

Multi-scale temporal patterns in stream biogeochemistry indicate linked permafrost and ecological dynamics of boreal catchments

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Abstract: Temporal patterns in stream chemistry provide integrated signals describing the hydrological and ecological state of whole catchments. However, stream chemistry integrates multi-scale signals of processes occurring in both the catchment and stream. Deconvoluting these signals could identify mechanisms of solute transport and transformation and provide a basis for monitoring ecosystem change. We applied trend analysis, wavelet decomposition, multivariate autoregressive state-space modeling, and analysis of concentration-discharge relationships to assess temporal patterns in high-frequency (15 min) stream chemistry from permafrost-influenced boreal catchments in Interior Alaska at diel, storm, and seasonal time scales. We compared catchments that varied in spatial extent of permafrost to identify characteristic biogeochemical signals. Catchments with higher spatial extents of permafrost were characterized by increasing nitrate concentration through the thaw season, an abrupt increase in nitrate and fluorescent dissolved organic matter (fDOM) and declining conductivity in late summer, and flushing of nitrate and fDOM during summer rainstorms. In contrast, these patterns were absent, of lower magnitude, or reversed in catchments with lower permafrost extent. Solute dynamics revealed a positive influence of permafrost on fDOM export and the role of shallow, seasonally dynamic flowpaths in delivering solutes from high-permafrost catchments to streams. Lower spatial extent of permafrost resulted in static delivery of nitrate and limited transport of fDOM to streams. Shifts in concentration-discharge relationships and seasonal trends in stream chemistry toward less temporally dynamic patterns might therefore indicate reorganized catchment hydrology and biogeochemistry due to permafrost thaw.

Highlights:

- Storms flushed nitrate and dissolved organic matter from a high-permafrost catchment
- Storms diluted nitrate in a stream draining a low-permafrost catchment

- Diel oscillations were minor compared to seasonal trends and storm-scale variation

Introduction

Temporal patterns in stream chemistry provide integrated signals describing the hydrological and ecological state of whole catchments (Likens and others, 1970). Seasonal trends, storm responses, and diel fluctuations in stream chemistry can reveal the distribution of hydrologic flowpaths, dominant biogeochemical processes, and phenology of water and energy exchange within catchments, as well as biotic processes in receiving streams (Kirchner and others, 2004; Hall and others, 2016; Burns and others, 2019). Deviation or long-term change in these characteristic patterns might therefore indicate ecosystem responses to changing climate and disturbance regimes (e.g., Czikowsky and Fitzjarrald, 2004; Keller and others, 2010; Jones and others, 2017; Fork and others, 2020). However, gaining mechanistic insights from records of stream chemistry requires deconvoluting signals that arise from both terrestrial and aquatic processes and occur on multiple time scales (Bernhardt and others, 2005; Brookshire and others, 2009).

Stream chemistry in the boreal forest of Alaska reflects the influence of spatially discontinuous permafrost (ground that remains frozen for ≥ 2 years) on hydrological and ecological processes. Forests underlain by permafrost have cold, moist soils and low rates of organic matter decomposition and nutrient cycling (van Cleve and others, 1991). Organic matter accumulates on the forest floor where it insulates permafrost, fosters growth of spruce forests, and provides a source of dissolved organic matter (DOM) (Johnstone and others, 2010; Jorgenson and others, 2010; Ping and others, 2010). Ice-rich permafrost confines hydrologic flowpaths to shallow, seasonally thawed soils (i.e., the active layer) of relatively high organic

content (Haugen and others, 1982, Jorgenson and others, 2010), resulting in DOM export to streams (Petrone and others, 2006; Harms and others, 2016). In contrast, permafrost-free soils are warmer and drier and favor establishment of deciduous forests that store organic matter in aboveground biomass, rather than in the forest floor (Johnstone and others, 2010; Alexander and Mack, 2016). Here, deeper flowpaths through well-drained hillslopes interact with mineral soils and fractured bedrock aquifers (Yoshikawa and others, 2003), which support little nitrate (NO_3^-) uptake (Harms and Jones, 2012) and are therefore sources of NO_3^- (Verplanck and others, 2003).

Permafrost and resulting patterns in forest composition thus impart biogeochemical signals that are transported by hydrologic flowpaths to streams. Identifying the mechanisms that generate temporal patterns in stream chemistry of permafrost-influenced catchments might provide new approaches for tracking catchment-scale permafrost distribution and understanding its influence on ecosystems. Spatially distributed analyses have demonstrated that streams draining catchments of lower permafrost extent or deeper active layer depth have lower DOM and higher NO_3^- concentrations (Petrone and others, 2006; Harms and others, 2016). These patterns are hypothesized to result from deeper flowpaths intersecting mineral soils, compared to shallow organic soils. On seasonal time scales, this phenomenon could manifest as season-long directional trends in solute concentrations as groundwater flowpaths migrate from organic to mineral layers as the active layer thaws (Harms and Jones, 2012; Table 1: H1). During storms, the timing, magnitude, and duration of concentration-discharge relationships could indicate the influence of permafrost on the depth and length of catchment flowpaths as well as the distribution and size of solute pools within catchments (Williams, 1989; Petrone and others, 2007; Burns and others, 2019). For example, where permafrost is present, flowpaths are shallow (Koch and others, 2013) and expected to rapidly deliver pulses of DOM from organic-rich soils

to streams during storms (Table 1: H2 and H3). Where permafrost is absent, well-drained soils yield longer transit times (Yoshikawa and others, 2003) and flowpaths intersect mineral soils, which could produce delayed flushes of inorganic N into streams (Table 1: H2 and H3).

Biotic processes in both streams and catchments also generate temporal signals in stream chemistry. Vegetative uptake regulates NO_3^- export from streams draining nitrogen-limited temperate forests (Likens and others, 1970) and could contribute to declines in NO_3^- concentration in streams during periods of peak growth in boreal forests (Table 1: H4). Terrestrial productivity and evapotranspiration can also generate diel variation in stream discharge, flowpath depth, or soil redox conditions that cause corresponding patterns in stream chemistry (Czikowsky and Fitzjarrald, 2004; Nimick and others, 2011). For example, evapotranspiration during the day can lower water tables and promote nitrification by oxygenating soils (Duncan and others, 2015). At night, a higher water table might foster denitrification and decreased NO_3^- export (Flewelling and others, 2013) or elevate DOM export as shallower flowpaths intersect organic-rich soils. Such patterns caused by terrestrial productivity would result in synchronous diel changes in stream discharge and chemistry (Table 1: H5). Alternatively, diel patterns might arise from metabolism in streams, whereby primary producers draw down NO_3^- concentration and release DOM during daytime (Odum, 1956; Nimick and others, 2011; Hall and others, 2016). Biogeochemical signals imparted by in-stream biota would therefore occur in synchrony with aquatic photosynthesis and respiration (Table 1, H6). Such biotic-driven patterns could vary with permafrost and forest composition, or may need to be disentangled from stream chemistry records to identify other signals.

We examined the temporal patterns in catchment biogeochemistry emerging from catchments varying in spatial extent of permafrost. We collected high-frequency (15 min)

records of stream chemistry using sensors deployed in four headwater boreal catchments and used these data to test hypotheses regarding the processes generating temporal patterns and how they related to permafrost extent (Table 1). Specifically, we tested hypotheses distinguishing signals generated in catchments from those produced within streams at seasonal (H1), within-season (H2–H4), and diel time scales (H5–H6) using statistical approaches to deconvolute temporal patterns into characteristic frequencies and assess correlations of biogeochemical patterns with potential drivers.

Methods

Site description

We monitored four headwater streams in the Caribou-Poker Creeks Research Watershed (CPCRW), which is part of the Bonanza Creek Long-Term Ecological Research Program in the boreal biome of Interior Alaska. Mean annual air temperature is -3.6°C , with the lowest mean monthly temperature typically in January (-23.2°C) and highest in July (15.1°C). During the study period (May–Sept 2017), mean monthly temperatures were similar to long-term averages, with the exception of mean July temperatures approximately 2°C warmer than average. The CPCRW receives 351 mm precipitation annually on average, with 294 mm falling as rain. Mean monthly precipitation during the study period was higher in May (39.4 compared with 24.5 mm), lower in June (34.8 compared with 57.9 mm), and similar in July (104 compared with 96.9 mm) and August (83.1 compared with 82.0 mm) compared to long-term monthly averages.

In the study area thin (approximately 1 m) soils are underlain by a fractured bedrock aquifer of weathered schist, dispersed carbonate-bearing lithologies, bands of gravel and loess, and spatially discontinuous permafrost (Haugen and others, 1982; Yoshikawa and others, 2003).

Permafrost underlies valley bottoms and north-facing slopes. North-facing slopes are dominated by black spruce (*Picea mariana*) with mosses in the understory (e.g., *Sphagnum* spp., *Hylocomium* spp.). South-facing slopes and ridges are dominated by mixed hardwoods (*Betula neoalaskana* and *Populus tremuloides*) and develop little understory groundcover.

The monitored streams drain catchments of varying aspect and therefore spatial extent of permafrost (Fig. 1). Though aspect is not replicated, the study design captured a gradient in permafrost extent from minimal to nearly spatially continuous permafrost extent within the drainage network of each catchment. A south-facing catchment (C2, hereafter “low permafrost”) is underlain by permafrost in approximately 4.5% of its total area and 10% of its drainage network area, defined as the stream network and likely areas of zero-order channels and shallow subsurface flow (see Appendix 1: *Geospatial analysis*). A southeast-facing catchment (C4; hereafter “low-mid permafrost”) is underlain by permafrost in 25% of its total area and 30% of its drainage network area. In contrast, a northwest-facing catchment (P6; hereafter “mid permafrost”) is underlain by permafrost in 38% of its total area and 66% of its drainage network area, and a north-facing catchment (C3; hereafter “high permafrost”) is underlain by permafrost in 57% of its total area and 97% of its drainage network area. The catchments also vary in fire history. An experimental fire burned 28% of the C4 catchment in 1999 (Hinzman and others, 2003) and a wildfire burned 65% of the P6 catchment in 2004 (Betts and Jones, 2009). Streams draining catchments are narrow (<1 m width) and have closed canopy or are well shaded (Betts and Jones, 2009; Mutschlecner and others, 2018)

Data collection

We installed *in situ* sensors adjacent to an existing flume for gauging discharge in each catchment and monitored stream chemistry at 15 min frequency from May 23–Aug 31, 2017 (Fig. 1). We instrumented the high- and low-permafrost catchments with a Submersible Ultra-Violet Nitrate Analyzer (SUNA, Satlantic LP, Seabird Scientific, Halifax, NS, Canada) to measure NO_3^- concentration (LOQ = 0.014 mg N L⁻¹, short-term precision = 0.004 mg N L⁻¹). Nitrate and organic N comprise the majority of dissolved N in the study streams and contribute nearly equally to total N loads, on average (Petrone and others, 2006). We instrumented all four streams with photosynthetically active radiation (PAR) loggers (Odyssey, Christchurch, New Zealand) at water level and EXO2 multi-parameter water quality sondes (YSI, Yellow Springs, OH, USA) equipped to measure fluorescent dissolved organic matter (fDOM), turbidity, specific conductivity (SPC), and temperature. The EXO sondes installed in the high- and low-permafrost catchments were additionally equipped with optical dissolved oxygen (DO) sensors. Before deployment, we verified that SUNAs met the manufacturer specification of ± 0.028 mg N L⁻¹ in nanopore water and we calibrated SPC, fDOM, and turbidity sensors to externally-verified standards. Following manufacturer's specifications, we calibrated DO sensors in moist air. Once deployed, we recalibrated all EXO sensors monthly, checked EXO and SUNA sensors against nanopure water for drift and biofouling weekly, and cleaned optical read windows weekly. Sensors collected a burst of 9–15 measurements within 30 s at each 15-min interval, and the mean and variance of these measurements were used to detect faulty measurements and estimate uncertainty in subsequent analyses.

Water samples were collected weekly for validation of sensor data. Water samples were filtered in the field (0.7 μm glass fiber filters), transported on ice to the laboratory, and frozen

until analysis. Nitrate was analyzed by ion chromatography on a Thermo ICS-2100 with AS-18 columns (LOQ = 0.0017 mg N L⁻¹). Dissolved organic carbon (DOC) was analyzed on a Shimadzu Total Carbon analyzer by combustion followed by non-dispersive infrared gas analysis (LOQ = 8 µM). NO₃⁻ concentration measured by each SUNA was corrected to laboratory-measured values with site-specific calibration curves (low-permafrost n = 11, slope = 0.94 [95% CI = 0.56, 1.32], R² = 0.75; high-permafrost n = 12, slope = 0.91 [95% CI = 0.82, 1.01], R² = 0.98). EXO-measured fDOM was corrected for temperature and turbidity as described in Watras and others (2011) and Downing and others (2012). Corrected fDOM concentration was correlated with lab-determined DOC concentration across all sites (R² = 0.67, n = 30). However, the strength of this relationship varied among sites (site-specific R² range = 0.029–0.95) and we report fDOM in quinine sulfate units (QSU).

We gathered hydrological and meteorological attributes (lter.uaf.edu/data) to evaluate the influence of storms (Table 1: H2–H3) and terrestrial primary productivity (Table 1: H4) on stream chemistry. First, discharge was monitored using pressure transducers installed at flumes adjacent to sensor installations in each catchment. In-stream pressure was corrected for barometric pressure and converted to discharge with rating curves calculated from weekly measurements of discharge by salt dilution. Second, we obtained hourly precipitation and air temperature data from high- and low-elevation meteorological stations in the CPCRW (Fig. 1) and used mean values of the two stations. We calculated daily growing degree units (GDU), which quantify heat accumulation above a threshold temperature (5°C for boreal regions following Ruostenoja and others (2016) and references therein). GDU (estimated as $[(\text{Temp}_{\text{MAX}} + \text{Temp}_{\text{MIN}}) * 2^{-1}] - 5^{\circ}\text{C}$) were used to approximate growing conditions for

terrestrial vegetation and provided a proxy for terrestrial productivity (Bugmann and others, 2001; Ueyama and others, 2013; Rollinson and others, 2017).

To evaluate potential changes in the sources of water and NO_3^- coincident with seasonal migration of flowpaths (Table 1: H1), we compared the isotopic composition of NO_3^- ($\delta^{15}\text{N}$ and $\delta^{18}\text{O}$) and water ($\delta^{18}\text{O}$) in streams to previously observed values in groundwater and precipitation (lter.uaf.edu/data; T. Douglas, unpublished data). Stable isotopes of NO_3^- were measured using the denitrifier method (Casciotti and others, 2002) at the UC Davis Stable Isotope Facility. Ratios of $^{15}\text{N}:^{14}\text{N}$ and $^{18}\text{O}:^{16}\text{O}$ are reported relative to N_2 in air for $\delta^{15}\text{N}_{\text{NO}_3}$ and to Vienna Standard Mean Ocean Water for $\delta^{18}\text{O}_{\text{NO}_3}$ with analytical precision of $\pm 0.4\text{‰}$ for $\delta^{15}\text{N}$ and $\pm 0.5\text{‰}$ for $\delta^{18}\text{O}$. Samples with NO_3^- concentration $\ll 0.02 \text{ mg N L}^{-1}$ were below the LOQ and could not be analyzed. Isotopic composition of H_2O was measured by Cavity Ringdown Spectroscopy on a Picarro L2120i and is reported relative to Vienna Standard Mean Ocean Water with an analytical precision of $\pm 0.2\text{‰}$ for ^{18}O .

Statistical analysis

To test hypotheses linking stream chemistry to ecological and hydrologic processes influenced by permafrost, we examined temporal patterns at diel to seasonal time scales. We assessed seasonal trends (Table 1: H1) and correlates of daily variation in solute chemistry (Table 1: H2–H4) using autoregressive state-space models. Storm-scale analysis of concentration-discharge relationships tested hypotheses regarding the influence of permafrost on solute supplies (Table 1: H2) and depth of hydrologic flowpaths (Table 1: H3). Finally, wavelet decomposition was applied to evaluate the relative magnitude of temporal variation across scales and coherence of solute chemistry with processes occurring within the catchment (Table 1: H5)

or stream (Table 1: H6). To facilitate analyses of time series, gaps of <24 h were filled by applying a Kalman filter to forecasts from an ARIMA model. All data analyses were performed in R version 4.0.2 (R Core Team 2020).

Stream metabolism— To test possible effects of in-stream biotic processes on stream chemistry (Table 1: H6), we modeled rates of gross primary productivity (GPP) and ecosystem respiration (ER). We used the BAYesian Single-station Estimation of Metabolism (BASEmetab) package (Grace and others, 2015; Giling and Bond, 2020) because it provides both daily and instantaneous GPP and ER rates and we required instantaneous rates for testing hypotheses about diel variation in stream chemistry. Details related to metabolism model fitting and validation are provided in Appendix 2: *Stream metabolism*.

Seasonal and within-season dynamics— We tested hypotheses explaining variation in stream chemistry at seasonal (Table 1: H1) and within-season (Table 1: H2–H4) time scales using univariate and multivariate autoregressive state-space (MARSS) models, respectively (Holmes and others, 2018). The MARSS framework partitions variation among observation error, autoregressive processes, and covariate effects when included. First, to test the hypothesis that seasonal deepening of flowpaths influences stream chemistry where permafrost was present (Table 1: H1), we evaluated season-long trends in stream solutes by fitting univariate autoregressive state-space models to solute concentrations summarized by weekly means. Weekly means reduced the effect of autocorrelation but still captured season-long trends. We fit models to datasets reflecting three different time periods: a) the longest available time series for each variable in each catchment, b) excluding the influence of snowmelt (dates <June 15), and c) excluding a sudden change in discharge and chemistry that occurred in some catchments in late summer (dates >Aug. 14). See Appendix 3: *MARSS models* for details on model selection and fit.

To test hypotheses linking stream chemistry to storm-driven, terrestrial, and in-stream processes (Table 1: H2–H6), we used multivariate models to quantify correlations between NO_3^- or fDOM concentration and covariates reflecting these hypothesized processes. Daily mean values were $\log_{10}$ -transformed, centered, and standardized (mean = 0, standard deviation = 1) to meet model assumptions and facilitate comparison of covariate effects. Precipitation was evaluated as a covariate capturing storm effects (Table 1: H2–H3) and was included as total precipitation on the same day as measured stream chemistry as well as lagged by one day because discharge can lag precipitation by 4–48 h (McGuire and McDonnell, 2006; Koch and others, 2013). The potential effect of terrestrial primary production (Table 1: H4) was evaluated by inclusion of GDU lagged by one day as a covariate. Potential influence of in-stream processes on solute concentrations (Table 1: H6) was quantified by estimating effect sizes associated with daily max PAR and mean GPP and ER. Models also accounted for a sudden change in discharge and chemistry that occurred in some catchments by including a dummy variable corresponding to Aug 14–15. Covariates were uncorrelated with one another (Pearson's $r < |0.4|$). Additionally, GDU, GPP, ER, and precipitation were uncorrelated with day of year (Pearson's $r < |0.4|$), whereas PAR was negatively correlated. We fit a separate multi-state model for each response variable (NO_3^- or fDOM concentration) and used the longest available time series for each combination of response and covariate data. In-stream GPP and ER were available only for the high- and low-permafrost catchments in June and July. Thus, we explored these potential in-stream effects by fitting a second set of models to the shorter time series. For all models, we tested the hypothesis that each catchment represented a unique ecosystem state by comparing AIC scores of multi-state models to a model representing all catchments as a single state. Further details on fitting multivariate MARSS models are provided in in Appendix 3: *MARSS models*.

Storm-scale dynamics— To test hypotheses related to the pool sizes of solutes available for transport to streams (Table 1: H2) and the activation of flowpaths (Table 1: H3) during storms, we analyzed storm-scale concentration-discharge relationships. This analysis entailed delineating individual storms and identifying the rising and falling limbs of discharge responses to precipitation. Discharge in streams draining the high- and mid-permafrost catchments (Fig. 1) responded strongly to precipitation whereas responses in the low- and low-mid-permafrost catchments were subtle. However, solute concentrations responded to precipitation in all catchments. We identified storms as events when discharge exceeded 41.9 L s^{-1} and 64.3 L s^{-1} in the high- and mid-permafrost catchments, respectively. These thresholds correspond to twice the mean base flow for each catchment, which we calculated using a recursive digital filter approach from the EcoHydRology package (Fuka and others, 2018). In catchments with weak discharge responses, we identified storms within the time windows of events identified in the nearest “responsive” catchment, rather than using a discharge threshold. In all catchments, we delineated the beginning of each storm as the inflection point when discharge began to rise and the end point when discharge returned to the pre-event level or when another storm began.

To compare storm dynamics across catchments and events, we calculated a hysteresis index (HI) and flushing index (FI) for NO_3^- , fDOM, turbidity, and SPC during each storm (Lloyd and others, 2016; Vaughan and others, 2017). The HI indicates the magnitude and direction of hysteretic responses of solute concentration to rising and falling discharge and was calculated at every 2% interval of normalized discharge of each event. The HI at each Q interval i was calculated as $HI_i = C_{RL} - C_{FL}$ where C_{RL} and C_{FL} are normalized concentrations on the rising and falling limb, respectively. The median of all HI_i summarizes the central tendency for each event and bootstrapped 95% CIs around the median HI describe uncertainty. Median HI values range

between -1 and 1 , with negative and positive values indicating anti-clockwise and clockwise hysteresis, respectively. The FI quantifies solute dilution or flushing during each storm as the difference between mean solute concentration at the onset of an event compared to peak discharge. Mean solute concentrations at storm onset and at peak discharge were calculated as the mean of the burst of measurements ($n=9-15$ within 30 s) collected by the sensor at each time point. Bootstrapped 95% CI of the difference between these means provided an estimate of uncertainty for each FI.

Multi-scale temporal variation— We conducted complementary analyses in the frequency domain to characterize the relative variation in NO_3^- and fDOM concentrations across time scales and quantify correlations with indicators of terrestrial and in-stream processes (Table 1: H2–H6). Using wavelet analysis, we decomposed 15-min resolution time series into power spectra across a range of frequencies ($0.25-32$ d), quantifying power as the squared amplitude of oscillations localized in time. We used a ‘morlet’ mother wavelet, executed with the WaveletComp package (Roesch and Schmidbauer, 2018) and determined the significance of wavelet power using an autoregressive (“red”) background noise spectrum and 100 simulations. Wavelet analysis requires continuous time series and datasets with >12 h gaps were therefore analyzed separately. We used wavelet coherence to quantify correlations between NO_3^- or fDOM concentrations and indicators of storms (Table 1: H2), terrestrial processes (Table 1: H5), or in-stream processing (Table 1: H6). Wavelet coherence is a measure of localized correlation and phase angles between time series across time-frequency space. Phase angles are in the interval $[-\pi, \pi]$ and describe whether two time series at a given frequency oscillate in synchrony (in-phase = absolute values less than $\pi/2$, e.g., NO_3^- is high when discharge is high), anti-synchronously (out-of-phase = absolute values greater than $\pi/2$, e.g., NO_3^- is high when

discharge is low), while the sign of the phase angle describes which time series is leading in the relationship (Roesch and Schmidbauer, 2018).

Results

Seasonal dynamics

Yields of NO_3^- and fDOM contrasted among catchments at a seasonal timescale (Fig. 2). The high-permafrost catchment (C3) exported 32% less NO_3^- and 75% more fDOM, on average, than the low-permafrost catchment (C2; Fig. 2). The low-mid-permafrost catchment (C4) exported the least fDOM of all catchments studied and the mid-permafrost catchment (P6) exported the most (Fig. 2).

A season-long, positive trend in NO_3^- concentration occurred in the high-permafrost catchment, consistent with the prediction that a seasonal increase in NO_3^- concentration would occur where permafrost is present (Table 1: H1-P1a; Fig. 3). The positive trend was significant both before ($0.007 \text{ mg N L}^{-1} \text{ wk}^{-1}$, 95% CI: 0.001–0.01) and including a rapid increase in NO_3^- concentration that occurred late in the monitoring period ($0.02 \text{ mg N L}^{-1} \text{ wk}^{-1}$, 95% CI: 0.002–0.03), and also when the snowmelt period prior to June 15 was excluded ($0.02 \text{ mg N L}^{-1} \text{ wk}^{-1}$, 95% CI: 0.0002–0.04). No other measured solutes exhibited significant trends in any catchment (Table S1).

A rapid, sustained increase in discharge and solute concentrations punctuated the seasonal records of the high- and mid-permafrost catchment. In the high-permafrost catchment, this “step-change” entailed an increase of $\sim 0.2 \text{ mg NO}_3^- \text{-N L}^{-1}$ starting on Aug 14 that was sustained through the remaining period of monitoring (Fig. 3). The increase in NO_3^- concentration was accompanied by increased discharge and a decline in SPC, though the increase

in NO_3^- concentration was disproportionately large (fDOM data were unavailable in this catchment during this period; Fig. 3). A similar increase in NO_3^- concentration was observed in weekly samples in the mid-permafrost catchment and was accompanied by a sustained increase in fDOM of ~ 6 QSU (Fig. 3). Analogous patterns were not observed in the low- or mid-low-permafrost catchment.

The step-change in solute concentrations was accompanied by a change in isotopic composition of NO_3^- (Fig. 4). In general, $\delta^{18}\text{O}_{\text{NO}_3}$, $\delta^{15}\text{N}_{\text{NO}_3}$, and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ of streams were bounded by values observed in groundwater from springs and wells, and the range of $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ values we observed were similar to those observed in the nearby Nome Creek catchment (Koch and others, 2014). Stream $\delta^{15}\text{N}_{\text{NO}_3}$ significantly increased after the step-change by 0.71‰ and 1.35‰ in the high- and mid-permafrost catchments, respectively (Fig. 4), as indicated by non-overlapping bootstrapped 95% CIs of mean values before and after the step-change. Similarly, mean stream $\delta^{18}\text{O}_{\text{NO}_3}$ significantly increased across the same catchments and dates by 0.83‰ and 0.79‰ in the high- and mid-permafrost catchments, respectively. Stream $\delta^{18}\text{O}_{\text{NO}_3}$ and $\delta^{15}\text{N}_{\text{NO}_3}$ in catchments where no step-change occurred and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ in all catchments showed little seasonal variation (Fig. 4).

Within-season dynamics

Daily variation in stream chemistry within the period of measurement was strongly correlated with precipitation, as indicated by significant covariate effects in MARSS models. NO_3^- and fDOM concentrations were significantly correlated with same-day and/or the previous day's precipitation in all catchments (Fig. 5a–f). Previous-day precipitation had a strong positive effect on NO_3^- concentration in the high-permafrost catchment (Fig. 5a). However, both same-

and previous-day precipitation had strong negative effects on NO_3^- concentration in the low-permafrost catchment (Fig. 5b), a pattern counter to that predicted to result from larger stores of mobilizable NO_3^- where permafrost is absent (Table 1: H2-P2c). Finally, same- and previous-day's precipitation were positively correlated with fDOM in all catchments (Fig. 5c–f), which is also inconsistent with the hypothesis that differential stores of soil OM would result in contrasting temporal dynamics among catchments (Table 1: H2-P2b).

Solute concentrations were also correlated with indicators of in-stream and catchment biological activity on a daily timescale. NO_3^- concentration was negatively correlated with previous day's growing degree units (GDU), an indicator of terrestrial productivity, in both the high- and low-permafrost catchments; this was a predicted result of the hypothesis that uptake of N by terrestrial vegetation influences temporal patterns of NO_3^- concentration in streams (Table 1: H4; Fig. 5a–b). GDU was not correlated with fDOM concentration (Fig. 5c–f). NO_3^- concentration was also negatively correlated with in-stream GPP in the high-permafrost catchment (Fig. S2a), consistent with the predicted result of biotic retention of NO_3^- within streams (Table 1: H6; Fig. 5a–b). fDOM concentration was positively correlated with PAR in the low-permafrost catchment and low-mid-permafrost catchments (Fig. 5d–f), a pattern suggestive of the role of in-stream production in generating DOM (Table 1: H6). In the high-permafrost catchment, fDOM was negatively correlated with in-stream ER (Fig. S2c), providing mixed support for the role of in-stream biota in generating temporal patterns in DOM (Table 1: H6). In all MARSS models, estimating a unique state process for each catchment improved model fit ($\text{AIC}_{\text{single-state}} \gg \text{AIC}_{\text{multi-state}}$), indicating statistically distinguishable temporal patterns among the headwater catchments.

Storm-scale dynamics

Storm dynamics indicated differential patterns of solute sources and transport across catchments varying in spatial extent of permafrost (Fig 6). In the high-permafrost catchment, four of five storms resulted in increased NO_3^- concentration (positive FI) and counterclockwise hysteresis (negative HI; Fig. 6b). The smallest increase occurred during the first storm observed (July 2), which was followed by dilution of NO_3^- (negative FI) during the next storm (July 7) and flushing of NO_3^- during all storms thereafter (Fig. 6b). In the low-permafrost catchment, all storms resulted in dilution of NO_3^- , and all but one storm produced clockwise hysteresis (positive HI; Fig. 6b). These patterns are opposite to those predicted to result from larger stores of NO_3^- in deep, permafrost-free soils (Table 1: H2). Though all catchments and storms resulted in flushing of fDOM, only the low- and low-mid-permafrost catchment produced clockwise hysteresis for fDOM. This pattern is opposite of that expected if shallow flowpaths dominate during storms of high, but not low spatial extent of permafrost (Table 1: H3). Temporal dynamics of SPC and turbidity were similar across all catchments, with dilution and clockwise hysteresis of SPC in nearly all storms (Fig. 6d) and heterogeneous patterns in turbidity across storms (Fig. 6e).

Multi-scale temporal variation

Wavelet analysis corroborated the significant influence of storms and discharge variation on NO_3^- and fDOM concentrations previously described in the time domain. At frequencies of 2–8 d and times corresponding to storms, oscillations of NO_3^- and fDOM were significantly greater than expected of an autoregressive (red noise) spectrum in all catchments (Fig. S3, S4). In the high-permafrost catchment, NO_3^- concentration was coherent with discharge at temporal

frequencies and times associated with storms, typically varying in-phase with NO_3^- following discharge (Fig. 7a–b), a pattern indicating flushing of NO_3^- during storms and which contradicts the prediction that NO_3^- flushing would only occur where there were larger stores of NO_3^- in deep, permafrost-free soils (Table 1: H2). NO_3^- concentration was also coherent and in-phase with discharge at 4–16 d frequencies during the time period of the step-change in the high-permafrost catchment. At a diel frequency, NO_3^- concentration was coherent with discharge in both the high- and low-permafrost catchments, as predicted to result from the influence of terrestrial transpiration (Table 1: H5). Out-of-phase coherence with NO_3^- leading discharge tended to occur at the diel scale in the high-permafrost catchment during the early thaw season, whereas NO_3^- varied in-phase, following discharge during the later season in the high-permafrost catchment and throughout the record in the low-permafrost catchment (Fig. 7a–b). This corresponded to discharge peaks ~03:00 and NO_3^- peaks ~04:00 throughout the records of both catchments, with the exception of NO_3^- peaks ~07:00 in the early thaw season of the high-permafrost catchment. During inter-storm periods, NO_3^- was coherent and consistently out-of-phase with GPP, with NO_3^- following GPP in both the high- and low-permafrost catchments (Fig. 7c–d). During these periods, GPP peaked ~14:00, a few hours prior to NO_3^- minima, consistent with predicted effects of in-stream N uptake (Table 1: H6). Wavelet power spectra indicated no diel pattern of variation in fDOM concentrations (Fig. S4).

Discussion

Rapid climate warming at high latitudes is thawing permafrost, which in turn is altering cycling of carbon and nitrogen, elements that directly and indirectly participate in feedbacks to climate warming (Wolken and others, 2011; Schuur and others, 2015; Salmon and others, 2016).

We identified indicators of linked permafrost and ecological dynamics using multi-scale statistical approaches to compare daily to seasonal temporal patterns in stream chemistry of boreal catchments that vary in spatial extent of permafrost. We observed patterns consistent with shallow, seasonally dynamic flowpaths in a catchment with high permafrost extent, including a seasonal increase in NO_3^- concentration and flushing of NO_3^- during storms. In contrast, dilution of NO_3^- during storms and lack of seasonal trends in a low-permafrost catchment was indicative of drainage by deep flowpaths. Overall, seasonally dynamic flowpaths and contrasting solute storage produced distinct temporal patterns in stream chemistry among catchments varying in spatial extent of permafrost that can be used to monitor ecosystem change.

Solute export reflects catchment state

Contrasts in the specific yields of NO_3^- and fDOM from catchments varying in permafrost extent (Fig. 2) reflected different sizes of solute pools accessible to hydrologic flowpaths. Specifically, greater yield of NO_3^- and smaller yields of fDOM from the low- and low-mid-permafrost catchments compared to high- and mid-permafrost catchments indicated flowpaths that traversed deeper soils of greater inorganic N and lower OM content (Harden and others, 2012; Melvin and others, 2015). Previous studies documented similar patterns in the same study catchments (Petrone and others, 2006; Balcarczyk and others, 2009) and high-frequency data further revealed variance in these general patterns due to storms (Fig. 2). In addition, the mid-permafrost catchment that burned in a wildfire 13 years prior to this study exported more fDOM than all other catchments, on average (Fig. 2). This contrasts with a decline in DOM export observed following the fire (Betts and Jones 2009) and indicates recovery of OM pools connected to the stream in the decade following fire.

Interactions of hydrologic flowpaths with solute sources

Seasonal-scale patterns in solute chemistry supported the hypothesis that flowpaths in permafrost-influenced catchments intersect deeper soils as the thaw season progresses (H1). Specifically, a significant increase in NO_3^- concentration in the high-permafrost catchment suggested seasonal changes in flowpath depth, whereas no trend in a low-permafrost catchment reflected the dominance of deep groundwater supporting streamflow there (Bolton and others, 2004). Importantly, the trends reported here are robust to type I errors caused by features of autoregressive data (Kirchner and Neal, 2013) because stream chemistry was modeled as an autoregressive process in the models we applied to estimate trends.

Several mechanisms might have contributed to the observed increase in NO_3^- concentration at a seasonal timescale, including lower biological N demand or greater rate of nitrification at depth (Harms and Jones, 2012; Melvin and others, 2015), increased rates of N mineralization and nitrification as soils warm (Klingensmith and Van Cleve, 1993), or increasing contribution of NO_3^- -rich groundwater that occurs in the region (Verplanck and others, 2003). High biotic demand for N likely limits hydrologic export of N mineralized under seasonal warming (McFarland and others, 2002; Lavoie and others, 2011) and sustained export of fDOM from the high-permafrost catchment throughout the monitoring period (Fig. 3) suggests that major flowpath contributions to streamflow likely remained within a relatively shallow active layer, rather than incurring substantial input from deep groundwater. Thus, we conclude that downward migration of shallow flowpaths is responsible for the observed season-length trend in NO_3^- concentration. This temporal trend is complementary to a positive correlation of active layer depth with NO_3^- concentration previously documented in streams across Interior Alaska

(Harms and others, 2016). The seasonal rate of NO_3^- increase estimated here provides a baseline against which to assess effects of continued warming, fire, or changes in precipitation regime on deepening of the active layer.

A large, sustained increase in NO_3^- , fDOM, and discharge that was accompanied by a decline in SPC in the high- and mid-permafrost catchments indicated rapid activation of hydrologic flowpaths in August (Fig. 2). This unexpected step-change was associated with prolonged precipitation in late summer, which can cause rapid transport of water and solutes along the permafrost boundary as well as rapid thaw of frozen soils (Koch and others, 2013; James and others, 2019; Douglas and others, 2020) and therefore could reflect mobilization of NO_3^- from mineral or previously frozen soils. However, the concurrent flushing of fDOM and decline in SPC do not support deepening of flowpaths as the origin of the step-change because OM content declines and ions contributing to SPC increase with soil depth (Kokelj and Burn, 2005; Keller and others, 2007; Gough and others, 2008; Ping and others, 2010). Neither was precipitation the likely source of increased NO_3^- observed in the stream because $\delta^{18}\text{O}_{\text{NO}_3}$ did not reflect the enriched values characteristic of atmospheric deposition (Kendall and others, 2007) and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ also did not shift toward heavier values more similar to rain following the step-change (Fig. 4).

We hypothesize that sustained precipitation enhanced lateral hydrologic connectivity and that solutes mobilized from within the active layer caused the observed step-change in solute concentrations and discharge. The isotopic composition of NO_3^- became more enriched in the catchments where the step-change occurred, suggesting that it was derived from a different source, or that precipitation and/or transport stimulated N processing. Isotopic enrichment of NO_3^- occurs during denitrification, though increased denitrification is inconsistent with the

observed increase in NO_3^- concentration. Instead, an increase in $\delta^{18}\text{O}_{\text{NO}_3}$ that remained greater than $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ of water sources within the catchment suggested export of recently nitrified NO_3^- that had not yet undergone complete exchange of $\delta^{18}\text{O}$ with local water (Boshers and others, 2019). Seasonal decline in terrestrial evapotranspiration might have additionally contributed to enhanced lateral hydrologic connectivity and solute mobilization during this time by raising the water table and decreasing plant N uptake (Yi and others, 2009; Iwata and others, 2012). Hillslope tracer studies in the nearby Nome Creek headwater catchment similarly found that shallow subsurface storage zones are rapidly flushed during large rain events (Koch and others, 2017), and smaller step changes in NO_3^- concentration followed late summer and early autumn rain events in an adjacent, larger river (Douglas and others, 2013), suggesting that solute dynamics of headwater streams might propagate to larger rivers. Absence of a step-change in the low- and low-mid-permafrost catchments underscored the role of infiltration and deep drainage in structuring hydrology and biogeochemistry in boreal catchments where permafrost is absent.

Storms mobilized solutes and activated flowpaths, revealing contrasts in solute stores (Table 1: H2) and depth of dominant flowpaths (Table 1: H3) among catchments varying in spatial extent of permafrost. Specifically, storms caused flushing of NO_3^- and fDOM from the high-permafrost catchment, but dilution of NO_3^- and smaller flushes of fDOM in the low-permafrost catchment (Fig. 6). Flushing of NO_3^- on the falling limb of storms in the high-permafrost catchment suggests that NO_3^- was delivered to the stream from distal locations in the catchment. A positive correlation of daily mean NO_3^- concentration with the previous-day's precipitation in the high-permafrost catchment (Fig. 5) corroborates the notion of longer-range transport of NO_3^- delivered to the stream on the falling limbs of storms. Together, these patterns underscore the role of permafrost in maintaining shallow flowpaths that connect soils to streams.

In contrast, dilution and clockwise hysteresis of NO_3^- in the low-permafrost catchment during most storms was counter to our prediction that larger stores of NO_3^- there would generate flushing during storms (Table 1: H2-P2c). Dilution of the NO_3^- -rich groundwater that supports baseflow likely occurred when storms increased flow through shallow flowpaths, such as the NO_3^- -depleted soil water present in the riparian zone (Rinehart and others, 2015). This pattern resembles storm responses observed in agricultural catchments encompassing NO_3^- -enriched groundwater (Dupas and others, 2016; Lloyd and others, 2016). Simultaneous flushing of fDOM and turbidity in the low-permafrost catchment is consistent with activation of shallow flowpaths during storms. Smaller flushes of fDOM in the low- and mid-low-permafrost catchments (Fig. 3) supported our expectation of smaller DOM stores there (Table 1: H2-P2b), and instances of clockwise hysteresis of fDOM in these catchments (Fig. 6c) indicated that DOM supplies were sometimes exhausted during storms (Williams, 1989). Overall, however, patterns on the timescale of storms emphasized that the distribution of solute supplies relative to locations of dominant flowpaths, rather than differences in total C and N stocks in soils, influences the biogeochemical signals of permafrost observable in streams.

Biotic influence on temporal variation in stream chemistry

Both potential growth of terrestrial vegetation (Table 1: H4–H5) and in-stream metabolism (Table 1: H6) covaried with solute concentrations in streams, though these effects were secondary to the influence of precipitation. The significant negative correlation of NO_3^- concentration with conditions favorable to terrestrial vegetation productivity (GDU) at a daily resolution (Fig. 5) suggests a potential effect of biota on N export within both the high- and low-permafrost catchments, a pattern also observed in other forested catchments (Pardo and others,

1995; Lucas and others, 2016). GDU peaked in early July and declined by late summer, making it unlikely that this effect was related to the season-long increase in NO_3^- observed in the high-permafrost catchment. Wavelet transforms also revealed coherence in oscillations of NO_3^- concentration and discharge at a diel timescale (Fig. 7), which might occur due to the effects of terrestrial evapotranspiration on water table elevation (Flewelling and others, 2013; Schwab and others, 2016). A higher water table enhances capacity for denitrification in the riparian zone (Harms and Jones 2012), whereas oxygenated soils above the water table might support nitrification (Duncan and others, 2015). However, this mechanism is plausible only in the high-permafrost catchment early in the thaw season, when daily peaks in NO_3^- concentration occurred when discharge was low (Fig. 7). Later in the thaw season, and throughout the record in the low-permafrost catchment, NO_3^- and discharge oscillated synchronously on a diel timescale, or peak daily NO_3^- concentration followed peak daily discharge, which is counter to expectations resulting from fluctuation of the water table, but consistent with N uptake by either in-stream or terrestrial biota. Negative coherence in oscillations of NO_3^- and in-stream GPP at a diel timescale in both catchments (Fig. 7) as well as a negative correlation of daily in-stream GPP with NO_3^- concentration in the high-permafrost stream (Fig. S2) further suggested that stream biota also contributed to diel variation in NO_3^- concentration. Lack of diel variation in fDOM and inconsistent or non-significant correlations of either solute with daily max PAR suggest that in-stream photodegradation did not play a major role in DOM processing and was not coupled to diel NO_3^- variation, as expected in these narrow well-shaded streams.

Implications

Permafrost generated biogeochemical signals that were observable as multi-scale temporal patterns in solute chemistry of headwater streams. Distinct signals of spatially extensive permafrost included seasonal trends, responsiveness to precipitation, counterclockwise hysteresis of concentration-discharge relationships during storms, and increasing influence of biota on diel oscillations through the growing season. Importantly, biogeochemical signals observed in streams resulted from interaction of permafrost with hydrologic flowpaths contributing to streamflow. Quantitative metrics describing these signals could be applied to detect degradation of permafrost at catchment scales, which is occurring due to climate warming, fire, and increased precipitation (Douglas et al. 2020, Brown et al. 2015).

Further, dominance of precipitation as a driver of catchment biogeochemistry in permafrost-influenced boreal headwaters, as demonstrated in this study, portends significant changes to solute export because precipitation is expected to become more variable at high latitudes (Bintanja and others, 2020). Based on the patterns observed here, we predict that increased variation in precipitation will generate an initial increase in temporal variability of solute export, followed by a decrease in variability as permafrost thaw deepens hydrologic flowpaths. In particular, a shift toward clockwise hysteresis of concentration-discharge relationships during storms, decreased discharge and solute responses to storms, and increase in mean NO_3^- or decreased fDOM concentrations are expected indicators of deepening hydrologic flowpaths that accompany permafrost loss.

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