Surprises and lessons from the 1988 Yellowstone fires

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Understanding forest fire dynamics is becoming increasingly important as scientists, policy makers, and the public grapple with wildfires every year. Wildfires burn as much as 3.5 million ha annually in the US (NIFC 2003), and fire seasons during the past two decades have been among the most severe and costly on record in western North America. The causes and consequences of these fires remain the subject of considerable discussion. Some experts argue that decades of fire suppression, resulting in extensive and unnatural buildup of fuels, are to blame, while others suggest that variation in climate, particularly the occurrence of persistent and severe drought, is largely responsible.

Each view is correct with respect to some fire events and some ecosystems, but natural fire regimes and human impacts on those regimes are complex and variable from place to place. Furthermore, the interaction between climate change and fire may potentially be as important as the direct effects of global warming on forests (Dale et al. 2001). Thus, as policy makers and managers develop and implement measures to reduce future wildfire damage, it is critical that we incorporate our best understanding of the inherent variability in forest fire dynamics, as well as the importance of natural disturbance in ecological systems.

In this article we synthesize the results from 14 years of studying vegetation and ecosystem processes following the 1988 fires in Yellowstone National Park (YNP), WY, highlighting key ecological surprises and lessons learned about the role of fire.

In order to understand the 1988 Yellowstone fires, it is necessary to consider the broader context of fire regimes. Natural forest fire regimes vary widely, yet this complexity is often overlooked and underappreciated. The distinction between natural understory fire regimes and stand-replacing fire regimes is particularly important (Brown 2000). Understory fires burn at relatively low intensity and are typically characterized by short return intervals (from years to several decades). These fires burn along the ground, consuming woody fuels and understory vegetation, reducing tree regeneration, and maintaining an open forest structure. Historically, understory fire regimes dominated the once extensive long-leaf pine (Pinus australis) forests of the southeastern coastal plain, the open ponderosa pine (Pinus ponderosa) forests of the southern Rocky Mountains, and the ecotone between the forest and the prairie biomes. Because these low-intensity fires can be readily suppressed, many areas of the US that had been characterized historically by an understory fire regime have experienced increased fuel buildup and connectivity during the 20th century, raising the risk of high-intensity, stand-replacing fires. In these areas, past fire suppression is a major factor contributing to the increased occurrence and severity of wildfire (Covington and Moore 1994; Allen et al. 2002).

Recent wildfire mitigation programs at national and state levels have focused on forests formerly dominated by understory fire regimes. Unfortunately, some of these programs have failed to take into account that fire plays a very different role in many other forest types. Northern and subalpine conifer forests are typically characterized by

In a nutshell:

- Many northern conifer forests such as those in Yellowstone National Park are characterized by natural stand-replacing fires that are infrequent but severe
- Climate is the primary driver of stand-replacing fire regimes, while variation in fuels is less important
- Disturbances of this type and magnitude may influence plant population structure, genetics, and evolution
- Spatial heterogeneity resulting from this type of fire is the rule, not the exception
- Fires and fire regimes are not all the same, and a “one size fits all” approach to wildfire management is likely to be misguided

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in frequent, high-severity fires that kill most of the canopy, either via intense surface fire or flames spread through the crowns of the trees. Fire return intervals are long, ranging from about 60 years in jack pine (Pinus banksiana) to several centuries in some spruce–fir (Picea, Abies spp) communities. Stand-replacing fire regimes dominate the extensive boreal forests of North America, Europe, and Asia, as well as the forests of the northern Rocky Mountains, including YNP (Turner and Romme 1994). Climatic conditions, particularly severe regional drought, set the stage for occasional years of extensive conflagrations, and these few large fires account for most of the cumulative area burned over a long period of time (Johnson 1992; Johnson and Wowchuck 1993; Bessie and Johnson 1995; Flannigan and Wotton 2001). Under such extreme fire-weather conditions, variation in fuel structure is often of little importance, and fire suppression appears to have had little influence on recent fire behavior during these big fire years.

A stand-replacing fire regime characterizes the extensive subalpine forests of YNP. Thus, the lessons and surprises gleaned from intensive study of the 1988 fires can help us interpret the effects of recent and future fires in other ecosystems with a similar natural fire regime. Our Yellowstone findings are probably not very applicable to other ecosystems with a similar natural fire regime. Our focus here is on this forested subalpine plateau and not the lower elevation ecosystems. Tree-ring evidence (Romme 1982; Romme and Despain 1989) and studies of pollen and charcoal accumulation over millennia (Millspaugh et al. 2000) have clearly shown that YNP has experienced large-scale fire throughout the Holocene Epoch (the most recent 10,000 years). Fire return intervals have varied between 100 and 500 years.

Fire suppression was instituted in Yellowstone in 1886, but was not consistently effective prior to about 1945 (Schullery 1989). In response to a growing recognition of the ecological importance of fire, a natural fire program was initiated in 1972, in which lightning-caused fires in remote areas were allowed to burn without interference under proscribed conditions. Of over 200 such fires observed in the park between 1972 and 1988, 83% went out by themselves before burning more than 0.5 ha, and in 1981, the largest fire year prior to 1988, a total of 3300 ha was burned in 28 natural fires (Renkin and Despain 1992). During the summer of 1988, severe fires burned in YNP under conditions of extreme drought and high winds (Christensen et al. 1989; Renkin and Despain 1992). This surprised both managers and researchers, and focused worldwide attention on wildfire. It was reported in the media that the nation’s “crown jewel” was being destroyed by wildfires and would never recover (Smith 1996). The fires were indeed large, the most extensive since the park was established, and ultimately affected over 250,000 ha in the Greater Yellowstone Region. However, post-fire research has clearly demonstrated that the park’s important features were not “destroyed”.

Analyses of the 1988 fires strongly suggest that climate, rather than fuels, was primarily responsible for the extent of the fires (Romme and Turner in press). In June and July, the fires responded in similar fashion to what had been observed between 1972 and 1987 (Renkin and Despain 1992), and during these months, 19,555 ha were burned in subalpine portions of YNP. Full fire suppression efforts began in late July. As severe weather conditions persisted and intensified, fire activity increased. Between August 20 and September 15, another 157,726 ha was added to the area within the burn perimeter. These later fires spread rapidly through all forest successional stages and were influenced primarily by wind speed and direction. Even large natural firebreaks, such as the Grand Canyon of the Yellowstone, did not impede the spread of these late-season fires. Burning embers were carried 1 to 2 miles ahead of the main fire front, initiating new spot fires. Despite full suppression efforts, the fires ultimately stopped only with the arrival of the mid-September snowfall. Thus, it was weather that largely controlled the 1988 fires; landscape heterogeneity had minimal influence, and historic fire suppression probably had little effect.

The fiery summer of 1988

Established in 1872 as the world’s first national park, Yellowstone is famous for its unique thermal features, natural beauty, and abundance and variety of wildlife. Encompassing 9000 km² in the northwest corner of Wyoming, YNP is primarily a high, forested plateau. Approximately 80% of the park is covered with coniferous forests dominated by lodgepole pine (Pinus contorta var latifolia). Our focus here is on this forested subalpine plateau and not the lower elevation ecosystems. Tree-ring evidence (Romme 1982; Romme and Despain 1989) and studies of pollen and charcoal accumulation over millennia (Millspaugh et al. 2000) have clearly shown that YNP has experienced large-scale fire throughout the Holocene Epoch (the most recent 10,000 years). Fire return intervals have varied between 100 and 500 years.

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Fires do not homogenize the landscape

The public, and even some ecologists, feared that the 1988 fires had produced a desolate area of uniform devastation, but this was not the case. Instead, one of the most striking features of the 1988 fires was the post-fire heterogeneity of the burned landscape (Christensen et al. 1989). The fires created a spatially complex mosaic of unburned and burned patches, encompassing a wide range of burn severities (Figure 1). Understanding the effect of fire on landscape heterogeneity is important because the kinds, amounts, and spatial distribution of burned and unburned areas may influence the reestablishment of plant species on burned sites. A map of post-fire burn severity derived from Landsat Thematic Mapper imagery was used to quantify the isolation of burned areas and the relationship between heterogeneity and fire size (Turner et al. 1994). The majority of severely burned areas were within 50–200 m of unburned or lightly burned areas, suggesting that few burned sites were very distant from potential sources of off-site propagules. Spatial heterogeneity was the rule, not the exception, and a synthesis across differ-
ent disturbance types suggests that this characterizes many large, infrequent disturbances (Foster et al. 1998).

Post-fire vegetation

Plant cover and composition

The forests that burned in 1988 have since been repopulated by extensive stands of lodgepole pine and a wide variety of native herbaceous plants and shrubs. We have sampled the post-fire vegetation in nine patches — a small (1 ha), moderate (70–200 ha), and large (500–3600 ha) burned patch at each of three geographic locations — annual from 1990 to 1993 and again in 1996 and 2000. Post-fire plant reestablishment in most areas of YNP was rapid, with substantial increases in plant cover during the 4 years following the fires (Turner et al. 1997, 1999). We were surprised to find that most post-fire colonization occurred from plant parts that survived within the burned areas and then produced seed rather than by dispersal from unburned sites (Turner et al. 1997). Perennial herbs, graminoids (grasses and grasslike plants), and shrubs resprouted in 1989 and flowered profusely the following year, and widespread, abundant seedling establishment occurred in 1991 (Figure 2). The mean depth to which soil was charred was <14 mm, even in areas affected by severe surface and crown fires (Turner et al. 1999), suggesting that deeper roots and rhizomes had a high chance of survival. Therefore, although the fires were very large and severe, many plants survived and quickly refilled the burned forests. This rapid response of native species may also have provided a natural defense against invasion by exotic species that, surprisingly, did not increase in extent or abundance after the fires.

Our analyses of the vegetation data revealed substantial effects of fire severity and patch size on early post-fire plant cover and species richness (Turner et al. 1997, 1999; Romme and Turner in press). Some of these effects are now diminishing, but small patches still have greater forb, graminoid, and total cover compared to large patches, and more severely burned sites still have lower cover of forbs, graminoids, and shrubs compared to less severely burned sites (Figure 3). However, the effects of environmental variation on species richness and community composition are becoming more pronounced (Figure 3), and post-fire communities are similar in composition to nearby forests that did not burn. The influence of the abiotic template on vegetation is, therefore, becoming more evident as succession proceeds.

Lodgepole pine

Lodgepole pine reestablished well following the 1988 fires. Adult lodgepoles do not survive fire, except where fire intensity is low, and post-fire regeneration depends entirely on seedling establishment which often generates dense, even-aged stands (Johnson and Fryer 1989). Lodgepole pine is serotinous, meaning that trees may produce cones that remain closed at maturity and only open in response to high temperatures (Figure 4). Trees may retain closed cones for 40 years or longer, and the canopy...
may therefore contain millions of seeds per ha (Clements 1910; Lotan and Jensen 1970). Seed stored in serotinous cones can remain viable for 50–75 years (Clements 1910), and the seedlings thrive in the high-light environment of a recently burned forest.

We found significant influences of fire severity, patch size, and geographic location on post-fire pine seedling densities (Figure 5). Large patches had more pine seedlings than smaller patches, and areas with severe surface burns had more seedlings than areas with crown fires or less severe burns (Figure 5). In areas with severe surface burns, temperatures were probably sufficiently high to break the resin seal on the serotinous cones, but not so high as to reduce viability of the seed; thus, local variation in burn severity (Turner et al. 1994) resulted in fine-scale variation in post-fire pine density. However, we did not expect such large variation in pine seedling density between the three geographic locations of our study patches. We discovered that our three “replicate” locations had unexpectedly captured a strong and previously undescribed gradient in pre-fire serotiny (Table 1).

Although lodgepole pine is a serotinous species, not all trees produce this type of cone. The spatial and temporal controls on expression of serotiny are not well understood. Measurements of serotiny in unburned lodgepole pine forests in YNP revealed that it is affected by both elevation and stand age (Tinker et al. 1994; Schoennagel 2002; Schoennagel et al. in press). At high elevations (>2300 m), where estimated fire return interval is ~300 years, serotiny is low. However, at low elevations (<2300 m), where estimated fire return interval is ~180 years, serotiny is low in young stands but high in stands >70 years old (Schoennagel 2002; Schoennagel et al. in press). These patterns are consistent with our observations of geographic variation in both serotiny and post-fire pine recruitment in our study patches.

After observing the variation in post-fire pine density in our nine study patches, we were curious about the spatial heterogeneity in pine density across the broader landscape. Using a combination of aerial photography and extensive field studies, we found post-fire pine densities ranging from 0 to 535 000 stems/ha (Turner et al. in press). Collectively, our analyses described a matrix of larger patches of low-density lodgepole pine randomly interspersed with smaller patches of high-density pine. The six orders of magnitude
variation in post-fire lodgepole pine density was astounding, and also generated considerable spatial variation in aboveground (Reed et al. 1999, Turner et al. in press) and belowground (Litton 2002; Litton et al. 2003a, b) net primary production. For example, post-fire annual net primary productivity in 11-year-old lodgepole pine stands ranged from 0.04–15.12 mg/ha/yr, spanning much of the range of variation expected through successional time (Turner et al. in press). Our results suggest that the factors influencing post-fire stand density – namely, pre-fire serotiny and fire size, severity, and pattern – also influence post-fire ecosystem function.

How long is this post-fire spatial variation likely to persist on the landscape? Variation in stand density will probably be reduced through time as high-density stands self-thin and low-density stands continue to have gradual recruitment (Kashian 2002). However, the initial post-fire mosaic will probably influence the structure and function of the YNP landscape for at least a century. Analyses of basal area increment in lodgepole pine stands spanning a range of age classes and densities indicate that this measurement varies with density until the trees are about 125 years old (Kashian 2002). Clearly, the fire-generated patterns in forest structure have left a legacy that will persist for many decades in the YNP landscape.

Aspen

The post-fire dynamics of aspen (*Populus tremuloides*) in YNP provided yet another ecological surprise. As graduate students, we had absorbed the conventional wisdom that the aspen clones in the northern Rocky Mountains (Figure 6) have persisted since the Pleistocene Epoch and do not reproduce by seed. Therefore, although we anticipated root sprouting of burned aspen clones where they were present before the fires (Romme et al. 1995), we did not expect aspen on the plateau. During our 1989 field season, we encountered small seedlings in the burned lodgepole pine forests that we could not identify definitively. In 1990, these seedlings were becoming woody, and they keyed out to aspen. Indeed, seedling aspen established successfully throughout the burned pine forests and well beyond the pre-fire range of aspen.

We have documented seedling aspen occurring as far as 15 km from the nearest known aspen clone, and most are well over 5 km away (Romme et al. 1997; Turner et al. 2003). If the statistical models of post-fire aspen seedling occurrence as measured along infrequently used foot trails (Turner et al. 2003) are extrapolated spatially to the landscape, we can see areas of YNP in which aspen seedlings have a higher likelihood of occurrence (Figure 7). Genetic analyses of seedling aspen populations found greater variation than that observed among pre-fire clones (Tuskan et al. 1996), and most variation occurred within rather than among aspen seedling populations (Stevens et al. 1999). Even geographically distant seedling populations shared nearly 90% of the same markers (Stevens et al. 1999). Within plots, the amount of genetic variation decreased slightly with increasing percentage of vegetative cover and mean size of the aspen seedlings, suggesting that competition between aspen and other plants might already be eliminating some genotypes.

Aspen is well known as the preferred forage for elk, and the role of ungulate browsing on the dynamics of aspen throughout the Rocky Mountains has been the subject of some controversy (NRC 2002). Nearly all the aspen seedlings that we have observed in the burned pine forests have been browsed and average 30 cm in height (Turner unpublished; Figure 6, top). However, recent studies suggest a potentially complex interaction between fire, post-
fire coarse woody debris, and aspen recruitment. Many of the conifers that were killed in the 1988 fires have now fallen to the ground, creating in some areas a thicket of fallen trees that is difficult to traverse, even for elk. Ripple and Larson (2001) observed tall aspen ramets (individual members of a clone) in areas where the plants were protected from browsing by natural piles of fallen trees. Therefore, although most aspen seedlings remain small and are browsed, post-fire coarse woody debris may be creating “safe zones” for aspen, in which new clones appear to be establishing (Figure 6, bottom). We suggest that recruitment of new aspen genets in this region is episodic, is usually associated with large severe fires like those in 1988, and contributes considerably to population genetic structure and diversity. Numerous perennial herbs of the forest floor also appear to persist primarily via asexual sprouting during fire-free intervals, but to recruit new genetic individuals after fire (Romme et al. 1993).

Coarse woody debris

The 1988 fires in Yellowstone killed millions of lodgepole pines, creating an estimated 25 million metric tons of standing and downed dead wood, known as coarse woody debris (CWD) (Tinker and Knight unpublished). This wood is one of the most important legacies of the pre-fire forest. CWD in the form of snags, logs, and stumps may cover as much as 24% of the forest floor immediately following an intense fire, and this percentage increases to almost 60% by 50 years after the fire, as the snags fall (Tinker and Knight 2001), modifying the forest floor environment. The decaying logs, roots, and stumps are also a primary source of organic matter inputs to Yellowstone's generally infertile forest soils, which may help maintain higher levels of productivity. Surprisingly, very little of the CWD that was present on the forest floor at the time of the fires was consumed during 1988. A small proportion (about 8%) was consumed and an equal amount was converted to charcoal (Tinker and Knight 2000). This persistent and important pool of organic material will affect forest floor and soil structure and function for decades after stand-replacing fires.

Lessons from the fires

The 1988 Yellowstone fires produced spatially complex patterns of succession in what is often considered a relatively simple system dominated by lodgepole pine. The post-fire spatial variability in forest structure in turn influences – perhaps even dominates – ecosystem function across the landscape, and its imprint will probably persist for at least a century. Long-term study is essential for understanding natural disturbance dynamics, particularly when events are both large and infrequent (Turner et al. 1998, 2003). The 1988 fires continue to offer irreplaceable opportunities to address basic questions in ecology and to study natural ecosystem structure, function, and dynamics.

The significance of the 1988 Yellowstone fires extends far beyond the park. Large fires will probably occur again in YNP and in many other areas of western and northern North America, where extensive coniferous forests are characterized by large, infrequent blazes. Many of these areas are prized for their aesthetic beauty and environmental amenities, and are experiencing extensive rural development (Hansen et al. 2002; Romme et al. in press). This combination may inadvertently set the stage for increasing future conflicts between competing values, similar to those which have developed in flood- or storm-prone regions. Neither complete fire exclusion nor conversion of the natural stand-replacing fire regime to an
understory fire regime appears feasible, either operationally or ecologically (Romme et al. in press). Although local thinning to create defensible space around individual homes may be practical, fire management in ecosystems with stand-replacing fire regimes must accept and incorporate large, severe, infrequent fires throughout much of the landscape (Romme et al. in press).

What lessons learned from the 1988 fires may be relevant elsewhere? First, distinctions must be clearly drawn between natural stand-replacing fire regimes such as YNP and other areas where past fire suppression has shifted the fire regime from understory to stand-replacing. Interventions such as low-intensity prescribed burning or mechanical thinning to restore natural fire dynamics are not appropriate for crown-fire systems. Second, large stand-replacing fires are an important source of landscape heterogeneity. Even though such fires create patches of even forest age across the landscape, unburned patches are interspersed throughout, and there is tremendous spatial variation in both forest structure and function within the burned area.

Third, natural processes restored plant cover and composition in the burned area quite rapidly, with no deleterious consequences observed (Romme and Turner in press). Thus, active post-fire rehabilitation is probably not needed, and may even be counterproductive. For example, exotic plant species introduced deliberately to expedite revegetation may retard forest recovery. The CWD created by the 1988 fires played an unexpected role in the establishment of new aspen clones—a role that never would have been recognized had the park quickly salvaged the fire-killed trees. Finally, episodic large-scale disturbances may play a key role in the population structure, genetics, and evolution of long-lived clonal plant species such as aspen and many forest floor herbs. Continuing research on the effects of the 1988 Yellowstone fires will lead to additional insights and new surprises about the dynamics of systems characterized by stand-replacing fire regimes.

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