Present Effects of Past Wildfires on the Diets of Stream Fish

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This document was originally published by Brigham Young University in *Western North American Naturalist*. Copyright restrictions may apply. DOI: 10.3398/1527-0904(2007)67[429:PEOPWO]2.0.CO;2
Stream ecosystems are intimately tied to the catchments they drain (Hynes 1975, Van-note et al. 1980). Any disturbance in the catchment can influence the structure and function of the stream’s biotic components. For example, clearing of forested and riparian areas can directly affect fish assemblages by increasing stream temperatures and discharge (Hetrick et al. 1998b). Indirectly, reduction of terrestrial vegetation may alter in-stream cover by reducing inputs of large woody debris, and by potentially altering the prey base (Hetrick et al. 1998a, Keith et al. 1998).

In the western United States, wildfire is a commonly occurring landscape-level disturbance which has pronounced effects on both terrestrial and aquatic systems. Immediate effects of fire on stream ecosystems have been well documented (Minshall et al. 1989, 2004, Gresswell 1999, Benda et al. 2003, Robinson et al. 2005), but these effects usually wane within a decade (Gresswell 1999, Minshall et al. 2004). Wildfire’s indirect impacts on the stream, usually mediated through changes in terrestrial vegetation, have been less studied. By removing streamside vegetation, fire may influence fish and invertebrates not only by raising water temperatures and escalating substrate scouring, but also by decreasing transported organic material and increasing the importance of autotrophic energy pathways (Minshall et al. 1989, McIntyre and Minshall 1996, Rieman and Clayton 1997).

In landscapes where wildfire is a recurring disturbance, fish populations tend to recover quickly (Rinne 1996, Rieman et al. 1997, Gresswell 1999). These populations can evolve strategies that incorporate the fire regime into their overall life histories (Reiman and Clayton 1997). Thus, the effect of wildfire on some fish populations may be evident for only a short time (Rinne 1996, Gresswell 1999). However, recent studies suggest that indirect effects, which are more subtle and longer-lasting, can occur between a catchment’s terrestrial vegetation and the stream that drains it (e.g., Nakano et al. 1999, Fausch et al. 2002, Dunham et al. 2007). In the long term (tens to hundreds of years), the response of the stream system to fire may be tied to the recovery of the riparian vegetation (Minshall et al. 1989, 2004). While fish assemblages tend to recover quickly, do wildfires still influence fish ecology a decade after the burn? Diet composition in fish may be 1 variable that displays lasting effects due to fire. The diets of fish (hereafter “fish diets”) are, in part, linked to the riparian canopy (Wootton 1992, Gerking 1994), and wildfire has

**ABSTRACT.—**We investigated present indirect effects from a decade-old burn on the diets of stream fish. Based on soil instability and burn patterns from a 12–14-year-old wildfire complex, we separated 9 streams into 3 conditions: unburned, burned, and burned/scoured (i.e., experiencing a massive scour event 1 year post-burn). In the streams, we measured light levels and water temperatures, and we estimated fish density and biomass. We removed the digestive tracts from 9–15 rainbow trout (*Oncorhynchus mykiss*) from each stream and used a gravimetric procedure to analyze gut contents. Canopy cover development may dictate the composition of dietary items. Greater amounts of aquatic invertebrates and inorganic material were found in trout from streams with reduced overhead canopy. Further, these streams had lower amounts of terrestrial invertebrates and organic materials in the diets of fish. Although trout abundance was not different among the stream treatments, fire-induced, indirect effects on fish diets were still evident more than a decade after the burn. This suggests that recovery rates for trout assemblages may take longer than predicted and may depend on riparian recovery.

**Key words: wildfire, rainbow trout, diet composition, riparian canopy, streams.**

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PRESENT EFFECTS OF PAST WILDFIRES ON THE DIETS OF STREAM FISH

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long-term impacts on the recovery of that canopy. Based on these relationships, we hypothesized that fire-induced change that alters riparian vegetation should be expressed in the diet composition of stream fish. We collected fish from unburned streams and streams burned in 1992–1994. We compared fish stomach contents using gravimetric procedures, and found differences in gut content items that were correlated to fire-altered riparian vegetation.

**STUDY AREA**

Our study was conducted on nine 2nd- and 3rd-order tributaries of the Boise River catchment, located in the southern forested subregion of the Idaho batholith ecoregion (44°10′N–44°20′N, 115°30′W). Catchments in the Boise River Basin, located in the Boise National Forest, have hillside gradients ranging from 20° to 40° and are highly dissected. Upland vegetation is dominated by stands of ponderosa pine (*Pinus ponderosa*) and Douglas-fir (*Pseudotsuga menziesii*), while riparian vegetation is dominated by willow (*Salix* spp.), cottonwood (*Populus* spp.), and water birch (*Betula occidentalis*), each type with an associated understory of mixed shrubs, grasses, and forbs. The regional climate is continental with an average precipitation of 500 mm per year (Benda et al. 2003). The bedrock substrate in the basin is dominated by moderate to well-weathered coarse-grained quartz monzonite typical of much of the Idaho batholith. Soils are lithosols with weakly developed A horizons ranging from 5 to 25 cm thick (Megahan and Ketcheson 1996). These granitic, droughty soils have limited fertility and are highly erodible when vegetation is removed (McGarth et al. 2001).

**METHODS**

This research was part of ongoing investigation examining the effects of wildfire on stream trophic structure. Our present study was conducted from 7 July through 20 July 2005. Using aerial photographs, GIS layer maps, and considering a stream fire study design developed by the USDA FS Rocky Mountain Research Station (see Dunham et al. 2007), we tried to choose streams for which we could match catchment area, slope, gradient, elevation, and discharge (Table 1). Of the 9 study streams, 6 were burned in the 1992–1994 Foothills fire and Idaho City complex of wildfires (85%–100% of catchment; Dunham et al. 2007). Due to soil instability after vegetation removal, 3 of these burned systems experienced a massive scouring (erosion and redeposition of bed material throughout the stream’s reach) the following year. Subsequently, these 3 streams continued to experience moderate scour and redeposition of bed material after minor storms. Following Dunham et al. (2007), we grouped the study streams into 3 conditions based on wildfire patterns and soil instability. These stream conditions were (1) unburned: streams that were not burned by the fires; (2) burned: streams that burned above ground but retained much of the belowground vegetative structures; and (3) burned/scoured: streams that, less than a year after the fire, experienced major erosion, movement, and deposition of

**Table 1. Physical characteristics of the 9 streams sampled during July 2005.**

<table>
<thead>
<tr>
<th>Stream/condition</th>
<th>Order</th>
<th>Elevation (m)</th>
<th>Mean width (m)</th>
<th>Gradient (%)</th>
<th>Catchment area (km²)</th>
<th>Discharge (m³ s⁻¹)</th>
<th>Location (latitude, longitude)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Burned/scoured</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SF Sheep Creek</td>
<td>2</td>
<td>1137</td>
<td>4.2</td>
<td>2.8</td>
<td>31.7</td>
<td>0.19</td>
<td>43°40'926&quot;N, 115°38'811&quot;W</td>
</tr>
<tr>
<td>Trapper Creek</td>
<td>3</td>
<td>1479</td>
<td>2.6</td>
<td>4.7</td>
<td>9.1</td>
<td>0.06</td>
<td>43°53'543&quot;N, 115°26'918&quot;W</td>
</tr>
<tr>
<td>Wren Creek</td>
<td>3</td>
<td>1363</td>
<td>2.4</td>
<td>10.2</td>
<td>9.4</td>
<td>0.29</td>
<td>43°52'435&quot;N, 115°30'990&quot;W</td>
</tr>
<tr>
<td><strong>Burned</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cottonwood Creek</td>
<td>3</td>
<td>1296</td>
<td>3.0</td>
<td>4.5</td>
<td>24.1</td>
<td>0.19</td>
<td>43°40'538&quot;N, 115°49'417&quot;W</td>
</tr>
<tr>
<td>Hungarian Creek</td>
<td>3</td>
<td>1286</td>
<td>1.8</td>
<td>6.4</td>
<td>11.5</td>
<td>0.07</td>
<td>43°49'039&quot;N, 115°32'127&quot;W</td>
</tr>
<tr>
<td>Lost Creek</td>
<td>2</td>
<td>1368</td>
<td>3.3</td>
<td>5.3</td>
<td>14.4</td>
<td>0.05</td>
<td>43°51'586&quot;N, 115°31'737&quot;W</td>
</tr>
<tr>
<td><strong>Unburned</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Beaver Creek</td>
<td>2</td>
<td>1332</td>
<td>1.9</td>
<td>4.6</td>
<td>14.2</td>
<td>0.05</td>
<td>43°53'186&quot;N, 115°30'564&quot;W</td>
</tr>
<tr>
<td>Lost Man Creek</td>
<td>2</td>
<td>1599</td>
<td>2.7</td>
<td>2.1</td>
<td>13.3</td>
<td>0.07</td>
<td>43°44'971&quot;N, 115°26'264&quot;W</td>
</tr>
<tr>
<td>Trail 2 Creek</td>
<td>2</td>
<td>1559</td>
<td>2.2</td>
<td>4.1</td>
<td>17.2</td>
<td>0.06</td>
<td>43°45'623&quot;N, 115°08'881&quot;W</td>
</tr>
</tbody>
</table>
bed material, which removed most of the riparian belowground vegetative structures (see Benda et al. 2003 and Dunham et al. 2007 for fire history of this area).

We used the amount of light potentially penetrating the riparian vegetation and reaching the stream's surface as a proximate measure of riparian development. After walking along 1–2 km of each stream, we chose a 100-m segment as a representative reach. To estimate riparian canopy, we randomly chose 50 points along and across the 100-m reach in each stream. At each point, we measured the amount of photosynthetically active radiation (PAR) using a LI-COR quantum sensor (model LI-192SA) and a LI-COR light meter (model LI-250). We placed the sensor 3 cm above the water's surface and measured PAR on cloudless days between 1200 hours and 1300 hours.

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Fish assemblages in the tributaries of the upper Boise River catchment are comprised of rainbow trout (*Oncorhynchus mykiss*), brook trout (*Salvelinus fontinalis*), and bull trout (*S. confluentus*), mountain whitefish (*Prosopium williamsoni*), and shorthead sculpin (*Cottus confuses*). However, 98% of the fishes collected in the study streams were *O. mykiss*. Consequently, we limited our study of fish variables and gut content comparisons to this species. We sampled *O. mykiss* by backpack electrofishing immediately after we measured PAR and discharge. Block nets (mesh size: 5 mm) were set at the upstream and downstream ends of the sampling segment. Two sequential passes were made (in all streams, probability of fish capture equaled or exceeded 0.80, necessitating only 2 passes; Armour et al. 1983). We estimated fish density using a standard depletion-curve method (Youngs and Robson 1978, Armour et al. 1983, Brower et al. 1997). All *O. mykiss* collected were identified and weighed, and the total lengths were recorded. Because we collected no young-of-the-year from 1 of the burned/scoured streams, we omitted all individuals <70 mm from this study. Of the fishes collected, we removed the digestive tracts from 9–15 fish from each stream and stored them on ice for transport back to the laboratory.

In the laboratory, we dissected each digestive tract and removed the contents. With the aid of a dissecting microscope, we separated gut contents into 4 categories: aquatic invertebrates, terrestrial invertebrates, organic material (leaf and bark particles, algae, etc.), and inorganic matter (sand, small pebbles, etc.). We identified invertebrates to family, but in most cases we could also recognize genera. All stomachs that we examined contained material that could be placed in 1 (or more) of our gut content categories; no empty stomachs were encountered. Once stomach contents were removed, identified, and sorted into categories, the materials were oven-dried for 2 days at 80°C to stabilize their weights and to decrease sample variability (Hyslop 1980, Bowen 1996), and then the samples were weighed. Additionally, we estimated the condition coefficient of *O. mykiss* collected in each stream using the following equation (Everhart and Young 1981):

\[
C = \left( \frac{W}{L^3} \right) \times 10^5,
\]

where \(C\) = condition coefficient, \(W\) = fish biomass (g), and \(L\) = fish length (mm).

To determine PAR and water temperature differences among our 3 stream conditions, we analyzed the data using an analysis of variance (ANOVA) with stream condition as the treatment and PAR or temperature as response variables. These datasets were log\((x + 1)\)-transformed to achieve linearity and homoscedasticity. We also used ANOVA to identify differences in fish density, total length, biomass, and condition among the stream conditions. Because each of our gut content categories was recorded as a percentage of the sample’s total weight, we applied an arcsine transformation to the data to normalize their distribution (Zar 1999). We used the nonparametric Kruskal-Wallis ANOVA (Siegel and Castellan 1988) to determine if stream condition affected the percent composition of fish diet. If significance was found (\(P < 0.05\)), we then used a 2-sample Kolmogorov-Smirnov test as a post hoc
pairwise comparison between the stream conditions within that gut content category (Siegel and Castellan 1988).

RESULTS

Noticeable visual differences in riparian canopy were evident between the 3 stream conditions. These field observations were confirmed by significant differences in both photosynthetically active radiation (PAR) measured at the streams' surface ($F_{2, 441} = 36.07, P < 0.001$) and by stream water temperatures ($F_{2, 432} = 50.41, P < 0.001$; Table 2). Our burned/scoured streams had the highest mean values for PAR (1200 µmol · m$^{-2}$s$^{-1}$) and water temperature (16.3°C). Measurements for these variables were significantly greater ($P < 0.05$) than those recorded in our unburned streams (PAR 292 µmol · m$^{-2}$s$^{-1}$; temperature 11.6°C). For both PAR and water temperature, the burned streams were intermediate between the other conditions (Fig. 1).

Average *O. mykiss* density was highest in burned streams (density = 34 individuals · 100 m$^{-2}$, $s_{x} = 4.0$) while density was the lowest in burned/scoured streams ($x = 18$ individuals · 100 m$^{-2}$, $s_{x} = 0.4$); however, these differences were not significant ($F_{2, 6} = 1.14, P = 0.38$). Significant differences did occur in fish total length and biomass among the stream conditions (Fig. 2). Mean total length and biomass were largest in the burned streams, but smallest in the unburned ones (total length: $F_{2, 6} = 3.47, P = 0.04$; biomass: $F_{2, 6} = 3.07, P = 0.05$). Regardless of differences in fish length and biomass, fish condition was nearly equal in all 3 stream conditions ($F_{2, 6} = 2.01, P = 0.14$).

Gravimetric comparisons of gut contents did show significant differences among the 3 designated stream conditions (Fig. 3). Both aquatic invertebrate (Kruskal-Wallis ANOVA: $H = 6.04, P = 0.049$) and inorganic matter (Kruskal-Wallis ANOVA: $6.36, P = 0.04$) categories displayed the same trend, with proportionately more of these materials ingested by fish in the burned/scoured streams than in either of the other 2 stream conditions. Although aquatic invertebrates were highest in *O. mykiss* guts collected from the burned/scoured streams, these values did not vary significantly from values for guts collected from the burned systems ($P = 0.37$), but both were greater than the values for trout guts collected from unburned streams ($P < 0.05$). Likewise, trout from burned/scoured systems had a higher proportion of ingested inorganic matter in their stomachs than trout in the other streams ($P < 0.04$), while material from fish in the burned and unburned streams was not significantly different.

In contrast, the organic material category displayed an opposite trend, with more material ingested by *O. mykiss* from the unburned streams when compared to the other streams, with the least organic material in fish from the burned/scoured systems (Kruskal-Wallis ANOVA: $H = 9.29, P = 0.01$). Material in the organic category consisted of leaf and bark particles as well as some algae (frequently occurring in stomachs from fish in the burned/scoured systems). There were no proportionate differences of organic material in trout among the burned and unburned conditions, but each was significantly greater than that found in fish from the burned/scoured systems ($P < 0.04$). Although the terrestrial invertebrate category showed the same pattern as that of organic material, the pattern was not significant.

DISCUSSION

The immediate effects of wildfire on stream biotic structure and function have been well documented (Minshall et al. 1989, Minshall...
and Brock 1991, Rinne 1996, Rieman and Clayton 1997, Gresswell 1999, Spencer et al. 2003). Less clear are the mid- to long-term patterns in biotic response (but see Robinson et al. 2005 and Dunham et al. 2007 for physical data). At the community level, biotic variables have shown large variation with little correlation to predicted response signatures (Minshall et al. 2004). For example, fish abundance and community structure were predicted to attain preburn densities 10–12 years after the fire (Minshall et al. 1989, Gresswell 1999). But fish assemblages in streams of the Boise River Basin have returned to prefire numbers within 3 years post-burn (Rieman and Clayton 1997, Rieman et al. 1997). Lack of correlation to predicted patterns and wide variation have led some to suggest that studying population-level responses might be more appropriate for detecting long-term change and recovery (Mihuc 2004, Mihuc and Minshall 2005).

Though we found no differences in rainbow trout population densities or fish condition among the study streams, our results do suggest that wildfire continues to affect biotic
processes in the long-term, albeit indirectly, through alterations of the riparian canopy cover. While trout densities were not different among our streams, we did find significant differences in the gravimetric comparisons of fish stomach contents. Fish diet appears to be influenced by the rate of riparian recovery. Due to a lack of vegetative overhang in our burned/scoured systems, fewer terrestrial invertebrates may have fallen into these streams. As a result, aquatic invertebrates made up a large portion of the fish diet. During the process of feeding on the benthic fauna, fish may have engulfed sand and small stones from the substrate itself. Inorganic materials from ingested case-dwelling invertebrates were also present in fish from this stream condition. We found several intact cases from the caddis larvae *Dicosmoecus* within fish stomachs from the burned/scoured streams. Indeed, inorganic material was significantly greater in fish stomachs from the burned/scoured streams when compared to fish stomachs from the other 2 stream conditions. Others have interpreted the presence of organic and inorganic matter in salmonid gut analyses as material that was...
ingested accidentally during epibenthic feeding (Tippets and Moyle 1978, Angradi and Griffith 1989).

The opposite was true of fish from the unburned systems. With a thick overhanging canopy present, the input of terrestrial invertebrates into these streams may have been high. In addition, the thick canopy cover reduced the amount of solar radiation available to the primary producers of unburned streams, potentially limiting the amount of autotrophic energy available to the aquatic invertebrate community (Minshall and Brock 1991, Minshall 2003). The combination of these 2 factors may result in a larger percentage of terrestrial insects found in the stomachs of fish from the unburned sites. In feeding on the terrestrially-derived invertebrates, fish may ingest plant material as well. We found higher amounts of organic matter in the stomachs of trout from these unburned streams. Most of this material was composed of leaf, needle, and bark particles. Vascular plant material may supplement the energy requirements of some fish (Bowen et al. 1995), but it is likely that this material was accidentally ingested during water-column feeding.

Our study supports some of the predictions originally made by Minshall et al. (1989) immediately after the 1988 Yellowstone fires.
The return of stream ecosystems to prefire levels of structure and function appear to be closely tied to the recovery of the catchment’s forest and understory vegetation (Rieman and Clayton 1997). The highly erodible nature of the granitic soils (common to the Idaho batholith region) underlying our streams coupled with fire intensity dictate the recovery time of the catchment’s vegetation. Within 1 or 2 years after the fires, some of the study streams experienced a massive scouring, erosion, and redeposition of bed materials (Benda et al. 2003), removing much of the belowground vegetative structures. Consequently, streams like these may be prone to substrate-scouring events at moderate levels of precipitation (Megahan 1983, Megahan and Ketcheson 1996). In turn, this continued scouring could further delay riparian recovery. By using the amount of solar radiation reaching the water’s surface as a proxy for canopy coverage, we concluded that the riparian vegetation for the burned/scoured streams (when compared to the unburned ones) had not yet attained prefire levels 12–14 years after the burn. The sparse amounts of riparian overhang directly affected stream water temperatures (Minshall et al. 1989, 2004, Dunham et al. 2007). In turn, many biotic components of stream ecosystems—algal standing crop, invertebrate growth rates and community composition, fish growth and production—are partially governed by water temperature (Allan 1995, Giller and Malmqvist 1998). Indeed, in 1 of the burned/scoured streams (Wren Creek) we found no young-of-the-year of any fish species. Thus, long-term, indirect effects of the 1992–1994 wildfires may be evident in other biotic parameters within these streams.

The intensity of the fire coupled with vegetation removal and the resulting soil instability may alter the recovery trajectory of the burned/scoured streams (sensu Drake 1990). Streams with eroded catchments and severe streambed scour may experience prolonged recovery times, or they may move to a new steady-state final configuration (Matthaei et al. 2004). In severe disturbance regimes, such as repeated scouring of the stream channel over a long time period, new lower levels of biotic richness and abundance may be established (Minshall et al. 1989).

In conclusion, wildfire may have subtle, long-lasting effects on the functioning of stream ecosystems. The duration of these effects are dependent upon riparian recovery and soil stability. Too often, studies of stream catchments in fire-prone basins are classified as only burned or unburned, leading to a wide variation in long-term response signatures (e.g., Robinson et al. 2005). Our study supports the notion that a 3rd category, burned/scoured streams, should be considered based on catchment soil types, channel scouring frequency, and riparian vegetation recovery. Identifying the type and frequency of post-fire disturbance (sensu Lake 2000) may reduce the variation evident in long-term fire recovery studies and lead to a better understanding of wildfire’s effects on long-term stream dynamics.

ACKNOWLEDGMENTS

This study was supported by the National Research Initiative of the USDA Cooperative State Research, Education and Extension Service, grant number 2003-35101-13682, and by the Dan Montgomery Foundation for ecological study. We would also like to thank C. Valdivia, T. Krause, and A. Fonner for field and laboratory assistance; D. Nagel (USDA Forest Service) for GIS layer maps; and A. Rosenberger (USDA Forest Service) and J. Dunham (USDA Forest Service / USGS) for many thoughtful discussions on study site selection and the role of fire in forested ecosystems.

LITERATURE CITED


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2007


ROBINSON, C.T., U. UEHILINGER, AND G.W. MINSHALL. 2005. Functional characteristics of wilderness streams...


Received 4 April 2006
Accepted 8 January 2007