



Influence of Drying and Wildfire on Longitudinal Chemistry Patterns and Processes of Intermittent Streams

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Stream drying and wildfire are projected to increase with climate change in the western United States, and both are likely to impact stream chemistry patterns and processes. To investigate drying and wildfire effects on stream chemistry (carbon, nutrients, anions, cations, and isotopes), we examined seasonal drying in two intermittent streams in southwestern Idaho, one stream that was unburned and one that burned 8 months prior to our study period. During the seasonal recession following snowmelt, we hypothesized that spatiotemporal patterns of stream chemistry would change due to increased evaporation, groundwater dominance, and autochthonous carbon production. With increased nutrients and reduced canopy cover, we expected greater shifts in the burned stream. To capture spatial chemistry patterns, we sampled surface water for a suite of analytes along the length of each stream with a high spatial scope (50-m sampling along ~2,500 m). To capture temporal variation, we sampled each stream in April (higher flow), May, and June (lower flow) in 2016. Seasonal patterns and processes influencing stream chemistry were generally similar in both streams, but some were amplified in the burned stream. Mean dissolved inorganic carbon (DIC) concentrations increased with drying by 22% in the unburned and by 300% in the burned stream. In contrast, mean total nitrogen (TN) concentrations decreased in both streams, with a 16% TN decrease in the unburned stream and a 500% TN decrease (mostly nitrate) in the burned stream. Contrary to expectations, dissolved organic carbon (DOC) concentrations varied more in space than in time. In addition, we found the streams did not become more evaporative relative to the Local Meteoric Water Line (LMWL) and we found weak evidence for evapoconcentration with drying. However, consistent with our expectations, strontium-DIC ratios indicated stream water shifted toward groundwater-dominance, especially in the burned stream. Fluorescence and absorbance measurements showed considerable spatial variation in DOC sourcing each month in both streams, and mean values suggested a temporal shift from allochthonous toward autochthonous carbon sources in the burned stream. Our findings suggest that the effects of fire may magnify some chemistry patterns but not the biophysical controls that we tested with stream drying.

Keywords: intermittent streams, fire, carbon, nitrogen, spatiotemporal patterns, allochthonous and autochthonous, evaporation, groundwater

INTRODUCTION

Intermittent streams, those streams experiencing periods of disconnected surface water flow (Larned et al., 2010), currently constitute ~30% of the total river length and discharge of the world river network and about half of the United States (US) network (Datry et al., 2014). Despite the ubiquity of stream drying, perennial streams have historically dominated our understanding of stream chemistry patterns. Indeed, study and understanding of intermittent streams have lagged behind perennial ones (Costigan et al., 2016; Allen et al., 2020; Busch et al., 2020). For example, traditional stream chemistry studies have often focused on temporally intensive measurements at the outlet of a catchment and assumed that downstream patterns are representative of the upstream segment (e.g., Fisher and Likens, 1973; Schiff and Aravena, 1990; Boyer et al., 1997). However, this assumption may not be valid and the approach is unlikely to capture the spatiotemporal variability of intermittent streams (Godsey and Kirchner, 2014; Costigan et al., 2016). To address this assumption, the scope of intermittent stream studies, which is determined by the extent (area or time-period over which measurements are conducted) and the grain (frequency of measurement through space or time) need to be adjusted (Schneider, 2001; Fausch et al., 2002). Indeed, high spatial-scope studies adopting a spatially continuous sampling approach (fine-grain) carried out over larger stream segment lengths (greater extent) have revealed previously undetected spatial patterns along the length of perennial streams and stream networks (Fausch et al., 2002; Likens and Buso, 2006; McGuire et al., 2014). Stream intermittence is characterized by contractions and disconnections that occur with drying at the scale of meters with consequences for stream structure at segment to network scales (Stanley et al., 1997; Dent and Grimm, 1999; Zimmer et al., 2013; Hale and Godsey, 2019; Jensen et al., 2019). Thus, temporally repeated high-spatial scope investigations seem necessary to understand patterns and mechanisms associated with stream intermittence.

Stream intermittency is projected to increase in extent, frequency, and duration as climate changes (Datry et al., 2014), with potentially important consequences for stream chemistry. In the western US, 80% of streams experience drying (US Geological Survey USGS, 2006)¹, and by 2050, intermittent streams and ephemeral streams (those that only experience event-based flow) are projected to increase by 5–7% (Döll and Schmied, 2012). In basins with moderate relief and elevation, such as the Snake, Great Salt Lake, and Oregon Closed Basins, snowmelt-fed streams are particularly susceptible to future stream drying as these areas may experience 100% loss of snow by 2050 (Klos et al., 2014). In most of the western US, projections include earlier snowmelt (Stewart et al., 2004; Abatzoglou et al., 2014). Although watershed geomorphology and vegetation determine hydrologic responses to precipitation shifts (Tague and Grant, 2009; Godsey et al., 2014), potential climate change impacts to streams include early-season drying (Datry et al., 2014; Jaeger et al., 2014; Tennant et al., 2015), lower base flows, and increased

groundwater sourcing (Tague and Grant, 2009; Godsey et al., 2014). Stream surface water chemistry patterns will likely reflect and respond to these anticipated changes, but the patterns and processes dominating these responses are complex because of potential feedbacks.

Coincident with changing snowmelt and low-flow patterns, wildfire is also increasing in its frequency and spatial extent in the western US (Abatzoglou and Williams, 2016; Parks et al., 2016), which alters land cover and influences stream chemistry (Hauer and Spencer, 1998; Mast et al., 2016). In particular, fire combusts vegetation and biotic soil components and releases mineral forms of nutrients and ions into soils (Murphy et al., 2006; Rau et al., 2007) that can be transported into streams (Spencer et al., 2003; Mast et al., 2016). Fire impacts to the water budget of a catchment can be substantial, especially in forested systems (Kinoshita and Hogue, 2015; Wine and Cadol, 2016; Atchley et al., 2018). Reduced terrestrial vegetation can lower evapotranspiration (ET) rates (Poon and Kinoshita, 2018), which can increase stream flow (Kinoshita and Hogue, 2015; Costigan et al., 2016; Wine and Cadol, 2016; Atchley et al., 2018), increase sediment yields (Subiza and Brand, 2018; Vega et al., 2020), and decrease terrestrial carbon inputs (Bixby et al., 2015; Cooper et al., 2015). Moreover, fire opens up the stream canopy increasing radiation, temperature, and wind (Naiman and Sedell, 1980; Poole and Berman, 2001; Bixby et al., 2015), which can increase evaporation (Poole and Berman, 2001; Maheu et al., 2014) and in-stream primary production (Davis et al., 2013; Rugenski and Minshall, 2014; Cooper et al., 2015). To date, most studies have focused on forested watersheds with less attention paid to seasonally snow-dominated mountain areas where sagebrush steppe vegetation often predominates. In these ecosystems, fire regimes are shifting with the spread of invasive *Bromus tectorum* (cheatgrass; Bradley, 2009; Bradley et al., 2016) and increasing fire frequency and fuel availability (Link et al., 2006; Abatzoglou and Kolden, 2011). The few studies conducted in these ecosystems have rapid recovery of ET losses due to increased grass and herbaceous cover with only marginal and short-term effects on streamflow (Flerchinger et al., 2016; Fellows et al., 2018). Because stream drying and fire are accelerating, there is a critical need to understand how both phenomena will affect spatiotemporal patterns of stream surface water chemistry in these contexts.

Natural drivers of stream drying and stream chemistry patterns include increases in surface water evaporation (Brooks and Lemon, 2007; Gallo et al., 2012) or ET losses (Poon and Kinoshita, 2018; Warix, 2020) and/or changes in subsurface connectivity (Brooks et al., 2015; Costigan et al., 2016); fire may have consequences for these processes as well (Kinoshita and Hogue, 2015; Wine and Cadol, 2016; Atchley et al., 2018; Poon and Kinoshita, 2018). For example, increased in-stream evaporation may cause evapoconcentration, or an increased concentration of solutes as the amount of water decreases as found in hot desert streams at low flows (Brooks and Lemon, 2007; Gallo et al., 2012). Under low-flow conditions, evapoconcentration may occur in patches of open canopy stream that experience particularly high radiation, and contribute to overall patchiness in stream solute patterns. Fire may promote

¹<http://nhd.usgs.gov/index.html>

such processes by opening stream canopies (Cooper et al., 2015), which can increase evaporative processes in streams by increasing radiation and wind (Maheu et al., 2014) or increase streamflow by reducing evapotranspiration (Kinoshita and Hogue, 2015; Costigan et al., 2016; Wine and Cadol, 2016; Atchley et al., 2018; Poon and Kinoshita, 2018). Alternatively, patchiness may arise from subsurface processes like changing patterns of hillslope connectivity and water residence times that influence groundwater inputs of water and solutes (Zimmer and McGlynn, 2017; Dohman, 2019; van Meerveld et al., 2019). Dynamic dissolved organic carbon (DOC) patterns in streams have been linked to springtime upland snowmelt and lateral flushing in alpine catchments of the Rocky Mountains (Boyer et al., 1997). However, the flushing process in the headwaters at Reynolds Creek Critical Zone Observatory (RC CZO) located in southwest Idaho was found to be slightly more complex (Radke et al., 2019) than Boyer et al.'s (1997) lateral flushing paradigm. At RC CZO, geophysical and hydrochemical evidence pointed to vertical and then lateral springtime flushing. The prior year's soil-water and DOC flushed vertically to the saprolite during snowmelt and then laterally along this interface to the stream. DOC in the stream was primarily allochthonous, presumably sourced from DOC that had accumulated in the soil matrix during the dry summer rather than from in-stream processes (Radke et al., 2019).

As streams dry, connectivity to hillslope carbon sources may shift and in-stream primary production may also be patchy. The resulting stream chemistry reflects the relative magnitudes of these processes and whether these changes occur synchronously or not. Both autochthonous (in-stream) and allochthonous (terrestrially produced) sources of carbon may shift as hydrologic and biologic inputs change seasonally, with stream size, or following fire (Minshall et al., 1989; Cooper et al., 2015). For example, high DOC concentrations can result from autochthonous carbon production during low flows in desert streams in Arizona (Jones et al., 1996; Brooks and Lemon, 2007). Similarly, in warm, humid systems in Tennessee, low flows and open canopy in autumn are associated with high DOC concentrations driven by increased autochthonous processing (Mulholland and Hill, 1997). Immediately following fire, stream algal blooms can result from elevated nutrient levels and open canopy that increase sunlight while also reducing the amount of available allochthonous carbon (Cooper et al., 2015). The chemical signatures of the organic carbon in the dissolved organic matter (DOM) can characterize organic carbon as autochthonous or allochthonous in origin. Fluorescence Index (FI) is a metric of spectral properties of fulvic acids component of DOM and absorbance coefficient ($a_{254}[\text{m}^{-1}]$) is a metric of carbon aromaticity; both tools help to evaluate autochthonous or allochthonous sourcing (McKnight et al., 2001; Brooks and Lemon, 2007; Inamdar et al., 2012). The connectivity of streams to the terrestrial surroundings may impact the variability of DOM sourcing. We expect that smaller headwater catchments may be more hydrologically connected to proximal terrestrial carbon (Hornberger et al., 1994; Brooks et al., 1999) and thus, more impacted by terrestrial processes (Creed et al., 2015). However, hillslope connections may also vary more in low-flow conditions and across the seasonal streamflow recession following snowmelt.

In this exploratory study, we measured spatial patterns of biogeochemistry in two mountainous, intermittent headwater streams as they dried. We sampled along the length of each stream, hereafter referred to as the longitudinal sampling or patterns. Sampling occurred following snowmelt through mid-summer in both streams: one unburned stream and one recently burned stream. We used a spatially continuous sampling approach over 2.5 km repeated over 3 months during the growing season to assess changes in solute sourcing patterns and processes. We asked three questions: (1) How does surface stream chemistry vary longitudinally and temporally in intermittent streams as they begin to dry? (2) What are potential driving processes that can explain these patterns? and (3) How do these patterns and processes change following fire? We hypothesized stream chemistry would vary spatially downstream as a function of different physical and biological processes, that spatial patterns would shift temporally as a function of seasonal drying, and that these shifts would be more pronounced following fire. Specifically, we hypothesized that carbon concentrations would increase owing to in-stream evapoconcentration, groundwater connectivity, and in-stream primary production (Table 1). On the other hand, we hypothesized that nutrient concentrations would decrease owing to increased uptake with drying, with higher nitrogen losses following fire in the burned stream owing to an open canopy and high nitrogen availability (Table 1). To test our hypotheses, we performed an initial cluster analysis to evaluate similarity and dissimilarity amongst chemical properties including solute concentrations, water isotopes, and spectral indices. We then evaluated spatial and temporal patterns and tested hypotheses about associated processes using a combination of these properties that emerged as relevant to our questions.

METHODS

Experimental Design

Study Site

Our study took place at the Reynolds Creek Experimental Watershed (RCEW) and RC CZO, a 239 km² watershed located southwest of Boise, ID, USA. The USDA Agricultural Research Service (ARS) established RCEW in 1960 as an experimental watershed representative of the Intermountain West region in the United States (Marks et al., 2011), and the ARS has monitored long-term precipitation and stream discharge trends (Seyfried et al., 2000, 2018). The RCEW extends over a steep climatic gradient with mean annual precipitation varying from 250 to 1,100 mm/yr and mean annual temperatures from 5.5 to 11°C. This climatic variability is driven by the nearly 1,000 m elevation range. At lower elevations, rain is the dominant form of precipitation in the RCEW whereas snow is dominant at the highest elevations (Nayak et al., 2010; Kormos et al., 2014). Peak stream discharges across the watershed are driven by snowmelt patterns (Pierson et al., 2001). Vegetation includes Wyoming sagebrush steppe in the lower elevations, transitioning to mountain big sagebrush (*Artemisia tridentata*), antelope bitterbrush (*Purshia tridentata*), rabbitbrush (*Chrysothamnus viscidiflorus*), western juniper (*Juniperus occidentalis*), aspen (*Populus tremuloides*) and coniferous forest [mostly Douglas fir

TABLE 1 | Outlines the study framework and aims (left two columns under “Study aims” in bold outline), expectations, and rationale for stated expectations.

Study aims		Expectations as streams dry (April to June) for unburned and burned streams		Rationale
Patterns observed	Spatial	Analytes will group into those that behave more conservatively and those that behave more reactively. Distinct longitudinal patterns will be observed, and these will differ between groups of analytes. <i>Approach:</i> solute concentrations, isotope, and spectral indices collected at high-spatial scope (50-m intervals over 2,500 m). Initial cluster analysis.		Source/sink dynamics will become more pronounced due to less surface water connectivity and increased local in-stream production and/or unique chemical signature of deeper groundwater sources with distinct flow path chemistry; this will be greater following fire when nutrients are high and uptake/biological production can be highly heterogeneous in the stream.
	Temporal	Changes in the longitudinal spatial patterns and increases in conservative concentrations; decreases in bio-reactive solutes with greater shifts in burned stream. <i>Approach:</i> repeated solute concentration measurements across 3 months (April, May, and June) as stream shifts from snowmelt to low flow. Initial cluster analysis.		Higher conservative solutes concentrations during lower water flow due to decreased snowmelt and increased groundwater sourcing. Lower observed bio-reactive solute concentrations due to in-stream biotic uptake.
Processes and mechanisms tested	Physical hydrological	Evaporation, evapoconcentration, dilution	Temporal shifts with heavier stable water isotopes later in the season and increased evidence of in-stream evapoconcentration, greater shifts in burned stream. <i>Analyses:</i> stable water isotopes slopes, source/sink chloride dynamics.	As ambient air temperature increases, evaporative processes will increase including evapoconcentration, especially in burned stream where an open canopy offers no sun or wind protection.
		Groundwater influence	Increases in cation concentrations, shifts toward groundwater sourcing in water isotope intercept. <i>Analyses:</i> stable water isotopes and conservative solutes end members.	As snow melt diminishes, water will shift from shallow subsurface water to deeper groundwater.
	Biological	Autochthonous or allochthonous carbon sourcing	Shifts toward autochthonous sourcing, greater shifts in burned stream. <i>Analyses:</i> FI and abs coefficient $a_{245}[\text{m}^{-1}]$.	As growing season progresses and water velocity flows, nutrients will be more labile and in-stream production will contribute.

The approach and method employed to observe patterns and test processes are described under each expectation.

(*Pseudotsuga menziesii*), at higher elevations (Seyfried et al., 2018). The site is underlain by Cretaceous granites and volcanic rocks such as the Miocene Salmon Creek Volcanics (McIntyre, 1972).

To evaluate spatial and temporal variation in stream chemistry, we studied two headwater streams within the RCEW known to be intermittent, Johnston Draw (basin area 1.83 km²) and Murphy Creek (basin area 1.32 km²; Seyfried et al., 2000; Pierson et al., 2001; Patton et al., 2018). Johnston Draw was ideal for this study owing to the availability of streamflow data measured at a dropbox v-notch 90° weir (Seyfried et al., 2000; Godsey et al., 2018). We opportunistically studied Murphy Creek to evaluate possible extremes in spatial and temporal

variation in stream chemistry following wildfire. Specifically, the Soda Fire burned 68 km² of RCEW in August 2015, including Murphy Creek (Vega et al., 2020). Briefly, the fire was classified as moderate severity in the study area and consumed nearly all above-ground live vegetation resulting in more than 60% bare ground (bare soil, ash, and rock), with high sediment delivery to the stream (Vega et al., 2020). In October 2015, a dropbox v-notch 90° weir with pressure transducer for discharge measurements was re-activated in Murphy Creek. Hereafter we refer to Johnston Draw as “unburned JD” and Murphy Creek as “burned MC.”

Unburned JD and burned MC are comparable in basin area, discharge, elevation, mean annual precipitation (MAP), and

aspect (Seyfried et al., 2000; **Table 2**). Annual water yields for 2016 were lower than the long-term average for both watersheds (Glossner, 2019). Though some studies find increased water yield after fire due to loss of vegetation (Atchley et al., 2018), in burned MC, the 2016 water year represented the third lowest

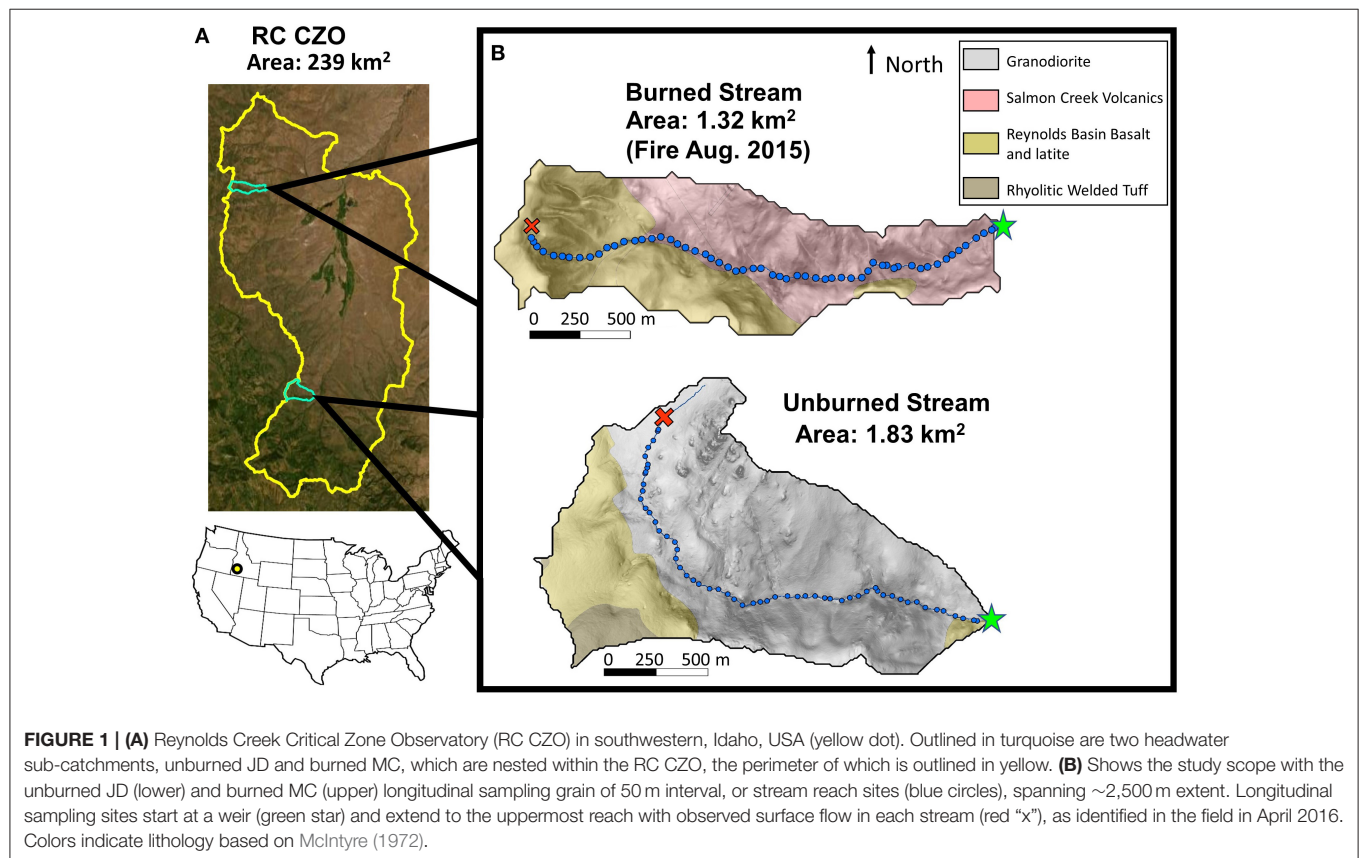
TABLE 2 | Characteristics of the unburned JD and burned MC sub-catchments.

	Unburned JD	Burned MC
Drainage area (km ²)	1.8	1.32
Streamflow direction	East with upper ~500 flowing south	East
Mean annual discharge (Q, m ³ /s)	0.007	0.007
Weir type	Dropbox v-notch 90°	Dropbox v-notch 90°
Elevation (m)	1,490–1,850	1,383–1,822
MAP (mm/year)	550	500–600
Lithology	Granodiorite	Salmon Creek Volcanics and basalt
2015 Fire	No	Yes
Primary vegetation	Mountain big sagebrush, juniper, alder, aspen, bunch grasses	(pre fire) Mountain big sagebrush, antelope bitterbrush, rabbitbrush, bunch grasses

streamflow year out of 12 years (1968–1977 and 2016–2017) of RC CZO data (Vega et al., 2020). The major differences between the two catchments are recent fire history and lithology (**Table 2**). The lithology of unburned JD is predominantly granodiorite with minor lithologic discontinuities including a combination of quartz latite and rhyolite flows underlying the high plateau at the top of the catchment, and a small olivine-rich basalt flow that extends into the outflow (McIntyre, 1972). Basalt and Salmon Creek Volcanics, underlies the burned MC. Unburned JD is described in more detail by Godsey et al. (2018) and Patton et al. (2018, 2019) and burned MC is described in more detail in Seyfried et al. (2000), Pierson et al. (2001), and Vega et al. (2020).

Sampling Design

To investigate spatiotemporal chemistry patterns in unburned JD and burned MC, we used a high-spatial scope approach: temporally repeated high-spatial density measurements at the km extent. In April, May, and June 2016, we sampled each stream at 50 m intervals (the grain of our study) over ~2,500 m distance (the extent of our study; **Figure 1**). Each 50 m site is referred to as a reach site (blue dots in **Figure 1**) and the reach sites span over ~2,500 m distance, the extent, which we refer to as the stream segment (**Figure 1**). In each stream segment, sampling started at the weir and continued to the uppermost observed surface flows (**Figure 1**) to include the entire sub-catchment. We refer to the suite of 50 m reach sites collectively as the stream longitudinal profile, and the spatial patterns identified using these sites as



longitudinal patterns. Water samples were collected in pools at the stream thalweg following Dent and Grimm (1999).

Biogeochemical Characterization of Intermittent Streams

Physical and Chemical Field Properties

We characterized each stream reach site by measuring a suite of *in situ* properties including temperature ($^{\circ}\text{C}$), pH, dissolved oxygen (DO mg/L), and estimated canopy cover. Water temperature ($^{\circ}\text{C}$) and pH were measured at each stream reach site using an Oakton pH 110 Series probe (Vernon Hills, IL) calibrated with 4, 7, and 10 pH standard solutions; DO was similarly measured at each reach site with a YSI dissolved oxygen probe (Burlington, VT). Canopy cover was visually estimated in July at half of the stream sites using upward-oriented fisheye lens photos at pool height. Maximum canopy coverage was estimated at alternating sites from photos taken during full leaf-out. Canopy coverage was categorized based on the proportion of the photo covered by foliage (0–25, 25–50, 50–75, and 75–100%). Discharge (m^3/s) was calculated from continuous stage measurements using a stage-discharge rating curve only at the outlet weir of each stream segment (Figure 1B). We acknowledge that direct measures of discharge and associated water balance at stream reach site would have augmented our understanding of stream dynamics and local water balance, but this was beyond this study. Accurately measuring discharge throughout a stream network at low flows is methodologically challenging and time-consuming. Instead, we focus here on concentration patterns and use indirect and environmental tracer approaches described below to understand inputs to and outputs from each reach site.

Surface Water Collection

To establish biogeochemical patterns in each stream, we collected surface water samples in the field and then filtered and analyzed them in the lab for the concentrations of a suite of chemical constituents. In the field, samples were collected in 250 mL amber high-density polyethylene bottles (HDPE) bottles, which had been rinsed three times and pre-leached in 18.2 MOhm distilled (DI) water. Bottles were then rinsed three times in the field with the stream sample and then filled to eliminate headspace. Samples were transported by backpack out of the steep terrain (~ 35 –60 total per stream per sample period), and the water samples were kept refrigerated (4°C) until filtered and analyzed. For carbon and total nitrogen (TN) analysis, water samples were filtered within 72 h (typically within 24 h) in the lab using vacuum filtration through pre-combusted $0.7\ \mu\text{m}$ Whatman glass fiber filters (GFF) into DI- and sample-rinsed 60 mL amber HDPE bottles. Amber HDPE bottles were used instead of glass owing to transport hazards (Sanderman et al., 2009). The remaining sample was syringe-filtered through a $0.45\ \mu\text{m}$ Puradisc nylon filter into four 60 mL clear HDPE bottles for analysis of nutrients, anions, cations, isotopes, and fluorescence index. Samples were collected moving from downstream to upstream to minimize disturbance.

Laboratory Analysis

Stream chemistry was analyzed for a suite of chemical constituents, including dissolved inorganic carbon (DIC), DOC, TN, nutrients including ammonium-N, nitrate (NO_3^- -N), orthophosphate (PO_4^{3-} -P), anions including chloride (Cl^-) and sulfate (SO_4^{2-}), and cation concentrations including base cations and rare earth elements such as strontium (Sr). In addition, we analyzed stream water for stable water isotopes ($\delta^{18}\text{O}$ and $\delta^2\text{H}$), fluorescence index (FI), and absorbance coefficient (a_{254} [m^{-1}]). FI and a_{254} methods are discussed in the DOC sourcing section. DIC, DOC and TN concentrations were measured on a Shimadzu TOC-V CSH (Columbia, MA, USA) equipped with an ASI-V autosampler and TNM-1 chemiluminescence detector for TN. Errors <2 –3% were accepted for concentrations $>1\ \text{mg C/L}$. High DIC concentrations (~ 10 –30 mg DIC/L as C) resulted in difficulty using the non-purgeable organic carbon method (NPOC). Thus, we calculated DOC as the difference between TC and DIC and propagated the errors associated with this calculation (~ 0.3 –0.9 mg/L). Cross validation of our DOC values with the Perdrial Environmental Biogeochemistry Lab at the University of Vermont (Burlington, VT) showed reasonable agreement ($n = 140$, $r = 0.65$). Owing to the higher error associated with DOC by difference, we approached DOC concentrations with some caution but retained them because DOC and spectral characteristics showed similarly variable patterns, and DOC provided an interesting contrast to other nutrients and DIC. Nutrients were measured on a Westco Discrete Analyzer (Unity Scientific, Brookfield, CT, USA), an automated chemical spectrophotometer. We accepted a $<10\%$ error for NO_3^- -N and PO_4^{3-} -P concentrations $<1\ \text{mg/L}$ and a 20% error for NH_4^+ -N $<0.10\ \text{mg/L}$. For most of the analyses, we utilize TN because both NO_3^- and NH_4^+ were below detection limit, with the exception of early season samples and those from burned MC. Anions Cl^- and SO_4^{2-} were analyzed on a Dionex ion chromatograph (ICS-5000, Sunnyvale, CA, USA) with AS18 column 4 X 250 mm, and we accepted $<10\%$ error for sample $<1\ \text{mg/L}$ and 2–3% for samples $>1\ \text{mg/L}$. Cations were measured by the Center for Archaeology, Materials and Applied Spectroscopy (CAMAS) lab (Idaho State University, Pocatello, ID) on a Thermo X-II 283 series Inductively Coupled Plasma Mass Spectrometer (ICP-MS) equipped with a Cetac 240-position 284 liquid autosampler (ThermoFisher Scientific). Dilutions were 1:10 sample to de-ionized water. Some cations were below the detection limit of 10 ppb. We reliably measured Sr, Ba, Na, Mg, Al, Si, K, Ca, Sc, Ti, Fe, and Zn for all 3 months. Lastly, stable water isotope samples were analyzed at ISU/CAMAS Stable Isotope Laboratory on a Thermo Scientific, High Temperature Conversion Elemental Analyzer (TC-EA) interfaced to a Delta V Advantage mass spectrometer. Precision for both $\delta^{18}\text{O}$ and $\delta^2\text{H}$ was better than $\pm 0.2\text{‰}$ and $\pm 2.00\text{‰}$ respectively.

Lastly, autochthonous and allochthonous DOC sourcing was investigated by analyzing the spectral characteristics of DOM in the lab, specifically Fluorescence Index (FI) and absorbance. Completed by the Perdrial Environmental Biogeochemistry Lab at the University of Vermont (Burlington, VT), these characteristics were measured using the Aqualog Fluorescence

and Absorbance Spectrometer (Horiba, Irvine, CA, USA). The excitation (EX) wavelength range spanned from 250 to 600 nm (increment 3 nm) and emission (EM) ranged from 212 to 619 nm (increment 3.34 nm). All excitation emission matrices (EEMs) were blank-subtracted (nanopure water, resistivity 18 M Ω cm⁻¹), corrected for inner filter effects, and Raman-normalized (Ohno, 2002; Miller et al., 2010). All samples were diluted to absorbance values below 0.3 and we computed relevant indices, including the FI (calculated as the intensity at Emission 470 nm divided by the intensity at Emission 520 nm for Excitation at 370 nm (Cory and McKnight, 2005). Absorbance (a) at 254 nm can also be used as a direct measure of aromaticity of DOM. We report the absorption coefficient (a_{254} [m⁻¹]) which is calculated independently of DOC concentrations and using Equation 1 (Green and Blough, 1994; Inamdar et al., 2012) below:

$$a_{254}[\text{m}^{-1}] = (\text{UV absorbance at 254 nm}) \times 2.303 \times 100. \quad (1)$$

Even though specific UV absorbance (SUVA₂₅₄) is the most commonly used indicator for DOM aromaticity (Weishaar et al., 2003), we refrained from its use due to the high error in DOC measurements and report a_{254} [m⁻¹] instead.

Spatiotemporal Pattern Analyses

Initial Clustering of Water Properties and Biogeochemistry

To assess possible common synchronous or asynchronous drivers of biogeochemistry, we initially identified chemical variables within each stream that were strongly and weakly related over time using cluster analysis. We explored highly correlated groupings among chemical variables described above using cluster variable analysis for each month in order to understand a given variable's relationship with other variables over time. The loaded variables were the chemical properties and analyte concentrations measured in the field and lab including temperature, pH, DO, carbon (DIC and DOC), anions (Cl⁻), nutrients (TN and PO₄³⁻), all cations measurable across the 3 months (Sr, Ba, Na, Mg, Al, Si, K, Ca, Sc, Ti, Fe, and Zn), and stable water isotopes ($\delta^{18}\text{O}$ and $\delta^2\text{H}$). The cluster variable analyses were conducted in JMP (version 14.2) which first grouped variables into cluster components determined by first and second principal components and then ranked variables within cluster components based on the strength determined by a cluster algorithm (SAS Institute Inc., 2017). This algorithm iteratively clusters variables based on a combination of strength of the principal component and relative eigenvalue strength. The order in which the analytes and chemical properties are reported within each cluster is based on the highest Pearson correlation coefficient (r) within the cluster. The results of these analyses informed our choice of analytes used for longitudinal pattern reporting and testing our specific process hypotheses. We chose analytes that juxtapose each other by selecting both highly ranked variables within clusters of strongly temporally correlated variables as well as analytes that were not strongly clustered. We did this in order to represent a spectrum of more conservative solutes such as Cl⁻ to more biological reactive ones such as TN in

each stream. The above criteria supported reporting longitudinal patterns for DIC, Cl⁻, Sr, DOC, TN, and PO₄³⁻. However, we assessed temporal patterns from April to June in all chemical properties and solutes as well.

Longitudinal Spatial Patterns and Temporal Tendencies

To explore the biogeochemical patterns both along the length of the stream and monthly changes as the stream dried, we plotted the longitudinal patterns of all solutes for each month, but focused on those identified above in the cluster analysis: DIC, Cl⁻, Sr, DOC, TN, and PO₄³⁻. These analytes also represent more conservative analytes and more reactive analytes as defined by Baker and Webster (2017). Stream data collected longitudinally from non-perennial streams presented challenges. First, samples collected 50 m apart in a stream are not statistically independent, second as streams dried April to June, it was impossible to sample in some locations and stream sample sizes changed, and third underlying longitudinal patterns complicated typical methods of quantifying spatial heterogeneity (details in statistical challenges of stream drying discussion section). Overall temporal comparisons of chemical properties, solutes, and cations from high to low flow were analyzed using mean differences and propagated errors from April to June within each stream. We determined differences to be substantial when they were larger than errors. As detailed above (Table 1), we expected to observe longitudinal and temporal similarities in both streams from April to June and predicted higher concentrations of carbon with evapoconcentration, lower concentrations of nutrients as in-stream primary production seasonally increased, and increasing cation concentrations as both streams became increasingly groundwater-dominated. However, we expected differences between the two streams like higher TN due to fire and higher cation loss due to volcanic lithology in burned MC.

Explaining Patterns: Testing Hypothesized Processes

Evaporative Processes

The potential evaporative signature of stream water was assessed by plotting stable water isotopes $\delta^{18}\text{O}$ and $\delta^2\text{H}$ against the Global Meteoric Water Line (GMWL) as well as the Local Meteoric Water Line (LMWL; $\delta^2\text{H} = 7.1(\delta^{18}\text{O}) - 6.3$), with data derived by Tappa et al. (2016). We compared monthly $\delta^{18}\text{O}$ and $\delta^2\text{H}$ data within each stream to the LMWL and calculated the monthly best-fit slopes based on simple linear regression and the LMWL-intercepts. To test for statistical differences between months and streams, we ran *t*-tests on the slopes of the regressed isotopic relationships. We hypothesized that the streams would undergo evaporation with drying in this semi-arid system (Fritz and Clark, 1997), and thus, that streams' isotopic data would fall below the LMWL (Gat, 1996). In addition, we predicted both streams would exhibit an increasingly evaporative signature as the streams dried, with a stronger shift in burned MC.

We evaluated the role of in-stream evapoconcentration in explaining chemistry patterns associated with stream drying by comparing the behavior of $\delta^{18}\text{O}$ to Cl⁻ between upstream and downstream sites. We calculated predicted downstream

values of Cl^- concentration ($[\text{Cl}_D^-]$) based on ratios of $\delta^{18}\text{O}$ from upstream (subscript “U” in Equation 2) and downstream (subscript “D”) samples and the observed upstream Cl^- ($[\text{Cl}_U^-]$) values (Mulholland and Hill, 1997; Gallo et al., 2012):

$$[\text{Cl}_D^-] = \frac{[\text{Cl}_U^-][\delta^{18}\text{O}_D]}{[\delta^{18}\text{O}_U]} \quad (2)$$

Because both $\delta^{18}\text{O}$ and Cl^- are expected to be conservative tracers (Baker and Webster, 2017), Equation 2 should lead to accurate predictions of Cl^- and each downstream site unless evapoconcentration or flushing were dominant processes. Therefore, we compared the predicted $[\text{Cl}_D^-]$ and observed Cl^- concentrations for each downstream site. If evapoconcentration were driving patterns in the water samples, we would expect to observe differences in the predicted vs. observed Cl^- concentrations. To explain any differences, we would need to parse out the role of flushing vs. evapoconcentration (Gallo et al., 2012).

Groundwater Influence

We used water isotopes and the Sr/DIC ratio to evaluate the potential influence of groundwater as streams dried. First, we calculated unburned JD's and burned MC's isotopic intercepts with the LMWL (Tappa et al., 2016) using each monthly regression (Fritz and Clark, 1997) and evaluated temporal shifts. We then compared the stream's LMWL-intercepts to the mean rain, snow, and groundwater samples collected by Radke et al. (2019) in a nearby but higher elevation watershed within RC CZO (Reynolds Mountain East ~2,100 m elevation). In this way, isotopic means of rain ($\delta^2\text{H} -66.5 \pm 12.7$ and $\delta^{18}\text{O} -8.4 \pm 2.4$, mean $\pm$ SE), snow ($\delta^2\text{H} -129 \pm 10.2$ and $\delta^{18}\text{O} -16.8 \pm 1.3$, mean $\pm$ SE) and groundwater samples ($\delta^2\text{H} -121 \pm 1.04$ and $\delta^{18}\text{O} -16.4 \pm 0.18$, mean $\pm$ SE) were assumed to be reasonable endmembers for our sites as well. As the streams dried, we expected the stream-LMWL intercepts to shift toward the groundwater signature later in the season if groundwater made up a larger component of the streams' surface water.

We also plotted Sr against DIC for each month because a previous study showed Sr to be a good indicator of groundwater at the RC CZO (Radke et al., 2019). We calculated RC CZO groundwater and precipitation (rain and snow) averages from samples collected by Radke et al. (2019) in late summer (August 2017), and used these values as endmembers in our analysis. Sr averaged 142.74 ± 8.77 ppb (mean $\pm$ SE) and DIC averaged 21.20 ± 0.55 mg C/L (mean $\pm$ SE). Rain and snow water samples had average values of 1.37 ± 1.63 ppb (mean $\pm$ SE) and 1.54 ± 1.80 mg C/L (mean $\pm$ SE) for Sr and DIC, respectively (Radke et al., 2019). We compared the upstream/downstream Sr-DIC relationship by comparing the uppermost 1,000 m sites with those in the lowest 1,000 m. We expected to observe a shift toward the groundwater ratio reflected in the late summer Sr/DIC ratios in unburned JD and burned MC that would be consistent with the shift in isotopic signatures.

Autochthonous or Allochthonous DOC Sourcing

Allochthonous carbon can be spectrally distinguished from autochthonous due to the structural carbon components in terrestrial vegetation, like lignin, which are also more aromatic. FI and a_{254} [m^{-1}] measure these spectral components and help classify carbon sources as allochthonous or autochthonous (McKnight et al., 2001; Inamdar et al., 2012). We compared FI to published ranges of these values associated with more autochthonous or allochthonous sources (Inamdar et al., 2012). Based on these values, we interpreted FI values between 1.2 and 1.5 as reflecting an allochthonous signature whereas FI above 1.7 reflects an autochthonous/microbial signature (McKnight et al., 2001). Furthermore, we report the variability using coefficient of variation (CV) because FI had no underlying longitudinal patterns in either stream (see discussion section, “Statistical Challenges of Stream Drying Data”). However, in our calculations, we controlled for sample size in each stream by including only those sites that had water flowing all 3 months so as not to bias any one month.

Aromatic and humic DOM is a product of terrestrial vegetation so that a higher aromatic a_{254} signature reflects a more allochthonous DOM with structural carbon. Thus, the relationship between FI and a_{254} is inverted, such that lower a_{254} result from less aromatic DOM, more autochthonous/microbial derived carbon and higher a_{254} result from more aromatic, more allochthonous derived carbon (Inamdar et al., 2012). Due to increased in-stream primary production as both streams dried, we expected to observe FI to increase reflecting shifts from more allochthonous to autochthonous carbon, and a_{254} to decrease reflecting diminished aromatic properties with a more pronounced shift in burned MC.

RESULTS

Biogeochemical Characterization of Intermittent Streams

Stream Hydrologic Conditions

Seasonal drying in both intermittent headwater streams was similar, with weir discharges (Q) decreasing in April, May, and June 2016 (Figure 2A). Drying occurred in both streams between April and June. For unburned JD, all of the April sampling sites ($n = 57$) had surface flow, 89% persisted in May ($n = 51$), and only 65% sustained surface flows in June ($n = 37$). Drying occurred in reaches interspersed throughout the unburned JD stream. Similarly, all of the reach sites in burned MC were flowing in April ($n = 59$), but only 73% in May ($n = 43$), and 64% sustained flow in June ($n = 38$). Unlike in unburned JD, drying in burned MC occurred from the top of the stream to the bottom as though the entire stream contracted longitudinally (though subsequent work showed this pattern did not persist later in the summer; Warix, 2020). Thus, by June, 35% of unburned JD's and 36% of burned MC's reach sites had dried, with the highest proportion of sites drying from May to June in unburned JD stream and from April to May in burned MC stream. Over the sample period, unburned JD stream discharge exhibited a slightly greater relative change (April Q $\sim 0.022 \text{ m}^3/\text{s}$ to June Q < 0.005

m^3/s) than burned MC (April Q $\sim 0.017 \text{ m}^3/\text{s}$ to June Q $\sim 0.005 \text{ m}^3/\text{s}$; **Figure 2A**). Burned MC also exhibited flashier stream flow as spikes in discharge, presumably responding to event-based precipitation or snowmelt inputs (**Figure 2A**) that did not occur as strongly in unburned JD.

Physical and Chemical Field Properties

The streams were similar in physical and chemical properties measured in the field but as expected, canopy cover differed between the unburned stream and burned stream. Stream temperature showed strong longitudinal and increasing temporal shifts in both streams, whereas pH and DO were more variable longitudinally and temporally (**Figures 2B–G** and **Supplementary Table 1**). The two streams differed in canopy cover in that 44% of unburned JD had $>25\%$ canopy cover during full leaf-out whereas 100% of sites sampled in burned MC had $<25\%$ canopy cover, and most sites fell very close to 0% canopy cover even in July 2016 (**Figures 2B,C**). Although some burned MC sites had $>0\%$ canopy cover in July, the post-fire vegetation differed from the vegetation in the unburned JD. The riparian vegetation of burned MC in July was dominated by fast-growing, non-woody shrubs and small, young willows because all vegetation, namely sagebrush and bunch grasses were charred to the ground (Vega et al., 2020). This contrasted with the overhanging riparian alder, willow, and juniper in unburned JD.

Biogeochemical Pattern Analysis Initial Clustering of Water Properties and Biogeochemistry

Correlations between stream analytes varied between streams and these relationships changed as each stream dried. We identified groups of solutes that behaved similarly or dissimilarly for each stream at different moments in the season. For example, in unburned JD, we observed positive correlations between Sr, Mg, Ca, K, Na, Ba, Fe, DOC, temperature (Temp), DIC, $\delta^{18}\text{O}$, TN, Cl^- , Sc, and Si in April. Ti, Al, DO, pH, PO_4^{3-} , and Zn were negatively correlated with these analytes and chemical properties, but positively correlated with each other (**Figure 3A**). By June, Mg, Fe, Cl^- , K, Temp, TN, $\delta^{18}\text{O}$, Sc, and Si were positively correlated with each other, but shifted to being negatively correlated with other analytes or chemical properties. From April to June, shifts from negatively to positively correlated analytes were observed between Ti, DO, pH, and PO_4^{3-} and other analytes or properties in unburned JD (**Figure 3E**). We observed similar (though not identical) patterns within burned MC (**Figures 3B,D,F**). Strongly correlated (either positively or negatively) analytes were placed into clusters for each month and stream (**Supplementary Table 2**). Based on these results, we selected representative analytes to illustrate different clusters and behaviors for the longitudinal analysis, including DIC, DOC, TN, PO_4^{3-} , Sr and Cl^- .

Longitudinal Spatial Patterns

Common longitudinal patterns were evident in both unburned JD and burned MC streams (**Figure 4**) for many of the more conservative analytes (e.g., DIC, Sr, and Cl^-) than the biologically reactive ones (e.g., DOC, TN, and PO_4^{3-}). As expected, mean concentrations generally increased for DIC, Sr,

and Cl^- downstream (**Figures 4A–F**). The exception to this pattern were consistently high concentrations at the top of unburned JD (relative to downstream concentrations) at reach sites that were south-facing and meadow-like. Concentrations of more reactive analytes, DOC, TN, and PO_4^{3-} , tended to decrease downstream in unburned JD whereas DOC and PO_4^{3-} were relatively invariant longitudinally in the burned MC stream. TN was exceptional in that it varied in the uppermost portions of both streams with patterns that changed each month of the sampling campaign.

Temporal Shifts

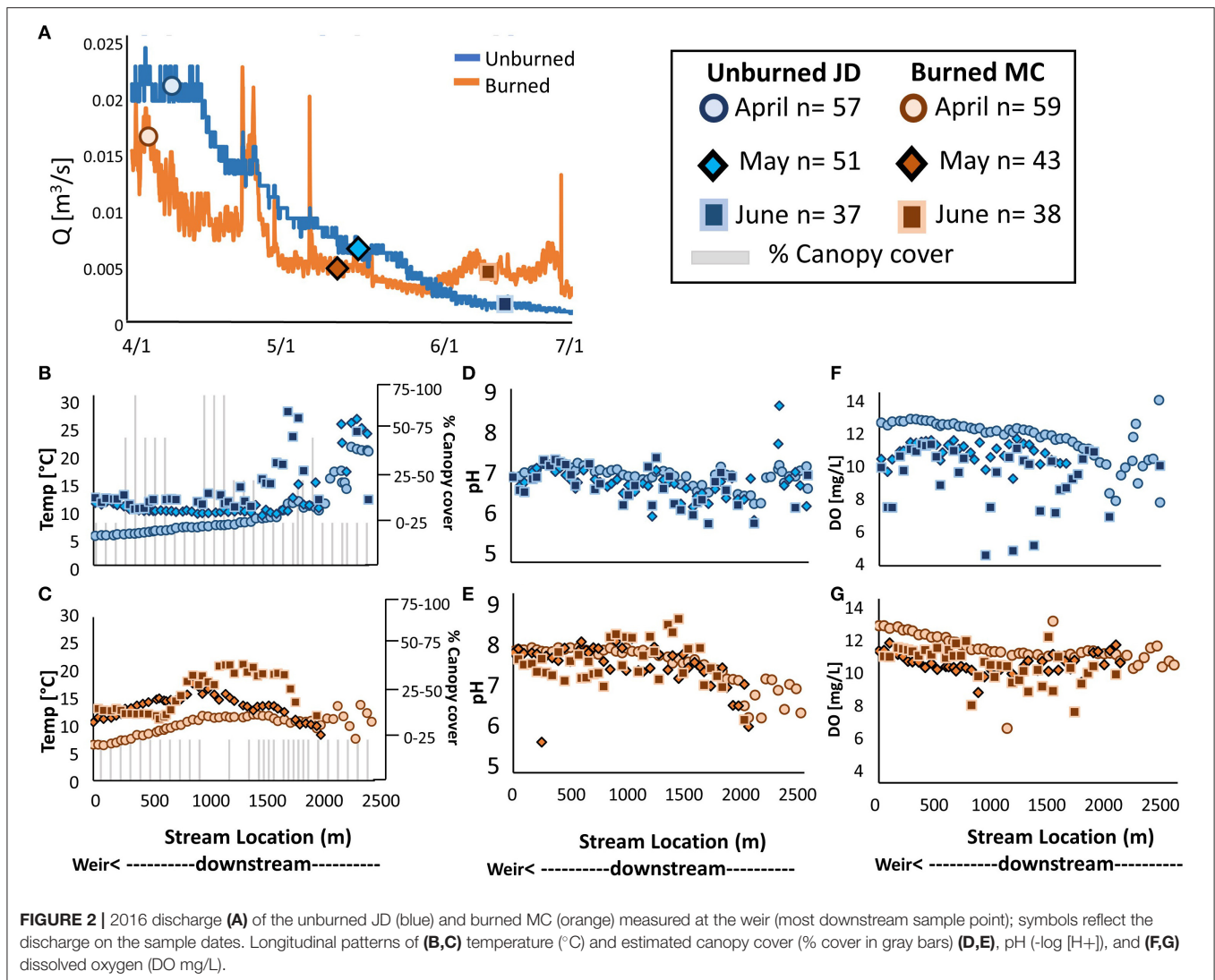
As expected, mean differences in concentrations between April and June showed that TN concentrations substantially decreased over the growing season in each stream, and by an order of magnitude in burned MC (**Figure 5B**). Mean TN concentration decreased by $>80\%$ from April to June (**Figure 5B**) in burned MC stream (1.14 ± 0.03 to $0.19 \pm 0.02 \text{ mg N/L}$, mean $\pm$ SE); TN was composed mostly of NO_3^- ; even in April, NH_4^+ was largely below our instrument detection limit. We observed a similar but less pronounced decrease in TN in unburned JD (**Figure 5A**), where a 39% decrease occurred over the same months (0.36 ± 0.04 to $0.22 \pm 0.03 \text{ mg N/L}$, mean $\pm$ SE). Unlike TN, PO_4^{3-} decreased only slightly in the burned MC stream over the growing season and did not exhibit a measurable change in unburned JD (**Figures 5A,B** and **Supplementary Table 3**). In contrast to our expectations, DOC exhibited no clear temporal shift from April to June in unburned JD and slightly decreased in burned MC (**Figure 5**).

In both streams, we observed similar temporal patterns in the average differences from April to June for base cations and other solutes (**Figures 5C,D**). Like DIC and Cl^- , most cations increased from April to June in both streams as indicated by negative difference values between April and June mean concentrations (**Figures 5C,D**). However, in unburned JD, some cation concentrations decreased as indicated by positive difference values as was the case for Si, Fe, and Ba. As expected, burned MC exhibited three to ten times larger cation differences than unburned JD for several cations including Na, Mg, Si, K, and Ca (**Figures 5C,D** and **Supplementary Table 4**).

Explaining Patterns: Testing Hypothesized Processes

Evaporative Processes

Isotopic signatures for both streams showed that evaporation was occurring, which we expected. However, counter to our predictions, we did not find evidence of temporal shifts toward increasing evaporation with stream drying in either stream. The regression slopes of the $\delta^{18}\text{O}$ - $\delta^2\text{H}$ relationships for all months for both unburned JD and burned MC fell below the GMWL and LMWL (Tappa et al., 2016), indicating an evaporative signature in these two streams compared to local precipitation (**Figures 6A,B**). In the unburned JD, we observed a lower-sloped evaporative trend with drying as April's slope ($\pm$ SE) was 2.7 ± 0.16 , May's was 3.6 ± 0.27 , and June's slope 4.7 ± 0.30 (**Table 3**). All months within unburned JD were statistically different so that April differed from May ($t = -5.2, p = < 0.0001, \text{df} = 103$), May differed from June ($t = -4.4, p = 3.7\text{E-}05, \text{df} = 81$), and



April from June ($t = -14.14$, $p < 0.0001$, $df = 84$). In the burned MC stream, April's slope ($\pm$ SE) was 4.3 ± 0.30 , May's was 3.0 ± 0.26 , and June's was 4.5 ± 0.23 (Table 3). In contrast to unburned JD, burned MC slopes for April and June did not statistically differ from each other ($t = -0.51$, $p = 0.61$, $df = 91$), although both differed from May ($t = -2.6$, $p = 0.01$, $df = 95$ and $t = -5.1$, $p < 0.0001$, $df = 76$, respectively), which had the most evaporative signature over the study period. Between the two streams, slopes were only significantly different from each other in April ($t = -10.24$, $p < 0.0001$, $df = 108$) and not in May ($t = 1.0$, $p = 0.32$, $df = 89$) or June ($t = 0.6$, $p = 0.56$, $df = 70$). We generally did not observe longitudinal patterns of evaporation (Figures 6A,B), with the exception of the uppermost portion of unburned JD. There, the aspect and vegetative cover of the stream was distinct from downstream reaches: these south-facing meadow sites had a strong evaporative signature in April (Figure 6A) and had dried by May.

In-stream Evapoconcentration or Dilution Influences on Solute Concentrations

To investigate potential effects of evaporation or dilution on the concentration of solutes as streams dried, we evaluated source-sink dynamics of chloride compared to $\delta^{18}\text{O}$. In both streams, most measured Cl^- fell within the error of the predicted Cl^- based on $\delta^{18}\text{O}$, indicating the two analytes behaved conservatively, with no major new gains or losses of water. Given this finding, we did not further investigate spatial or temporal deviations due to either evapoconcentration or dilution (Figures 6C,D). Although we hypothesized that in-stream evapoconcentration would be a driver of increased analyte concentrations, we found few sites in either stream that showed concentrations outside of the $\delta^{18}\text{O}$ -based prediction intervals. Additionally, counter to our predictions, we observed no differences between streams despite their difference in canopy cover.

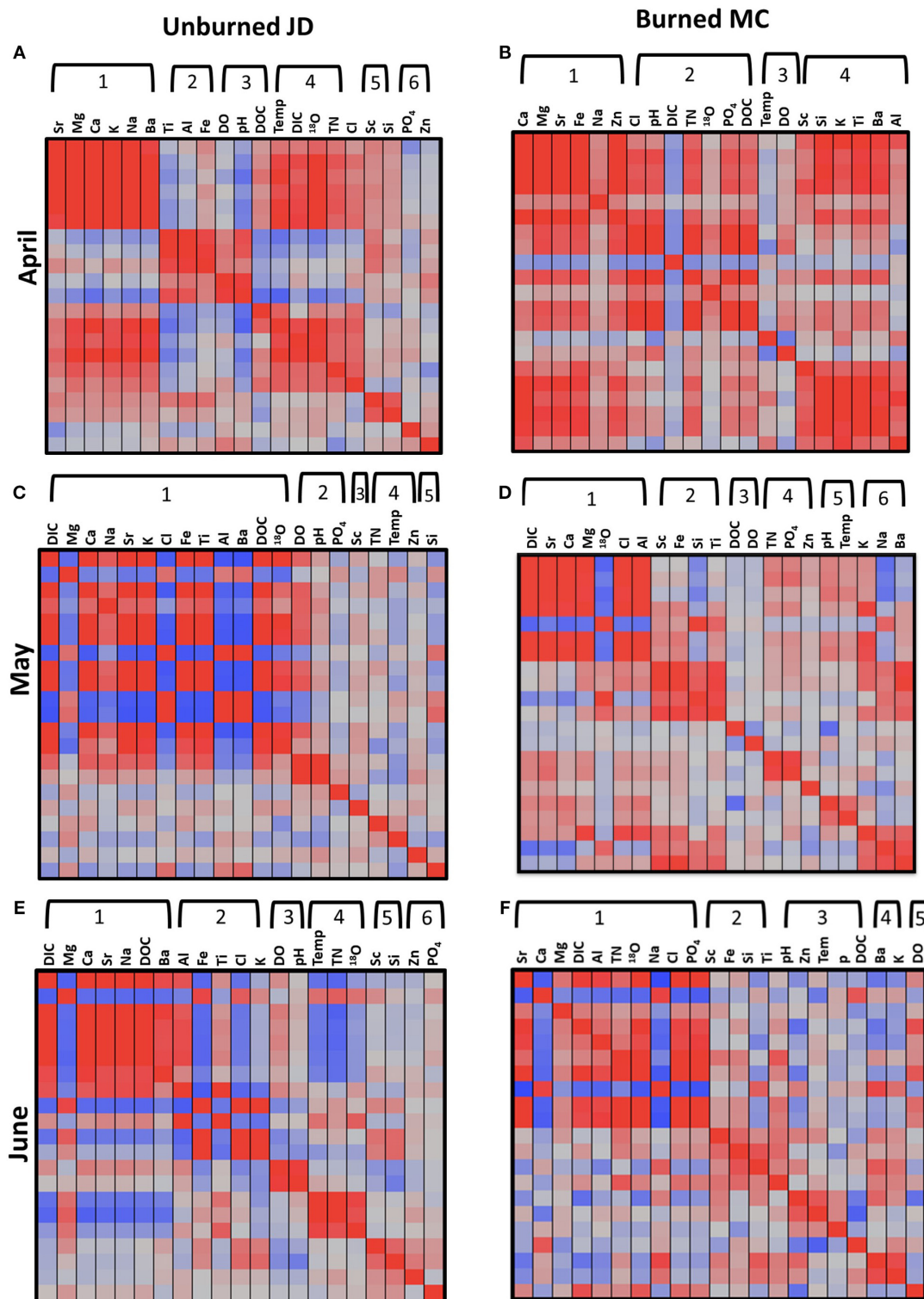


FIGURE 3 | Cluster analysis shows correlations between analytes by rows for April (**A,B**), May (**C,D**), and June (**E,F**) for unburned JD (left column) and burned MC (right column). Analytes include: temperature, or Temp ($^{\circ}\text{C}$), DO (mg/L), pH, DIC (mg C/L), Cl^{-} (mg Cl/L), DOC (mg C/L), TN (mg N/L), PO_4^{3-} (mg P/L), $\delta^{18}\text{O}$, and cations (ppb): Sr, Ba, Na, Mg, Al, Si, K, Ca, Sc, Ti, Fe, and Zn. Analytes are labeled across the top of each x-axis and are listed in the same order along the y-axis in each panel. Analyte order is based on cluster analysis groupings where analytes within each cluster are more strongly correlated (higher Pearson correlation coefficient or r -value) to each other than to other groups of analytes. The r -values are colored by saturation where the higher the r -value, the more saturated the color, and gray represents a low r -value and low correlation. Red indicates positive correlations and blue negative correlations.

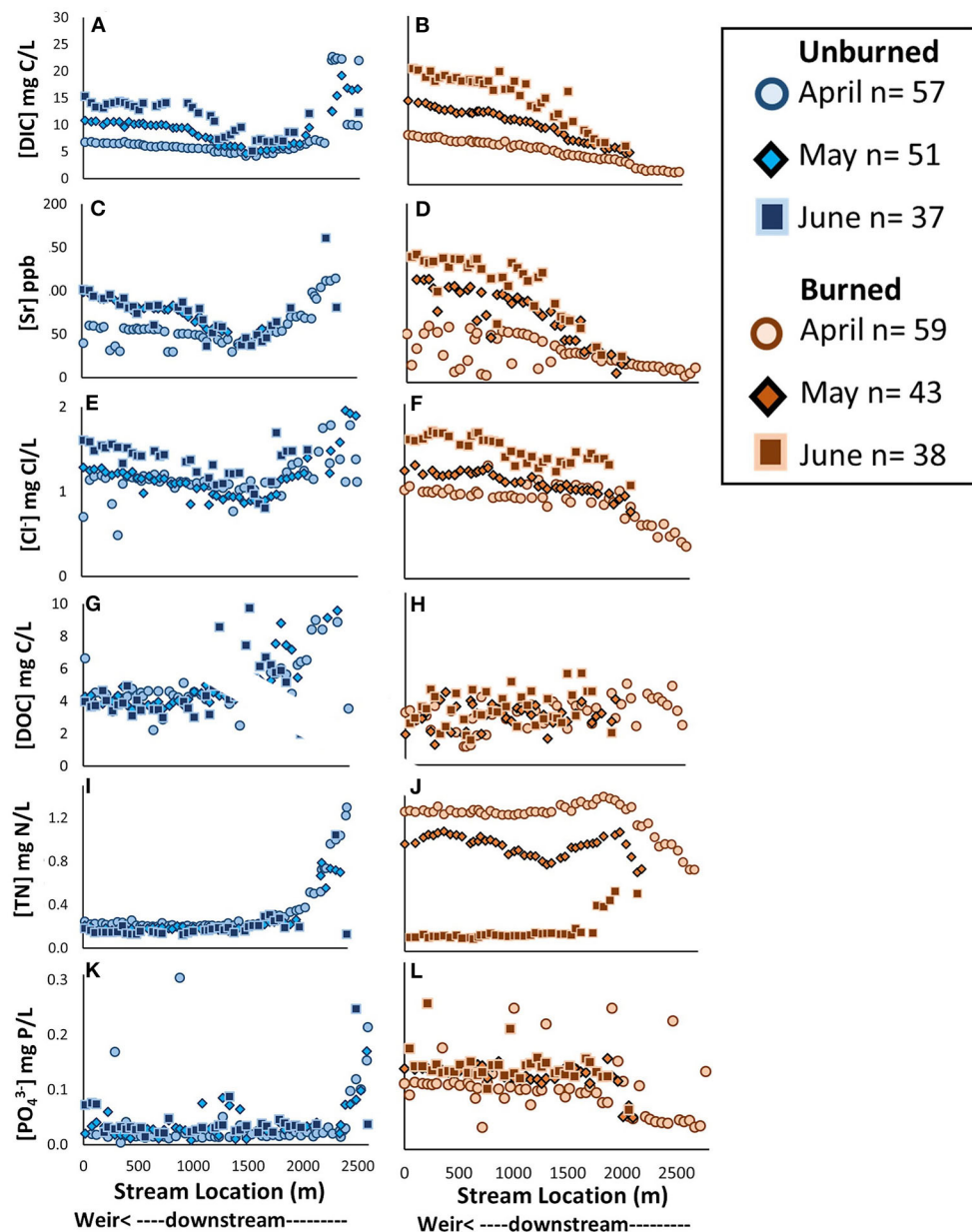


FIGURE 4 | Longitudinal patterns of (A,B) dissolved inorganic carbon (DIC mg C/L) (C,D), strontium (Sr ppb) (E,F), chloride (Cl⁻ mg Cl/L) (G,H), dissolved organic carbon (DOC mg C/L) (I,J), total nitrogen (TN mg N/L) (K,L), phosphate (PO₄³⁻ mg P/L) for unburned JD (blue, left column) and burned MC (orange, right column) streams. Values from April are represented by circles, May sites are diamonds, and June sites are squares along the length of the stream where 0 m is the weir and ~2,500 m is the upper stream reach.

Groundwater Influence

The isotopic signature of both streams showed surface water sources to be mixed contributions of snowmelt and rain: each stream's isotope values fell between the snow and rain mean values reported by Radke et al. (2019; **Figures 6A,B**). Based on intercepts, water in both streams had dominant contributions from snow/groundwater; snow sources for unburned JD reflected a larger rain contribution as indicated by isotopic-intercepts with the LMWL (~ -111 $\delta^2\text{H}$ per mil) whereas the LMWL-intercept

for burned MC stream (~ -120 $\delta^2\text{H}$ per mil) reflected a larger snow and/or groundwater contribution (**Figures 6A,B**). Because the groundwater and snow isotopic signatures were similar, we could not distinguish between the two sources with isotopic tracers. Counter to our expectations, no clear temporal shifts in water sources emerged in the isotope data in either stream (**Table 3**).

However, temporal shifts in groundwater-precipitation mixing were apparent by pairing the isotopic patterns with

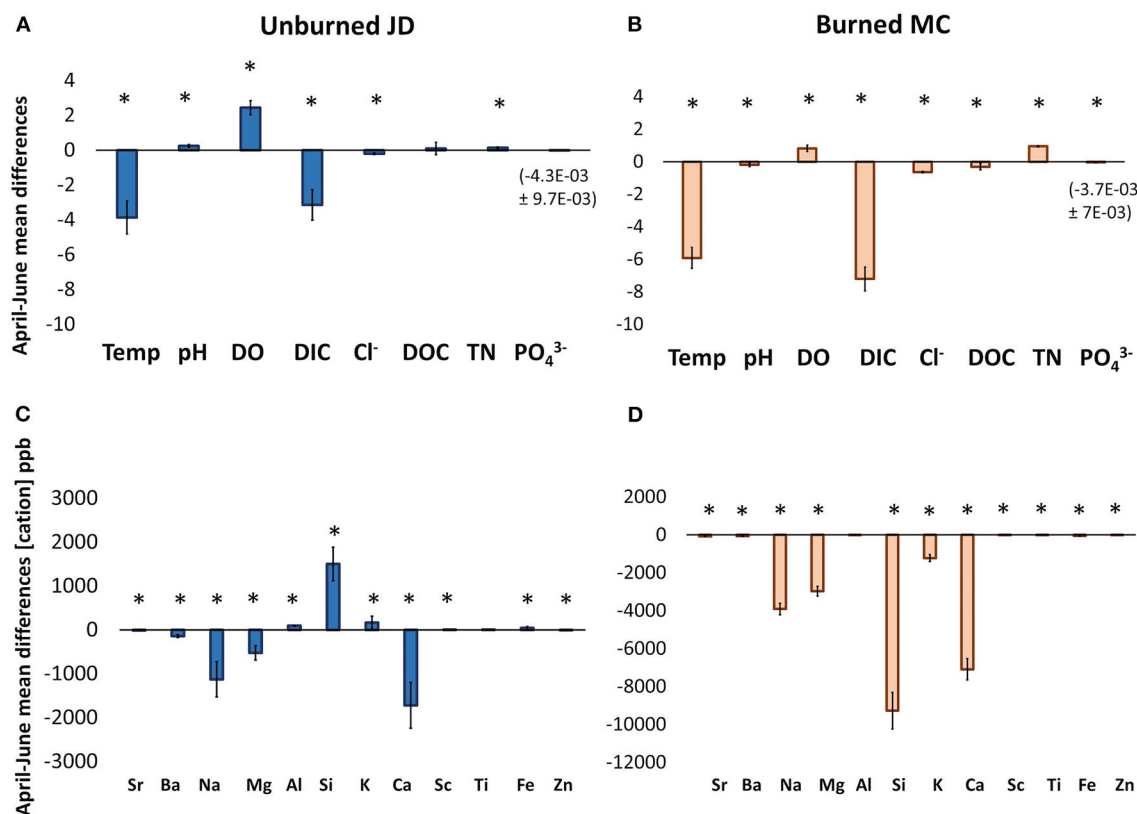
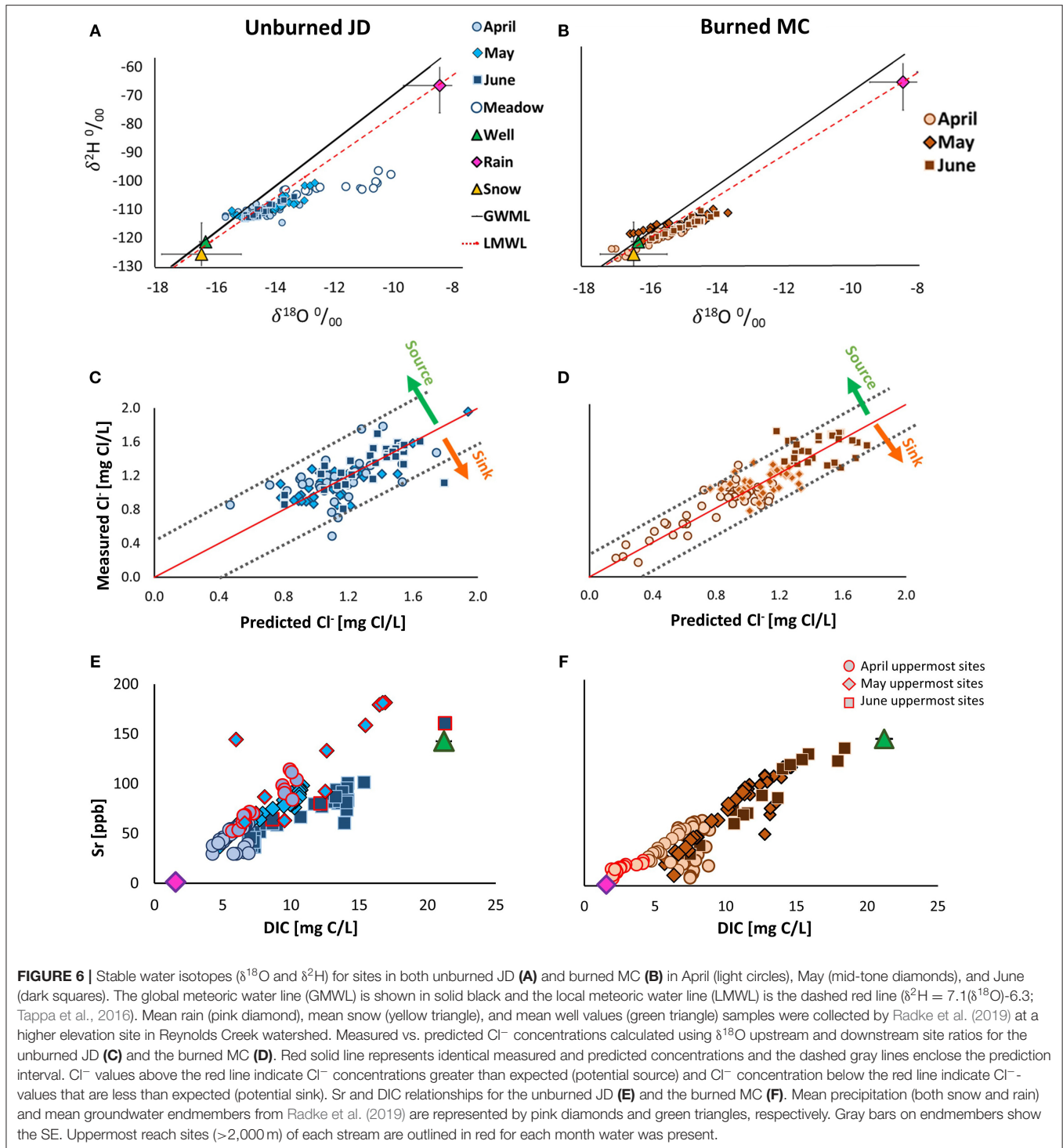


FIGURE 5 | Mean solute differences from April to June are shown for **(A)** unburned JD and **(B)** burned MC for temperature, or Temp, (°C), pH (-log [H⁺]), DO (mg/L), DIC (mg C/L), DOC (mg C/L), TN (mg N/L), PO₄³⁻ (mg P/L), Cl⁻ (mg Cl/L), DOC (mg C/L), TN (mg N/L), PO₄³⁻ (mg P/L). Mean cation differences (ppb) from April to June are shown for unburned JD **(C)** and burned MC **(D)**. Error bars show propagated error and asterisks indicate where differences are greater than the propagated error. Positive differences indicate decreasing in concentration from April to June and negative differences indicate increasing concentration from April to June (arrows for reference). Note pH measures are representative of log scale, so small changes are not equivalent to raw concentrations of other solutes. Additionally, please note the difference in axes scales between panel **(C,D)**.

observed shifts in DIC and other analyte concentrations. For example, we observed that DIC and Sr were strongly correlated (Figure 3). When compared to the average precipitation (rain and snow) and late summer groundwater Sr/DIC endmembers, stream water was sourced from precipitation in April and shifted toward groundwater sourcing in June in each stream (Figures 6E,F). In contrast to isotopes of water, rain and snow precipitation had similar DIC and Sr signatures and these signatures were substantially different from groundwater DIC and Sr signatures (Radke et al., 2019). We likely did not capture an additional endmember in unburned JD in the upper portion of the catchment (Figure 6E) whereas most of the variation in the burned MC was captured by two endmembers, potentially with exception of a few downstream sites (Figure 6F). The uppermost reach sites (>2,000 m) of each stream (flowing in April and May) had different signatures; the signatures from the upstream reaches in unburned JD fell close to groundwater and burned MC upstream reach sites fell close to a precipitation signature (Figures 6E,F and Supplementary Figure 1).

Organic Carbon Sourcing: Autochthonous and Allochthonous Processes

DOM sources and spectral characteristics varied along the profile of each stream during each month; FI varied by similar amounts in both streams whereas a_{254} [m⁻¹] varied more in unburned JD (Figure 7). Many FI values fell between the 1.5 allochthonous derived threshold and the 1.7 autochthonous/microbially derived threshold (Figures 7A,B), indicating mixed carbon sources in both streams. Unburned JD had a slight but surprising longitudinal FI pattern in April (mean 1.59 ± 0.01 SE) when we observed a more autochthonous/microbial FI signature downstream, while the upper reach of the stream had a more allochthonous signature (Figure 7A, circles). May generally had an allochthonous signature (mean 1.49 ± 0.003 SE) as did June (mean 1.50 ± 0.007 SE), and this similarity between seasons was counter to our predictions. Interestingly, while May and June mean FI values were similar, June exhibited greater variation (CV 2.8) than May (CV 1.2), with points spanning the thresholds between autochthonous and allochthonous sources in both months (Supplementary Table 5).



In burned MC (Figure 7B), we observed a mixed autochthonous and allochthonous FI signature in April (mean 1.55 ± 0.005 SE), and a similar but more varied FI signature in May (mean 1.50 ± 0.009 SE). June's FI varied along the stream profile with an overall downstream increase toward autochthonous/microbial signature (mean 1.62 ± 0.02 SE). This tendency matched our expectations of increasing

in-stream productivity, which we expected would be more pronounced in the burned stream. In addition, burned MC's higher mid-summer FI variation was reflected in the relatively high June FI CV (7.5) compared to that of April (1.0) and May (3.5; Supplementary Table 5). Despite the overall autochthonous tendency in June, there were also sites for which the FI value fell below or close to the allochthonous threshold

(1.5, **Figure 7B**, squares). We did not expect to observe this high variation.

We expected that as streams dried, DOC aromaticity (high in terrestrially derived carbon) would be decreasing due to increased in-stream productivity, especially in burned MC, and thus, we expected decreases in a_{254} values. Overall, unburned JD had higher a_{254} values that were more longitudinally and seasonally variable than burned MC (**Figures 7C,D** and **Supplementary Table 5**). We expected higher overall aromaticity in unburned JD. Though we expected to see higher FI values coupled with low a_{254} values, we did not observe these relationships consistently. In April, unburned JD had higher a_{254} values (mean 44.5 ± 1.4 SE) coupled with higher FI values (**Figure 7E**). The unburned JD maintained higher a_{254} values in May (mean 48.1 ± 3.0 SE) and dropped in June (mean 38.5 ± 2.9 SE); this decrease in aromaticity suggests less terrestrial carbon and followed our expectations. Compared to unburned JD, burned MC a_{254} values shifted less from April (mean 26.2 ± 0.48 SE) to May (mean 27.2 ± 0.54 SE) and June (mean 29.1 ± 0.53 SE, **Figure 7D**). The differences in a_{254} temporal patterns between the streams were illustrated by the relationship of FI and a_{254} (**Figures 7E,F**) where the spread of FI values was greater in the burned MC, but the spread of a_{254} was greater in the unburned JD. We hypothesized that increased in-stream productivity would result in shifts of increased FI and decreased a_{254} values. However, the spatial patterns we observed in both streams were more longitudinally varied and had less clear seasonal shifts than expected.

DISCUSSION

Stream intermittency and fire regimes are changing in the Intermountain West and cold montane shrubland ecosystems that characterize the northern Rocky Mountains and Great Basin are especially vulnerable to these changes. We hypothesized stream chemistry patterns would change seasonally and longitudinally in response to both drying and fire, and that process shifts would explain these biogeochemical patterns. Specifically, we hypothesized that with drying, in-stream evaporative processes, increased groundwater influence, and shifts in DOC sourcing from allochthonous to autochthonous would contribute to differences in these longitudinal stream chemistry patterns. In a recently burned watershed, we expected the impacts of these processes would be amplified.

Stream Drying: Non-linear Spatiotemporal Biogeochemical Patterns

Both streams exhibited non-linear longitudinal patterns in chemistry that changed with stream drying. Spatially variable and distinct patterns among analytes were revealed by this study's high-scope sampling approach and breadth of measured chemical constituents. For instance, we identified potential springs/deeper groundwater inputs where temperature or solute concentrations remained stable despite seasonal shifts at other locations longitudinally along the streams (e.g., around $\sim 1,500$ m

TABLE 3 | The slope $\pm$ standard error and the LWML-intercept extracted from the data presented in **Figures 6A,B**.

Isotope analyses		Slope	$\pm$ SE	LWML intercept (per-mil)
Unburned JD	April	2.7	0.16	-111.47
	May	3.6	0.27	-111.09
	June	4.7	0.30	-114.04
Burned MC	April	4.3	0.30	-122.36
	May	3.0	0.26	-117.04
	June	4.5	0.23	-121.73

in unburned JD and $\sim 2,000$ m in burned MC). In contrast to our expectations, we found similar longitudinal patterns and processes driving these patterns in both streams. This provides some evidence for emerging regional behavior in the subsurface processes (i.e., groundwater contributions) driving stream chemistry. Our findings collectively point to the power of high-scope sampling to capture both the temporal and spatial variability in intermittent stream processes, and they contribute empirical data on how spatiotemporal patterns and processes may change with stream drying in headwater streams like these.

Streams Similarities: DIC Dominance With Drying

Given the differences in fire history and lithology, the similarity of chemistry patterns and shifting processes between these two streams was surprising. In both unburned JD and burned MC, DIC and many cations exhibited clear longitudinal patterns that increased in mean concentration with stream drying. Though we did not directly analyze concentration-discharge relationships longitudinally at each 50 m reach site, chloride source/sink dynamics did not suggest strong spatial inputs or outputs in any month. Monthly increases in DIC and other conservative solute concentrations were consistent with dilution behavior of concentration-discharge observed over the water year at the weir of each stream (Glossner, 2019). While conservative solutes in larger streams often display chemostatic behavior (Godsey et al., 2009), our data is consistent with chemodynamic concentration-discharge relationships documented in smaller catchments (Hunsaker and Johnson, 2017) and intermittent streams (Bernal et al., 2019). Instead of evapoconcentration or dilution, our findings point to deeper groundwater sources driving increased concentrations as the streams dried. Our findings are consistent work by Wlostowski et al. (2020), whose findings describe loamy and sand-rich soils and relatively shallow bedrock in unburned JD resulting in more baseflow-dominated streamflow with less subsurface water storage compared to CZOs with clay-rich soils. Distinct from other baseflow dominated streams, groundwater contributed to low flow in our streams, not high flow as observed in the Jemez River Basin Critical Zone observatory in New Mexico (McIntosh et al., 2017). Our results build off of those reported by Radke et al. (2019) at a higher elevation RC CZO catchment during snowmelt, where deeper flow paths rather than shallow subsurface flow

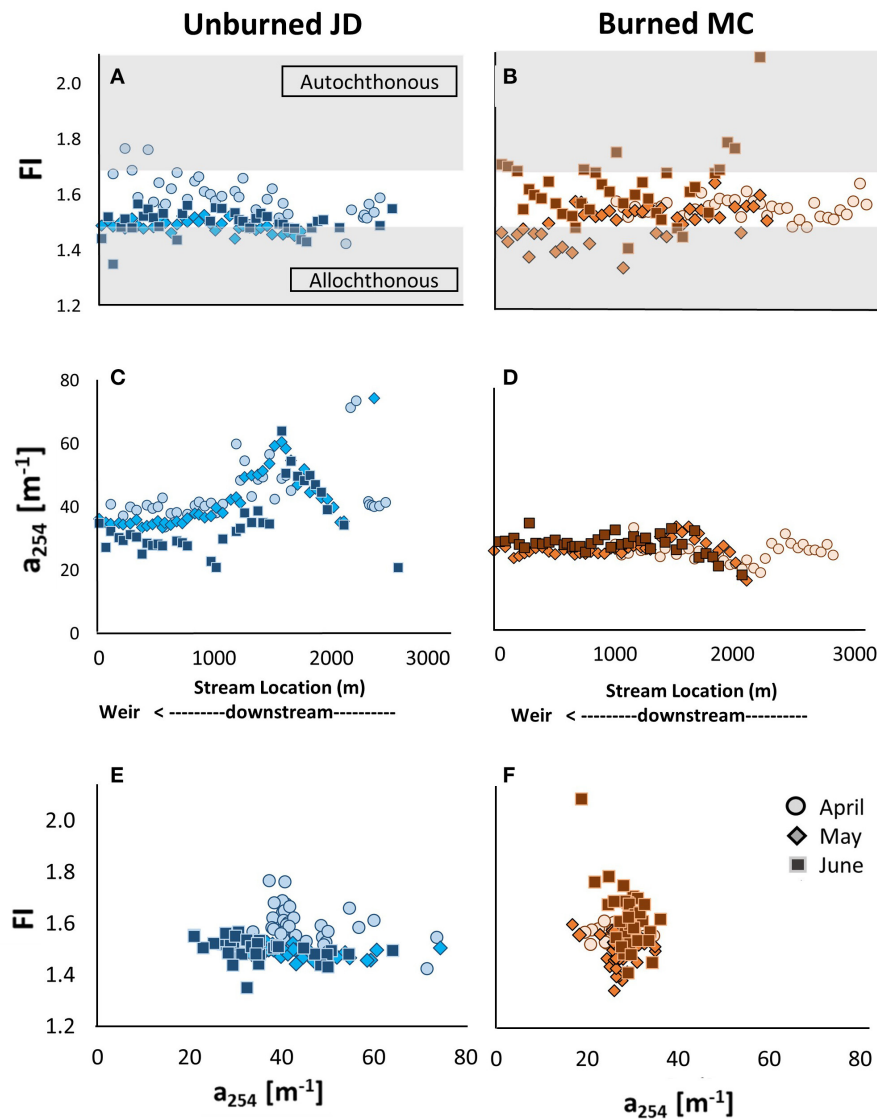


FIGURE 7 | (A,B) Fluorescence index (FI) (C,D), absorbance coefficient (a_{254} [m⁻¹]), and (E,F) the relationship of FI to a_{254} for stream profile of unburned JD (blue, left) and burned MC (orange, right); shape, shade, and color of months are described in **Figure 2**. Gray shaded boxes (A,B) indicate the 1.5–1.2 allochthonous-derived threshold and the 1.7–2.0 autochthonous/microbial-derived threshold.

paths contributed to stream DOC as observed by Boyer et al. (1997). Deeper flow paths also contribute to increasing ion concentration patterns at low flow in Dry Creek Experimental Watershed near Boise, ID (McNamara et al., 2005). Our findings, along with these earlier works suggest a regional occurrence of limited near-surface storage capacity due to loamy soils and the importance of deeper groundwater for stream water.

In contrast to DIC, more biologically reactive solutes TN, DOC, and PO_4^{3-} either decreased or were relatively invariant with stream drying. Mean TN concentrations decreased throughout the growing season, and DOC showed relatively high variability and no clear longitudinal pattern or large

seasonal shift in either stream. With respect to TN, we expected temporal decreases because nitrogen is biologically reactive and often subject to uptake, especially during low flows and low concentrations (Moatar et al., 2017). On the other hand, many studies of stream carbon dynamics have found clear seasonal patterns in DOC (Hornberger et al., 1994; Boyer et al., 1997; Mulholland and Hill, 1997; McGuire et al., 2014; Hale and Godsey, 2019). However, we did not observe a strong DOC pattern with drying in either stream. Rather, we found strong temporal shifts in DIC, and DIC concentrations that were 2–3 fold higher than DOC in unburned JD and burned MC. Thus, DIC, rather than DOC, appeared to dominate stream carbon dynamics at our sites.

Non-linear spatial patterns in longitudinal stream chemistry spatial patterns have been observed in other studies that employed high spatial-scope measurements, but few have examined these longitudinal patterns in the context of stream drying or across such a broad suite of biologically reactive and conservative analytes. In one such example, within snowmelt-driven, deciduous forested streams at Hubbard Brook, NH, Likens and Buso (2006) sampled at 100 m intervals over a network and observed non-linear spatial patterns in a wide suite of constituents including DOC, DIC, K, Na, SO_4^{2-} , NO_3^- , Sc, dissolved Si, and Ca (also see McGuire et al., 2014; Zimmer and Lautz, 2014). Similarly, Dent and Grimm (1999) found that NO_3^- -N and soluble reactive phosphorus (SRP or PO_4^{3-}) varied longitudinally in the hot desert at Sycamore Creek, AZ. Moreover, based on 25 m intervals over 10 km of stream, they observed that NO_3^- -N decreased and SRP increased with succession following monsoonal flooding. These findings agree with our observations of decreasing TN concentrations and relatively invariant (unburned JD) or slightly decreasing PO_4^{3-} (burned MC) with drying.

Stream Differences: Lithology and Fire Impacts

We found differences between the two streams we studied in cation concentrations, the proportion of open canopy, and magnitude of seasonal decrease in TN, organic carbon and water sourcing. Lithological differences likely account for higher cation concentrations in volcanic burned MC compared to granitic unburned JD stream (Meybeck, 1987; Ibarra et al., 2016). It is also likely that fire impacts increased cation concentrations through ash and sediment inputs and/or mineralization of these inputs, especially during snow melt. At the watershed scale, sediment losses were high during winter and snowmelt in the year following fire, a low flow year, and sustained in 2017 (>450 compared to $20 \text{ g m}^{-2} \text{ yr}^{-1}$ mean sediment yield in Johnston). Sediment losses were low during the drying summer months ($<5\%$ of annual sediment yield) indicating low contributions of particulates during this study period (Glossner, 2019); mineralization of these ash-derived products warrants further study.

Elevated nitrogen levels in April and May in burned MC decreased by $>80\%$ in June. These TN results supported our hypothesis that nitrogen levels would be elevated immediately following wildfire but that decreases in TN would be observed as burned MC dried due to potentially increased nutrient uptake. Other studies such as Murphy et al. (2006) and Rau et al. (2007) have documented that there is often a lag in nitrate uptake following fire due to the time required to re-establish uptake by denitrifying bacteria populations and vegetation. In some cases, high nitrogen levels have been documented in streams for years following fire (Hauer and Spencer, 1998; Mast et al., 2016), potentially related to low DOC from terrestrial carbon losses (Rodríguez-Cardona et al., 2020). However, TN levels in burned MC decreased relatively quickly to concentrations similar to those of unburned JD by June. Finally, DOM sourcing followed different temporal tendencies in the two streams: burned MC shifted as expected from more allochthonous to autochthonous whereas unburned JD showed more mixed contributions. These

findings will be discussed further below in the “Statistical Challenges of Stream Drying Data”.

Fire can have a large effect on water budget and cause increases in stream flow as catchment ET diminishes following vegetation losses (Kinoshita and Hogue, 2015; Atchley et al., 2018), but we did not find strong evidence of this influencing stream chemistry drying patterns. This in part may be because burned MC experienced a relatively low water year in 2016 (Glossner, 2019; Vega et al., 2020) and rapid growth of grasses and herbaceous was observed. Rapid fire recovery (within a year) has been observed previously at RC CZO with regards to gross ecosystem production (Fellows et al., 2018) and ET (Flerchinger et al., 2016), with statistically marginal short-term increases to stream flow (Flerchinger et al., 2016). Moreover, ET rates have been shown to be relatively low in sagebrush in RC CZO (Sharma et al., 2020). Isotopic LMWL-intercepts indicated differing water sources between the streams and suggested unburned JD was more rainfed than burned MC, which was fed more by groundwater/snow. Given the vegetation differences, it is possible unburned JD had more shallow subsurface storage of rainwater than burned MC. However, as vegetation regrew in burned MC, there was no clear LMWL-intercept shift from April to June suggesting changes in soil water storage capacity. Alternatively, water source differences may have reflected pre-fire differences and further study would be required to parse this out. As mentioned, we observed stream chemistry responses to fire in some instances like elevated TN, but we did not detect increased stream flow. These observations are consistent with the weak evidence of monthly changes in evapoconcentration or dilution processes and burned MC's lack of shifts in water isotopic water sources or evaporative slopes. In addition to the similarity of stream patterns indicating the importance of deeper subsurface processes on stream chemistry, the similarity may indicate muted effects of fire on stream chemistry in the burned watershed.

Processes Explaining Spatiotemporal Patterns

Our findings highlight the utility of stream chemistry to elucidate both hydrological processes (evapoconcentration or dilution and water sourcing) and the in-stream biological processes (potential autochthonous carbon-sourcing) that contribute to spatiotemporal patterns as streams dry seasonally and following wildfire. A strength of our data was that we were able to evaluate surface-groundwater interactions, which has been identified as a key research priority to understanding intermittent stream controls (Costigan et al., 2016). Our inferences are only made possible by the high spatial scope sampling approach we employed.

Evaluating Hydrological Processes: Shift Toward Groundwater With Drying

By quantifying evaporation and groundwater influence, we could distinguish between chemistry drying patterns driven by surface processes and chemistry drying patterns driven by subsurface hydrologic processes in these streams. Isotopic slopes showed both streams were evaporative; however, we observed no temporal shifts toward increased in-stream evaporation with

drying. Contrary to our expectations and despite differences in canopy cover, both streams showed similar evaporative trends in water isotopes with no statistical difference in slope between the two streams in May or June. Furthermore, we found only weak evidence of in-stream evapoconcentration or dilution on the spatial stream chemistry patterns in either stream during any month. In contrast, Sr-DIC ratios provided strong evidence that stream surface water shifted toward a deeper groundwater signature with drying. These low-flow groundwater contributions differ from the lateral input dominance of snowmelt described by Boyer et al. (1997). The Sr-DIC relationships varied longitudinally and were better captured by precipitation and groundwater endmembers in burned MC than unburned JD. In April, the Sr-DIC signature of the uppermost portion of unburned JD led us to reason that either that these sites are spring-fed or that our sampling missed the shift from precipitation to groundwater sourcing. In general, increased contact time between water and soil/bedrock may explain elevated cation and DIC concentrations in groundwater (Fritz and Clark, 1997; Richter and Billings, 2015; Olshansky et al., 2019), each of which increased with drying in both streams. Although isotope intercepts did not show strong shifts in water sourcing between months as Sr-DIC ratios did, the isotopic signatures of snow and groundwater were indistinguishable. This suggests two potential scenarios. First, as snowmelt recharges deeper groundwater, snowmelt increases its solute concentrations. Alternatively, shallow subsurface evaporation could cause evapoconcentration (Radke et al., 2019). Quantifying subsurface processes such as those that would affect the residence times of water (Brooks et al., 2015), carbon in groundwater, or potential evapoconcentration in soils (Radke et al., 2019) may be important for understanding stream dynamics at RC CZO and warrants further regional study.

Evaluating Biological Processes: Carbon Sourcing Is Longitudinally Variable

Rather than exhibiting distinct seasonal shifts over time, biologically active solutes, such as organic carbon, showed a mixed signature of allochthonous/autochthonous sourcing and composition in both streams each month. We found DOC sourcing (FI) and aromatic properties a_{254} varied in both streams, but burned MC was less aromatic overall (lower a_{254}). Longitudinally, unburned JD had mixed sourcing (FI values) and a wide spanning aromatic signature (a_{254}) even in June, though there was a tendency toward being less aromatic. Although unburned JD FI observations were similar to summer 2017 samples from higher elevation streams at RC CZO (Radke et al., 2019), they were counter to our expectations. Canopy cover in the unburned JD was heterogeneous: leaf out may have decreased sunlight availability to the stream, limiting productivity and contributing more allochthonous material throughout the summer. Burned MC more closely matched our expectations of increased in-stream productivity with a more autochthonous FI in June and an overall less aromatic signature, but FI was nevertheless unexpectedly variable. More in-stream productivity may have been supported by high (~20-fold more) nitrogen availability in the early season and an

open canopy from April to June, combined with the generally reduced vegetation as allochthonous carbon sources post fire. While there were temporal differences between the streams, the high variability that characterized DOM sourcing and spectral characteristics reflected a mixed organic carbon signature in both streams. This may be typical of small-area catchments that are often hydraulically connected to surrounding terrestrial uplands (Creed et al., 2015). Nevertheless, burned MC showed clearer, and potentially less complex patterns of increasing groundwater influence and autochthonous DOC. Though it is unclear how characteristic these patterns may have been before the fire in burned MC, it is possible the fire's reduction of canopy cover and increase of TN may have contributed to more pronounced temporal shifts toward autochthonous DOC.

Study Approach: Pros and Cons Power of High Spatial-Scope Sampling and Integrated Methodology

Collectively, our findings have contributed to recognizing and understanding stream chemistry patterns that cannot be addressed with traditional sampling approaches that rely on a small number of spatially discrete locations, such as stream outlets (e.g., Fisher and Likens, 1973; Boyer et al., 1997; Mulholland and Hill, 1997; Hood et al., 2006; Jaffe et al., 2008 etc.). For instance, Jaffe et al. (2008) found FI to vary from 1.15 to 1.75 at the landscape to biome scale. We found comparable variation both longitudinally down the 2.5 km stream segments at a 50 m grain and temporally across months we studied. This finding suggests that traditional coarse-grain spatial sampling of DOM sources (1) may not capture heterogeneity of the streams above or below a sampling point and (2) may lead to inappropriate inferences about a given stream network. It is unclear if network convergence would occur downstream in DOM sourcing, as has been observed in DOC concentrations (Asano et al., 2009; Hale and Godsey, 2019), and if so, at what scales (Creed et al., 2015). Our findings provide evidence to support the Creed et al. (2015) hypothesis that DOM sourcing dynamics within a single catchment may be more complex than thought and may have consequences for the chemical composition and quality of organic matter being transported downstream.

Stream chemistry studies have traditionally either focused on hydrological processes (Brooks et al., 2015) or on in-stream biological processes (e.g., Minshall et al., 1989; Dent and Grimm, 1999), and study scope determines at what scale(s) patterns can be detected or processes can be assessed (Schneider, 2001; Fausch et al., 2002). In this observational study, our aim was to explore both hydrological and biological processes that may drive stream chemistry patterns using integrated methodology at a high spatial scope as streams dry. Taking this spatial approach required that we balance trade-offs between distance covered (extent) and sampling interval (spatial grain); we did not study a large stream network, but focused on the headwater stream segment, where we expected to see the most pronounced drying and biogeochemical patterns and variation. Although we also used repeated monthly sampling to investigate temporal patterns, we focused our efforts at the beginning of the drying season with moderate temporal

resolution (monthly) over a limited temporal extent (3 months at the beginning of one drying season).

Due to their small size, headwater streams are highly variable with regards to stream chemistry (Zimmer et al., 2013; Creed et al., 2015; Hunsaker and Johnson, 2017), and thus they are likely to be more susceptible to seasonal and spatial regime shifts such as increased stream intermittency and fire, both promoted by climate change. The particulars of these shifts in headwaters are important to understand in various biomes for study design as well as water quality monitoring (Abbott et al., 2018). Future work might include spatially distributed streamflow measurements to accompany stream chemistry observations because it is possible that the local water balance differs seasonally and longitudinally, affecting in-stream evapoconcentration/dilution dynamics (e.g., Glossner, 2019; Warix, 2020). Lastly, we balanced sampling intensity and number of streams studied. Although our inferences are limited by the fact that we studied only two streams, the similarities between the two suggest some of the patterns we observed may be expected throughout the RC CZO and region. Our analysis of a wide range of chemical constituents at 50 m reach intervals complimented the work of Hale and Godsey (2019) who measured DOC at 200 m intervals at the network scale in southeastern Idaho, USA, showing that additional analytes can improve process interpretation. We suggest that exploratory studies, such as this one, are important to tease apart which processes are effectively uniform throughout a stream reach, and which are likely to be spatially heterogeneous and seasonally dynamic.

Statistical Challenges of Stream Drying Data

Additional challenges existed applying typical statistical approaches used to describe perennial stream chemistry patterns to spatiotemporal drying patterns. It was particularly challenging to describe temporal patterns, compare the longitudinal spatial variation between months as streams dried, and to quantify the spatial patterns of a given month. Much of this difficulty stemmed from the loss of data points as streams dried, impacting the extent and resolution and effectively reducing the sample size each month and leaving few perfect statistical approaches. Some authors have reported maximums and minimums to capture seasonal differences within a network (Likens and Buso, 2006; Brooks and Lemon, 2007; Zimmer et al., 2013), but this limits comparisons between variables, particularly of different magnitudes or different units of measure. Coefficient of variation (CV; calculated by the standard deviation, SD, divided by the mean) is a unitless tool for quantifying variation and is therefore appealing to compare different variables. As streams dry, CV has been used to quantify temporal patterns (Likens and Buso, 2006), spatial patterns at varying scales (Dent et al., 2001), and spatiotemporal stream chemistry patterns (Hale and Godsey, 2019) to compare across varying monthly means and sample sizes. However, CV is sensitive to changes in the mean, as Dent and Grimm (1999) point out, as well as the sample size which makes CV potentially problematic for drying streams. For example, in our data if April and June solute concentrations

have the same SD (representing spatial variation) but June has a higher mean, June will have a lower CV. This could reflect the loss of sampling locations rather than changes in spatial variability of the sites that remained wet in June. Dent and Grimm (1999) acknowledged the limitation and emphasized CV only in specific instances.

We also faced difficulties in quantifying spatial patterns in our streams due to underlying, non-linear longitudinal patterns, which changed with stream drying. Brooks and Lemon (2007) addressed this issue by reporting means and SD for an upper, middle, and lower region of the San Pedro River, AZ. To control for differences in the mean, sample size and underlying longitudinal patterns, we calculated CV in instances for which underlying longitudinal patterns did not change with drying, which occurred only for FI. More sophisticated approaches have included constructing semivariograms to quantify spatial patterns (McGuire et al., 2014) and spatiotemporal patterns (Dent and Grimm, 1999). The data from our study required an in-depth approach to detecting and quantifying spatiotemporal patterns which was beyond the scope of this paper, and we more thoroughly addressed quantifying patterns in a separate, forthcoming publication (MacNeille, 2020). Continued development of robust statistical methods and conceptual models that incorporate drying (changes in sample size) is merited. This is a vital step to developing more appropriate approaches and tools to understand patterns and mechanisms of stream drying (Jensen et al., 2019).

CONCLUSIONS

Understanding intermittent stream ecosystem structure and processes requires a shift in stream conceptual models to include stream drying (Allen et al., 2020) and testing controls on drying like surface water-groundwater interactions (Costigan et al., 2016). Exploring patterns using a high-spatial scope approach, biogeochemical data enabled us to test several hypotheses about processes associated with stream drying and following fire. We found headwater streams had distinct chemical patterns that shifted longitudinally with seasonal drying and showed similar patterns despite fire history differences. In this study, an unburned and a burned stream exhibited similar temporal patterns with stream drying and similar shifts in processes. We argue that detecting shifts in the processes that influenced these patterns required measurements at a fine grain over an intermediate extent. DIC dominated carbon dynamics in both streams, with increasing concentrations with drying. We observed similar drying shifts in increasing Sr and Cl^- , relatively invariant DOC, decreasing TN, and relatively invariant PO_4^{3-} . Cations generally increased from April to June with more substantial losses in burned MC potentially due to volcanic bedrock. The most striking difference in temporal patterns between the unburned JD and burned MC was the dramatic loss of TN from April to June in the burned MC. As the streams dried, surface water chemistry patterns could be influenced by surface evaporative processes, dilution, and/or from subsurface

processes with shallow and deep groundwater influence. During stream drying, the snow-fed, semi-arid mountainous catchments we studied were both characterized as evaporative streams, but with weak evidence for in-stream evapoconcentration or dilution as either stream dried. There were not substantial in-stream evaporative differences despite canopy cover differences between unburned JD and burned MC. Instead, stronger evidence for deeper groundwater processes influencing stream chemistry patterns with longitudinal distinctions were observed in these streams, with more clear chemistry shifts with drying in burned MC. The similarities between streams may suggest that subsurface structure mutes the effects of fire and lithology differences at RC CZO. Organic carbon sourcing was mixed and highly longitudinally varied in both streams, but with an overall shift toward autochthonous in burned MC. These findings contribute to understanding streams in the sagebrush steppe by exploring spatiotemporal stream chemistry patterns and processes shifts as stream drying and following wildfire. Such empirical data are critical to the development of more sophisticated stream models, which are especially needed as climate changes across the western U.S.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found at: <http://dx.doi.org/10.18122/td/1332/boisestate/reynoldscreek/16>, Boise State ScholarWorks.

AUTHOR CONTRIBUTIONS

RM: contributed to project conception, experimental design, primary researcher conducting sampling, lab analysis, statistical analysis, manuscript writing and editing, and

major concepts/framing. KL: contributed to project conception and experimental design, primary supervisor of sampling, lab analysis, statistical analysis, manuscript editing, major concepts/framing, Reynolds Creek Critical Zone Observatory PI, and funding. SG: data and statistical analysis consultant, manuscript editing, and contributed to major concepts/framing. JP: supervisor for laboratory analysis of organic carbon and fluorescence data, data analysis consultant, manuscript editing, and major concepts/framing. CB: project conception, experimental design, data analysis consultant, manuscript editing, and major concepts/framing. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/frwa.2020.563841/full#supplementary-material>

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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