouts (greenleaf manzanita) are from 3 to 5 years old in the photo, and are a common understory response to opening of the canopy. Nearby stands with the same treatment history that burned in the Angora Fire suffered only 5 to 15 percent overstory mortality. The site on the right (panel B) is about 500 m SW of panel A, and was not treated before the date of the fire. The site is characterized by large Jeffrey pine and red fir in the overstory, with white fir and lodgepole pine dominating the smaller size classes. Nearby stands with the same forest structure that burned in the Angora Fire suffered 80 to 100 percent overstory mortality.
Plate title: Standard mixed-conifer stand, Sierra Nevada west slope

Plate Notes: Near Camp Nelson, Greenhorn Mountains, Sequoia National Forest, Tulare County, 1770 m (5,800 ft). This area was selectively logged more than once in the 19th and 20th centuries. The stand is dominated by ponderosa pine, incense cedar, white fir, sugar pine and black oak. There was some western pine beetle damage in the stand when the photo was taken in 2015; by 2016 there was widespread pine mortality. No fire has burned here in at least 100 years.
Plate title: Dry yellow pine-black oak forest

Plate Notes: Beaver Creek Pinery, Ishi Wilderness, Lassen National Forest, Tehama County, 870 m (2,850 ft), a well-known natural range of variation reference site. This low-elevation forest is dominated almost completely by ponderosa pine and black oak. The open canopy and low surface fuels of this site and other nearby unlogged “pineries” contrast strongly with nearby lands that were heavily logged and fire suppressed, which are now dominated by dense stands of shade-tolerant species (especially Douglas-fir). This site burned in 1903, 1924, 1990, and 1994. Recent fires in the area were suppressed. The current 22-year gap in the fire record has led to some fuels accumulations, and there is concern that the site could be lost to wildfire if prescribed burns are not instituted.
Plate title: Gap regeneration following fire

Plate Notes: Illilouette Creek Basin, Yosemite National Park, Mariposa County, 2200 m (7,200 ft). This stand is near the upper limits of yellow pine and mixed-conifer forests on the west slope of the Sierra Nevada. The stand is dominated by white fir and red fir with some Jeffrey pine, as well as lodgepole pine in wet undrained areas. This is a good example of the regeneration that occurs in forest gaps, in this case caused by a small lightning fire about 20 years previous to the photo.
Plate title: Sparse Jeffrey pine.

Plate Notes: Owens River Valley, Inyo National Forest, Mono County, 2410 m (7,900 ft). Pure Jeffrey pine stands occur at the eastern edge of its distribution in the Sierra Nevada rain shadow. Soils here are on volcanic rocks. Selective logging occurred in this area in the late 19th and early 20th centuries. When the photo was taken, no fire had burned here in at least 100 years. Then two fires burned the area in the summer of 2016. The photo foreground experienced very high tree mortality in a wind-driven fire after three years of drought.
Plate title: Mixed-conifer stand on a serpentine soil boundary

Plate Notes: Along Concow Road, Plumas National Forest, Butte County, 850 m (2,800 ft). A geological contact between ultramafic rocks (peridotite) and metasediments runs vertically through the middle of the photo. Site history is identical across the contact. The soils that develop on ultramafic rocks (often called “serpentine” soils) are usually critically low in macronutrients like nitrogen, phosphorus, and potassium, and enriched in magnesium, iron, and a number of heavy metals. As a result biomass accumulation is slow, woody plants are often stunted and much older than they appear, and there is high plant species endemism. Both soils support yellow pine and mixed-conifer forest, with the serpentine forest on the left dominated by gray pine and a thin and stunted complement of species from the more fertile soil to the right (Douglas-fir, sugar pine, incense cedar, ponderosa pine). Hardwoods are uncommon on serpentine soils (except where they have been hydrothermally altered). Photo taken in 2005 after more than 100 years without fire; the entire area of the photo burned in the 2008 BTU Lightning Complex Fire. Because of fuel differences, the dense forest to the right burned with nearly complete overstory mortality, the serpentine forest suffered about 30 percent mortality.
Plate title: Dense understory, mesic mixed conifer stand.

Plate Notes: Private property, North San Juan, Nevada County. 640 m (2,100 ft). The understory “jungle” that is common in mesic stands on unmanaged/unburned lands at low elevations on the Sierra Nevada west slope. This stand is dominated by ponderosa pine, incense cedar, Douglas-fir, black oak, interior live oak, and madrone (abundant in the understory), with some invasion by Spanish broom (*Spartium junceum*). This area was heavily logged in the mid to late 19th century and has not experienced a fire in at least 100 years.
Plate title: Low-elevation ponderosa pine-oak woodland

Plate Notes: Near Priest Reservoir, Tuolumne County, 760 m (2,500 ft). At very low elevations, ponderosa pine occurs as scattered individuals or small groves within landscapes dominated by oak species (interior live oak, blue oak, canyon live oak), other hardwoods, and gray pine. Chaparral stands are also common. The ponderosa pine in this photo had been attacked by western pine beetle that year (2015) and are now dead. This area was logged in the late 19th century.
Plate title: Open mixed-conifer forest.

Plate Notes: Illilouette Creek Basin, Yosemite National Park, Mariposa County, 2070 m (6,800 ft). This mixed-conifer stand is dominated by Jeffrey pine and white fir. The stand structure is an example of open forest maintained by frequent surface fire. This stand burned at low severity in 1981 and 2004.
Plate title: Open dry mixed-conifer forest

Plate Notes: Sierra de San Pedro Mártir National Park, Baja California, 2600 m (8,540 ft). This high-elevation stand is dominated by Jeffrey pine, with white fir, lodgepole pine, and sugar pine (white fir and sugar pine mostly occur on north slopes). Aspen is found along the drainage and also on north slopes. A few oak species, including canyon live oak, are also found in this stand. This site last burned in 1948, a previous fire occurred in 1929. There is also moderate grazing pressure here. Most of the national park, including this site, has never been logged. Since the 1980s, the Mexican Park Service has been aggressively suppressing fires in the park, and recent science is showing that fuel loads are increasing as a result.
Plate title: Sugar pine stand

Plate Notes: D.L. Bliss State Park, Lake Tahoe Basin, Eldorado County, 2070 m (6,800 ft). Sugar pine often dominates areas such as this with thin soils and rocky outcrops but decent precipitation. There are also white fir and some Jeffrey pine in this stand. White pine blister rust is killing many of the trees in the stand. This stand was one of few to escape the clearcut logging that pervaded the Lake Tahoe Basin in the late 19th century. No fire has burned here in at least 100 years.
Plate title: Fire suppressed mixed conifer forest

Plate Notes: Plumas National Forest, Butte County, 1220 m (4,000 ft). Standard moist mixed-conifer stand after late 19th century logging and a century of fire suppression. Stand is dominated by Douglas-fir, with individuals of ponderosa pine, sugar pine, black oak, canyon live oak, incense cedar, and madrone also present. This is in the area of the highest conifer growth rates in the assessment area. No fire has burned here in at least 100 years.
Plate title: Regeneration in the absence of fire

Plate Notes: Buck Meadows, west of Yosemite National Park, Stanislaus National Forest, Mariposa County, 915 m (3,000 ft). Regeneration of ponderosa pine, incense cedar, and black oak is occurring in a gap left by fallen beetle-killed pine. No fire has burned here in at least 100 years; however, the area was logged in the early 20th century.
Plate title: High-elevation Jeffrey pine

Plate Notes: Big Pine Creek, Inyo National Forest, Inyo County, 2560 m (8,400 ft). Yellow pine stands follow drainages in these high-elevation, rain shadowed sites on the east side of the Sierra Nevada. The stand is dominated by Jeffrey pine, with a minor component of lodgepole pine and aspen. This is near the upper elevation of Jeffrey pine at this latitude, with lodgepole pine the dominant species at higher elevation. No fire has burned here in at least 100 years.
Plate title: Fire-suppressed moist mixed-conifer/white fir

Plate Notes: Illilouette Creek drainage, Yosemite National Park, Mariposa County, 2100 m (6,900 ft). Moist mixed-conifer stands such as this are typical in bottomlands and cold air drainages at higher elevations. This stand is almost entirely white fir and lodgepole pine, with a few scattered Jeffrey pine. None of the Illilouette Creek Basin was logged. No fire has burned here in at least 100 years.
Plate title: Invaded low-elevation ponderosa pine-oak woodland

Plate Notes: City property, Nevada City, Nevada County, 820 m (2,700 ft). The overstory in this stand is dominated by ponderosa pine with some Douglas-fir and black oak; the understory is heavily invaded by Spanish broom (*Spartium junceum*), a Mediterranean legume originally introduced as an ornamental. The species is actively invading yellow pine and mixed-conifer forests at lower elevations near seed sources and areas of disturbance, especially along roads and near urban areas. There are many other noxious weeds in this stand as well, including Himalayan blackberry (*Rubus armeniacus*), and a variety of Mediterranean grasses. This area was heavily logged in the mid to late 19th century. No fire has burned here in at least 100 years.
Plate title: Open Jeffrey pine forest

Plate Notes: Golden Trout Wilderness, Inyo National Forest, 2440 m (8,000 ft), Tulare County. Jeffrey pine dominates this high-elevation site, which last burned in a surface fire 5 years before the photo was taken. This area is subject to wildland fire use. The site is also within an active grazing allotment, with low levels of livestock use.
Plate title: Moist mixed-conifer stand

Plate Notes: Station Creek Research Natural Area, Eldorado National Forest, Eldorado County, 1650 m (5,400 ft). This moist mixed-conifer stand is dominated by white fir and sugar pine, with some Douglas-fir, ponderosa pine, and black oak. Soils are thin, over steep granitic bedrock, keeping forest densities somewhat lower than in nearby drainage bottoms. There was some logging here in the late 19th century. No fire has burned here in at least 100 years.
Plate title: Large legacy ponderosa pine

Plate Notes: Emerald Bay State Park, Lake Tahoe, Eldorado County, 1905 m (6,250 ft). This property is one of few uncut old-growth stands at lake level; the dominant trees are ponderosa pine, which is rare in the Lake Tahoe Basin. Other conifers include (in order of decreasing basal area) Jeffrey pine, incense cedar, white fir, and sugar pine. Ponderosa pine in the foreground are 100 to 150 cm diameter at breast height (d.b.h.) (40 to 60 inches), some ponderosa pine in this stand reach 220 cm d.b.h. (87 inches). No fire has burned here in at least 100 years.
Plate title: Low-density Jeffrey pine stand on serpentine soils

Plate Notes: North-facing slopes south of North Fork Feather River, near Red Hill, Plumas National Forest, Plumas County, 1250 m (4,100 ft). Ultramafic “serpentine” soils, on peridotite. Site is dominated by Jeffrey pine, with some incense cedar, Douglas-fir, and sugar pine. Although Jeffrey pine is normally a higher elevation species in the assessment area, it is a better stress tolerator than ponderosa pine, and it supplants ponderosa pine on serpentine soils as low as 1100 m. At the time of the photo, this site had not experienced a fire in over 100 years, but it burned in the 2008 Rich Fire. Windy conditions led to surprisingly high mortality, up to 50 percent in some stands in this area.
(1999a) included fire in their simulations, but only surface fires were modeled, so fire played little role in directly causing biomass loss. Miller and Urban’s (1999a) results are thus probably conservative with respect to the velocity of change in biomass.

Bachelet et al. (2001) modeled the potential effects of climate warming on carbon budget using climate data from seven different GCMs. Their results varied widely, depending on the GCM used, the vegetation/carbon model used, the degree of future warming, and precipitation. Forested areas were especially sensitive to changes in temperature. Bachelet et al. (2001) found that most scenarios predicted increases in forest cover and forest biomass in the assessment area under moderate warming (+2 to 4 °C); their vegetation model, which included fire, resulted in less biomass gain. Their projections included a relatively large increase in precipitation, however; according to Dettinger (2005), the most common prediction among more recent GCMs (which are considerably more complex and realistic than the models available in the late 1990s) is temperature warming by about 5 °C by 2100, with precipitation remaining similar or slightly reduced compared to today. The newer GCMs thus cast some doubt on Bachelet et al.’s (2001) results. Under more extreme warming scenarios (>4 °C), Bachelet et al. (2001) found that increases in forest biomass were reversed. They referred to this temperature-driven pattern as the “early green-up, later browning” hypothesis. Overall, Bachelet et al. (2001) suggested that a temperature increase above a threshold of about 4.5 °C would result in more carbon loss than sequestration across the United States. In a subsequent modeling effort, Bachelet et al. (2007) projected that biomass consumption by wildfire would increase by 25 to 67 percent across most of the assessment area over current (1961 to 1990) rates.

Lenihan et al. (2008) modeled vegetation distribution and productivity in California under three future climate change scenarios. Under the most moderate warming scenario, which also featured similar precipitation to today, the extent of tree-dominant vegetation in the assessment area increased substantially (although some losses of conifer forest occurred), and California as a whole was modeled to be a significant carbon sink by the end of the 21st century. Under the two drier and warmer scenarios, significant loss of conifer forest occurred (much of it to fire), and broadleaf expansion was not sufficient to offset a large net loss of ecosystem carbon by 2099.

Loudermilk et al. (2013) modeled carbon sequestration in the Lake Tahoe basin under two future climate scenarios that included interactions with fire. Because the basin was heavily logged at the turn of the 19th and 20th centuries, their modeling suggested that forests in the basin would remain a carbon sink until the end of the 21st century under all modeled scenarios. Including the effects of fire, however (constrained to burn in similar fashion to current fires and under continued fire suppression), under the more extreme climate scenario, resulted in 30 to 40 percent less C storage.
Canopy cover—
Canopy cover is a measure of the percentage of the ground surface that is shaded from directly above by trees. A tree’s “canopy” is a polygon whose perimeter coincides with the outer edge of the tree’s foliage. The canopy is assumed to be whole, and gaps within the tree canopy perimeter are not subtracted. In this section, we refer to “relative” canopy cover, where overlapping tree canopies are not double-counted (or triple-counted, etc.), so the maximum canopy cover in a plot is 100 percent. Canopy cover is usually an overestimate of the amount of shade on the underlying ground because it ignores intercanopy gaps. “Canopy closure” is better correlated with actual light availability, but it must be measured on a plot-by-plot basis and cannot be easily modeled or estimated from plot tree data (see North and Stine 2012).

NRV—As indicated in the “General forest structure” section, observations of forest cover in the 19th and early 20th centuries are overwhelmingly characterized by remarks about the general openness of the canopy of YPMC forests in and near the assessment area, but it is clear from descriptions that conditions differed across the landscape (e.g., Brewer 1930, Greeley 1907, King 1871, Larsen and Woodbury 1916, Leiberg 1902, Muir 1894, Sudworth 1900; also see Pearson [1923]) for descriptions of yellow pine forests in Arizona). Recently, a number of researchers have undertaken plot-based reconstructions of stand structural conditions in the late 19th century, and data are also now available from a number of contemporary reference sites where logging did not occur and—in many cases—where fire was not completely suppressed during the 20th century. From these studies, we can now obtain a more quantitative idea of the levels of canopy cover supported by YPMC forest stands before timber harvest and fire suppression altered much of the landscape.

Historical datasets and reconstructions of presettlement or prefire suppression conditions do not generally report canopy cover. However, most of these datasets provide data on size class distributions of trees in the sampled plots. Where such data were available, we used two methods to generate canopy cover estimates. First, we used species-specific equations for crown-width (Keyser 2010: table 4.4.2.1) from the FVS (Dixon 2002) to estimate crown area, using the midpoint of the size class in our calculations, then multiplying by the number of trees in that size class. Percentage of cover was obtained by summing across the size classes and standardizing to square meters, then dividing the result by 10 000 (number of square meters in a hectare) and multiplying by 100 for the percentage. Second, cover was adjusted by applying a formula that assumes the random spatial location of the trees in the sampled area and subtracts overlapping canopies from the maximum cover value (Crookston and Stage 1999). In table 7, we report both the adjusted value and the
Table 7—Percentage of tree cover in historical and contemporary reference yellow pine and mixed-conifer (YPMC) stands in the assessment area and comparable reference sites, arranged from north to south

<table>
<thead>
<tr>
<th>Site</th>
<th>Mean</th>
<th>Standard error</th>
<th>Method</th>
<th>Observations</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>North and central assessment area</td>
<td>48.5 (79.9) a</td>
<td>1.2</td>
<td>FVS modeled estimate^b</td>
<td>1930s: Forest Service vegetation mapping inventory, YPMC</td>
<td>Dolanc (2014b)</td>
</tr>
<tr>
<td>Central assessment area</td>
<td>69.1 (89.1) a</td>
<td>4.1</td>
<td>FVS modeled estimate^b</td>
<td>1899: timber inventory, mixed conifer</td>
<td>Sudworth (1900)</td>
</tr>
<tr>
<td>Lake Tahoe Basin</td>
<td>16.6 (18.1) a</td>
<td>4.1</td>
<td>FVS modeled estimate^b</td>
<td>1870—1900: stump counts, Jeffrey pine forest, east shore</td>
<td>Taylor (2004)</td>
</tr>
<tr>
<td>Lake Tahoe Basin</td>
<td>35.7 (44.1) a</td>
<td></td>
<td>FVS modeled estimate^b</td>
<td>1893: plot-based reconstruction, mixed conifer, west shore</td>
<td>Taylor et al. (2014)</td>
</tr>
<tr>
<td>Lake Tahoe Basin</td>
<td>45.3</td>
<td></td>
<td>Line intercept</td>
<td>Modern: uncut old-growth, YPMC, white fir, red fir</td>
<td>Barbour et al. (2002)</td>
</tr>
<tr>
<td>Stanislaus National Forest</td>
<td>36/45^c</td>
<td></td>
<td>Tree canopy area calculations</td>
<td>1889/1929: plot-based reconstruction, mixed conifer</td>
<td>Lydersen et al. (2013)</td>
</tr>
<tr>
<td>Yosemite National Park</td>
<td>22</td>
<td></td>
<td>FVS modeled estimate</td>
<td>1911: timber inventory, mixed conifer</td>
<td>Collins et al. (2011)</td>
</tr>
<tr>
<td>Yosemite National Park</td>
<td>41 (52.7) a</td>
<td></td>
<td>FVS modeled estimate^b</td>
<td>1899: plot-based reconstruction, moist mixed conifer</td>
<td>Scholl and Taylor (2010)</td>
</tr>
<tr>
<td>Sierra National Forest</td>
<td>33.9 (41.4) a</td>
<td></td>
<td>FVS modeled estimate^b</td>
<td>1865: plot-based reconstruction, moist mixed conifer, Teakettle Experimental Forest</td>
<td>North et al. (2007)</td>
</tr>
<tr>
<td>Sequoia National Forest</td>
<td>12.1</td>
<td>5.1</td>
<td>FVS modeled estimate</td>
<td>1911: timber inventory, ponderosa pine</td>
<td>Stephens et al. (2015)</td>
</tr>
<tr>
<td>Sequoia National Forest</td>
<td>25.2</td>
<td>7.1</td>
<td>FVS modeled estimate</td>
<td>1911: timber inventory, mixed conifer</td>
<td>Stephens et al. (2015)</td>
</tr>
<tr>
<td>Assessment area (mostly southern and central)</td>
<td>43.7</td>
<td>1.5</td>
<td>Densiometer</td>
<td>Modern: YPMC forest, uncut, with frequent or recent non-stand-replacing fire</td>
<td>Lydersen and North (2012)</td>
</tr>
<tr>
<td>Sierra de San Pedro Mártir</td>
<td>24.9</td>
<td>2.4</td>
<td>Ocular estimate</td>
<td>Modern: Jeffrey pine, mixed-conifer forests, Baja California</td>
<td>Dunbar-Irwin and Safford (2016)</td>
</tr>
<tr>
<td>Sierra Juarez</td>
<td>35.8</td>
<td>7.6</td>
<td>Ocular estimate</td>
<td>Modern: Jeffrey pine, Baja California</td>
<td>Safford, in preparation</td>
</tr>
</tbody>
</table>

^a The first modeled canopy cover value reduces crown cover by removing canopy overlap under the assumption of random distribution of stems (Crookston and Stage 1999). The value in parentheses is the total cover of all trees in the sampled plot, assuming that no trees grow beneath others. This might be taken as a rough estimate of the mean maximum canopy cover in the analyzed dataset.

^b Forest Vegetation Simulator (FVS) modeling carried out post facto on size-class distribution data provided in the cited paper.

^c Values refer to 1889 and 1929 reconstructions.
value before adjustment, which assumes that no trees in the stand grow beneath any others and may therefore be seen as a rough estimate of the maximum possible canopy cover in the sampled stands.

Direct measurements of tree canopy cover in contemporary reference YPMC forests, and modeled canopy cover from historical datasets both strongly substantiate the oft-stated notion that presettlement forests were generally more open than modern forests in the assessment area (table 7). Average canopy cover values ranged from around 17 percent to less than 50 percent, except in the Sudworth (1900) dataset (16 YPMC plots from his table 1), which averaged almost 70 percent. The much higher canopy covers in Sudworth’s data support the notion that his data were not a random sample of forest conditions (Bouldin 1999, Stephens 2000, Stephens and Elliott-Fisk 1998). Uncorrected modeled cover values suggested that values as high as 70 to 90 percent were possible in very dense forest stands (table 7).

With respect to the FVS modeled values, it should be noted that a few studies have found that FVS techniques tend to estimate lower cover values than field-based methods, especially in areas of higher canopy cover. Fiala et al. (2006), in Douglas-fir/western hemlock forests in western Oregon, found that FVS estimated lower canopy cover than four field-based methods by 11 to 27 percent in high canopy cover mature and old-growth forests. The sampled stands were almost entirely >60 percent cover, however, and the difference between FVS and the most precise field methods became nearly zero below 60 percent canopy cover. Because YPMC forests in the assessment area were highly heterogeneous and mostly open canopied, FVS modeling probably provides a reasonable “ballpark” estimate of canopy cover for historical and reconstruction plots. Another issue is that most of the modeled FVS values in table 7 were carried out on summary data that had been organized into size classes. This lessens the variability of the input data and could affect the calculated mean in either direction, depending on the distribution of tree sizes within the size class categories.

Comparison to current—Data from the compiled FIA plots (USDA FS 2013a) show that mean modern canopy cover in YPMC forests is 45.7 percent (±21.8 SD; west-side mean = 50 percent, east–side mean = 38 percent). The average from the 15 reference values in table 7 is 34.4 percent (±14.3 SD) when including Sudworth, 34 percent when excluding Sudworth. The FIA canopy cover values were generated using the same FVS algorithm that produced most of the values in table 7. The modern average is about 33 percent higher than the presettlement average from table 7, but, as with all of the quantitative comparisons using the FIA data, it is difficult to statistically compare the two data sources given their different scopes and sample sizes. Generally, however, the data lend strong support to the idea that current canopy cover is higher on average than under presettlement conditions.
Future—Assuming that current trends in forest densification and fire exclusion continue, canopy cover will probably continue to increase across the YPMC forest belt. This will be offset to some extent if forest mortality agents, such as fire, insects, and disease, increase.

Forest gaps and tree clumps—

NRV—A “forest gap” is an opening or hole that exists within the forest canopy, or put another way, an area within a forest stand that is unoccupied by mature trees. Forman (1995) argued that gaps are small entities that exist within a mosaic of larger landscape “patches” (or what foresters refer to as “stands”), where a patch is a relatively large and homogeneous area that differs from its surroundings in its structure and composition. Essentially, patches are a component of coarse-grained heterogeneity, whereas gaps are a component of fine-gained heterogeneity. In assessment-area YPMC forests, the historical rarity of large-scale severe disturbances resulted in a very fine-grained structural heterogeneity that does not lend itself well to the recognition of landscape patches (Franklin and Fites-Kaufman 1996, USDA FS 2001). Given the generally low canopy cover and clumpy nature of YPMC forest stands, many presettlement forest stands were as dominated by gaps as by trees.

When gaps are created in the canopy, enhanced light is provided to the forest understory, which often results in a dynamic ecological response from both plants and animals. Gaps are important locations for tree, shrub, and herb regeneration, and they also serve a variety of roles in providing cover, forage, and other habitat for animals. Gaps are created through disturbance processes (“disturbance patches”), or they can also arise as a consequence of the physical habitat, e.g., soil type, rocks, and so on (“environmental patches”). The sizes and distributions of canopy gaps in forested ecosystems are important ecological variables (Agee 1993, Connell 1989, Spies and Franklin 1989), and over the past few decades it has become clear that the creation and maintenance of canopy gaps are important forest management objectives. Because open gaps in the forest canopy in a reasonably productive forest will quickly fill with regeneration, the size of “regeneration patches” can be roughly equated to the sizes of the original disturbance patches that led to the regeneration event. After growth to adult size, the surviving members of the regeneration patch will form an aggregation of trees, or a “tree clump” (Agee 1993). In general, canopy gaps are somewhat larger that the regeneration patches that arise within them, and mature tree clumps somewhat smaller than the regeneration patches that gave rise to them (Fry et al. 2014, Stephenson 1999).

Larson and Churchill (2012) outlined three components of local pattern in frequent-fire YPMC forests: tree clumps, widely spaced individual trees (usually
large), and openings. Tree clumps could be composed of similar-aged overstory
trees or multiaged aggregates, or they could be regeneration patches. In the studies
that Larson and Churchill (2012) reviewed, the average number of trees in a clump
ranged from 2 to 44 trees, and the average number of clumps per hectare ranged
from 10 to 27. Fry et al. (2014) compared spatial clustering within reference
YPMC forests in the SSPM of Mexico and forests in the Sierra Nevada of Califor-
nia, and found that the mean number of trees per clump ranged from 5.8 to 7.4 in
Mexico and from 8.1 to 11.2 in California, with a larger proportion of trees found
in large patches in California than in Mexico.

A number of studies in and around the North American Mediterranean climate
zone have used stand-mapping techniques to estimate typical sizes of canopy gaps,
regeneration patches, and tree clumps in YPMC forests. Table 8 provides a sum-
mary of the range of published gap-patch-clump sizes in the literature. Minimum
sizes averaged about 0.04 ha, and maximum sizes averaged about 0.3 ha, with a
maximum range from 0.07 ha to 1.17 ha. Larson and Churchill (2012) conducted an
independent summary of tree clump sizes in YPMC forests across the Western
United States, and found a range of 0.003 to 0.4 ha per clump. Their summarized
range for regeneration patches was somewhat wider, but with a similar mean: 0.001
to 0.64 ha.

<table>
<thead>
<tr>
<th>Site</th>
<th>Minimum size</th>
<th>Maximum size</th>
<th>Observations</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Central Oregon</td>
<td>?</td>
<td>0.16</td>
<td>Ponderosa pine; tree clumps</td>
<td>Youngblood et al. (2004)</td>
</tr>
<tr>
<td>Eastern Oregon</td>
<td>0.025</td>
<td>0.35</td>
<td>Ponderosa pine; tree clumps</td>
<td>Morrow (1985)</td>
</tr>
<tr>
<td>Lassen National Forest</td>
<td>0.003</td>
<td>0.14</td>
<td>Ponderosa pine-black oak; tree clumps</td>
<td>Taylor (2010)</td>
</tr>
<tr>
<td>Lassen National Forest</td>
<td>?</td>
<td>0.18</td>
<td>Ponderosa pine; tree clumps</td>
<td>Youngblood et al. (2004)</td>
</tr>
<tr>
<td>Lake Tahoe basin</td>
<td>0.03</td>
<td>0.07</td>
<td>Jeffrey pine-white fir; tree clumps</td>
<td>Taylor (2004)</td>
</tr>
<tr>
<td>Stanislaus National Forest</td>
<td>0.03</td>
<td>0.21</td>
<td>Mixed conifer; canopy gaps</td>
<td>Knapp (2012)</td>
</tr>
<tr>
<td>Kings Canyon National Park</td>
<td>0.014</td>
<td>0.16</td>
<td>Sequoia-mixed conifer; tree clumps</td>
<td>Bonnicksen and Stone (1980)</td>
</tr>
<tr>
<td>Sequoia National Park</td>
<td>0.16</td>
<td>1.17</td>
<td>Sequoia-mixed conifer; canopy gaps</td>
<td>Demetry (1995)</td>
</tr>
<tr>
<td>Northern Arizona</td>
<td>0.02</td>
<td>0.29</td>
<td>Ponderosa pine; tree clumps</td>
<td>White (1985)</td>
</tr>
<tr>
<td>Eastern Arizona</td>
<td>0.06</td>
<td>0.26</td>
<td>Ponderosa pine; regeneration patches</td>
<td>Cooper (1960)</td>
</tr>
<tr>
<td>Sierra de San Pedro Mártir,</td>
<td>0.001</td>
<td>0.07</td>
<td>Jeffrey pine-mixed conifer; regeneration</td>
<td>Stephens and Fry (2005)</td>
</tr>
<tr>
<td>northern Baja California</td>
<td></td>
<td></td>
<td>patches</td>
<td></td>
</tr>
</tbody>
</table>
Lydersen et al. (2013) described reference and current variability in canopy
gaps and tree clumps in plots in a mixed-conifer forest in the central Sierra Nevada
(same area studied by Knapp et al. [2012, 2013]). They found that, between 1889
and 2007–2008, the areal proportion of the studied stands occupied by canopy gaps
fell from about 50 percent to nearly zero, and gap density dropped from 6.2 gaps/
ha to 0.2 gaps/ha. Lydersen et al. (2013) also found that the average number of
trees found in a definable clump rose from 5.2 trees/clump to 26.3 trees/clump over
the same period, and the maximum number of trees per clump rose from 27 trees/
clump to 543 trees/clump. Overall, the modern stands were extremely homogeneous
in canopy cover, and completely lacked canopy gaps greater than 250 m² in area.

**Future**—Increased forest densification will continue the trend of reducing forest
openings/gaps. This will be offset to some extent if forest mortality agents continue
to increase.

**Snags and coarse woody debris**—

**NRV**—As he left the Sierra Nevada after 3 years of travel and mining, Peter
Decker, in his diary from 1849 to 1851, wrote that dead trees were seldom seen and,
“it is very rarely that timber is laying rotting” (Giffen 1966: 232). Arno (2000: 100)
stated that, in western YPMC forests characterized by understory fires, “shrubs,
understory trees, and downed logs were sparse, as testified to by dozens of histori-
cal photographs and narrative accounts.” In areas characterized by more moderate/
mixed-severity fires, surface fuels, snags, and coarse woody debris (CWD) tended
to be heterogeneously distributed, with concentrations found in scattered patches
burned so often, “It is unlikely that much large woody material survived fire long
enough to decompose fully in fire regimes that preceded the fire-suppression era.”

Stephens (2004) reported on fuel loads and snag densities in unlogged YPMC
forests in the SSPM, which also have seen only recent suppression of fires. Stephens
(2004) found that snag density in the sampled forests (all stems ≥2.5 cm d.b.h.)
averaged 3.5/ha before a major drought, and 5.1/ha after a multiyear drought.
Heterogeneity in snag distribution was very high: the modal snag density in the first
sample (before drought) was 0 snags/ha (35 percent of plots), in the second sample
(after drought) it was about 2.5/ha (27 percent of plots had no snags). The average
snag measured 59 cm d.b.h. Coarse woody debris, measured as the 1,000-hr fuels
(branches and logs >7.6 cm d.b.h.), averaged 15.8 tons/ha, but more than a third
of plots had no CWD and only 20 percent of the plots supported 75 percent of the
CWD. Eighty-one percent of the CWD was in a rotten state, and 19 percent was
sound (Stephens 2004).
Stephens et al. (2007) reported on CWD over 15 cm in diameter and ≥1 m long from the SSPM. Thus defined, CWD averaged 15.7 tons/ha, with a large range (0 to 154.5 tons/ha). The median was only 1.4 tons/ha, and about half of the plots sampled contained no CWD of this size or above. Overall, the average density of CWD was about 108 pieces/ha (±16.3 SE). Most sampled logs were rotten.

Lydersen and North (2012) sampled assessment-area YPMC forests that had not been logged and had experienced at least two fires in the 65 years before the field work commenced. They found very high variability in snag and CWD measurements. Snag density for snags greater than 50 cm d.b.h. averaged 9.1/ha, and snag volume averaged 123 m³/ha (±20.1 SE), with the highest densities and volumes of snags occurring on lower, northeast-facing slopes. Lydersen and North (2012) provided two measurements of CWD: logs ≥50 cm in diameter averaged about 10.3 pieces/ha (no measurement of variability given), while 1,000-hr fuels (>7.6 cm diameter) averaged 32.3 tons/ha (±3.4 SE).

Agee (2002), in a paper discussing the general relationship between fire regime and the creation and persistence of snags and CWD, suggested that a Fire Regime I forest with an undisturbed fire regime might typically support around five snags/ha, with the average snag size about 75 cm d.b.h. According to Agee (2002), CWD derived from the same average size of tree might total around 5 tons/ha. Because of very frequent fire in these forest types, both snags and CWD would fall or decompose at a higher rate than under conditions free of fire. For so-called mixed-severity fire regimes, Agee (2002) estimated that snag densities and CWD mass would be higher, as fires were more severe, which creates more snags and CWD, and less frequent, which allows them to persist longer on the landscape. Agee cited Wright (1998), who found an average of 40 tons/ha of snags and 55 tons/ha of CWD (ranging from 20 to 250 tons/ha) in Douglas-fir forests. Agee (2002) noted that modern fire-suppressed forests that used to experience frequent, low-severity fire might now support (much) more CWD than they did under presettlement conditions. Concerning these Fire Regime I forests, Agee (2002) stated that:

Burning prescriptions designed to retain most coarse woody debris can produce a “non-window”: duff moisture levels so high that such moisture contents are rarely if ever attained on these sites. Constraints to preserve all or most coarse woody debris effectively eliminate the use of fire for restoration purposes and leave the dry forest types at risk for stand-replacing fire. When such high-severity fire occurs, it brings with it the “boom and bust” coarse woody debris dynamics of the high severity fire regimes. This is a
classic case of the fine-filter (log preservation) trumping the coarse-filter (restoring the natural process), and in the long run is likely to result in a failed conservation strategy.

Harrod et al. (1998) used stand reconstruction and modeling techniques in an attempt to derive historical reference conditions for snag densities for an area of dry ponderosa pine/Douglas-fir forest in eastern Washington. They determined that, depending on the size classes included, snag densities under presettlement conditions probably ranged from about 14.5 to 34.6 snags/ha. Agee (2002) noted that Harrod et al. (1998) did not incorporate the effects of fire on snag-fall rates, implying that their numbers were probably biased upward.

Youngblood et al. (2004) studied three areas of old-growth ponderosa pine forest in eastern Oregon and northern California. All sites had experienced about a century of fire exclusion, although parts of the California site had recently experienced a prescribed fire. Youngblood et al. (2004) found that snag densities ranged from 8 to 12/ha (trees ≥15 cm d.b.h.), and CWD (≥15 cm diameter and 1 m long) averaged about 47 pieces/ha (±5.3 SE). Eighty-five percent of the CWD was between 15 and 65 cm in diameter, and most pieces were rotten (i.e., old). Youngblood et al. (2004) noted that the long-term demography of both snags and CWD in Western frequent-fire forests is largely unknown, with the current lack of fire in these stands a particular impediment. They suggested that, with respect to their ability to discern presettlement reference conditions from their data, fire exclusion over the previous century had probably biased their measurements of CWD (and snags) upward.

**Comparison to current**—Stephens (2004: 109) provided a table (his table 4) summarizing snag densities for seven studies in areas managed under fire suppression during the 20th century, two of which had also experienced timber harvest. The minimum sizes of measured snags differ from study to study, but the average snag density from these seven sites is 47.9 snags/ha, an order of magnitude higher than found in Stephens’ (2004) study of the SSPM, and also much higher than two other studies of unlogged YPMC forests also summarized in the table (Savage 1997, Stephens 2000).

Dunbar-Irwin and Safford (2016) carried out a direct statistical comparison of snag densities and CWD in the SSPM versus an array of forest plots sampled in analogous YPMC forests on the Sierra Nevada east slope. In SSPM, Dunbar-Irwin and Safford (2016) sampled from a much broader geographic area than Stephens (2004) or Stephens et al. (2007). They found that snag densities (trees
>10 cm d.b.h.) averaged 12/ha in SSPM (±2.8 SE), while the fire-suppressed eastern Sierra Nevada plots averaged 31/ha (±8.7 SE). Coarse woody debris (1,000-hr fuels) was not significantly different between the two regions, and averaged 28.9 tons/ha in SSPM vs. 25.1 tons/ha in the eastern Sierra Nevada; the median CWD loading was higher in the eastern Sierra Nevada (5.9 tons/ha vs. 4.8 tons/ha in SSPM). Note that Dunbar-Irwin and Safford sampled 16 years after Stephens (2004), during which time SSPM suppressed every lightning-ignited fire in the park. This would partly explain the higher snag and CWD values found in the more recent study.

Studies of the effects of prescribed fire on CWD and snags in the assessment area provide more evidence that levels of both were probably lower on average in presettlement forests than in current, unlogged (but fire-excluded) forests. Kauffman and Martin (1989) summarized the results of 60 prescribed fires conducted at three experimental sites in the northern Sierra Nevada in the mid-1980s. Fires were ignited at four different times of year: early and late spring, and early and late fall. We summarize the results from late spring and early fall burns, as they best represent the natural fire season in the assessment area: fires reduced the mass of sound CWD (>7.6 cm diameter, i.e., 1,000-hr fuels) by 62 percent on average, and the mass of rotten CWD by 87 percent, resulting in an overall postfire mean of 11.3 tons/ha; CWD averaged about 28 tons/ha before fire.

Various other studies of the effects of prescribed fires on CWD and snags corroborate the Kauffman and Martin (1989) results in indicating that frequent fire in these forests would quickly consume most of the snags and CWD that have resulted from a century of fire exclusion. For example, Kilgore and Sando (1975) measured 15 tons/ha of 1,000-hr CWD after fire (down from 97.5 ton/ha prefire); Stephens and Finney (2002) measured 3.4 tons/ha (from 42.1 tons/ha prefire); and Knapp et al. (2005) reported 15 tons/ha postfire (from 66.2 tons/ha prefire) for their fall burn. Kilgore (1973b) measured the change in log (>15 cm diameter) mass after prescribed fire, finding a reduction to 2.8 tons/ha from 12.8 tons/ha prefire. Knapp et al. (2005) also measured losses in log mass (>15 cm diameter) to a fall prescribed fire: postfire = 7.4 tons/ha, prefire = 47.2 tons/ha. Kilgore (1973b) reported that snags were almost entirely reduced to charred stumps and cavities in the prescribed fire he studied. Uzoh and Skinner (2009) investigated the effects of prescribed fire on CWD (>7.6 cm diameter) at Blacks Mountain Experimental Forest in the northern subregion of the assessment area. A single fall burn reduced CWD mass by 1.5 to two times more than the mechanical treatment effects, which also reduced CWD. In the burned plots, overall reductions of CWD ranged from 57 to 90 percent, with higher losses in the more decayed CWD classes. Uzoh and Skinner (2009) con-
cluded that “restoration goals that include large amounts of decayed, CWD do not appear to be appropriate for these dry pine forests of northeastern California.”

An important point is that these large reductions in CWD (and snags) were the result of single prescribed fires under moderate conditions at times of the year when naturally ignited fires are uncommon because of higher fuel moisture. They were also all carried out in stands that had not experienced fire for 50 to 100 years or more. Average fire return intervals in presettlement assessment-area YPMC forests were between 10 and 20 years, and such frequent fire would likely have rapidly reduced snag densities and CWD loadings.

The most recent FIA data compilation (USDA FS 2013a) gives an average of 23 tons/ha (±12.9 SD ) for CWD measured as 1,000-hr fuels (>7.6 cm diameter) in contemporary YPMC forests. The average of Kauffman and Martin (1989), Kilgore and Sando (1975), Knapp et al. (2005), Lydersen and North (2012), Stephens (2004), and Stephens and Finney (2002), all of which we interpret to broadly represent reference conditions, is 15.5 tons/ha (±9.4 SD ). The FIA cutoff for measurement of logs is a diameter of 25 cm, which does not perfectly correspond to any of the studies we found in the scientific literature. Nonetheless, the FIA average density for logs ≥25 cm in diameter is 42.6 pieces/ha, which is comparable to the density of 44.8 pieces/ha (logs ≥30 cm diameter) found by Stephens et al. (2007) in the SSPM.

FIA data give an average snag density (≥15 cm d.b.h.) of 37.2 snags/ha (±69.5 SD; mixed-conifer mean = 47.7 snags/ha, yellow pine mean = 20.2 snags/ha) for contemporary assessment-area YPMC forests. For the same size categories, Stephens (2004) found 4.4 snags/ha in the reference forests in the SSPM, and Youngblood et al. (2004) found 8 to 12 snags/ha in the old-growth pine forests they studied. Harrod et al.’s (1998) estimate of presettlement yellow pine forests in eastern Washington, which did not account for fire consumption of dead trees (see above), was in the range of 14 to 36 snags/ha.

Based on these data sources and considerations, we conclude that the levels of CWD and snags in modern YPMC forests in the assessment area are somewhat higher on average than in the average presettlement YPMC forest stand. Note that this statement does not pertain to the area of intensive beetle mortality that has occurred since 2014–2015. In this area, snag densities and CWD are both orders of magnitude higher than under average presettlement conditions.

**Future**—Continuation of current trends toward more snags and CWD seems likely, especially if nonfire forest mortality agents increase substantially. Increases in fire frequency will increase snags and CWD initially, but will reduce them as areas are reburned.
Forest Understory and Nonforest Vegetation

Few of the early observers of assessment-area YPMC forest provide more than passing reference to nonforest vegetation, except with respect to seedling and sapling densities and an occasional reference to shrubs.

Tree seedlings and saplings—

NRV and comparison to current—The very low average density of tree seedlings and saplings in YPMC forests alarmed early foresters in the assessment area, but it was clear to them that recruitment potential was high and fire was the main factor reducing recruitment (e.g., Greeley 1907, Leiberg 1902, Show and Kotok 1924, Sudworth 1900). Sudworth (1900) wrote that:

The frequent open spaces in yellow-pine forests are sooner or later covered with dense patches of young trees, but these thickets may in turn be swept off by fire. So continuous and widespread are these forest fires that… they keep a very large percentage of the seedling growth down… The forest floor looks clean swept. But the remarkable productive power… is seen only in localities where fences and the exclusion of fire have protected the incoming seedlings. Here the stand is so dense as to be quite impenetrable.

Where seedlings and saplings successfully survived (usually because of longer intervals without fire), shade-tolerant species tended to dominate. Greeley (1907) wrote:

The very large proportion of fir and cedar in its reproduction is the worst feature of the west Sierra Forest. Both of these species are prolific seed bearers… Except at lower elevations and on very warm exposures, where yellow pine grows in pure stand, dense thickets of fir and cedar crowd the young pine down to one fifth or less of the reproduction.

Bonnicksen and Stone (1982), in one of the first modern stand reconstruction studies, noted that the 1890s landscape at their study site supported much more open ground with seedlings and saplings than the current forest. Around 50 percent of the area covered by seedling and sapling “aggregations” was dominated by white fir, which surprised the authors, as they had expected shade-intolerant species to dominate recruitment into gaps.

What emerges is an ecosystem in which conditions were greatly determined by the frequency and nature of fire. Frequent fire substantially reduced reproduction, and the resulting bare mineral soil and open forest conditions favored species like yellow pine and black oak, which could better survive low-intensity burns and did

Modern reference sites like the SSPM or Beaver Creek Pinery (Lassen National Forest) show very high heterogeneity in seedling densities, which is similar to the conditions described by early observers. The mean seedling density in the SSPM was measured by Stephens and Gill (2005) at 125 seedlings/ha, with about 50 percent of the sampled plots supporting <100 seedlings/ha and 10 percent supporting >300 seedlings/ha. Sixteen years of fire exclusion later, Dunbar-Irwin and Safford (2016) found mean seedling densities in SSPM of 420 per hectare, but the median was zero; the coefficient of variation for seedling density in SSPM was more than twice as high as in the eastern Sierra Nevada plots that were compared to it. Mean seedling density in the Beaver Creek Pinery was measured by Taylor (2010) at about 660 seedlings/ha, including ponderosa pine and black oak. More than two-thirds of the ponderosa pine seedlings were found in high light conditions in forest gaps and variation in densities was very large, ranging from 37 to 1,408 seedlings/ha. It was observed that surviving seedlings after fire in the Beaver Creek Pinery tended to be associated with open canopy gaps that did not burn because of the lack of surface fuels, whereas seedlings were killed where they grew in gaps with some canopy cover that supported sufficient needle litter to carry fire (see footnote 3).

Compilation of the most recent FIA data (USDA FS 2013a) gives a mean of 1,820.6 seedlings/ha (± 3.973 SD) for yellow pine and mixed-conifer plots. The coefficient of variation (CV) is 2.2, which indicates high variance in seedling density among plots. We conclude from the data and from inference based on successional processes that current mean seedling densities are almost certainly higher than mean seedling densities under presettlement conditions.

Future—Seedling composition will largely reflect the overstory composition (van Mantgem et al. 2006). The very high proportion of shade-tolerant species in current YPMC forests in the assessment area suggests that future forest recruitment will be dominated by species like white fire and incense cedar. Even where repeated, prescribed fires have been employed in order to enhance pine and oak recruitment, the persistence of mature shade-tolerant species in the forest canopy has resulted in a strong dominance of these species in postfire seedling densities (Webster and Halpern 2010). If an increase in recruitment of fire and drought-tolerant species is desired, removal of shade-tolerant species from the overstory or direct planting of fire-tolerant seedlings may be necessary. If fire-tolerant species are already present at sufficient numbers in the overstory, the restoration of a frequent fire regime may accomplish this same objective.
Shrubs—

NRV and comparison to current—Like tree cover, most evidence is that shrub cover in presettlement YPMC forests was relatively low but highly heterogeneous on the landscape. For example, Fitch (1900) described forests in the Yosemite area as: “…remarkably free from undergrowth… and only along streams, in the bottom of gulches, and on rocky southern slopes is the brush so thick as to impede progress.” Marshall (1900), surveying the higher elevations of Yosemite National Park for the U.S. Geological Survey, which included some areas of moist mixed conifer, stated that the forest was “everywhere open without undergrowth of any kind.” Decker, in his journal, wrote of the forests in the upper Mokelumne River drainage that there was “generally no underbrush” (Giffen 1966). On the other hand, Bruff, in his journal from the same time period refers to both open forests and forests with dense, bushy undergrowth (Read and Gaines 1949).

Leiberg (1902) stated that much of the forest understory in his survey area was more or less bare. At the same time, he noted that burned areas supported a “great amount” of undergrowth, mostly dominated by species of Ceanothus, which he ascribed to recent anthropogenic fire activity. According to Leiberg (1902), burned areas in YPMC forest often supported dense growths of montane chaparral, from 1.2 to 1.5 m in height. He was convinced that montane chaparral came only from fire in previously forested stands and counted such areas as representing destroyed forest even where there wasn’t any evidence of previous forest (Miller and Safford 2017). Greeley (1907), referring primarily to fires set by Euro-Americans, decried the loss of valuable timberlands to montane chaparral. Contemporary data from the Plumas National Forest and Illilouette Creek basin in Yosemite confirm that fire-created chaparral patches are more likely to reburn at high severity, which can result in inhibition of forest succession and a (semi-) permanent conversion to chaparral (Coppoletta et al. 2016, van Wagendonk et al. 2012).

Mitchell (1913) noted that the quantity of brush (shrubs) in yellow pine forest tended to be minimal, but it was much higher in mixed-conifer stands (and even higher in red fir forests). Given that forest openings were much more prevalent in yellow pine stands, the difference was probably due to more frequent fire (which reduced shrub cover), less intense fire (which would result in less fire-induced germination of shrubs like Ceanothus or manzanita), and lower site productivity.

By using the section-line information recorded by crews doing GLO surveys, Baker (2014) found that >90 percent of the area surveyed had some mention of shrubs, mostly Ceanothus and Arctostaphylos. Surveyors neglected, however, to record entry or exit information from shrub patches on almost 30 percent of northern Sierra section-lines, and more than 55 percent of southern Sierra section-lines,
so an estimate of shrub cover is not possible. Stephens et al. (2015) found that 54 percent of belt transects from a 1911 timber survey in the Greenhorn Mountains of the Sequoia National Forest contained shrub patches, and the average shrub cover within a given transect was 25 percent.

Show and Kotok (1924) decried the loss of forest to severe fires set by humans, especially in situations where logging slash was heavy. Severe forest fires in the assessment area normally result in postfire succession to shrublands (usually some form of chaparral), especially on the west side. In the absence of further fire, such shrublands will last for decades before succession to forest occurs. In the presence of further fire, such stands can become quasi-permanent features of the landscape, because chaparral generally burns at high intensity and kills most young trees (Nagel and Taylor 2005, Skinner and Taylor 2006). Show and Kotok (1924) produced a table providing summaries of the areas of the 10 northern national forests (minus the Modoc) that supported “brushfields” in the early 1920s. In their interpretation, these brushfields had developed in previously forested stands after fires set by Euro-American settlers. We reproduce that table for the six national forests in the assessment area in table 9, and compare it to modern areas of chaparral on the same national forests. Percentages of national forest area in brushfields in the early 1920s ranged from 7 percent on the Eldorado National Forest, to over 16 percent on the Stanislaus National Forest; the overall average was about 11 percent. Modern data (from the most recent Forest Service existing vegetation mapping, data at http://www.fs.usda.gov/detail/r5/landmanagement/gis/?cid=STELPRDB5327836), using “mixed chaparral” and “montane chaparral” California Wildlife Habitat

Table 9—Estimated percentages of the area covered by seral shrub fields in 1924 and the early to mid-2000s on six national forests in the assessment area

<table>
<thead>
<tr>
<th>National forest</th>
<th>Seral shrub fields</th>
<th>1924</th>
<th>2000s</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Percent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lassen</td>
<td>11.8</td>
<td>8.5</td>
<td></td>
</tr>
<tr>
<td>Plumas</td>
<td>11.7</td>
<td>6.2</td>
<td></td>
</tr>
<tr>
<td>Tahoe</td>
<td>8.6</td>
<td>10.6</td>
<td></td>
</tr>
<tr>
<td>Eldorado</td>
<td>7.0</td>
<td>9.2</td>
<td></td>
</tr>
<tr>
<td>Stanislaus</td>
<td>16.2</td>
<td>9.9</td>
<td></td>
</tr>
<tr>
<td>Sierra</td>
<td>11.1</td>
<td>7.0</td>
<td></td>
</tr>
<tr>
<td>Average</td>
<td>11.1</td>
<td>8.6</td>
<td></td>
</tr>
</tbody>
</table>

Data are from table 19 in Show and Kotok (1924).

Data from most recent Forest Service existing vegetation maps, using “Classification and Assessment with Landsat of Visible Ecological Grouping” types of mixed chaparral and montane chaparral where growing in areas identified as having sufficient productivity to support forest. Imagery is from the early and mid-2000s.
Relations types where they occur in sites identified as having sufficient productivity to support forest as our definition of shrublands, show that the overall averages are slightly lower on average. Four of the national forests experienced a decrease in the area occupied by shrubs, and two experienced increases. According to our comparison, the Eldorado National Forest experienced the greatest increase (+31 percent), the Plumas National Forest the greatest decrease (-47 percent). Note that the most recent vegetation mapping on the northern Sierra Nevada national forests (Lassen, Plumas, and Tahoe) was completed some time ago, using imagery from 2000 and 2005. The values in table 9 thus exclude the large expanses of fire-caused shrubfields that have resulted from the high number of large, often very severe fires that have occurred in the northern Sierra Nevada and southern Cascade Range since 2005 (e.g., Rich, Butte Lightning Complex [BTU], Moonlight, Antelope Complex, Chips, Reading, Eiler). Including these numbers would raise the modern percentages substantially, especially on the Plumas National Forest. Note also that the techniques for measuring the areal extent of shrubfields in the two surveys were different (summary of ocular and land surveyed measurements in 1920s, versus remotely sensed imagery in 2000s), and the numbers in table 9 should therefore be viewed as approximations.

Modern studies of reference YPMC forests have found an average of about 15 to 25 percent relative shrub cover (but with high variability) in forests that have not been logged or have not experienced complete fire exclusion (table 10). Note that very large areas of chaparral would not have been sampled by these studies as they would not be considered forest for sampling purposes.

Contemporary FIA data on shrub cover are difficult to compare with other datasets, as shrub cover is computed by summing the species-specific covers without accounting for overlap (and therefore total cover can sum to more than 100 percent). Average absolute cover by shrubs in YPMC forest plots is 22.6 percent (±23.3 SD) (USDA FS 2013a). These values will be higher than the actual relative values.

Bonnicksen and Stone (1982) carried out a stand reconstruction in a small watershed of YPMC and giant sequoia forest in Kings Canyon National Park. They estimated that about three-fourths of the forest aggregations dominated by large and very large trees had some sort of shrub cover in their understory in 1890. They also found that about 19 percent of the watershed was covered by shrub-dominated “aggregations” in 1890 versus 11 percent in 1977, but these would have represented shrubfields rather than forest understory.

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Table 10—Percentages of shrub cover in reference yellow pine and mixed-conifer (YPMC) stands in the North American Mediterranean zone

<table>
<thead>
<tr>
<th>Site</th>
<th>Mean</th>
<th>Median</th>
<th>Mode</th>
<th>Standard error</th>
<th>Observations</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Northeastern assessment area</td>
<td>21.4</td>
<td>8</td>
<td>8</td>
<td>1.0</td>
<td>Yellow pine forests, mature and old-growth, uncut</td>
<td>Smith (1994)</td>
</tr>
<tr>
<td>Northern assessment area</td>
<td>20.8</td>
<td>11.5</td>
<td>10</td>
<td>3.9</td>
<td>Mixed-conifer forests, mature and old-growth, uncut</td>
<td>Fites (1993)</td>
</tr>
<tr>
<td>Assessment area (mostly southern and central)</td>
<td>16.9</td>
<td></td>
<td></td>
<td>1.9</td>
<td>YPMC forest, uncut, with frequent or recent non-stand-replacing fire</td>
<td>Lydersen and North (2012)</td>
</tr>
<tr>
<td>Stanislaus National Forest</td>
<td>28.6</td>
<td></td>
<td></td>
<td>5.5</td>
<td>YPMC forest on productive site, 40 years after most recent fire</td>
<td>Knapp et al. (2013)</td>
</tr>
<tr>
<td>Sierra Juarez, northern Baja California</td>
<td>16.3</td>
<td>17.5</td>
<td>0</td>
<td>3.5</td>
<td>Yellow pine forests, mostly uncut, only recent fire suppression</td>
<td>Safford, in preparation</td>
</tr>
<tr>
<td>Sierra de San Pedro Mártir, northern Baja California</td>
<td>17.4</td>
<td>5</td>
<td>0</td>
<td>2.9</td>
<td>Mixed-conifer forests, uncut, only recent fire suppression</td>
<td>Dunbar-Irwin and Safford (2016)</td>
</tr>
</tbody>
</table>

Vankat and Major (1978) found that, in general, shrub cover had dropped in YPMC forests in Sequoia & Kings Canyon National Parks over the previous half century or more. They ascribed the changes to more light competition from trees, increased browse by ungulate populations, or less fire. The most affected shrub genera were *Arctostaphylos* and *Ceanothus*, both of which support species with fire-cued germination. Vankat and Major (1978) also noted some areas in which shrub cover had increased.

Laudenslayer and Darr (1990), studying areas where timber harvest had occurred, stated that shrub cover had increased in most places owing to greater availability of light and the long-term lack of fire, which they suggested had reduced shrub mortality.

Comparison of aerial photos from 1930s and 1940s with photos from today sometimes shows greater cover of shrubland in the early photos. It is difficult to determine whether the early photos are NRV, however, as there were many severe fires set by settlers in the late 1800s. For example, Nagel and Taylor (2005) found that the average area of six chaparral stands in the Lake Tahoe basin retracted by 62.4 percent between 1939 and 2000. Note that these comparisons are mostly focused on large stands of chaparral, rather than patches of shrubs found within the forest matrix itself.
Thorne et al. (2008) compared Forest Service VTM maps from the 1930s (Wieslander 1935) with modern Forest Service vegetation maps in the lower elevations of Eldorado County, on the west side of the central assessment area. They found that the extent of montane chaparral stands had declined by more than 90 percent over the 60-year period. Thorne et al. (2008) noted that some of these stands had potentially transitioned to hardwood stands, but others were large patches of chaparral from earlier timber harvest and fires that were reoccupied by conifer forest after the institution of fire suppression. As above, the extent of chaparral on the landscape in the 1930s was probably notably enhanced over presettlement conditions by human activities.

Kauffman and Martin (1990) reported on the effects on shrubs of 60 prescribed fires in the northern Sierra Nevada. We report only those results from the late spring and early fall burns (n = 30), as they best replicate the seasonality of natural fires. Kauffman and Martin (1990) found that fires at these times of year tended to consume more fuel and (mostly) burned more intensely than fires in the early spring and late fall. One year after fire, an average (across the three study sites) of 25 percent of shrubs within the burn plots had survived the early fall burns, and 36 percent survived the late spring burns. Kauffman and Martin (1990) also tracked 2-year survival of black oak and tanoak (Lithocarpus densiflorus (Hook. & Arn.) Rehder) and found only 9 percent survival of black oak and 18 percent survival of tanoak in the early fall burn and, 28 percent and 21 percent survival, respectively, from the late spring burn. Survival was higher from the early spring and late fall burns. Seedling recruitment and sprouting from Ceanothus species was relatively strong, with hotter burns increasing mortality and reducing sprouting but increasing the number of seedlings. Kauffman and Martin (1990) concluded that intense burns could be used to reduce shrub cover on sites where that was a management goal.

Knapp et al. (2012) studied the shrub seed bank in 24 assessment-area YPMC sites. They found that seeds of Ceanothus were found at 88 percent of the unburned sites, at high densities (mean = 246 seeds/m² of soil surface), and as deep as 10 cm. Arctostaphylos and Prunus seeds were found at 64 and 45 percent of the unburned sites, respectively. The intrinsic potential for shrub response to fire is thus very high throughout assessment-area YPMC forests, and it has not been appreciably affected by fire exclusion.

Overall, considerations of shrub cover on assessment-area landscapes suggest that the overall portion of the YPMC landscape occupied by shrubs today is broadly similar to, but possibly somewhat lower than, the portion occupied at the beginning of the 20th century. Many early observers believed that shrub
cover had been increased after Euro-American settlement, so it may be that the current overall areal extent of shrublands is actually somewhat higher than when Euro-Americans began arriving in the mid-19th century, but we have no quantitative data to substantiate this inference. At the same time, fire suppression, logging, and other management practices since the beginning of the 20th century have greatly homogenized assessment-area YPMC forests, resulting in higher canopy cover and denser stands, and less optimal conditions for shrub survival in the forest understory. In summary, the landscape extent of shrubfields and other early seral vegetation within the YPMC forest belt is probably within NRV, but shrub cover in the forest understory has probably been generally reduced by increasing tree density and canopy cover and decreasing understory light availability.

**Future**—The increase in fire area, fire severity, and high-severity patch size in assessment-area YPMC forests over the last quarter-century (Mallek et al. 2013, Miller and Safford 2012, Miller et al. 2009b) has led to a recent increase in the extent of early seral montane chaparral stands, especially in Forest Service lands in the northern half of the assessment area. Most future models and analyses of paleo-data suggest that future disturbance cycles and warming climates will further increase the amount of early seral vegetation on assessment area landscapes (e.g., Cole 2010, Lenihan et al. 2008, McKenzie et al. 2004).

**Grass and forbs**—

**NRV and comparison to current**—William Brewer’s (1930) memoirs of his 1861–1864 travels with the Whitney Survey refer to the scarcity of good grass cover in the Sierra Nevada, and lush areas of grass are highlighted where they occur. Sudworth (1900) also referred repeatedly to the lack of good pasturage, and stated toward the beginning of his report that “…forage is exceedingly short on all the unfenced mountain ranges.” Later Sudworth suggests that in many places, the lack of herbaceous growth is due to the effects of sheep grazing, although it is important to note that his field surveys also coincided with one of the worst California droughts on record. Interestingly, Fitch (1900), working adjacent to Sudworth’s survey area, described “excellent pasturage” on high mountain slopes, meadows, and river valleys, as well as in open timber stands. None of Sudworth’s photos from the YPMC forests, except perhaps one from the lower boundary with the oak-foothill pine belt, which supports much annual grassland, show any obvious areas of grass cover. Leiberg (1902) noted that humus layers were rare to nonexistent in the forests he surveyed, and the forest floor was usually bare, perhaps with a layer of needles “rarely exceeding two inches in depth.”
Evett et al. (2007, 2006), studied soil phytoliths in mixed-conifer forests in the North American Mediterranean zone, looking to determine whether there was any support for hypotheses of substantial grass cover in presettlement YPMC forests. Phytoliths (also known as “grass opal”) are small structures of silica left behind in the soil after death of the parent plant, and grass phytoliths are morphologically unique. Evett et al. (2006) found that the grass phytolith content of soils in a giant sequoia–mixed-conifer forest in the southern Sierra Nevada indicated a long-term (at least centuries-long) lack of a substantial grass component at the study site. However, the study found evidence of somewhat higher grass cover in some places on the landscape, such as in ponderosa pine-dominant forest on ridge tops, and in areas near stream channels. Evett et al. (2007) carried out a similar study in a Jeffrey pine/mixed-conifer forest in the SSPM of northern Baja California. As above, they found that grass phytolith levels were too low to indicate extensive cover of grass over the previous centuries.

Takahashi et al. (1994) studied volcanic soil development under mixed-conifer forests in northern California. Presence of melanic epipedons (humus) in forest andisols has traditionally been ascribed to periods in which the soils supported grassy vegetation, but phytolith analysis of the California YPMC soils showed very little to no grass component could have been present during the formation of the soils, i.e., over many centuries. According to Takahashi et al. (1994), the occurrence of frequent fire in their study forests may have been the key factor in forming melanic epipedons in the andisols they studied.

Modern surveys of YPMC forests in the California Mediterranean zone do not suggest that grass or forb cover is particularly high, even in relatively undisturbed stands with low canopy cover. Smith (1994) described 45 associations of yellow pine forests in the northeastern assessment area, sampling only from uncut stands that were at least 100 years old. Few of the sampled sites were heavily grazed.15 Average overstory cover (relative) was about 53 percent (range = 13 to 82 percent), shrub cover 21.4 percent (range = 1 to 61 percent), forb cover 8.9 percent (range = 2 to 41 percent), and grass cover 6.5 percent (range = 2 to 20 percent). Fites (1993) conducted a similar survey of older and undisturbed mixed-conifer forests in the northern and southern Cascade Range. Canopy covers were much higher in these more productive sites (range = 25 to 96 percent), and forb and grass covers similarly low (relative cover means of 11.8 percent and 3.2 percent, respectively). Oliver (2000) described forest conditions

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at Black Mountain Experimental Forest on the Lassen National Forest. Perennial
grass and herb cover was very low both before and after forest treatment (less
than 4 percent total in both cases), although the cover of nonnative invasive
annual grasses increased notably after treatment, especially prescribed fire.
However, C. Skinner noted that grass cover has increased in the years since this
study (see footnote 3). Barbour et al. (2002) sampled uncut old-growth mixed-
conifer stands in the Lake Tahoe basin and found an average of only 0.5 percent
(range = 0 to 10 percent) relative herbaceous cover, even though overstory tree
cover was only 45 percent on average.

The contemporary FIA data provide measurements of herbaceous cover that are
generated by summing the cover percentages of the different herbaceous species,
thus total cover can sum to more than 100 percent. Comparison with relative cover
data (which is what is typically reported in scientific studies) should be done with
care, as the FIA summing practice ignores overlap among plants. The average
cover of herbaceous plants reported from the FIA plots is 11.1 percent (± 12.5 SD )
(USDA FS 2013a). Remember that the FIA dataset is a statistical sample of all stand
conditions, whereas the reference values measured above are all from relatively
undisturbed, old-growth-type stands.

The SSPM in northern Baja California has not been logged and has expe-
rienced more or less effective fire suppression only since the 1980s. Cattle are
present but spend relatively little time in the upland forest at any distance from
water sources and meadows (although it is true that low densities of cattle dung
are present throughout forested areas in SSPM). A recent study of the understory
vegetation in conifer stands in the SSPM found that relative forb cover averages
9 percent and grass cover less than 2 percent, under an average tree cover of 29
percent. Drier and lower elevation Jeffrey pine stands in the Sierra Juarez near
the U.S. border show similarly low levels of understory cover: averages of 6.6 and
11.8 percent for forbs and grasses, respectively, under tree cover of 36 percent on
average (see footnote 13).

Pearson (1942) described the effects of herbaceous vegetation on ponderosa
pine recruitment and survival, and explained how the substantial cover of under-
story grass in southwestern (Arizona and New Mexico) ponderosa pine forests was
highly dependent on the occurrence of monsoonal summer rain showers. He noted
that, “Shallow-rooted herbs are dependent on summer showers, whereas the deeper
rooted shrubs and trees are able to grow with little or no summer precipitation if the
soil is well saturated during the winter months… A pronounced midsummer rainy
season as well as relatively heavy soil favors the grasses.”
Swetnam and Betancourt (1998) showed evidence from Arizona and New Mexico that fires in both mixed-conifer forests and ponderosa pine forests in that region tended to occur in years of low precipitation, but their analysis also suggested that fire in the latter forest type was positively correlated with precipitation in prior years. Swetnam and Betancourt (1998) ascribed this pattern to the increased importance of fine fuels—both live (grass) and dead (needles)—to fire dynamics in the more open and drier ponderosa pine stands, whereas moister mixed-conifer forests are more characterized by woody fuels that build up gradually over years and are less responsive to annual changes in precipitation. California supports a Mediterranean climate that is similar to the southwestern climate (Arizona and New Mexico) in its overall aridity but very different in its distribution of rainfall. For example, July and August are the driest months in California, but the wettest in Arizona and New Mexico; in much of New Mexico, the driest months are December through February, which is the height of the rainy season in California. These differences affect herbaceous vegetation much more profoundly than woody vegetation, and the general paucity of rainfall during the growing season in the assessment area means that grass and forb production is usually severely water limited in upland sites.

Norman and Taylor (2003) carried out a study of fire history and fire-climate interactions in pine forests found along meadow edges on the Lassen National Forest in an area of relatively subdued topography. They found that the historical occurrence of widespread fires was partly dependent on moisture conditions 1 to 3 years prior, which suggested that grass production in and around the meadow systems was important to fire spread. It is difficult to extrapolate this study to the broader landscape, as most assessment-area YPMC forests are not located adjacent to extensive meadow complexes, and grass cover in upland forests is generally not sufficient to carry fire on its own (Fites 1993, Oliver 2000, Smith 1994).

In summary, there appears to be little basis for the idea that presettlement assessment-area YPMC forests supported abundant swards of grass in their understory, except possibly in areas of high soil moisture and relatively open canopy conditions, and at lower elevations in yellow pine-dominant stands, especially where these interfinger with oak woodland. Except in these places, surface fires may have been primarily fed by tree litter, dried forbs, and shrubs, with a variable component of grass. Variability in precipitation, especially in the spring and summer, certainly played a role in driving interannual differences in herbaceous and graminoid biomass, however.

Although grass cover in presettlement YPMC forests may not have been particularly high, many forbs are more shade tolerant, and overall herbaceous cover and local species richness has probably dropped as tree cover has increased with fire suppression. Parks (2009) noted that stem densities of understory herbaceous species in the Stanislaus-Tuolumne Experimental Forest have been reduced by
around 80 percent over the past eight decades as a result of increases in forest stand density and canopy cover.

Another wildcard is the increased presence of cheatgrass (*Bromus tectorum* L.) in assessment-area YPMC forests. In east-side and lower elevation west-side forests, this invasive grass can take advantage of disturbance to invade forest stands (Keeley and McGinnis 2007). Wet, warm years can increase cheatgrass cover substantially, and in recent wet years, we have seen yellow pine stands with substantial cheatgrass cover in their understories where little understory existed before.

**Future**—As with shrubs, it seems two trends are likely. On the landscape scale, it seems inevitable that current and projected future trends will lead to increased importance of (early seral) herbaceous vegetation on the YPMC landscape. On the other hand, in undisturbed forest stands, forest cover and density will continue to increase and herbaceous diversity and abundance will continue to be suppressed.

**Litter**—

**NRV and comparison to current**—There are no measurements of litter depth or cover that we know of from YPMC forests in the late 19th or early 20th centuries. However, many early observers remarked on the common occurrence of bare soil and the prevalence of thin layers of needle litter on the soil surface (e.g., Fitch 1900, Leiberg 1902, Sudworth 1900).

Modern litter + duff depth measurements from contemporary reference sites tend to average between 1.5 and 2 cm (e.g., Lydersen and North [2012]: 2.1 cm [±1.8 SD]; Stephens [2004], Sierra de San Pedro Mártir: 1.6 cm [no standard deviation given]; Safford, unpublished data, Sierra Juarez: 2.1 cm [± 1.6 SD]26).

The FIA data compilation (USDA FS 2013a) for YPMC forests gives an average of 3.4 cm (±4.5 SD), with mixed-conifer plots averaging 3.9 cm and yellow pine plots averaging 2.8. These estimates are consistent with measured data from modern fire-suppressed YPMC forests from Stevens et al. (2014), who found an average litter + duff depth of 3.47 cm (±2.17 SD). Van Wagendonk et al. (1998b) found that modern litter + duff layers ranged between an average of approximately 4 and 6 cm depending on the age of the stand, but there were strong differences in depth among different conifer species.

These are very few data points on which to base an ecosystem-wide conclusion. See the “Forest fuels” section below for more evidence relating to forest floor accumulations of dead plant matter. Van Wagendonk et al. (1998b) found that modern litter + duff layers ranged from approximately 4 to 6 cm depending on the age of the stand, but there were strong differences in depth among different conifer species.

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26 Unpublished data. On file with: Hugh Safford, USDA Forest Service, Pacific Southwest Region, 1323 Club Drive, Vallejo, CA 94592.
Forest fuels—

**NRV and comparison to current**—Stephens (2004), working in reference YPMC forests in the SSPM in Baja, California, found average surface fuel loads (1- to 1,000-hr summed) of 15.8 tons/ha (±3.9 SE; the 1-hr to 100-hr sum was 2.16 tons/ha) and ground (litter + duff) fuels of 8.7 tons/ha (±0.83 SE). Duff is almost non-existent in this system (Stephens 2004). Fuel loadings were extremely variable.

For example, surface fuel loads were below average on 73 percent of the plots, and above 36.8 tons/ha on 8 percent of plots. Thirty-seven percent of the plots sampled had no 1,000-hr fuels (branches and logs >7.6 cm diameter), and 20 percent of the plots supported 75 percent of the 1,000-hr fuels. Dunbar-Irwin and Safford (2016) sampled a broader area of the SSPM 16 years after Stephens and found that fuel loadings had increased, probably largely because of the strict fire suppression that has been instituted in the area since the 1980s. The 1- to 100-hr sum was 4.7 tons/ha, and the overall sum (including 1,000-hr fuels) was 33.6 tons/ha.

Lydersen and North (2012), in their study of old-growth stands in the assessment area that had experienced at least two fires over the previous 65 years, found an average of 40.2 tons/ha for all surface fuels (1- to 1,000-hr summed; 1- to 100-hr sum was 7.9 tons/ha). Variability was high, with the standard deviations for all of the component fuel types about equal to or greater than the mean. Fuel loadings were lowest on ridgetops and southwest-facing slopes.

Taylor et al. (2014) used three methods to estimate prefire exclusion fuel loads in reconstruction plots in YPMC forests in the Lake Tahoe basin. Average estimated fuel loads from the three techniques ranged from 4.2 to 6.1 tons/ha (1- to 100-hr summed) for Jeffrey pine forest and 4.4 to 8.3 tons/ha for mixed-conifer forest. This compared to contemporary ranges of 4.2 to 7.5 tons/ha for Jeffrey pine forest and 8.5 to 12.9 tons/ha for mixed-conifer forest.

Studies of prescribed fire in modern, fire-suppressed forest can provide insight into fuel loadings that may have characterized presettlement YPMC forests in the assessment area. Kauffman and Martin (1989) report results from 60 prescribed fires from three sites in the northern Sierra Nevada. We report only their early fall and late spring results (30 burns), as those fires are most likely to represent conditions like those during the natural fire season. After fire, Kauffman and Martin (1989) found an average of 1.5 tons/ha (range of site means 0.6 to 2.2) for 1- to 100-hr fuels, and an average of 12.8 tons/ha (5.8 to 18.7) for 1- to 1,000-hr summed fuels; ground fuels (litter + duff) averaged 11.6 tons/ha (fig. 21). Unburned forest supported over eight times more ground fuels by mass, more than five times more fuel in the 1- to 100-hr classes, and 2.8 times more fuel when the 1- to 1,000-hr classes are summed (fig. 21). Other studies of prescribed-fire effects on fuels provide remarkably similar results. The means of postfire 1- to 100-hr fuels, 1- to
1,000-hr fuels and ground fuels from Kilgore and Sando (1975), Knapp et al. (2005; fall burn), and Stephens and Finney (2002) are 1.4 tons/ha, 10.7 tons/ha, and 12.5 tons/ha, respectively. Average prefire fuels in these studies were higher than in the Kauffman and Martin study sites. We would expect average presettlement fuel loadings to be less than the postfire loadings reported here, as these modern prescribed fires were one-time events after many decades of fire exclusion, whereas presettlement YPMC forests were experiencing burns every 10 to 20 years on average.

The FIA plot data compilation (USDA FS 2013a) showed that fuel loadings in contemporary YPMC forest plots average 7.3 tons/ha for 1- to 100-hr summed fuels, and 30.3 tons/ha (±24.9 SD) for 1- to 1,000-hr fuels. Mixed-conifer forests support much heavier fuels (1- to 1,000-hr summed = 36.7 tons/ha) than yellow pine forests (1- to 1,000-hr summed = 20 tons/ha). This gives a good idea of the kind of transformation that occurs in the fuel profile of YPMC forests as they transition from pine-dominant stands to mixed-conifer stands under fire suppression. The average of the reference studies detailed above is about 3.6 tons/ha for 1- to 100-hr fuels, and 17.7 tons/ha for 1- to 1,000-hr fuels (the latter including the very high value from Lydersen and North [2012]). We conclude that modern-day fuel loadings in assessment-area YPMC forests are substantially higher than loadings under presettlement conditions.
Future—We know of no scientific attempt to project fuel levels in assessment-area YPMC forests into the future. It seems most likely that the same sorts of trends that have characterized YPMC forests for the past 50 to 100 years will continue into the future, perhaps at an accelerated pace, as climate warming and rising carbon dioxide levels increase plant growth and fuels accumulation.

Composition
Forest landscape composition

NRV and comparison to current—
Here, we refer to the distribution of forest types (rather than specific species) across the landscape. There are not many data available, as mapping or extensive plot networks are necessary to assess this indicator.

Many early observers noted how yellow pine-dominant forests (yellow pine and dry mixed conifer) were more common at lower elevations, on warm aspects (south and west), and in areas of thin or otherwise low-productivity soils. Forests with a notable shade-tolerant component (moist mixed conifer) were mostly in moist microsites, along streams, on north slopes, and at high elevations (e.g., Leiberg 1902, Sudworth 1900). These general trends are also apparent today, although the absolute area supporting yellow pine-dominant forest has dropped and the area supporting shade-tolerant-dominant forest types has increased (Barbour et al. 1993, 2007; Fites-Kaufman 1997; Sugihara et al. 2006; Vankat and Major 1978).

Fites-Kaufman (1997) developed environmental models of presettlement forest types in two watersheds on the western slope of the central assessment area. She concluded that 64 percent of the areas modeled would be dominated by Douglas-fir and white fir forest types in the absence of fire (i.e., on cool and moderate slopes, watered draws on warm slopes, and higher elevations), but 28 percent of that area was in intermediate areas likely historically dominated by ponderosa and sugar pine because of frequent fire. Overall, according to Fites-Kaufman (1997), about two-thirds of the area modeled was probably dominated by pine-dominant forest before Euro-American settlement.

Dolanc et al. (2014a), compared the vegetation types represented by 4,371 VTM forest plots (Wieslander 1935) from the 1930s and 1,000 FIA forest plots from the 2000s in the central assessment area. They found that 19.9 percent of the 1930s plots classified as ponderosa pine, versus 8.9 percent of the plots from the 2000s; 27.3 percent of the plots were classified as mixed conifer in the 1930s dataset, versus 37.1 percent in the 2000s dataset. Both east-side and west-side Jeffrey pine also declined as a proportion of the sampled vegetation between the 1930s and 2000s.
Dolanc et al. (2014a) suggested that much of the ponderosa pine sampled in the 1930s and subsequently lost probably succeeded to mixed-conifer (through infilling of fir, Douglas-fir, and incense cedar) or to montane hardwood forest, through expansion of oaks after disturbance.

Thorne et al. (2008) compared the 1930s Forest Service vegetation maps (Wieslander 1935) of the Placerville Quadrangle, in the center west of the assessment area, with the Forest Service vegetation map from 1996, to compare the extent of different vegetation types in the two periods. The vegetation type with the largest loss of area was ponderosa pine, which declined by 64 percent over the six decades between maps. In their study area, Thorne et al. (2008) documented that ponderosa pine forest had primarily transitioned to forests dominated by hardwoods (where disturbance had reduced or removed the dominant pines) or by Douglas-fir (where lack of disturbance allowed succession of shade-tolerant species). Thorne et al. (2008) did not find much change in their mixed-conifer forest type, but their classification (California Wildlife Habitat Relations types) splits a number of forest types we include in our general definition of YPMC forests.

Current Forest Service vegetation maps can also be compared to Show and Kotok’s (1929) summary of forest cover types in northern California in the late 1920s. The major changes are in yellow pine and mixed conifer. The former comprised 33.7 percent of Show and Kotok’s analysis area in the 1920s (Modoc National Forest south to the Sequoia National Forest, excluding the Inyo National Forest), but only 17 percent of the area in the most recent Forest Service mapping. Mixed-conifer forests covered 19.8 percent of the area in the 1920s, versus about 30 percent today. Other forest types were similar between the two periods.\(^7\)

**Future**—See the following section.

**Forest composition and species diversity**—

**Trees**—A number of excellent summaries of the species composition of assessment-area YPMC forests have been published (e.g., Barbour and Billings 1999, Fites 1993, Fites-Kaufman et al. 2007, Smith 1994, Sugihara et al. 2006, USDA FS 2001). We direct the reader to these and other sources for information on modern forest conditions. In this section, we both qualitatively and quantitatively describe the patterns of tree composition that characterized YPMC forests before significant Euro-American impact.

\(^7\) Data provided by Jay Miller, remote sensing specialist, USDA Forest Service, Pacific Southwest Region, 3237 Peacekeeper Way, Suite 101, McClellan, CA 95652.
NRV—The U.S. Geological Survey (USGS) carried out general surveys of Sierra Nevada forests at the turn of the 19th century, primarily to assess the status of the timber resource. Leiberg (1902) surveyed forest lands on what is today the Plumas and Tahoe National Forests and the northern Lake Tahoe basin. Sudworth (1900) surveyed the southern Lake Tahoe basin, the Eldorado National Forest, and northern portions of the Toiyabe and Stanislaus National Forests. Fitch (1900) and Marshall (1900) provided brief descriptions of forest conditions in and around Yosemite National Park. McKelvey and Johnston (1992) evaluated the USGS reports and provided a summary of their results. Stephens and Elliot-Fisk (1998) and Stephens (2000) summarized unpublished plot data from Sudworth that were entered into his field notebooks, including data from the southern Sierra Nevada in what is today Sequoia & Kings Canyon National Parks and the Sierra and Sequoia National Forests.

McKelvey and Johnston (1992) note that the composition of forest trees in these early surveys included all the same species we encounter today, but the order of their dominance in assessment area forest stands has changed. Leiberg (1902) divided his survey area into three predominant forest types: his “yellow pine” forest type includes what we call today “mixed conifer” and corresponds to the YPMC forest types covered in this chapter. According to Leiberg, yellow pine was the “most conspicuous and important” species in this forest type, but although it may have once been the dominant species of tree, owing to heavy selective logging throughout his survey area, it was no longer dominant by the turn of the end of the 19th century. Sugar pine was not a dominant species in YPMC forests, but like the yellow pine species, it was selectively harvested and its densities and overall volume had decreased greatly by the time the USGS surveys were made. Leiberg’s (1902) opinions are qualitatively supported by the forest observations of Joseph Bruff, who walked the same northern Sierra Nevada forests a half century earlier. Bruff occasionally made journal notes of forest structure and composition in his 1849–1851 travels, and almost all his observations of conifer trees refer to forests dominated by pines (Read and Gaines 1949).

Sudworth (1900) stated that pine species dominated lower elevation YPMC forests, but incense cedar and fir were mixed with the pine in approximately equal proportions at higher elevations. Sudworth (1900) rarely referred to hardwood species, but he did note that “a few small, unimportant broad-leaved trees” were common along canyon-bottom streams and also grew at times in with the upland coniferous forest. According to Sudworth, yellow pine was the most abundant tree in the YPMC belt, followed by white fir, then incense cedar, sugar pine, and Jeffrey pine. The amount of yellow pine varied considerably from place to place and along
the altitudinal gradient, it was most common on south, west, and east aspects, with some sites containing 80 to 90 percent yellow pine. Most forest stands included different mixes of the above species, with Sudworth estimating the typical ranges as being 45 to 50 percent yellow pine, 30 to 45 percent white fir, 20 to 30 percent incense cedar, 5 to 20 percent sugar pine, and 0 to 5 percent Jeffrey pine on the west slope (but a local dominant on the east slope); Douglas-fir, 2 to 5 percent; black oak, 5 to 10 percent of stands at lower elevations. Black oak was a close associate of yellow pine and incense cedar and could account for more than 50 percent of a stand in dry, thinly soiled locations on south and west slopes.

Sudworth and the other USGS surveyors were visiting forests that already had nearly a half century of Euro-American presence. Many of the stands they surveyed had been cut, usually selectively for pine. This had an impact on their estimates of species importance. For example, Sudworth (1900) noted that the best sugar pine had already been removed from much of the forest he surveyed, and the same had happened with the larger specimens of Douglas-fir. Leiberg (1902) described the effects of heavy logging on forests in the Lake Tahoe basin and noted that sugar pine, which had comprised 20 to 25 percent of the virgin stand, was likely to contribute only 2 to 3 percent of the secondary forest. Greeley (1907) lamented that many stands had been so thoroughly high-graded (valuable species and large trees selectively cut) that yellow pine averaged only 15 to 20 percent of the trees in much of the western Sierra. Leiberg and other early surveyors also noted that the contribution of incense cedar to the stands they visited was artificially high, as it had no commercial value and was left standing far more often than the other species.

Leiberg (1902) estimated that YPMC forests in his survey area at the time of his work were on average 25 percent yellow pine, 1 percent sugar pine, 25 percent Douglas-fir, 44 percent white fir, and 5 percent incense cedar. The relative proportions of the pines and fir had been markedly changed by logging, which focused heavily on the former. At the time of the USGS surveys, Leiberg’s survey area had been (much) more heavily cut than the area assessed by Sudworth. (Miller and Safford 2017).

Leiberg (1902) also estimated the standing volume of trees species in as yet uncut forests. Proportionally, yellow pine species (both ponderosa and Jeffrey) were dominant and comprised about twice the volume of Douglas-fir, sugar pine, or white fir (see McKelvey and Johnston 1992); incense cedar had no economic value at the time and was not listed. White fir regenerated strongly on formerly pine-dominant lands, and Leiberg estimated that it would amount to 60 to 75 percent of the secondary forest, up from 25 to 40 percent of the uncut stands. Sudworth (1900) also referred to the “thickets of seedlings and saplings” of white fir that covered many acres and often excluded all other species. Other observers made similar observations.
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Volume estimates from the Plumas National Forest in 1913 are very similar to Leiberg’s earlier volume estimates for the northern Sierra Nevada as a whole. As above, yellow pine species dominated, with 1.6 times the volume of white fir, 2.4 times the volume of sugar pine, and 2.8 times the Douglas-fir volume; other species like red fir and incense cedar comprised the remaining 7 percent of volume (McKelvey and Johnston 1992, Moore 1913).

Greeley (1907) described the forests of the western Sierra Nevada from the viewpoint of a traditional forester. Like the other early observers, he noted that the original forest had varied from pure yellow pine at the lower elevations and on dry exposures to nearly pure fir toward the upper limit of merchantable timber. Greeley outlined “three bad features of the virgin stand (that) quickly impress the forester.” Two of these three features pertain to structure (“the widespread over-maturity of the timber,” and “the large area on which the stand is open or has disappeared entirely”) and are discussed above. The third feature was the “large and apparently increasing proportion of inferior species.” Except at lower elevations, cutting had been sufficient to reduce the pines to where white fir comprised a third of merchantable timber, and it was a much higher proportion of the younger growth. Together with incense cedar, white fir outnumbered both of the pines in small size classes. Both white fir and incense cedar produce high volumes of seed, and their seedlings are much more tolerant of shade than the pines. Greeley (1907) noted that except at lower elevations and on very warm exposures, dense thickets of fir and cedar were already crowding the pine to “one-fifth or less of reproduction.”

Early GLO land surveying of the assessment area produced useful data on the composition of late 19th century forests. General Land Office crews sampled up to four trees at each section corner and two trees at quarter-section corners (see above). Because of the gridded sampling scheme, GLO samples of forest structure and composition are relatively robust, statistically speaking, although they did not restrict their activities to uncut ground, and they tended to prefer smaller mature trees when they were available, as they were assumed to have the greatest longevity (Hyde 2002). The very low density of GLO data (a maximum of eight points per square mile) means that spatial interpretation of these data is nearly impossible (which is not to say that people have not tried) (e.g., Baker 2012, 2014), but they are very valuable as tabular summaries from larger landscapes. GLO tree data have been summarized for the Lake Tahoe basin (Manley et al. 2000), the Eldorado National Forest (Fites-Kaufman 1997), and for the Stanislaus, Sierra, and Sequoia National Forests (Hyde 2002).

GLO data from Lake Tahoe suggest that yellow pine and fir were present in nearly equal numbers in lower montane forests, with the balance shifting from fir on the wetter west shore to pine on the drier east shore (Manley et al. 2000). Many
of the GLO surveys in the basin were conducted after the mid-1870s, however, and in the lower montane zone their section lines, especially on the east shore, crossed stands where selective logging of Jeffrey and sugar pine had already commenced (Lindström 2000). In addition, the data summarized in Manley et al. (2000) are missing many survey points. It seems likely that the GLO counts in the Lake Tahoe basin in lower montane forests may underestimate the presence of pine before logging, at least on the east shore. Data from Taylor’s studies in the basin support this viewpoint (Taylor 2004, Taylor et al. 2014). For example, the modern Jeffrey pine stands sampled by Taylor and colleagues on the east shore of the lake support an order of magnitude more Jeffrey pine than white fir, even a century after the institution of fire suppression and 130 years or more after the removal of all merchantable Jeffrey pine. Taylor et al.’s (2014) results from west shore mixed-conifer forest agree more closely with the GLO data, with white fir in the 1873 forest outnumbering Jeffrey pine by 1.5:1 (versus 5:4 in the GLO surveys). Almost half of all the white fir measured were smaller than 25 cm d.b.h., and Jeffrey pine was the dominant tree in the larger size classes (Taylor et al. 2014).

Hyde (2002) summarized compositional patterns in GLO data sampled in the middle and late 1800s from areas on the Stanislaus (37 300 ha sampled), Sierra (46 600 ha), and Sequoia National Forests (18 600 ha). Her study areas were chosen to represent elevational transects of approximately 2000 m, along a latitudinal gradient from the central to southern Sierra Nevada. Below 1000 m elevation, forest vegetation was dominated by oak species (more than three-fourths of stems sampled), with only about 8.5 percent of the trees measured being pines (although these contributed about one-fourth of the basal area). Between 1000- and 1500-m elevation, pine (45 percent of sampled stems) and oak (37 percent of sampled stems, mostly black oak) shared in dominance, with fir species contributing 8 percent of stems. Pines comprised about 60 percent of the basal area, and oaks about 24 percent (Hyde 2002). Between 1500- and 2000-m elevation, pines (mostly yellow pines and sugar pine) accounted for 58 percent of stems, fir about 26 percent, and oaks about 11 percent. Basal area was dominated by pines (49 percent), giant sequoia (29 percent), and white fir (16 percent). Above 2000 m, the fir component increased to 42 percent of stems sampled, with pines comprising about 57 percent; yellow pine, lodgepole pine, and sugar pine were the dominant pine species, in that order. Fir (probably mostly red fir at these elevations) contributed slightly more basal area to the stand than the pines (Hyde 2002). Pines dominated slopes of all aspects except northeast, which was dominated by oak species; fir species were surprisingly constant in their contribution to stem densities, comprising in all cases about 20 percent of stems (Hyde 2002).
Baker (2014) reported results from four large areas of GLO surveys in the northern and southern Sierra Nevada (total area sampled equals about 330,000 ha). Baker does not report results by species, but does report percentage of stems encountered by GLO surveyors that were pines, fir plus incense cedar plus Douglas-fir (“shade tolerant”), and oaks. In his northern sites, the proportions of stems in these three classes were 30:38:29; in the southern Sierra Nevada the proportions were 46:28:22.

Fites-Kaufman (1997) summarized the GLO data for two watersheds on the Eldorado National Forest. She found that GLO sites recorded “fir” (Douglas-fir or white fir) on only 26 percent of the surveyed sites. Although Fites-Kaufman (1997) did not report the overall results of the GLO data, her figure 2.11 gives the proportion of stems by species for GLO sites that contained “fir” and for sites that her modeling had identified as being within the Douglas-fir/mixed-conifer potential forest type (i.e., on cool slopes, higher elevations, etc.). The pines (ponderosa and sugar) accounted for 24 and 31 percent of the stems in the two subsamples, respectively; black oak accounted for 26 and 30 percent in the two subsamples. These fire-tolerant species accounted for much more of the stands sampled by the GLO across the rest of the landscape, but Fites-Kaufman (1997) did not provide a summary of those data.

In the VTM project, the Forest Service inventoried and mapped vegetation on much of the land under its jurisdiction in the early 1930s (Wieslander 1935). In the assessment area, thousands of 0.2-ac (809-m²) vegetation plots were sampled in YPMC forest types (ponderosa pine, mixed-conifer, and east-side Jeffrey pine forests). Figure 22 gives the relative densities of tree species in YPMC forests sampled by the VTM project in forest stands that had not experienced intensive logging (n = 2250) and compares them to the most recent available FIA inventory of assessment area forests (n = 510) (see Dolanc et al. 2014b). The VTM protocol sampled only trees 10 cm d.b.h. and above, so the relative densities of species sampled are biased away from the youngest cohorts of trees, which were dominated by white fir and incense cedar that were beginning to benefit from 2 to 3 decades of fire exclusion. Because of this, relative tree densities represented in the VTM dataset provide an at least approximate idea of dominance patterns among tree species before fire suppression (with the caveat that large- and medium-size sugar pine and yellow pine had been selectively logged through much of the assessment area by the time of VTM sampling). The values given in figure 22 are overall averages, and obviously patterns varied from west-side to east-side forests (e.g., Jeffrey pine strongly dominated east-side stands, and Douglas-fir and black oak...
were much more common on the west side), from north to south slopes, and so on. The most obvious pattern in figure 22 is the strong shift in overall dominance from shade-intolerant/fire-tolerant species (ponderosa pine, Jeffrey pine, sugar pine, black oak) in the VTM dataset to shade-tolerant/fire-intolerant species (white and red fir, incense cedar, Douglas-fir) in the FIA dataset. The relative proportions of shade-intolerant versus shade-tolerant species change from 60:40 in the VTM data to 35:55 in the FIA dataset (fig. 22) (Dolanc et al. 2014b).

Figure 23 shows historical stem densities by species for seven sites in the assessment area in which stand reconstruction studies were carried out. Ratios of shade-intolerant to shade-tolerant species range from 92:7 to 16:84 (fig. 23). Collins et al. (2015) analyzed forest inventory data from 1911 in the Tuolumne River drainage at the western border of Yosemite National Park and found that basal areas were divided 58:42 among shade-intolerant and shade-tolerant species (they did not report stem densities). Stephens et al. (2015) conducted an analysis of a similar

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Figure 22—Relative densities of tree species in yellow pine–mixed-conifer forests, comparing vegetation type mapping (VTM) (early 1930s) and Forest Inventory Analysis (FIA) (early 2000s) datasets in the central assessment area (Plumas, Tahoe, Eldorado, Humboldt-Toiyabe, Eldorado, Stanislaus, and Sierra National Forests; Lake Tahoe Basin Management Unit; and Yosemite National Park—VTM data were not collected north or south of this area). Only trees >10 cm diameter at breast height are included. ABCO = *Abies concolor* (white fir); ABMA = *A. magnifica* (red fir); CADE = *Calocedrus decurrens* (incense cedar); PIJE = *Pinus jeffreyi* (Jeffrey pine); PILA = *P. lambertiana* (sugar pine); PIPO = *P. ponderosa* (ponderosa pine); PSME = *Pseudotsuga menziesii* (Douglas-fir); QUKE = *Quercus kelloggii* (black oak); live oaks = canyon live oak and interior live oak. Data are from Dolanc et al. (2014b).
Figure 23—Reconstructions of historical stem densities at seven sites in yellow pine–mixed-conifer forest in the assessment area. Sites are arranged in order of proportional dominance by shade-intolerant species, from high (left) to low (right). (A) absolute densities; (B) relative densities (percentage of stems). 1 = Parsons and Debenedetti (1979) (yellow pine, ≥12 cm diameter at breast height [d.b.h.]); 2 = Taylor (2004) (≥10 cm d.b.h.); 3 = USDA FS (1911) (≥15.2 cm d.b.h.); 4 = North et al. (2007) (≥5 cm d.b.h.); 5 = Scholl and Taylor (2010) (≥10 cm d.b.h.); 6 = Taylor (2014) (≥5 cm d.b.h.); 7 = Parsons and Debenedetti (1979) (mixed conifer). ABCO = Abies concolor (white fir); ABMA = A. magnifica (red fir); CADE = Calocedrus decurrens (incense cedar); PIPO = Pinus lambertiana (sugar pine); PIJE = P. jeffreyi (Jeffrey pine); PIPO = P. ponderosa (ponderosa pine); PSME = Pseudotsuga menziesii (Douglas-fir); QUKE = Quercus kelloggii (black oak).
1911 dataset from the Greenhorn Mountains in the southwestern Sierra Nevada and found that basal areas were split 46:54 between shade-intolerant and shade-tolerant species. It can be appreciated that the relative densities of shade-intolerant and shade-tolerant species were not constant on the landscape, and in some circumstances yellow pines, which dominated much of the general landscape, were entirely absent.

In another forest reconstruction study based on “aggregations” of vegetation, Bonnicksen and Stone (1982) found that a YPMC watershed in Kings Canyon National Park had seen an increase in the area of the watershed covered by white fir-dominant aggregations from 27 to 37 percent between 1890 and 1977. Black oak was reduced from 10 percent of the study area to 6 percent during the same period.

Kercher and Axelrod (1984) developed a Monte Carlo-based model of YPMC forest succession (known as “SILVA”) in the Sierra Nevada. This model is further discussed in “Successional processes” above. Kercher and Axelrod (1984) used SILVA to compare forest succession after a simulated clearcut for 500 years at two different elevations, 1520 m and 1830 m. The lower elevation site is at the upper reaches of YPMC forests historically dominated by ponderosa pine; the upper site is nearer the upper limits of YPMC forests in the Sierra Nevada and historically included a significant component of fir species. The time-averaged results of the SILVA lower elevation simulation under frequent fire (mean FRI = 7 years) are given in figure 11. Ponderosa pine strongly dominated the modeled stand throughout the 500-year simulation. The relative proportions of shade-intolerant versus shade-tolerant species in figure 11 are about 66:33. The SILVA model for the higher elevation site projected a forest stand dominated strongly by white fir (which comprised an average of 60 percent of the total basal area over the 500-year simulation), even under frequent fire (Kercher and Axelrod 1984).

In summary, presettlement YPMC forests in the assessment area tended to be dominated by yellow pine and other shade-intolerant species, but shade-tolerant species could be locally dominant, especially at higher elevations, at northern latitudes, and in moist microsites. Most assessments of species densities were carried out after selective logging of yellow pine and sugar pine had already occurred. The actual proportions of shade-tolerant to shade-intolerant species varied across the presettlement landscape, but the data suggest that among mature trees, shade-intolerant species (pine and black oak) typically comprised most of the individuals in the average stand, especially the larger individuals (see below). Then, as now, in the absence of fire, shade-tolerant species dominated the smallest size classes.
Comparison with current—Yellow pine and sugar pine have notably decreased in importance in assessment-area YPMC forests, while shade-tolerant conifers and hardwoods (white fir, incense cedar, Douglas-fir, tanoak, canyon live oak) have increased (fig. 24). This is due to selective logging, increased resource competition under increasing stand densities, the difficulty of regenerating under the denser modern canopy, and—in the case of sugar pine—to the effects of white pine blister rust (Cronartium ribicola), an introduced fungal pathogen that is having major effects on five-needled pine populations across the Western United States (Agee 1993; Barbour et al. 1993, 2007; Sugihara et al. 2006; van Mantgem et al. 2004). FIA data show that the component of shade-intolerant species in assessment-area YPMC forests has dropped from an average of maybe two-thirds of the mature forest stand to around one-third of mature trees. The largest decrease has been in ponderosa pine, but all shade-intolerant species have decreased in density.

Geographic, topographic, and successional patterns in tree species composition are driven by ecological differences between species. Table 2 lists the tolerances of common YPMC tree species in the assessment area to shade, frost, temperature, drought, and fire. The dominance of the yellow pine species on south and west aspects, on droughtier soils, and in fire-prone locations is easy to predict from the information in table 2, as is the heightened presence of white fir on north slopes, and in moist and protected topographic locations; incense cedar is somewhat intermediate in most characteristics. White fir is the major successional climax species for most of the YPMC belt in the assessment area, although Douglas-fir plays this role at lower elevations. White fir’s shade tolerance and fire intolerance, its ability to survive for very long periods as a suppressed tree, its capacity to respond rapidly to release (increased light availability), and its prolific production of seed are dominant factors in forest succession in the assessment area (Barbour et al. 2007, Burns and Honkala 1990, USDA FS 2013b). In prefire suppression times, very frequent fire limited white fir populations in most of the assessment area, but it could dominate stands in areas where conditions existed (such as high fuel moisture) such that fire frequency was reduced, or where fire intervals were long enough or variable enough to permit adults to develop thick bark. Increasing annual precipitation and lower summer temperatures along the south-to-north axis of the assessment area also result in higher fir densities to the north. The same gradient occurs with elevation, and many early observers noted how YPMC forests were dominated by white fir at higher elevations. With the onset of fire suppression, major increases in white fir
and incense cedar began across the assessment area (Vankat and Major 1978). Early observers noted this tendency already in the late 19th and early 20th centuries (Greeley 1907, Leiberg 1902, Sudworth 1900). Today, with the general absence of fire, climate is the principal factor regulating species distributions, and species requiring frequent disturbance to retain their place in the forest canopy—principally species of pine—are in the decline, except in areas of low site productivity such as areas of thin or nutrient-poor soil or extreme microclimates.

Dolanc et al. (2014b) compared forest conditions as documented in the Forest Service forest mapping inventory of the early 1930s (Wieslander 1935) with the most recent compilation of the FIA plot network in the assessment area. Figure 24, using data from table 2 in Dolanc et al. (2014b), is a graphic summary of the relative changes in tree composition that have occurred in the assessment area.

![Figure 24](image_url)

Figure 24—Relative change in species frequencies (measured as the percentage of plots in which a given species is found) between the 1930s Forest Service vegetation mapping inventory (VTM) and the most recent compilation of Forest Inventory and Analysis (FIA) data (USDA FS 2013a). Species below the x-axis have dropped in frequency. Codes are from the first two letters of the genus plus the first two letters of the species. PISA = *Pinus sabiniana* (California foothill pine); QUWI = *Quercus wislizenii* (interior live oak); ARME = *Arbutus menziesii* (Pacific madrone); QUCH = *Q. chrysolepis* (canyon live oak); LIDE = *Lithocarpus densiflorus* (tanoak); QUKE = *Q. kelloggii* (black oak); PSME = *Pseudotsuga menziesii* (Douglas-fir); PIPO = *P. ponderosa* (ponderosa pine); CADE = *Calocedrus decurrens* (incense cedar); PILA = *P. lambertiana* (sugar pine); ABMA = *Abies magnifica* (red fir); PICO = *P. contorta* (lodgepole pine); PIMO = *P. monticola* (western white pine); TSME = *Tsuga mertensiana* (mountain hemlock); PIAL = *P. albicaulis* (whitebark pine). Illustration adapted from data in Dolanc et al. (2014b).
since the arrival of Euro-Americans. The YPMC belt begins where QUKE (black oak) is located on the x-axis and continues to PIJE (Jeffrey pine). The most obvious pattern in figure 24 is the reduced frequency (calculated as the percentage of plots in which the species in question is found) of shade-intolerant/fire-tolerant species like ponderosa (PIPO), sugar (PILA), and Jeffrey pine; and the increased frequency of shade-tolerant/fire-intolerant species like Douglas-fir (PSME), incense cedar (CADE), and white fir (ABCO). Changes have been driven primarily by fire suppression and timber harvest activities.

**Future—**

Miller and Urban (1999a) added a soil water routine and a fire model to the forest gap model ZELIG and simulated successional dynamics in Sierra Nevada forests along an elevational gradient in Sequoia & Kings Canyon National Parks under a number of future climate scenarios. Simulations were run for 800 years, with the first 200 years having no fire and the last 600 years under a “natural” fire regime for the elevation in question. In all future climate scenarios except the cool-wet scenario, tree species composition at the two lowest elevation sites (1800 m and 2200 m), which are at elevations currently dominated by YPMC forests, changed markedly. Major changes were not apparent in the models until about a century into the simulations, but fire in Miller and Urban’s (1999a) model is entirely surface fire and is internally driven (by fuel load and fuel moisture), not externally forced, and effects of disease or insect outbreak were not modeled, so the results are probably very conservative with respect to the velocity of change. At the lower two elevations, trees either completely or nearly disappeared by the final century of the simulation. Under the less extreme warming and drying scenarios, the forest at 1800 m saw the basal area of tree species change from 15:9:3:1:1 (m²/ha; ponderosa pine:incense cedar:Jeffrey pine:black oak:white fir) to 1:1:1:1:0. Under more extreme warming, trees disappeared at this elevation. At 2200 m, baseline basal areas of 45:4:2:1:0 (m²/ha; white fir:sugar pine:incense cedar:Jeffrey pine:ponderosa pine) were changed to an average of 4:0:12:2.5:1 under the less extreme warming/drying scenarios, and the site transitioned to a yellow pine–black oak–incense cedar woodland (total basal area only 8 m²/ha) under the more extreme scenarios. The 2600 m site, which currently supports a red fir forest, transitioned to a moist mixed-conifer forest dominated by white fir under the less extreme future scenarios, and to a dry mixed-conifer (yellow pine-white fir-incense cedar in almost equal proportions) under the more extreme scenarios (Miller and Urban 1999a).
The Nature Conservancy carried out future climate suitability forecasts for major tree and shrub species of the Sierra Nevada, in support of the northern and southern Sierra Nevada Partnerships. The partnerships are collaborative efforts, organized by The Nature Conservancy, the Sierra Business Council, and a variety of other national, regional, and local conservation organizations, to carry out all-lands conservation planning for important subregions of the Sierra Nevada (see: http://consbio.org/products/projects/southern-sierra-partnership, and http://northernsierrapartnership.org/). Climate suitability forecasts for the period 2045 to 2065 were derived using maximum entropy (MaxEnt) modeling, and based on 11 GCMs run under the IPCC’s A2 climate scenario. Models only incorporated climate variables. Three generalized future climate scenarios were developed (“warm-dry,” “hot-dry,” and “hot-wet”) and an ensemble projection was also developed, where the degree of consensus across models was mapped. In the ensemble outputs, areas of projected climate “stress” (possible loss in distribution), climate refugia, and possible expansion were mapped, along with a measure of the degree of model consensus. More information and the data outputs are available at http://app.databasin.org/app/pages/galleryPage.jsp?id=8c5db744f9fe4d3e9375b100dc695c4d. All the projections for California can be viewed at http://www.tncmaps.org/climate/species/.

Overall, The Nature Conservancy’s climate suitability forecasts suggest more climatic stability and less future climate stress in the southern Sierra Nevada than in the north (SSP 2010). This is due largely to much higher elevations and more accentuated topography in the south, which lead to less probability of overall habitat loss as climates warm and snowpack decreases. Nonetheless, the models suggest notable areas of climate stress along the lower, western margins of the YPMC forest belt, as well as the potential for expansion to higher elevations (fig. 25). Both Jeffrey and ponderosa pine are projected to experience increased climate stress in much of the northern and northeastern assessment areas, primarily because of the lower average elevations in those areas. Douglas-fir (not shown) is projected to experience relatively little climate stress, except along the western margins of the central Sierra Nevada, and climate suitability for the species may increase markedly at higher elevations throughout the central and northern Sierra Nevada. In their summary of the top threats to conservation in mixed-conifer forests, the Southern Sierra Partnership (SSP 2010) identified changed fire regimes as the highest threat, followed by climate change and a number of other factors.
Figure 25—Future climate suitability forecasts for four yellow pine–mixed-conifer tree species for the period 2045 to 2065; ensemble results from 11 GCMs under the A2 IPCC climate scenario. (A) Jeffrey pine; (B) ponderosa pine; (C) white fir; (D) black oak. Red and orange = areas in which climates are currently suitable but will not be in the future (climate “stress”); blue = areas in which current and future climates are suitable (refugia); purple = areas in which climates are not currently suitable but will be in the future (possible expansion). Mapped outputs of MaxEnt species distribution projections are from modeling by The Nature Conservancy. See text for details.
Forest understory and nonforest vegetation

**NRV and comparison to current**—

Early observers did not measure richness of diversity of species other than trees, although they made reference to certain shrubs (mostly those that competed with trees), and sometimes referred in passing to herbaceous species as well. For example, Muir (1894) noted how the open canopy of YPMC forests in the assessment area led to many understory species, especially shrubs and forbs, but the growth was “never so dense… as to prevent the traveler from sauntering where he will” (Muir 1894: chapter 3). A general ecological principle is that the diversity of understory species tends to be negatively correlated with overstory canopy cover, and positively correlated with light availability (Barbour et al. 1987). Presettlement YPMC forests were generally more open than modern forests, and the enhanced incidence of light at the forest floor likely led to higher local species richness in understory plants, but we can only theorize based on numerous modern demonstrations of the negative relationship between forest canopy cover and understory species diversity in forests in and around the assessment area (e.g., Agee and Biswell 1970, Battles et al. 2001, Keeley et al. 2003, Wayman and North 2007, Webster and Halpern 2010). These probable local effects notwithstanding, there is no evidence that we know of that demonstrates any notable change in regional richness (overall species richness across the assessment area) of understory plants in the assessment area since presettlement times.

Although many plant species in California have been demonstrated to require fire scarification of their seeds to induce germination (e.g., through heat, presence of certain volatiles, or chemicals contained in charcoal), such plants are rare in assessment-area YPMC forests (Keeley et al. 2003, 2012; Webster and Halpern 2010). This is probably due to the relative rarity of high-intensity fire over evolutionary time in these forests (Denslow 1980, Grubb 1977). As such, the large decrease in fire frequency and overall area burned in assessment-area YPMC forests since presettlement times has probably not had a highly deleterious effect on regional understory diversity, or at least as deleterious an effect as it might theoretically have in an ecosystem characterized by highly intense fire. Several widely distributed shrub species from the genera *Ceanothus* and *Arctostaphylos* do possess fire-promoted seed germination, and although these are typically shaded out by overstory trees over time, their soil seedbanks may persist for hundreds of years and they are not at regional risk of extinction as a result of fire suppression (Keeley et al. 2012, Knapp et al. 2012).

As with the canopy tree species, understory species composition varies across the assessment area, depending on underlying environmental conditions as well as the influence of the tree canopy layer. It seems probable that understory species
requiring higher light environments may have become less abundant (if not less species rich) in the assessment area as a result of fire exclusion. At the same time, shade-loving species have probably increased in abundance, at least locally. Whether such changes have led to overall changes in understory species diversity is difficult to say in any general fashion; however, there are a few recent studies that provide some insight. Knapp et al. (2013) took advantage of surveys done in 1929 on the Stanislaus National Forest and resampled the same plots in 2007/2008. They did not find evidence that overall species richness had changed between the surveys, but shrub and herb species more tolerant of shade and litter layers were much more abundant in the modern forest (dense, high canopy cover, after long-term fire suppression) than in the original surveys (e.g., *Chrysolepis sempervirens* (Kellogg) Hjelmqvist, *Pyrola picta* Sm., *Symphoricarpos mollis* Nutt., *Apocynum androsaemifolium* L.), while heliophilic plants were less abundant (e.g., *Arctostaphylos patula* Greene, *Ceanothus* spp., *Chamaebatia foliolosa* Benth., *Epilobium*, and *Hieracium* spp.). Stevens et al. (2015) found that these same heliophilic genera, which have biogeographic affinity to California and xeric regions in southern North America, increased in richness with fire severity, from unburned forest to low-severity wildfire to high-severity wildfire. The increase in these southern-xeric species was accompanied by a loss of the “Arcto-Tertiary” geoflora (Raven and Axelrod 1978) following fire, particularly in more productive mixed-conifer forests that previously had higher canopy cover and stand density. Stevens et al. (2015) also found that within-plot (alpha) diversity was higher in burned forest than in unburned forest 3 years after fire, and that between-plot (beta) diversity was higher in stands that had previously been treated for fuels reduction and burned at low to moderate severity, compared with previously untreated stands nearby that generally burned at high severity. Their estimates for stand-scale diversity, combining alpha and beta diversity estimates, indicated a peak in diversity at the intermediate disturbance level created by low- to moderate-severity wildfire burning through fuel treatments. The number of invasive, nonnative plant species was positively related to disturbance and fire severity.

**Future**

The Nature Conservancy forecasted future climate suitability for a number of YPMC forest shrub species using MaxEnt modeling. See “Trees” on page 161 for more detail on the modeling. Outputs are viewable at http://www.tncmaps.org/climate/species/. These models suggest notable loss of suitable climates for important species like Greenleaf manzanita (*Arctostaphylos patula* E. Greene) and mountain whitethorn (*Ceanothus cordulatus* (Hook.) Nutt.) in the northern and northeastern parts of the assessment area. As noted above, these models include only climate variables, and many other factors interact to drive species distributions.
If current trends in fire severity continue (Mallek et al. 2013, Miller and Safford 2012, Miller et al. 2009b), it seems likely that species of more mesic habitats will decrease in abundance and perhaps richness, while xeric species adapted to higher light and warmer conditions will expand (Stevens et al. 2015). Alien species richness is also likely to increase under this scenario.

See “Forest understory and nonforest vegetation” above for more information on understory vegetation.

Summary of Probable Deviations From NRV—and Conclusion

Based on our understanding of YPMC ecosystems in the bioregional assessment area, we have attempted to draw conclusions with regard to whether key ecosystem variables are currently within or outside of the NRV. Table 11 summarizes our conclusions and directs the reader to the areas of this report that discuss the ecosystem elements in question.

We finish by making the following general conclusions:

1. With regard to ecosystem composition of assessment-area YPMC forests, although overall plant species diversity across the assessment area has probably not changed much (except for the addition of nonnative species), there has been a major shift over the past century from dominance by shade-intolerant/fire-tolerant species to dominance by shade-tolerant/fire-intolerant species. This has happened in both the forest overstory and understory.

2. With regard to ecosystem structure, assessment-area YPMC forests are greatly changed from the presettlement period, so much so that people from the 18th or 19th centuries would probably not recognize the modern forest. For example:

   A. Mean adult tree densities are an average of two to four times higher today than during the presettlement period.
   B. Tree seedling densities are similarly much higher in the modern forest, and they are dominated by fire-intolerant/shade-tolerant species.
   C. The average tree in today’s forest is 40 to 50 percent smaller (in d.b.h.) than in the presettlement forest.
   D. Even though there are fewer large trees in today’s forest, the huge number of small trees has resulted in basal areas that are equal to or higher on average than in presettlement forests.
   E. Tree canopy cover averages about 33 percent more today than in the presettlement period.

If current trends in fire severity continue (Mallek et al. 2013, Miller and Safford 2012, Miller et al. 2009b), it seems likely that species of more mesic habitats will decrease in abundance and perhaps richness, while xeric species adapted to higher light and warmer conditions will expand (Stevens et al. 2015). Alien species richness is also likely to increase under this scenario.
Table 11—Summary of probable deviations from the natural range of variation (NRV) for yellow pine and mixed-conifer (YPMC) forests in the assessment area

<table>
<thead>
<tr>
<th>Ecosystem attribute</th>
<th>Indicator group</th>
<th>Indicator</th>
<th>Variable</th>
<th>Within NRV</th>
<th>Confidence</th>
<th>See discussion on page:</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>Disturbance</td>
<td>Fire</td>
<td>Fire regime</td>
<td>No</td>
<td>High</td>
<td>31</td>
<td>Shift from Fire Regime I to Fire Regimes III and IV.</td>
</tr>
<tr>
<td>Function</td>
<td>Disturbance</td>
<td>Fire</td>
<td>Fire frequency</td>
<td>No</td>
<td>High</td>
<td>34</td>
<td>Current frequency far below presettlement but rising.</td>
</tr>
<tr>
<td>Function</td>
<td>Disturbance</td>
<td>Fire</td>
<td>Fire severity</td>
<td>No</td>
<td>Medium to High</td>
<td>38</td>
<td>Current severity higher than presettlement and rising.</td>
</tr>
<tr>
<td>Function</td>
<td>Disturbance</td>
<td>Fire</td>
<td>Fire size</td>
<td>No</td>
<td>Medium</td>
<td>52</td>
<td>For fires &gt;4 ha, current mean and mean maximum fire sizes larger than presettlement mean.</td>
</tr>
<tr>
<td>Function</td>
<td>Disturbance</td>
<td>Fire</td>
<td>High severity patch size</td>
<td>No</td>
<td>Medium</td>
<td>51</td>
<td>Current high-severity patch sizes higher than presettlement mean and rising.</td>
</tr>
<tr>
<td>Function</td>
<td>Disturbance</td>
<td>Fire</td>
<td>Fire rotation</td>
<td>No</td>
<td>High</td>
<td>37</td>
<td>Fire rotations much longer today than presettlement.</td>
</tr>
<tr>
<td>Function</td>
<td>Disturbance</td>
<td>Fire</td>
<td>Fire season</td>
<td>No</td>
<td>Medium</td>
<td>56</td>
<td>Fire season is becoming longer, but general seasonal patterns are similar.</td>
</tr>
<tr>
<td>Function</td>
<td>Disturbance</td>
<td>Fire</td>
<td>Annual area burned</td>
<td>No</td>
<td>High</td>
<td>55</td>
<td>Current mean annual area burned is much lower than all estimates of presettlement area (but it is rising quickly).</td>
</tr>
<tr>
<td>Function</td>
<td>Disturbance</td>
<td>Insect outbreaks</td>
<td>Tree mortality from insects</td>
<td>unknown</td>
<td>69, 84</td>
<td>Little in the way of presettlement estimates. Some conclusions can be drawn from comparisons of range of current conditions.</td>
<td></td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Canopy cover</td>
<td>Percentage of cover</td>
<td>No</td>
<td>Medium</td>
<td>136</td>
<td>Modern mean canopy cover is above presettlement.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Coarse woody debris (CWD)</td>
<td>Pieces of CWD per unit area</td>
<td>No</td>
<td>Medium</td>
<td>141</td>
<td>Density of CWD is higher in contemporary forests.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Coarse woody debris</td>
<td>Mass of CWD per unit area</td>
<td>No</td>
<td>Medium</td>
<td>141</td>
<td>Average tons/ha of CWD is higher in contemporary forests.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Forest fuels</td>
<td>Tons/ha</td>
<td>No</td>
<td>Medium</td>
<td>158</td>
<td>On average, contemporary YPMC forests support much higher fuel loadings than presettlement forests, in both fine-fuel and coarse-fuel classes.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Functional groups/growth forms</td>
<td>Proportion of early/middle/late seral forest</td>
<td>No</td>
<td>Medium</td>
<td>87</td>
<td>Current lack of old-forest successional stages, perhaps some localized lack of early stages.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Gap size distribution</td>
<td>Gap size</td>
<td>No</td>
<td>Medium</td>
<td>139</td>
<td>Gap sizes are generally decreasing (in undisturbed forests), but also increasing in disturbed forests owing to more severe disturbance.</td>
</tr>
<tr>
<td>Ecosystem attribute</td>
<td>Indicator group</td>
<td>Indicator</td>
<td>Variable</td>
<td>Within NRV&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Confidence</td>
<td>See discussion on page:</td>
<td>Notes</td>
</tr>
<tr>
<td>---------------------</td>
<td>----------------</td>
<td>----------</td>
<td>----------</td>
<td>------------------------</td>
<td>------------</td>
<td>-------------------------</td>
<td>-------</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Grass and forb cover</td>
<td>Percentage of cover</td>
<td>Maybe</td>
<td>Low</td>
<td>153</td>
<td>Difficult to assess, little presettlement data. Overall herbaceous cover on landscape may be similar; cover within forest stands may be lower owing to fire suppression.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Overstory density</td>
<td>Number of trees per unit area</td>
<td>No</td>
<td>High</td>
<td>97</td>
<td>Current density higher on average than presettlement.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Overstory density</td>
<td>Number of large trees per unit area</td>
<td>No</td>
<td>High</td>
<td>97</td>
<td>Large tree density is lower in modern forests.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Shrub cover</td>
<td>Percentage of cover</td>
<td>Maybe</td>
<td>Low</td>
<td>148</td>
<td>Difficult to assess, little presettlement data. Overall shrub cover on landscape not much changed over time; cover within forest stands may be lower owing to fire suppression.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Snag density</td>
<td>Number of snags per unit area</td>
<td>No</td>
<td>Medium</td>
<td>141</td>
<td>Snag density is higher in contemporary forests.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Tree size class distribution</td>
<td>Tree size-class distribution</td>
<td>No</td>
<td>High</td>
<td>99</td>
<td>Major increases in small size classes, and general decreases in large size classes. Change in distribution shape from ± flat, hump-shaped, or weakly J-shaped in average pre-settlement forest to strongly J-shaped in average modern forest.</td>
</tr>
<tr>
<td>Structure</td>
<td>Physiognomy</td>
<td>Average tree size</td>
<td>Mean d.b.h. or quadratic mean diameter</td>
<td>No</td>
<td>High</td>
<td>99</td>
<td>Average conifer tree in modern YPMC forests about one-half the diameter of the average tree in pre-settlement forests.</td>
</tr>
<tr>
<td>Structure</td>
<td>Productivity</td>
<td>Tree basal area</td>
<td>Basal area</td>
<td>Yes</td>
<td>Medium</td>
<td>107</td>
<td>Basal area similar if a bit higher in modern forests; major difference is distribution of more biomass in small and medium trees in contemporary forest than in pre-settlement forest.</td>
</tr>
<tr>
<td>Composition</td>
<td>Functional diversity</td>
<td>Functional groups/growth forms</td>
<td>Proportion of shade tolerant vs. shade intolerant species</td>
<td>No</td>
<td>High</td>
<td>160</td>
<td>Major shift from dominance of shade-intolerant species to dominance of shade-tolerant species.</td>
</tr>
<tr>
<td>Composition</td>
<td>Species diversity</td>
<td>Species richness</td>
<td>Plant species richness</td>
<td>Yes</td>
<td>Medium</td>
<td>161</td>
<td>No evidence of major species loss.</td>
</tr>
</tbody>
</table>

<sup>d.b.h. = diameter at breast height. TBD = to be determined.</sup>

<sup>a</sup> NRV pre-settlement reference period is assumed to refer to 1500/1600 to 1850, unless otherwise indicated in notes. NRV for most indicators/variables also includes information from contemporary reference sites.

<sup>b</sup> As defined as the range of means from multiple sources.
F. Forest structure has been greatly homogenized, with the size and number of forest gaps decreasing almost to zero in many modern forest stands. In presettlement forests, many areas supported more canopy gaps than canopy.

G. Contrary to what many people think, an objective assessment of the evidence suggests that snag densities and coarse woody debris are not depleted in modern forests, and indeed most of the evidence suggests they are more abundant today than in the average presettlement forest stand.

H. Coarse woody debris is also a component of forest fuels, and modern fuel loadings are much higher today than in the presettlement forest. Our estimate is that fuel loadings in assessment-area YPMC forests have risen by an average of 70 to 100 percent over the past century or so.

I. Shrub cover in modern YPMC forests is probably not very different from presettlement conditions (maybe slightly lower today), but the distribution of shrub cover certainly is. Modern forests are more likely to support large areas of contiguous shrub fields but relatively low shrub cover within forest stands (owing to higher stand canopy cover today), whereas presettlement forests supported higher cover of shrubs within stands, as light incidence at the soil surface was much higher.

3. With regard to ecosystem function, the major change in YPMC forests has been in the role and behavior of fire. Specifically:

A. Fires have gone from representing a frequently recurring disturbance on the landscape (5 to 10 events per century on average) to an extremely rare event (75 percent of all YPMC forest has not seen a fire in the past 100+ years).

B. The average area of fire in the assessment area between 1984 and 2010 was only about 10 to 15 percent of the presettlement mean (±150 000 ha per year), but the past 6 years have seen much more area burn, with large areas experiencing nearly complete tree mortality.

C. When fire occurs today, it behaves very differently on average than in the presettlement forest because of differences in forest structure and fuels, and as a result of changing climate. The proportional area of fires burning at high severity today (severity is a measure of mortality caused by fire or biomass lost to fire) is 5 to 10 times greater than in the average presettlement period fire.
4. As such, the role of fire has changed from one of forest maintenance (of relatively open-canopy, low-fuel-accumulation conditions with dominance primarily by fire-tolerant species) to one of forest transformation, where dense stands of fire-intolerant species and heavy fuel accumulations are more likely to burn at high severity, resulting in major ecosystem changes.

For decades, the major ecological issue in the assessment area was thought to be the loss of dense-canopied, old-growth forest to logging, and threats to wildlife species that depend on such conditions (Duane 1999, Ruth 1996). Today the major threat is clearly the loss of forest—old growth or not—to severe wildfire and insect and direct drought mortality (Keeley and Safford 2016, McKenzie et al. 2004). The irony is that a primary cause of this major threat is the historical widespread focus on fire suppression, which was viewed as a necessary means to prevent forest loss. In light of new scientific information, such as the information presented in this report, these management views have been changing, and we are at a pivotal point in resource management in the assessment area. Current trends in climate, fire, human land use, economics, and federal budgets are not auspicious, but recent collaborative management efforts at large landscape scales, political developments in California, and more progressive national forest planning suggest that there is a broadening understanding of the necessary ecological role of fire in the Sierra Nevada bioregion. We hope that this assessment of past and current conditions in the broader Sierra Nevada bioregion will add to this growing understanding and support effective management that can conserve California’s “Range of Light.”

Acknowledgments

We thank three anonymous reviewers and our editor for their constructive comments and sapient criticism of our submission draft. We also thank Ramona Butz (Six Rivers National Forest), Jonathan Long (Pacific Southwest Research Station), Marc Meyer (Sierra National Forest), Jay Miller (Pacific Southwest Region Fire and Aviation Management), Malcolm North (Pacific Southwest Research Station and University of California–Davis), Carl Skinner (Pacific Southwest Research Station), and Scott Stephens (University of California–Berkeley) for reviewing various early drafts of this NRV assessment. Chris Dolanc (University of California–Davis) and Jay Miller also provided datasets and carried out some analyses. Deb Beardsley (Forest Service Remote Sensing Laboratory) compiled the most recent FIA inventories for the assessment area and provided summaries of the data.
English Equivalents

<table>
<thead>
<tr>
<th>When you have:</th>
<th>Multiply by:</th>
<th>To get:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Millimeters (mm)</td>
<td>0.0394</td>
<td>Inches</td>
</tr>
<tr>
<td>Centimeters (cm)</td>
<td>0.394</td>
<td>Inches</td>
</tr>
<tr>
<td>Meters (m)</td>
<td>3.28</td>
<td>Feet</td>
</tr>
<tr>
<td>Kilometers (km)</td>
<td>0.62</td>
<td>Miles</td>
</tr>
<tr>
<td>Hectares (ha)</td>
<td>2.47</td>
<td>Acres</td>
</tr>
<tr>
<td>Square kilometers (km²)</td>
<td>0.386</td>
<td>Square miles</td>
</tr>
<tr>
<td>Square meters per hectare (m²/ha)</td>
<td>4.357</td>
<td>Square feet per acre</td>
</tr>
<tr>
<td>Degrees Celsius (°C)</td>
<td>1.8 °C + 32</td>
<td>Degrees Fahrenheit</td>
</tr>
</tbody>
</table>

Metric Equivalents

<table>
<thead>
<tr>
<th>When you have:</th>
<th>Multiply by:</th>
<th>To get:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inches (in)</td>
<td>2.54</td>
<td>Centimeters</td>
</tr>
<tr>
<td>Feet (ft)</td>
<td>0.305</td>
<td>Meters</td>
</tr>
<tr>
<td>Miles (mi)</td>
<td>1.609</td>
<td>Kilometers</td>
</tr>
<tr>
<td>Acres (ac)</td>
<td>0.405</td>
<td>Hectares</td>
</tr>
<tr>
<td>Square miles (mi²)</td>
<td>2.59</td>
<td>Square kilometers</td>
</tr>
<tr>
<td>Square feet per acre (ft²/ac)</td>
<td>0.229</td>
<td>Square meters per hectare</td>
</tr>
</tbody>
</table>

Literature Cited


Barth, M. 2014. Use of a forest reconstruction model to assess changes to Sierra Nevada mixed-conifer forest conditions during the fire suppression era. Missoula, MT: University of Montana. 170 p. M.S. thesis.


Dunbar-Irwin, M.; Safford, H.D. 2016. Climatic and structural comparison of yellow pine and mixed-conifer forests in northern Baja California (México) and the eastern Sierra Nevada (California, USA). Forest Ecology and Management. 363: 252–266.


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Appendix: Pacific Southwest Region Natural Range of Variation Assessments for Forest Planning:
A Summary of the Process and the Products

Hugh Safford, regional ecologist, U.S. Department of Agriculture, Forest Service,
Pacific Southwest Region, hughsa@fs.fed.us, 707-562-8934, May 12, 2013

According to the Forest Service 2012 Planning Rule (36 CFR 219):

Assessments rapidly evaluate existing information about relevant ecological, economic, and social conditions, trends, and sustainability and their relationship to the land management plan within the context of the broader landscape. The responsible official shall consider and evaluate existing and possible future conditions and trends of the plan area, and assess the sustainability of social, economic, and ecological systems within the plan area, in the context of the broader landscape.

The Planning Rule identifies 15 topic areas that must be addressed in the assessment. Among these are (1) terrestrial ecosystems (and aquatic ecosystems and watersheds), and (2) system drivers, including dominant ecological processes, disturbance regimes, and stressors, such as natural succession, wildland fire, invasive species, and climate change; and (3) the ability of terrestrial and aquatic ecosystems on the plan area to adapt to change.

FSH 1909.12 Chapter 10, Section 12.11

Requires that “the responsible official should identify and evaluate the ecological integrity of... ecosystems within the plan area.”

Steps include:

1. Identify relevant ecosystems to be evaluated and the appropriate scale for the assessment

2. Rapidly evaluate available information about those ecosystems, including composition, structure, and function, by:
   A. Selecting key ecosystem characteristics for the evaluated ecosystems that will permit evaluation of ecological integrity (sustainability)
   B. Describe the natural range of variation (NRV) for key ecosystem characteristics, when such information is readily available
   C. (describes an alternative to NRV assessment)
   D. Describe current condition and trends of the key ecosystem characteristics

3. Identify and evaluate system drivers and stressors
4. Describe the projected future status of ecosystem integrity, using the key ecosystem characteristics, by:
   A. Describing the status of the key ecosystem characteristics by comparing NRV to current conditions, or
   B. (describes alternative where NRV assessment is not available)

5. Identify status of key ecosystem characteristics and determine whether they are “functioning in a way that contributes to ecosystem integrity and sustainability.”

Natural Range of Variation Assessments

These were carried out by Pacific Southwest Region Ecology Program. Each ecologist was assigned one or two chapters. Work began October 2012, and was completed in May 2013 (with some exceptions). Updates were made in 2015 and 2016.

Detailed steps:

1. Identify ecosystems to be addressed
   A. NRV assessments are focused on terrestrial ecosystems
   B. We used Barbour and Billings (2000) Sugihara et al. (2006), Barbour et al. (2007), MCV, Van de Water and Safford (2011), and California WHR classification, and then requested input from about 25 academic, NGO, and agency experts. Ecosystems had to be mappable, relatively well-studied, common in some form to all of the sources above, and represented through much of the bioregional assessment area. Settled on:
      i. Forest types
         a. Yellow pine
         b. Mixed conifer
         c. Red fir
         d. Subalpine
         e. Pinyon-juniper
         f. Hardwoods
            • Oak-dominated forests
            • Aspen
      ii. Shrub types
         a. Chaparral
         b. Sagebrush
      iii. Herbaceous types
         a. Montane meadows
      iv. Combined types
         a. Riparian vegetation
2. Determine appropriate scale
   A. Spatial:
      i. SNEP/SNFPA boundary
      ii. NRV reference data used from outside of area when necessary and reasonable
      iii. Assessments included both stand-level and landscape-level analyses where possible
   B. Temporal:
      i. Holocene (12,000 YBP to today), with period 1500–1850 set as principal historical reference period (this is a general HRV/NRV standard)
      ii. FSH 1902.12 guidance: should be before widespread Euro-American influence

3. Determine key ecosystem characteristics (= ecological indicators)
   A. Guidance from FSH 1902.12 Chapter 10
      i. “Key ecosystem characteristics include the dominant ecological characteristics that describe the composition, structure, function, and connectivity of terrestrial, aquatic, and riparian ecosystems that are relevant to addressing important concerns about the land management plan. Key ecosystem characteristics are identified, selected, and evaluated during the assessment phase, brought forward to inform the development of plan components, and may be useful for monitoring progress towards maintaining or restoring ecological integrity. Key ecosystem characteristics may be added or modified during the planning phase.”
      ii. Need to be characteristics that can be measured, are meaningful, and for which we have data both now and in the past, as well as in contemporary reference ecosystems. Also should be something which will respond to Forest Service management, or “indicates something about the limits to Forest Service authority or the inherent capability of the land.”
      iii. Some examples provided in FSH 1902.12 Chapter 10 (12.14 exhibit 01)
      iv. Key ecosystem characteristics include ecological processes (“drivers and stressors”)
B. Generated list of ecological indicators (= key ecosystem characteristics)

i. Ecological indicators: “measurable characteristics of the structure (e.g., genetic, population, habitat, and landscape pattern), composition (e.g., genes, species, populations, communities, and landscape types), or function (e.g., genetic, demographic/life history, ecosystem, and landscape disturbance processes) of ecological systems.” (Niemi and McDonald 2004)

ii. Based on nested hierarchy of:

a. Ecosystem attribute (highest level)
   - Composition
   - Structure
   - Function

b. Ecological hierarchy
   - Population/species
   - Community/ecosystem
   - Landscape/region

c. Indicator group (examples)
   - Species diversity
   - Physiognomy
   - Productivity
   - Biogeography
   - Nutrient cycling
   - Patchiness
   - Connectivity
   - Disturbance
     - Fire, grazing, logging, wind, floods, etc.

d. Indicators
   - Examples from fire regime indicator group
     - Fire frequency, fire rotation, fire size, fire severity, fire season…
   - Examples from physiognomy indicator group
     - Tree density, basal area/volume, canopy cover…

e. Variables and their units (lowest level)
   - Example from fire frequency indicator
     - Fire return interval (mean number of years between fires)
   - Example from tree density indicator
     - Number of trees per hectare

f. A spreadsheet of attributes, hierarchy, indicators and variables is available from the Regional Ecologist (but perusal of the chapters will also provide this list)
4. NRV assessments
   A. 11 Chapters, Introductory chapter to be written; intention is to publish as general technical report in 2013 or 2014
   B. NRV is based on historical and contemporary reference systems. Direct data used when available, inference used where necessary and justifiable
   C. Includes comparisons to current conditions and summary of literature re. possible future trends
   D. Focus is on peer reviewed publications, including papers in press or soon to be in press; government publications; Forest Service and other federal and state agency data; and in some cases academic theses or dissertations. Because information on the historical state of some ecosystems and ecological processes and patterns is scarce, in some cases we also refer to published anecdotal information from the mid-19th to early 20th centuries. We do not refer to anecdotal information from more recent times.
   E. Used Forest Service Rocky Mountain Region HRV assessments from early 2000s as rough templates (e.g., Dillon et al. 2005, Meyer et al. 2005, Veblen and Donnegan 2005)
   F. Basic outline
      i. Introduction
      ii. Methods
      iii. NRV descriptions
         a. Function
         b. Structure
         c. Composition
      iv. Summary of NRV deviations
      v. Literature cited
      vi. Tables and figures
   G. Timeline
      i. Dec 14, 2012: Preliminary bibliography assembled
      ii. Feb 11, 2013: Drafts due for internal review
      iii. Feb 15: Internal reviews due
      iv. Feb 25: Delivery of draft chapters to planning staff
      v. Mar 11: Final revised drafts due for external review
      vi. Apr 1: External reviews due
      vii. Week of Apr 8: Final draft NRV assessments
      viii. Week of May 12, 2013: Final NRV assessments posted to web
Official Business
Penalty for Private Use, $300