Mounting evidence suggests that biogeochemical processing of permafrost substrate will amplify dissolved inorganic carbon (DIC = Σ[CO$_2$, HCO$_3^-$, CO$_3^{2-}$]) production within Arctic freshwaters. The effects of permafrost thaw on DIC may be particularly strong where terrain subsidence following thaw (thermokarst) releases large amounts of sediment into fluvial networks. The mineral composition and chemical weathering of these sediments has critical yet untested implications for the degree to which streams represent a source of CO$_2$ to the atmosphere vs. a source of bicarbonate to downstream environments. Here, we experimentally determine the effects of mineral weathering on fluvial CO$_2$ by incubating sediments collected from three retrogressive thaw slump features on the Peel Plateau (NWT, Canada). Prehistoric warming and contemporary thermokarst have exposed sediments on the Peel Plateau to varying degrees of thaw and chemical weathering, allowing us to test the role of permafrost and substrate mineral composition on CO$_2$:HCO$_3^-$ balance. We found that recently-thawed sediments (within years to decades) and previously un-thawed tills from deeper permafrost generated substantial amounts of solutes and DIC. These solutes and the mineralogy of sediments suggested that carbonate weathering coupled with sulfide oxidation was a net source of abiotic CO$_2$. Yet, on average, more than 30% of this CO$_2$ was converted to bicarbonate via carbonate buffering reactions. In contrast, the mineralogy and geochemical trends associated with sediments from the modern and paleo-active layer, which were exposed to thaw over longer timescales than deeper permafrost sediments, more strongly reflected silicate weathering. In treatments with sediment from the modern and paleo-active layer, minor carbonate and sulfide weathering resulted in some DIC and net CO$_2$ production. This CO$_2$ was not measurably diminished by carbonate buffering. Together, these trends suggest that prior exposure to thaw and weathering on the Peel Plateau reduced carbonate and sulfide in upper soil layers. We conclude that thermokarst unearthing deeper tills on the Peel Plateau will amplify regional inorganic carbon cycling.
INTRODUCTION

Rapid climate warming and intensifying hydrologic cycles at northern latitudes (Serreze and Barry, 2011) are activating immense stores of organic carbon from perennially frozen permafrost soils into modern biogeochemical cycles (Biskaborn et al., 2019; Vonk et al., 2019). The biotic and abiotic processing of this relatively reactive carbon pool stands to amplify climate warming in the coming centuries by releasing large amounts of carbon dioxide (CO₂) from terrestrial and aquatic environments into Earth’s atmosphere (Schuur et al., 2015; Turetsky et al., 2015; Anderson, 2007). Half of the CO₂ produced by plant root respiration and microbial oxidation of mineral-rich soil layers and accelerating physical erosion enhances solute production by exposing previously unweathered mineral C into solution as H₂CO₃⁺ (Equation 5) or HCO₃⁻ (Equation 6) via carbonate weathering, or generates no dissolved C via silicate weathering (Equation 7) (Lerman et al., 2007):

\[
\text{FeS}_2 + 15/4\text{O}_2 + 7/2\text{H}_2\text{O} \rightarrow \text{Fe(OH)}_3 + 2\text{H}_2\text{SO}_4(4)
\]

\[
2\text{H}_2\text{SO}_4 + 2(\text{Ca}, \text{Mg})\text{CO}_3 \rightarrow 2(\text{Ca}^{2+}, \text{Mg}^{2+}) + 2\text{SO}_4^{2-} + 2\text{H}_2\text{CO}_3^*(5)
\]

\[
\text{H}_2\text{SO}_4 + 2(\text{Ca}, \text{Mg})\text{CO}_3 \rightarrow 2(\text{Ca}^{2+}, \text{Mg}^{2+}) + 2\text{SO}_4^{2-} + 2\text{HCO}_3^*(6)
\]

\[
\text{H}_2\text{SO}_4 + \text{H}_2\text{O} + (\text{Ca}, \text{Mg})\text{Al}_2\text{Si}_2\text{O}_8 \rightarrow (\text{Ca}^{2+}, \text{Mg}^{2+}) + \text{SO}_4^{2-} + 2\text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4(7)
\]

When sediments are more limited in carbonates than in sulfides, H₂SO₄ from sulfdioxide oxidation can lower pH and increase the CO₂ proportion of DIC (Equation 1), facilitating CO₂ release to the atmosphere (Stumm and Morgan, 1996). Half of the HCO₃⁻ transferred from streams and rivers into the marine carbon cycle returns to the atmosphere as CO₂ over geological timescales (Calmels et al., 2007). Thus, the balance between H₂CO₃⁺ and H₂SO₄ weathering controls the degree to which mineral weathering is carbon neutral (Equation 2) vs. a carbon sink (Equation 3) or source (Equations 5, 6) (Calmels et al., 2007; Tank et al., 2012; Hindshaw et al., 2014, 2017; Pogge von Strandmann et al., 2017). Constraining the sources of mineral weathering and the balance between CO₂ and HCO₃⁻ in northern fluvial networks can help to understand DIC cycling and the abiotic component of permafrost-climate feedbacks.

Local and regional variation in lithology, physical erosion, and temperature are primary controls on the sources and magnitude of mineral weathering (Gaillardet et al., 1999; Li et al., 2017). For instance, riverine DIC can be relatively high in catchments with carbonate lithologies (Blum et al., 1998; Tank et al., 2012) as carbonates weather more readily than silicates (Stumm and Morgan, 1996). River chemistry reveals that physical erosion enhances solute production by exposing previously unweathered material (Riebe et al., 2001; West et al., 2005; Calmels et al., 2007) and by increasing the potential reactivity of sediments (Anderson, 2007). DIC cycling in terrains with glacial activity can be rapid. For instance, H₂CO₃⁺ weathering of sediments during fluvial transport in pro-glacial streams can consume dissolved CO₂ to below atmospheric levels (St. Pierre et al., 2019). While much of Canada is no longer glaciated, terrain subsidence following permafrost thaw (thermokarst) across vast swaths of northwestern Canada is enabling large quantities of glaciogenic sediments that were emplaced by the former Laurentide Ice Sheet for decades to centuries. However, CO₂ consumption via carbonate buffering may partly counterbalance CO₂ production and release to the atmosphere. Regional variability in the mineral composition of permafrost, thaw history, and thermokarst intensity are among the primary controls on mineral weathering within permafrost carbon-climate feedbacks.

Keywords: carbonate, sulfide, carbon dioxide, thermokarst, retrogressive thaw slump
to be mobilized into fluvial networks (Kokelj et al., 2017a). In these ice- and sediment-rich glacial margin landscapes like the Peel Plateau (NWT, Canada), retrogressive thaw slump (RTS) thermokarst activity can amplify DIC cycling and increase $\text{HCO}_3^-$ in downstream environments (Zolkos et al., 2018). While $\text{H}_2\text{SO}_4$ carbonate weathering in thermokarst-affected streams on the Peel Plateau is thought to be a net source of $\text{CO}_2$ to the atmosphere over contemporary and longer timescales (Zolkos et al., 2018, 2019), the degree to which mineral weathering by $\text{H}_2\text{CO}_3$ consumes $\text{CO}_2$ in streams on the Peel Plateau remains untested. The balance between $\text{CO}_2$ and $\text{HCO}_3^-$ in thermokarst-affected streams has important implications for the Arctic inorganic carbon cycle, as hillslope thermokarst features like RTSs may affect 900,000 km$^2$ (5%) of the northern permafrost region (Olefeldt et al., 2016). Further, the chemistries of streams on the Peel Plateau reflect weathering from sediment sources with varying degrees of prior exposure to thaw and thus mineral composition (Lacelle et al., 2019). Determining the influence of permafrost mineral composition and weathering sources on the balance between $\text{CO}_2$ and $\text{HCO}_3^-$ in streams can thus help to understand thermokarst effects on DIC cycling and potential feedbacks to long-term climate.

Here, we refine understanding of mineral weathering contributions to inorganic carbon cycling on the Peel Plateau (NWT, Canada) (Zolkos et al., 2018), where accelerating thermokarst activity is mobilizing large volumes of mineral-rich glacial tills with varying degrees of prior thaw into fluvial networks (Kokelj et al., 2013; Segal et al., 2016a; van der Sluijs et al., 2018). We build on previous research that investigated the effects of RTSs on solutes, sediments, and organic carbon in fluvial networks (Kokelj et al., 2013; Malone et al., 2013; Littlefair et al., 2017), and Holocene environmental change on the composition of permafrost soil organic carbon and solution chemistry (Lacelle et al., 2019). To do this, we characterized the mineral composition of RTS sediments that commonly enter streams on the Peel Plateau and experimentally weathered these sediments to determine how thermokarst and thaw history affect DIC concentration and speciation (Segal et al., 2016a; Lacelle et al., 2019). Our results indicate that fluvial inorganic carbon cycling on the Peel Plateau is amplified by more intense thermokarst, which thaws deeper sediments containing a greater proportion of carbonate and sulfide. We conclude that thermokarst on the Peel Plateau and in similar Arctic environments may intensify inorganic carbon cycling in fluvial networks for centuries.

**MATERIALS AND METHODS**

**Study Area and Retrogressive Thaw Slump Sites**

The Peel Plateau is characterized by permafrost tills up to 50 m in thickness underlain by sandstone and siltstone bedrock (Norris, 1985; Kokelj et al., 2017b). These tills are mainly comprised of carbonates, sulfides, silicates, and gypsum (Malone et al., 2013; Zolkos et al., 2018) likely derived from regional carbonate and shale bedrock (Norris, 1985; Stott, 1991) that was integrated into the Laurentide Ice Sheet (LIS) during its westward expansion. The last glacial maximum in the Peel Plateau region at around 18,500 cal yr BP (Lacelle et al., 2013) was followed by retreat of the LIS and early Holocene warming at 12,000–8,500 cal yr BP (Kokelj et al., 2017b). This warming promoted thaw of the ground surface, mineral weathering, and the accumulation of organic matter in the upper few meters of soil (Burn, 1997; Lacelle, 2004; Malone et al., 2013). A cooling climate 7,000–5,500 cal yr BP supported permafrost aggradation and the preservation of this solute- and organic-rich Holocene layer above the deeper, ice-rich Pleistocene sediments that have been isolated from thaw and exposure (Lacelle et al., 2013; Kokelj et al., 2017b). The shallow Holocene and the deeper Pleistocene permafrost layers are geochemically distinct and distinguished stratigraphically by an undulating thaw unconformity typically encountered 1.5–3 m below the surface (Malone et al., 2013) (Figure 1A).

The modern-day Peel Plateau is within the continuous permafrost zone and is comprised of ice-rich hummocky moraine, glaciofluvial, glaciolacustrine, and colluvial deposits (Duk-Rodkin and Hughes, 1992). Incised valleys and sloping terrain host hundreds of RTSs (Figure 1) (Kokelj et al., 2013; Lacelle et al., 2015; Segal et al., 2016a). The upslope growth of RTS features is sustained by the ablation of a vertical permafrost headwall of exposed ground ice and the downslope evacuation of a thawed slurry of slumped materials via fluidized flows which are enhanced by rainfall (Kokelj et al., 2015) (Figure 1A). Meltwater from the thawing headwall and its interaction with thawing substrate produce sediment- and solute-rich runoff often conveyed through rill channels in the scar zone (henceforth “runoff”), and into streams. The thawed materials that flow from thawing slopes are deposited as debris tongues that can grow over time to reach >1 km in length and tens of meters in thickness (van der Sluijs et al., 2018). These features are reshaped by the addition of debris from thawing slopes and erosion by the adjacent streams (Kokelj et al., 2015) (Figure 1B). Sediments unearthed by RTSs can be broadly categorized into five sources of material with varying degrees of prior exposure to thaw and weathering: (i) the seasonally-thawed active layer; (ii) the relatively organic- and solute-rich paleo-active layer contained in Holocene-aged permafrost; (iii) the un-thawed Pleistocene-aged permafrost sediments derived from glacial tills; and the composite of these three components, that are either recently mobilized and are undergoing active transport as fluvial suspended sediment in runoff (iv) or deposited in recent years to decades within the debris tongue (v) (Figure 1). Active layer and Holocene permafrost sediments in RTSs on the Peel Plateau have been exposed to thaw for decades to centuries, whereas Pleistocene permafrost sediments were frozen upon their deposition and have remained sequestered within permafrost for millennia.

The three focal RTSs in this study are situated within the ∼1,100 km$^2$ Stony Creek watershed, west of Fort McPherson, NWT (Figure 1, Table 1). The RTS FM2 is among the largest active RTS features on the Peel Plateau (Lacelle et al., 2015) and has persisted for at least 70 years (Kokelj et al., 2015; van der Sluijs et al., 2018). The RTS HC is moderately-sized and estimates from the field suggest it is likely at least several decades old. The RTS HD scar zone was <1 ha at the time of sampling and visual
Field sampling
During summer 2017, we collected active layer and permafrost sediments from the vertical headwall exposure, in addition to runoff and debris tongue sediments at RTS features FM2, HC, and HD (Figure 1, Table 1). These three RTS features were selected to capture potential variability in sediment geochemistry associated with varying compositions of glaciogenic materials (Duk-Rodkin and Hughes, 1992). To characterize RTS morphology, we measured headwall height at the time of sampling using a Uineye HK1200 laser range finder (±30 cm/0.35° accuracy) (Zolkos and Tank, 2019). Holocene and Pleistocene permafrost samples were collected in each RTS feature using a mallet and stainless-steel chisel. Permafrost layers were differentiated in the field based on cryostratigraphy (Lacelle et al., 2013) and later in the laboratory using stable isotopes of water (δ18O), to ensure isotopic values reflected previously published values for Holocene (δ18O = −17 to 25‰) and Pleistocene (δ18O = −26 to −32‰) ground-ice (Michel, 2011; Fritz et al., 2012; Lacelle et al., 2013). Active layer and debris tongue samples were collected using a stainless-steel trowel. Active layer material was composited from two roughly equal plugs of material from the A and B horizons (below the organic layer and above permafrost table). Debris tongue samples comprised of thawed, colluviated materials were collected near the adjacent stream, distant from the active headwall, reflecting sediments that have been thawed for years to decades. For active layer, permafrost, and debris tongue samples, the outer 5–10 cm of material was removed and discarded prior to sampling. Samples were stored in 5 L Whirlpak bags. Runoff was collected in pre-cleaned, triple-sample-rinsed 2 L HDPE bottles and stored chilled (4°C). Permafrost was thawed, and thawed permafrost and runoff were allowed to settle in the laboratory for 24 h at 4°C. Prior to drying sediments at
20°C, supernatant was removed to limit the precipitation of secondary minerals.

**Sediment Mineralogy and Geochemistry**

Prior to mineral analysis by X-ray diffraction (XRD), dried sediments were gently disaggregated with a mallet, sieved to <250 µm, and visible organic matter was removed. Samples were not pretreated prior to analysis and XRD spectra were visually inspected to confirm minimal interference from organic matter (Mandile and Hutton, 1995). Sieved sediments were then finely ground with a ceramic mortar and pestle, and mineralogy was analyzed at the University of Alberta by XRD (Rigaku Ultimate IV), using Cobalt radiation at 0.6 s/step and 0.02° steps (detection limit = 3%). The geochemical composition of sediments was analyzed by inductively coupled plasma mass spectrometer (Perkin Elmer Elan 6000 Quadrupole ICP-MS) at the University of Alberta Canadian Center for Isotope Microanalysis, following Cooper et al. (2008). The coefficient of variation for Al⁺³, Fe²⁺, K⁺, Mg²⁺, Ca²⁺, Na⁺, and Sr²⁺ is <6%. Subsamples (0.2 ± 0.02 g) of the sieved sediments were dissolved via sequential digestions using HF + HNO₃ and then HCl + HNO₃. Sediments were mixed with five mL of each acid (trace metal-grade), heated to 130°C, and digested overnight until dry. Finally, 10 mL 8N HNO₃ were added, heated to 130°C, and a 1 mL aliquot was mixed with 0.1 mL HNO₃, 0.1 mL of internal standards (In, Bi, and Sc), and 8.8 mL of MilliQ deionized water.

**Mineral Weathering Experiment**

To determine how varying degrees of prior thaw and biogeochemical alteration of glaciogenic materials can influence DIC cycling on the Peel Plateau, we incubated active layer, Holocene, Pleistocene, runoff, and debris tongue sediments under controlled conditions. Following collection and sieving (see above), unground sediments were sterilized in precombusted aluminum trays at 200°C for 24 h in a forced-draft oven (Wolf and Skipper, 1994) to isolate the abiotic controls on DIC speciation from CO₂ production by microbial oxidation of organic matter. In our experiment this likely inhibited microbial sulfide oxidation (Singer and Stumm, 1970), which is a primary biotic mechanism of carbonate and silicate weathering in glaciated environments (Montross et al., 2013; Tranter and Wadham, 2014; Hindshaw et al., 2016), thus constraining possible sulfide oxidation to abiotic pathways involving Fe³⁺ (Rimstidt and Vaughan, 2003) and dissolved O₂ (Calmels et al., 2007). Sterilization was tested by mixing sterile MilliQ water (0.2 µm-filtered, ultraviolet light-treated) with the sterilized sediment, and extracting and incubating the supernatant on nutrient-rich agar plates (37°C, 7 d; Zuberer, 1994). Sterilization was confirmed by no observable growth compared to identical sediments that had not been sterilized. To evaluate the relative importance of sulfide oxidation and gypsum dissolution to SO₄²⁻ production, we assumed that 1 mol O₂ could yield 0.53 mol H₂SO₄ via sulfide oxidation (Equation 4) and calculated the concentration of dissolved S₀²⁻ in the bottles under equilibrium conditions (10°C) to be 348 µM. This approach allowed us to estimate that S₀²⁻ concentrations up to 184 µM could originate from sulfide oxidation via the dissolved O₂ pathway (Equations 5, 6). We interpreted higher S₀²⁻ concentrations and the absence of sulfides from XRD detection as indicators of relatively greater S₀²⁻ production from gypsum dissolution.

Sterilized sediments were incubated at total suspended solid concentrations (1,800 mg L⁻¹) and water temperatures (10°C) representative of stream reaches immediately downstream of RTS features during the 2015 sampling season (Zolkos et al., 2018). We inoculated 200 mL of MilliQ water with 3.6 g of sterilized sediment in acid-washed, precombusted (5 h, 500°C) 250 mL glass bottles and sealed them with two silicone-teflon septa (Skidmore et al., 2004). For each sediment source, 12 identical bottles were prepared and replicate bottles were terminated at 0.17, 1, 6, 24, 72, and 168 h. Two control bottles containing only MilliQ water were also terminated at 168 h. We enabled weathering to proceed under closed system conditions by flushing headspace in bottles with N₂ gas, enabling dissolved CO₂ and O₂ to contribute to carbonate weathering and sulfide oxidation. For RTS HD runoff, we had only enough sediment to run replicate bottles that were terminated at 168 h and we had no replicates for the other time points. To induce sediment suspension, all bottles were incubated on a rotary shaker set to 250 rpm during the course of the experiment, following Skidmore et al. (2004).

**Chemical Analyses**

At each time point, bottles were removed and water was filtered (0.45-µm, polyethersulfone, ThermoScientific) for DIC, pH, and conductivity measurements. Water was also equilibrated with nitrogen gas to extract dissolved gas for measurements of CO₂ concentration and stable carbon isotope analysis of CO₂ (δ¹³C-CO₂) (see below). Filtrate for DIC was stored without headspace in precombusted 12 mL exetainers, sealed with butyl septa, and refrigerated until analysis within 2 weeks (Apollo SciTech DIC analyzer). DIC standard curves were

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**TABLE 1** Characteristics of RTS features in this study.

| RTS | Lat (DD) | Lon (DD) | δ¹⁸O Holocene | δ¹⁸O Pleistocene | Height (m, max) | Debris tongue length (~m) | Scar zone (~ha) |
|-----|----------|----------|---------------|-----------------|-----------------|--------------------------|----------------|
| FM2 | 67.257   | −135.236 | −20.3         | −28.3           | 24.2            | 1,600                    | 27             |
| HC  | 67.328   | −135.900 | −21.6         | −29.2           | 18.3            | 180                      | 2.5            |
| HD  | 67.400   | −135.334 | −21.8         | −30.3           | 13.6            | 150                      | 0.6            |

All headwall height measurements were made during July 2017. Stable water isotope (δ¹⁸O) values are reported in ‰ Vienna Standard Mean Ocean Water. *Estimated from satellite imagery and/or field photos; excludes area of debris tongue.
built from certified reference material (Scripps Institution of Oceanography) and, when required for low-DIC samples, from 100 or 600 μM solutions prepared from a 1,000 ppm Na₂CO₃-NaHCO₃ DIC standard (ACCUSPEC). pH and conductivity were analyzed within 2 h of filtering. pH was analyzed with a Mettler Toledo FiveEasy benchtop meter calibrated daily using a two-point calibration curve (7.01 and 10.01 pH, NIST). The mean pH of replicate samples was calculated from first converting pH to H⁺.

Conductivity was analyzed with a Thermo Orion 115 A+ meter calibrated daily using 1,413 μS cm⁻¹ solution. Dissolved CO₂ was extracted by equilibrating 20 mL of sample with 40 mL of nitrogen gas in an air-tight polypropylene syringe (Hesslein et al., 1991) and measured immediately by infrared gas analyzer (IRGA; PP Systems EGM-4). IRGA drift was checked daily with 60 mL of 1,010 ppm CO₂ standard (Scotty Gases). CO₂ concentration (μM) and the partial pressure of CO₂ (pCO₂) were calculated using Henry’s constants corrected for water temperature (Weiss, 1974). For instances when DIC concentration could not be measured by the Apollo, DIC concentration was calculated using pH and pCO₂ as inputs to CO₂SYS (v.2.3; Pierrot et al., 2006).

At the final time point of each experiment, water was also filtered (0.45-μm, cellulose acetate, Sarstourius) to determine the concentrations of cations and anions. Cations were preserved with trace metal-grade HNO₃ and all ions were refrigerated until analysis at the University of Alberta Biogeochemical Analytical Services Laboratory (BASL, ISO/EIC 17025) following standard procedures. Equilibrated headspace gas samples for δ¹³C CO₂ were stored over-pressurized in bottles that were sealed with baked (12 h, 60°C) butyl stoppers and pre-evacuated. δ¹³C CO₂ was analyzed within 2 months by a Picarro isotope analyzer (G2201-I; < 0.2‰ precision) equipped with an injection module for discrete samples (SSIM). The Picarro analyzer was checked for drift against commercial and in-house δ¹³C CO₂ standards during each run.

δ¹⁸O of permafrost ice was analyzed by cavity ringdown spectroscopy (Picarro L2130-i) at the University of Alberta BASL. Calibration curves for water isotopes were built from U.S. Geological Survey secondary standards (USGS45 and USGS46) and in-house tap water, and calibrated by an International Atomic Energy Agency standard as a quality control.

**Data Analysis**

We report treatment replicates as mean ± standard deviation for DIC, CO₂, and conductivity and mean ± range for pH. We used the proportions of solute equivalent concentrations in a Piper diagram (Piper, 1944) to characterize potential mineral weathering reactions (see also Lehn et al., 2017). Calculations were made using the concentrations of cations and anions in the experiment bottles at 168 h, first correcting for the solute concentrations measured in the MilliQ control bottles (Table A1 and Supplementary Material). For the Piper diagram, we calculated HCO₃⁻ as Σ[HCO₃⁻, CO₃²⁻] to correct for HCO₃⁻ hydrolysis to CO₂. The concentrations of dissolved CO₂ measured at each time point were corrected for dissolved atmospheric CO₂ in the control bottles (15-26 μM). To test for multivariate similarity (i.e., geochemical likeness) between treatments, we first performed a hierarchical cluster analysis using the R software package pvclust (Suzuki and Shimodaira, 2015), which employs the R base function hclust to generate a bootstrapped (n = 1,000 iterations), Euclidean distance-based estimate of a stable configuration of measurement clusters. Variables were selected for the pvclust to yield insight about DIC concentration and speciation (HCO₃⁻, CO₂) and CO₂ source (δ¹³C CO₂); to help differentiate between silicate and carbonate weathering (Al³⁺); and to evaluate contributions from sulfide oxidation and gypsum dissolution (SO₄²⁻). We then evaluated the associations between these geochemical variables and the sample clusters defined by pvclust using the metaMDS function in the R software package vegan (Oksanen et al., 2013). metaMDS augments traditional non-metric multidimensional scaling (NMDS) by running multiple MDS iterations to find a stable solution, whereby proximity of measurements indicates similarity and vectors reveal how variables correlate with measurements. All statistics were performed in R software v.3.4.4 (R Core Team, 2018), with significance interpreted at α = 0.05.

**RESULTS**

**Sediment Mineral and Geochemical Composition**

Permafrost sediments, RTS runoff, and debris tongue deposits contained carbonates (calcite, dolomite, and rarely siderite), pyrite, gypsum, quartz, and various Na-, Mg-, and K-aluminosilicates (Table 2; note the XRD detection limit of 3%). Albite, clinochlore, muscovite, and quartz were detected in all sediment sources. The XRD detected carbonates, pyrite, and gypsum in at least half of the sediment sources and in at least one source at each RTS. Carbonates, sulfides, and gypsum were generally present (>3%) in Pleistocene permafrost sediments and absent from active layer sediments (below detection), whereas the presence of these minerals in Holocene, runoff, and debris tongue sediments varied (Table 2). The presence of sulfide and/or carbonate varied among sediment sources, and there were no sediments where carbonate was detected by XRD and sulfide was not.

The geochemical composition of sediments was characterized by—in decreasing order of concentration—Al³⁺, Fe³⁺, K⁺, Mg²⁺, Ca²⁺, Na⁺, and Sr²⁺ (Table 3). Because concentrations of Ca²⁺ relative to Na⁺ and Sr²⁺ are typically higher for carbonates than silicates (Keller et al., 2007), we used molar ratios to discern the relative proportion of carbonates (high Ca/Na and Ca/Sr) vs. silicates in sediments. Ca/Sr and Ca/Na were typically higher in sediments where carbonates were detected by XRD (Table 2, Figure 2). Some sediments with sulfides but not carbonates also had relatively high Ca/Na and Ca/Sr (e.g., FM2 Holocene and Pleistocene, HD debris tongue), perhaps reflecting the presence of carbonates below the XRD detection limit.

**Weathering Experiment: Trends in Conductivity, DIC, CO₂, and pH**

For all RTSs and sediment sources, conductivity increased most strongly during the first 10 min of the incubation, and increased marginally thereafter, indicating that the
majority of the most reactive component of the mineral pool weathered by 168 h (Figures 3A–C). At 168 h, conductivity in treatments with Pleistocene permafrost, runoff, and debris tongue sediments (mean = 264 µS cm$^{-1}$) was significantly higher than in treatments with active layer and Holocene permafrost sediments [mean = 138 µS cm$^{-1}$, excluding HC Holocene [552 µS cm$^{-1}$]] (t-test: $t_{18} = 3.26$, $p < 0.001$). Solutes at 168 h broadly reflected sulfide oxidation and sulfate salt (e.g., gypsum) dissolution (SSD) and relatively greater contributions from silicate and carbonate weathering by $\text{H}_2\text{SO}_4$ than $\text{H}_2\text{CO}_3$ (Figure 4). Across RTSs and sediment sources, $\text{SO}_{4}^{2-}$ concentrations at 168 h were often greater than would be expected from sulfide oxidation alone (section Mineral Weathering Experiment) (Table 4), suggesting that gypsum dissolution was a primary source of $\text{SO}_{4}^{2-}$ in our experiment.

**TABLE 2** | Minerals detected by X-ray diffraction (x) in non-sterilized sediments.

| RTS Source Cluster | Gypsum | Pyrite | Calcite | Dolomite | Albite | Orthoclase/sanidine | Clinochlore | Muscovite | Quartz | Other |
|--------------------|--------|--------|---------|----------|-------|---------------------|-------------|-----------|--------|-------|
| FM2 AL 3           |        |        |         |          |       |                     |             |           |        |       |
| FM2 HO 3           |        |        |         |          |       |                     |             |           |        |       |
| FM2 PL 2           | x      | X      |         |          | x     |                     |             |           |        |       |
| FM2 RU 1           | x      | X      | x       |          | x     |                     |             |           |        |       |
| FM2 DT 1           | x      | X      | x       |          | x     |                     |             |           |        |       |
| HC AL 3            |        |        |         |          |       |                     |             |           |        |       |
| HC HO 1            | x      | X      | x       |          | x     |                     |             |           |        |       |
| HC PL 1            | X      | x      |         |          | x     |                     |             |           |        |       |
| HC RU 1            | x      | x      |         |          | x     |                     |             |           |        |       |
| HC DT 1            | x      | X      |         |          | x     |                     |             |           |        |       |
| HD AL 3            |        |        |         |          |       |                     |             |           |        |       |
| HD HO 3            |        |        |         |          |       |                     |             |           |        |       |
| HD PL 1            | x      | x      | x       |          | x     |                     |             |           |        |       |
| HD RU 2            |        |        |         |          |       |                     |             |           |        |       |
| HD DT 2            | x      | X      |         |          | x     |                     |             |           |        |       |

AL, active layer; HO, Holocene; PL, Pleistocene; RU, runoff; DT, debris tongue. Other minerals: *Siderite: FeCO$_3$; †Lepidolite: KLi$_2$Al(Si$_4$O$_{10}$)(F,OH)$_2$.

**TABLE 3** | Geochemical composition of sediments, measured by ICP-MS. Concentrations are reported in $\mu$mol g$^{-1}$ of dry sediment.

| Site | Source | Cluster | $\text{Al}^{3+}$ | $\text{Fe}^{2+}$ | $\text{K}^+$ | $\text{Mg}^{2+}$ | $\text{Ca}^{2+}$ | $\text{Na}^+$ | $\text{Sr}^{2+}$ |
|------|--------|---------|------------------|------------------|--------------|------------------|------------------|--------------|--------------|
| FM2  | AL     | 3       | 1,665            | 650              | 435           | 319              | 159              | 218          | 1.15         |
| FM2  | HO*    | 3       | 1,454            | 635              | 376           | 291              | 234              | 194          | 1.21         |
| FM2  | PL*    | 2       | 2,458            | 684              | 665           | 480              | 153              | 152          | 1.60         |
| FM2  | RU**   | 1       | 1,584            | 627              | 420           | 395              | 271              | 246          | 1.31         |
| FM2  | DT**   | 1       | 1,601            | 629              | 467           | 388              | 256              | 208          | 1.34         |
| HC   | AL     | 3       | 1,811            | 599              | 464           | 302              | 53               | 206          | 1.08         |
| HC   | HO**   | 1       | 1,722            | 594              | 495           | 396              | 240              | 207          | 1.26         |
| HC   | PL**   | 1       | 1,324            | 526              | 341           | 284              | 188              | 201          | 1.17         |
| HC   | RU**   | 1       | 1,560            | 556              | 430           | 359              | 175              | 237          | 1.32         |
| HC   | DT**   | 1       | 1,586            | 593              | 441           | 371              | 169              | 246          | 1.24         |
| HD   | AL     | 3       | 1,625            | 600              | 494           | 302              | 31               | 198          | 1.15         |
| HD   | HO     | 3       | 2,010            | 605              | 511           | 336              | 49               | 196          | 1.37         |
| HD   | PL**   | 1       | 1,986            | 646              | 519           | 402              | 202              | 195          | 1.38         |
| HD   | RU     | 2       | 2,330            | 752              | 651           | 479              | 113              | 181          | 1.51         |
| HD   | DT*    | 2       | 1,799            | 649              | 506           | 380              | 142              | 208          | 1.36         |

Minerals detected by XRD: **Sulfides and carbonates, *sulfides.
For all treatments, DIC concentrations increased rapidly during the first 6 h and more gradually after 24–48 h. Treatments containing sediments with carbonates and sulfides detected by XRD generated more DIC (mean = 728 μM after 168 h) than those in which only sulfides were detected (mean = 345 μM), or neither carbonates nor sulfides were detected (mean = 121 μM) (Figures 3D–F, Table 4). Accordingly, treatments with active layer and Holocene sediments generally had the lowest DIC at 168 h (mean = 111 and 342 μM, respectively), whereas DIC in treatments with Pleistocene, runoff, and debris tongue sediments had higher concentrations (581, 556, and 654 μM, respectively) (Table 4).

For all treatments, CO₂ increased rapidly within 6 h (Figures 3G–I) and, in treatments containing sediments with both carbonates and sulfides detected by XRD, decreased by 16–39 μM thereafter. The decrease in CO₂ was equivalent to 19–51% (mean = 33%) of the maximum CO₂ reached during the experiment for these treatments. At 168 h, CO₂ in these treatments was lower (mean = 75 μM) than in other treatments. In contrast, CO₂ in treatments containing sediments with sulfides but not carbonates detected by XRD generally reached the highest concentrations of CO₂ by 168 h (mean = 212 μM). Most of the treatments with active layer and Holocene sediments, which did not have carbonates or sulfides detected by XRD, had variable but relatively moderate CO₂ (mean = 95 μM).

pH increased in all treatments except for those with FM2 active layer sediments, which decreased from 7.04 to 6.69 pH units after 24 h. By 168 h, pH was highest where carbonates and sulfides were detected by XRD (mean = 7.23 pH units), lower when sulfides but not carbonates were detected (mean = 6.62), and the lowest in treatments where neither carbonates nor sulfides were detected (mean = 4.70) (Figures 3J–L).

**Statistical Clusters of Treatments Based on Geochemistry**

The cluster analysis revealed that treatments grouped into three geochemical clusters which broadly reflected gradients of prior thaw and exposure to weathering, and also the intensity of sulfide oxidation relative to carbonate weathering (Figure A1). Clusters one and two mostly included samples with more limited prior thaw compared to cluster three, which contained most of the active layer and Holocene permafrost sediments (Figure 5). Cluster one included sediments characterized by carbonate weathering that buffered H₂SO₄ acidity from sulfide oxidation more effectively than the sediments in cluster two. Cluster three included sediments which had relatively limited effects on DIC upon weathering. These statistical clusters help to describe the influence of prior exposure to thaw and the balance between carbonate weathering and sulfide oxidation on the evolution of DIC and CO₂ during the experiment.

**Cluster 1—High DIC and pH, Low Al³⁺**

Treatments containing sediments with both carbonates and sulfides detected by XRD (Table 2) had consistently high DIC (mean = 728 μM), elevated pH (mean = 7.22 pH units), and low dissolved Al³⁺ (mean = 0.1 μM) (Figure 3, Table 4). In this cluster, mineral weathering resulted in rapid CO₂ production early in the experiment followed by a decrease after ~24 h. Yet, CO₂ was always higher than in the control bottles. δ¹³C₀₂ at 168 h varied between treatments, but was generally intermediate (~8.3 to −13.2‰, n = 7) relative to samples in the other clusters (Table 4). Treatments in this cluster include Pleistocene permafrost sediments or recently thawed/colluviated material (i.e., runoff or debris tongue sediments), but not sediments from the active layer or Holocene permafrost.

**Cluster 2—High SO₄²⁻ and CO₂, ¹³C-Enriched CO₂**

Bottles with FM2 Pleistocene, HD runoff, and HD debris tongue sediments had the highest CO₂ concentrations (Table 4), with no measurable decrease in CO₂ during the 168 h incubation. Generally, these treatments had ¹³C-enriched CO₂ (~4.7 to −7.8‰), high concentrations of SO₄²⁻ (mean = 1.307 μM), and had sulfides and gypsum detected by XRD (except for HD runoff), but not carbonates. DIC concentrations were intermediate between clusters one and three (mean = 279 μM).

**Cluster 3—Low DIC and SO₄²⁻, High Dissolved Al³⁺, and ¹³C-Depleted CO₂**

Bottles with active layer and Holocene sediments (except HC Holocene) had high concentrations of dissolved Al³⁺ (mean = 20.3 μM), consistently lower conductivity, moderate to low DIC (mean = 160 μM), low SO₄²⁻ (mean = 202 μM), and ¹³C-depleted CO₂ (~16.3 to −20.3‰, n = 5) at 168 h (Figure 3, Table 4). There were no Pleistocene permafrost, runoff, or debris tongue sediments in this cluster.

**DISCUSSION**

Our findings reveal several interacting controls from thermokarst on fluvial inorganic carbon cycling on the Peel Plateau. In
FIGURE 3 | Weathering experiment trends in (A–C) conductivity, (D–F) dissolved inorganic carbon (DIC), (G–I) dissolved CO₂, and (J–L) pH by RTS and sediment source. Error bars represent ± standard deviation for conductivity, DIC, and CO₂, and range for pH (bars do not appear where smaller than symbol). See Figure 1 for locations and Table 1 for site characteristics. (G–I) Dashed lines represent CO₂ concentration in control bottles.

the modern and paleo (Holocene) active layer, we find that prior thaw and exposure of sediments to chemical weathering depleted carbonates, sulfides, and gypsum in these upper soil layers (Lacelle et al., 2019). In contrast, Pleistocene permafrost and recently thawed and colluviated sediments within the RTS runoff and debris tongue contain more of these readily
FIGURE 4 | Piper diagram showing the chemical composition of MilliQ water treated with five different retrogressive thaw slump sediments (see legend) exposed to varying degrees of prior thaw. The composition of each point reflects the relative concentrations (in meq L\(^{-1}\)) of major cations (left), anions (right), and both together (center) as a percent. See details in main text for calculation of end-members: CACW, \(\text{H}_2\text{CO}_3\) carbonate weathering; SACW, \(\text{H}_2\text{SO}_4\) carbonate weathering; CASW, \(\text{H}_2\text{CO}_3\) silicate weathering; SASW, \(\text{H}_2\text{SO}_4\) silicate weathering; SSD, sulfate salt (gypsum) dissolution.

weatherable sediments. Thus, thermokarst which unearths Pleistocene permafrost sediments at depth is more likely to intensify DIC cycling and abiotic \(\text{CO}_2\) production by facilitating \(\text{H}_2\text{SO}_4\) carbonate weathering, often in the presence of sulfides (Zolkos et al., 2018). Broadly, these trends show that renewed deglaciation on the Peel Plateau is accompanied by associated mineral weathering and biogeochemical dynamics documented in actively glacierized regions of the circumpolar north (Hindshaw et al., 2016; St. Pierre et al., 2019; Urra et al., 2019). These findings help to contextualize the effects of permafrost thaw on mineral weathering and inorganic carbon cycling on the Peel Plateau (Figure 6) and potentially in ice-rich, glaciated terrains elsewhere in northwestern Canada, Alaska, and Eurasia (Segal et al., 2016a; Kokelj et al., 2017a; Rudy et al., 2017).

Implications of Thaw History and Prior Sediment Exposure to Mineral Weathering for Inorganic Carbon Cycling

The lower conductivity observed in most treatments with active layer and Holocene sediments (except HC Holocene; Table 2, Figure 5) suggests that prior thaw of the upper layers of permafrost on the Peel Plateau promoted chemical weathering of readily-weathered minerals within the upper few meters of soil (Malone et al., 2013; Lacelle et al., 2019). Except for HC Holocene, active layer, and Holocene sediments did not have carbonate detected by XRD and also often had lower Ca/Sr and Ca/Na ratios. The relatively high DIC production associated with HC Holocene sediments may reflect the local presence of colluviated substrate from former debris flows that was integrated into permafrost during its upward aggradation.
The greater solute production associated with debris tongue sediments (Figure 3) indicates relatively limited weathering prior to burial within debris tongues, where some protection from wetting within the interior may reduce weathering prior to sample collection. Where RTSs on the Peel Plateau expose the Holocene active layer, the effects on fluvial inorganic carbon cycling will likely be greater if the substrate contains paleo debris flow material (Lacelle et al., 2019). The more modest DIC generation associated with active layer and Holocene sediments that did not contain debris tongue material suggests that carbonates were present, but in amounts typically minor enough to not be detected by XRD (here, <3%). This is consistent with observations of modern and paleo-active layer soils in Alaska (USA) and the central Yukon Territory (Canada), where top-down thaw promoted chemical weathering and a reduction of carbonates relative to silicates in upper soil layers (Burn et al., 1986; Keller et al., 2010). These findings broadly demonstrate that prior thaw and local and regional heterogeneity in soil composition are among the primary abiotic controls on fluvial inorganic carbon cycling on the Peel Plateau and likely in permafrost terrains across the circumpolar north.

**Thermokarst Sediment Mineral Composition As a Driver of DIC Speciation in Streams**

The grouping of treatments into distinct geochemical clusters indicates that thaw history enables a differentiation in mineralogy that provides an overarching control on the abiotic component of inorganic carbon cycling in thermokarst-affected streams on the Peel Plateau. Sediments in cluster one, which contained the most carbonates and sulfides (both >3%) and had more limited prior exposure to thaw, yielded the highest DIC concentrations and showed a trend of decreasing CO$_2$ after ∼24 h (Figure 3). The decrease in CO$_2$, high DIC, and elevated pH (mean ± 0.7) associated with sediments in cluster one (Figure 5) suggests that H$_2$CO$_3$ carbonate weathering converted some of the initially-produced CO$_2$ to HCO$_3^−$. The decrease in δ$^{13}$C$_{CO_2}$ values expected from this pH buffering (fractionation ca. −11‰) (Clark and Fritz, 1997) and the intermediate δ$^{13}$C$_{CO_2}$ values for cluster one (−8 to −13‰) indicate weathering of carbonate sediments that were isotopically similar to regional bedrock (δ$^{13}$C carbonate = −0.7 to −5.6‰) (Hitchon and Krouse, 1972) that may be present within tills on the Peel Plateau. Together, these trends

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**TABLE 4** Measurements of water chemistry from the mineral weathering experiment at 168 h.

| Site | Source | Cluster | pH (µS cm$^{-1}$) | Cond (µM) | DIC (µM) | Ca$^{2+}$ (µM) | Mg$^{2+}$ (µM) | Na$^+$ (µM) | K$^+$ (µM) | SO$_4^{2-}$ (µM) | Cl$^−$ (µM) | Al$^{3+}$ (µM) | δ$^{13}$C$_{CO_2}$ (% VPDB) |
|------|--------|---------|------------------|-----------|----------|-------------|-------------|-----------|-----------|--------------|------------|--------------|------------------|
| FM2  | AL     | 3       | 6.69             | 132.8     | 215.9    | 642.2       | 114.4       | 13.9      | 38.6      | 90.3         | 3.4        | 7.9          | −16.3            |
| FM2  | HO     | 3       | 6.68             | 202.2     | 392.1    | 1009.8     | 165.0       | 10.9      | 37.1      | 157.5        | 2.3        | 8.7          | −16.3            |
| FM2  | PL     | 2       | 6.65             | 357.7     | 244.4    | 1093.6     | 496.2       | 79.6      | 103.6     | 1476.0       | 2.3        | 0.1          | −7.8             |
| FM2  | RU     | 1       | 7.25             | 404.2     | 760.4    | 1649.5     | 308.6       | 52.2      | 49.1      | 1471.1       | 2.5        | 0.2          | −12.2            |
| FM2  | DT     | 2       | 7.39             | 286.2     | 851.0    | 1152.0     | 190.5       | 61.3      | 67.8      | 783.4        | 1.4        | 0.3          | −13.2            |

Cluster means (standard error, range for pH)

|          | 1 | 2 | 3 |
|----------|---|---|---|
| HD AL    | 4.19 | 126.1 | 53.1 | 169.4 | 82.3 | 34.4 | 35.3 | 313.6 | 2.5 | 10.0 | −20.3 |
| HD HO    | 4.12 | 158.5 | 75.8 | 283.7 | 102.9 | 75.3 | 35.0 | 381.1 | 2.8 | 49.5 | −19.1 |
| HD PL**  | 6.98 | 337.9 | 898.1 | 1212.4 | 337.4 | 36.1 | 67.0 | 1041.6 | 1.4 | 0.1 | −9.9  |
| HD RU    | 5.51 | 320.4 | 195.0 | 725.1 | 421.3 | 349.7 | 99.0 | 1153.0 | 3.4 | 16.1 | −6.9  |
| HD DT*   | 6.28 | 363.9 | 398.0 | 1047.7 | 509.8 | 128.6 | 76.0 | 1291.3 | 1.7 | 2.9 | −4.7  |

Source means (standard deviation, range for pH)

|        | AL  | HO  | PL  | RU  | DT  |
|--------|-----|-----|-----|-----|-----|
| 1      | 4.6 (1.2) | 5.0 (1.3) | 6.9 (0.3) | 6.0 (0.9) | 6.7 (0.6) |
| 2      | 7.21 (10.1) | 5.9 (0.6) | 3.2 (1.2) | 3.6 (1.2) | 3.2 (1.2) |
| 3      | 4.7 (1.2) | 6.9 (0.3) | 6.0 (0.9) | 6.7 (0.6) | 6.4 (0.9) |

*AL*, active layer; *HO*, Holocene; *PL*, Pleistocene; *RU*, runoff; *DT*, debris tongue. See text for details. Minerals detected by XRD: **Sulfides and carbonates, *sulfides.*

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for the cluster one treatments suggests that, when carbonates are sufficiently abundant, carbonate weathering by $H_2CO_3$ (Equation 2) and pH buffering (Equation 1) can fix some $CO_2$ as $HCO_3^-$ (Brantley et al., 2014) and thereby partially offset the high rates of $CO_2$ production and efflux within RTSs on the Peel Plateau (Zolkos et al., 2019). Although we did not observe it in our treatments, sediments with abundant carbonate and limited or no sulfides would have even stronger $CO_2$ consumption (Stiegl et al., 2007; St. Pierre et al., 2019). In contrast, the striking $CO_2$ production in treatments with a greater proportion of sulfide (cluster two) suggests that carbonate weathering was insufficient to noticeably temper dissolved $CO_2$ concentrations. Rather, more intense sulfide oxidation than carbonate weathering likely sustained a relatively low pH (mean $= 5.9$, Table 2) and enabled more complete dissolution of trace carbonates, resulting in $\delta^{13}C_{CO_2}$ (−16 to −20‰) reflective of an abiotic $CO_2$ source. In cluster three, the relatively more $^{13}C$-depleted $\delta^{13}C_{CO_2}$ values (−16 to −20‰) may partly reflect the abiotic breakdown of organic matter from the modern and paleo Holocene active layer soils in a relatively acidic environment (Bao et al., 2019). For all clusters, concentrations of $SO_4^{2−}$ were typically greater than would be expected from sulfide oxidation alone (section Mineral Weathering Experiment), indicating that gypsum dissolution generated a large proportion of the $SO_4^{2−}$ that we observed in our experiment. This suggests that even a small proportion of $SO_4^{2−}$ derived from sulfide oxidation can have critical implications for DIC cycling.

These findings build on recent work showing that RTSs exposing Pleistocene permafrost sediments can amplify inorganic carbon cycling (Zolkos et al., 2018) by revealing that the chemical weathering of these sediments on the Peel Plateau is most likely to be a net source of abiotic $CO_2$. This abiotic $CO_2$ flux from Peel Plateau thermokarst streams to the atmosphere—which occurs at high rates, but over a small area—is likely minor relative to $CO_2$ release from biotic processes occurring within recently thawed permafrost substrate (Littlefair and Tank, 2018; Zolkos et al., 2019) and from thickening of the active layer (Stiegl et al., 2005). As thermokarst intensifies on the Peel Plateau, RTSs may increase fluvial export of $HCO_3^−$ into marine environments, where climatic effects would unfold over geological timescales as carbonate precipitation reactions release $CO_2$ to the atmosphere (Calmels et al., 2007).

**Figure 5** | metaMDS results showing similarities between measurements (proximate = more similar) and correlations between measurements and original observations (black text). Note RTS IDs included above measurements. Ellipses constrain statistical clusters in Figure A1 (see section Results in main text).

**Figure 6** | Refined conceptual model of RTS effects on inorganic carbon cycling on the Peel Plateau (A) in the presence of carbonates and absence of sulfides and (B) in the presence of both carbonates and sulfides. (C) Depicts how this paper builds on previous understanding. Adapted from Zolkos et al. (2018).
tongue material reveals that these recently thawed sediments can augment fluvial inorganic carbon cycling for at least years to decades as RTSs develop. Initially, we reasoned that debris tongue sediments would have a limited effect on DIC, because this material is thawed and exposed at the surface environment prior to burial by subsequent debris flows (Kokelj et al., 2017b). Contrary to our hypothesis, RTS debris tongue sediments were associated with consistently strong DIC production, indicating greater protection from weathering and more limited depletion of carbonate and sulfide than the active layer and Holocene permafrost sediments. Mineral weathering in debris tongue environments may be enhanced as vegetation colonizes stabilized areas (Cray and Pollard, 2015) and as sediments erode into streams (van der Sluijs et al., 2018). From an inorganic carbon cycling perspective, this suggests that previously un-thawed sediments that are redistributed following thaw may be an important legacy of thermokarst on the Peel Plateau, particularly if increasing thaw and rainfall facilitate exposure and the transport of RTS sediments into stream networks (Kokelj et al., 2015; van der Sluijs et al., 2018). These interpretations align with observations of increasing downstream geochemical signals of carbonate weathering and sulfide oxidation (Zolkos et al., 2018) in conjunction with intensifying regional thermokarst in recent decades (Segal et al., 2016a). As future warming and increasing rainfall on the Peel Plateau increase RTS activity, the thawing of minerals in deeper permafrost can thus be expected to accelerate inorganic carbon cycling for decades to centuries.

**Implications for the Role of Mineral Weathering Within Permafrost Carbon Feedbacks**

Across the pan-Arctic, increasing riverine solute fluxes in recent decades signal a thickening of active layers, deepening flow paths, and enhanced mineral weathering in concert with climate warming and intensifying precipitation (Tank et al., 2016; Toohey et al., 2016; Drake et al., 2018a; Zolkos et al., 2018). While increasing riverine alkalinity fluxes in North America (Peel, Mackenzie; Tank et al., 2016; Zolkos et al., 2018) and Siberia (Ob’, Yenisey; Drake et al., 2018a) reflect a broad scale acceleration of inorganic carbon cycling, stronger coupling between carbonate weathering and sulfide oxidation in northwestern Canada diminishes the weathering CO₂ sink compared to other Arctic regions (Beaulieu et al., 2012). From an Arctic carbon cycling perspective, regional variability in lithology, hydrology (Tank et al., 2012), and microbial processes (Hindshaw et al., 2016) influence the mineral weathering component of carbon-climate feedbacks. Recent work suggests that biotic processes drive DIC cycling in ice- and organic-rich (syngenetic) regions affected by thermokarst (Drake et al., 2018b). Working in a relatively more mineral-rich permafrost terrain, we show that intensifying thermokarst stands to amplify abiotic processes, but that the presence of sulfide oxidation is a primary control on the degree to which mineral weathering represents a CO₂ sink. Despite increasing recognition that mineral weathering is an important component of the northern carbon cycle (Calmels et al., 2007; Beaulieu et al., 2012; Tank et al., 2012, 2016), the contribution of thermokarst to these processes remains poorly constrained.

This work indicates that thermokarst will have strong effects on inorganic carbon cycling where glaciated terrains coincide with carbonate lithologies (Zolkos et al., 2018; St. Pierre et al., 2019) and perhaps most strongly where sediments remain unmodified at the near surface (e.g., Rudy et al., 2017; Lewkowicz and Way, 2019; Ward Jones et al., 2019).

Thermokarst can expose large volumes of sediments to mineral weathering and fluvial transport (Kokelj et al., 2013; van der Sluijs et al., 2018), with unknown effects on CO₂ exchange between streams and the atmosphere as HCO₃⁻ is exported downstream. These effects are likely to be particularly strong across glacial deposits, where carbonates and sulfides are often highly reactive (Blum et al., 1998; Anderson et al., 2000; Torres et al., 2017) and even trace amounts can profoundly affect inorganic carbon cycling. Quantifying the fluvial CO₂: HCO₃⁻ balance across broader scales in thermokarst-affected fluvial networks will help to trace and better understand the effects of permafrost thaw on inorganic carbon cycling (Drake et al., 2018a).

**CONCLUSIONS**

In this study, we found evidence to suggest that thermokarst activity on the Peel Plateau will amplify inorganic carbon cycling where RTSs facilitate the exposure of deeper, previously unweathered permafrost sediments. Across the circumpolar north, future increases in inorganic carbon cycling will be strongest where thermokarst unearths carbonate-bearing tills with limited prior thaw and modification (Kokelj et al., 2017b). In these regions, the progression of—and balance between—H₂CO₃ and H₂SO₄ weathering will determine the degree to which DIC in freshwaters represents a CO₂ sink (Calmels et al., 2007; Tank et al., 2012, 2016; Drake et al., 2018a). By investigating the composition and reactivity of mineral substrate in permafrost regions elsewhere, particularly where thermokarst may expose and mobilize vast stores of minerals into fluvial networks (e.g., Olefeldt et al., 2016), future work will advance understanding of the role of mineral weathering within contemporary inorganic carbon cycling and long-term feedbacks within Earth’s climate system (Schuur et al., 2015; Zolkos et al., 2018).

**DATA AVAILABILITY STATEMENT**

All datasets generated for this study are included in the article/Supplementary Material.

**AUTHOR CONTRIBUTIONS**

SZ and ST designed the study. SZ led data analysis and interpretation and manuscript writing. ST contributed to data interpretation and manuscript writing.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/feart.
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Supplementary Material | The Supplementary Material for this study includes geochemistry data produced from the mineral weathering experiment.

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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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APPENDIX

TABLE A1 | The chemical composition of MilliQ de-ionized water in the control bottles at 168 h.

| Site | Conductivity (µS cm⁻¹) | DIC (µM) | CO₂ (µM) | SO₄²⁻ (µM) | Cl⁻ (µM) | Ca²⁺ (µM) | Mg²⁺ (µM) | Na⁺ (µM) | K⁺ (µM) | Fe (µM) | Al³⁺ (µM) |
|------|-----------------------|----------|----------|------------|---------|----------|----------|---------|---------|---------|----------|
| FM2  | 1.9 ± 0.6             | 23.4 ± 0.2 | 14.5 ± 1.3 | bd        | bd      | 0.8      | 0.4      | 13.1    | 0.3     | bd      | 0.1      |
| HC   | 1.5 ± 0.4             | 26.8 ± 2.2 | 26.0     | bd        | 1.1     | bd       | bd       | bd      | bd      | bd      | bd       |
| HD   | 1.6 ± 0.6             | 25.1 ± 2.4 | 24.9 ± 1.0 | bd        | bd      | bd       | bd       | bd      | bd      | bd      | bd       |

Conductivity, dissolved inorganic carbon (DIC), and CO₂ are reported as mean of replicates ± standard deviation. bd, below detection.

FIGURE A1 | Geochemical clusters from the pvclust analysis. Rectangles denote clusters where p < 0.05. au, approximately unbiased p-value; bp, bootstrapped probability.