Anonymous Reviewer #1

Zolkos et al., present a high quality characterisation of running water chemistry in a sub-catchment of the Peel River where the aim was to determine the effect of retrogressive thaw slump (RTS) on DIC sources and export. The study design, incorporating three transects at different spatial scales (1. retrogressive thaw slump (RTS) runoff water, 2. an intermediate size catchment with direct fed with RTS runoff, 3. A large catchment fed indirectly with RTS runoff through its tributaries), is an interesting and innovative sampling approach. The dataset, including a large number of key variables, is also of very high quality. The research question is also highly relevant to our understanding of the permafrost climate-feedback. The author rightfully stated that changes in carbonate alkalinity export in response to permafrost degradation has been far less studied that those for organic carbon and carbon dioxide. The study is also taking place in a relatively understudied region, which makes it even more valuable. While the subject and design of the study is of high quality, I find that the discussion and conceptualisation of the result need significant improvement. In short, the paper does not make full use of its potential.

The novelty of the paper lies in the approach of scales on the effect of RTSs on DIC export. At the moment, these three scales are taken separately, into three almost individual studies. Is the message simply that we can perceive the RTS effect at each of those three scales or is there a greater interpretation of how these effects integrate with increasing spatial scale and decreasing land-water connectivity? The paper would have more impact if the author could conceptualise these results and formulate how RTSs affect DIC cycling across scales rather than simply testifying that it has an influence. For example, does the “RTS effect” amplifies, is conserved, accumulates or becomes diluted with increasing scale. The author already documented that RTSs alter riverine DIC cycling in a previous publication in GRL (Zolkos et al, 2018). At present, this manuscript adds little to this state of knowledge, but this could be remediated by conceptualising further the effect of scales.

I have made a few suggestions to improve the presentation of results and conceptualisation of the discussion.

Thank you for the thoughtful and constructive feedback. Regarding your questions and comments above: Good question about the nature of RTS effects across scales (e.g. is it amplified, conserved, diluted, etc.). Effects on DIC are clearly amplified in headwaters. While CO₂ diminishes rapidly within headwaters, the HCO₃⁻ signal is preserved at broader scales. We use circumstantial evidence to infer the latter point in Zolkos et al. (2018), which focuses on biogeochemical effects immediately downstream of RTSs and leverages long-term Peel River data to assess changes over time. Our current study investigates processes occurring between headwater streams and the larger Peel River. We therefore contend that our current manuscript builds on our 2018 GRL paper and improves understanding of RTS effects on C cycling. We believe our edits help to clarify these points, in part by addressing your helpful suggestion to better conceptualize the effects of scale. Below, please find replies to your comments.
The sampling design is interesting and valuable, but arguably difficult to communicate to the reader. Having the Results and Discussion section together makes it even harder for the reader to put together the key results, and follow the discussion points that are mixed through the text. I recommend to separate these two parts. The first part of the results and discussion section details the changes in water chemistry in each of the three transect. The first part of the Result and Discussion section (section 3.1, 3.2 ad 3.3) details the water chemistry patterns in each transect with discussion points mixed through the text. Having this structure increases the impression that this study actually involves three separate studies rather than one. I suggest to also structure the results by water chemistry variable rather than sites. The study measured a large number of important and interesting water chemistry variables. Each should be presented clearly in the result section for the reader to identify. Each sections should provide, among other things, the overall range in values for the whole study, compare this range between each of the three scales and within each transect.

Great points. Thank you for the helpful suggestion. We restructured the manuscript so that Results and Discussion are presented separately. In the revised Results section we discuss results by hydrochemical parameter and present ranges of values, as you suggest.

Rather than naming the sites by their official river name, why not call them with a more conceptual name that represents the idea behind the sampling design. I like that the symbols in figures have numbers to indicate their position along the transect, but RTS FM2 site DC and SC has little meaning for the reader. I have also provided a few suggestions below to format the figures in a more visually telling way.

We appreciate the suggestion, but will use the official names for continuity with the literature. Our edits were made with this in mind, to improve demarcation between the different watershed scales.

Playing with your dataset I attempted to trace the d13CO2 source with the miller-trans plots, I found a clear difference in that value for the RTS runoff site (d13Csource -12‰ and the DC and SC rivers (-22.9‰. This suggest two predominant CO2 sources in this catchment and those end-members could potentially be used for calculations. The RTS site is consistent with a geogenic CO2 source, while the rivers have a predominant biogenic soil CO2 source. Would it be worth including this kind of approach to your results?

Thanks for the suggestion. We explored Miller-Tans plots, but do not have enough samples from undisturbed headwaters (n = 3) to trace CO2 sources with this approach. We agree that δ³¹C-CO₂ values reflect at least two predominant CO₂ sources in the catchment. Together, our δ³¹C-CO₂ values and δ³¹C-DIC vs. pH plot reflect spatial trends in DIC and CO₂ sources (atmospheric, soil biotic, mineral weathering). Building on this plot and following your suggested changes to the conceptual diagram, we clarify the relative importance of biotic versus mineral weathering sources for CO₂ and DIC, and how sources may change with movement downstream.
The effect of scales, with a varying degree of terrestrial connectivity, is only discussed in the context of circumpolar region with, but I believe that the study should also be put in the context of the broader literature, including lower latitude catchments, where many studies have also examined the effect of changing land-water connectivity with size.

*Yes, good point. We added brief text towards the end of the manuscript which considers this.*

I find it interesting to see the downstream changes in HCO3 concentration. It could be worth mentioning that studies modelling stream CO2 evasion based on d13CO2 value assume that carbonate alkalinity is conserve in river networks (Polsenaere 2012 Geochimica et Cosmochimica Acta and Venkiteswaran 2014 PLOS one).

*Good point. Though, this assumption will not hold true where changes in catchment lithology, groundwater inputs, etc. influence carbonate alkalinity. Nevertheless, it is interesting, so we have added brief text considering this.*

The term thermokarst and retrogressive thaw slumps are used interchangeably, I suggest to stick to retrogressive thaw slumps since this was the focus of the study and the findings may not be applicable to all kinds of thermokarst disturbances.

*Thank you for the suggestion. We have replaced thermokarst with retrogressive thaw slump where we discuss trends specific to the Peel Plateau. We retain “thermokarst” when we are discussing these effects from a broader perspective.*

The term “abiotic-inorganic processes” is a vague term to me, what are the processes included in that? Carbonate equilibrium reaction and CO2 evasion? Could you be more specific? Either a more specific term should be used or the choice of terminology should be justified in the text.

*Clarified.*

Abstract: Line 21:26: The results/discussion section of the abstract list changes in water chemistry in each of the three transect. I believe this section, and other relevant section in the main text, should come up with a more unified message of the RTS effect across scales, rather than at individual scales.

*Thank you for the suggestion. Please see our reply to your major comment, above.*

Line 30: I have trouble with the word “prevalent” here since it implies that one is larger than the other, while such calculations have not been done in the study.
Revised.

Introduction: Lines 32-33: This first sentence would introduce better the second paragraph where the source/sink relationship of DIC is detailed. This first paragraph discusses how the arctic landscape is changing and what is known of its influence on DIC export.

Thanks for the helpful suggestion. Revised.

Lines: 41-44: Since we include CO2 in this pool, the increase in soil respiration and/or increase in aquatic DOC degradation should also be part of this list. The citations be separated to indicate which if the listed mechanisms have been highlighted by each study.

Revised.

Lines 47-64: This is mostly textbook material and could be synthesised. I think what you want to express is why it’s important not only to account for the mass of DIC exported but also to identify its sources DIC. Without this we can’t determine whether this is a new sink or source of C in the short and long term perspective. I would suggest to move some of this information in section 2.5 of the methods and rework this paragraph to emphasise the simply the importance of source separation for DIC.

Fair point. We have modified the text but choose to discuss much of the original concepts, because our manuscript intends to target permafrost biogeochemists, who typically consider processes over contemporary timescales and processes associated with organic matter decomposition. So, this “textbook” information is provided – in part – to emphasize to this community the importance of incorporating weathering processes into consideration of the short- and long-term effects of permafrost thaw on biogeochemical (especially carbon) cycling.

Line 65: Could you state the representativeness of this study, which areas across the circumpolar north could be similar to your studied location?

Revised.

Line 94: This is a very important point which I think should be stated earlier in the introduction and given more emphasis.

Text added earlier in the Introduction to elaborate on this point.

Methods Line 133: How deep are those rivers? Can stream order be provided somewhere too.
We were not equipped with instruments to safely measure the depth of the rivers and unfortunately this information is not known. We have added stream order to Appendix Table A1.

Section 2.3 This is a nice way to work around flow measurement scarcity in this region. But should this section come after section 2.6?

Yes, this is more closely related to statistics and logically could be placed just before Sec. 2.7. Revised.

Line 212: What threshold was used in the flow accumulation to delineate the stream lines and catchment boundary? Was it validated with the areal photos mentioned on line 155 or something else?

We have clarified these methods in our ‘Geospatial Analyses’ section. Briefly, geospatial data were validated by modifying as needed to align with stream networks visible in moderate-resolution (10 m) Sentinel-2 satellite imagery, following St. Pierre et al. (2018).


Section 2.4, shouldn’t this section be called water chemistry analysis?

Changed to “Hydrochemical Analyses”.

Line 245: Why not model the full carbonate alkalinity pool (HCO3+ CO3)? Arguably the CO3 pool is small at this pH range and shouldn’t make a much difference to the model, but I find this conceptually more appropriate.

We agree that using \[HCO_3^- + CO_3^{2-}\] is conceptually more appropriate. We re-ran the model using carbonate alkalinity and yes, it is essentially identical to using only HCO3-. Nevertheless, we revised the results (Table 2) and text to reflect the updated model.

Results and Discussion

Section 3.1, 3.2 and 3.4 starts with a sentence stating how the author interprets DIC sources and cycling in each transect. This seems odd to me. I would rather the author takes me to that conclusion by presenting the results first.

Thank you for the suggestion. We believe that our revisions, which include splitting the Results and Discussion, address your suggestion.
The Miller-tans analysis supports that as well. Again, this statement comes before presenting the key results.

*Although we have not done a Miller-Tans analysis, for the reasons described above, our edits were done with an eye to elucidating the processes and sources which contribute to DIC in this system.*

Is the term “geogenic” more appropriate?

*Revised.*

Probably right, but do you have any measurements or estimates of CH4 concentration at the source - in the groundwater? The CH4 might have evaded already, but its imprint on d13CO2 values might still be there.

*Unfortunately, we do not have any measurements of [CH4] in groundwater.*

That is also supported with the Miller-tans. But this biogenic soil CO2 source seems to prevail in other sites as well albeit with some mixing with the geogenic source. Could mixing between the H2SO4 weathering and biogenic soil CO2 be back calculated?

*n = 1 in the Dempster Creek headwaters, so it is supported by the Miller-Tans analysis when data from all sampling points are considered. Also, [CO2] and stable isotopes alone would provide only a very rough estimate of the proportions of biogenic CO2 and abiotic CO2, the latter of which may also be sourced from H2CO3 carbonate weathering ~ DIC speciation reactions, and atmospheric CO2. As we lack the measurements (e.g. 14C of CO2) to make more robust estimates of CO2 contributions from these varied sources, we do not attempt this.*

This is also demonstrated in headwater streams at other latitudes and should be mentioned as well.

*Yes, good point. Elaborated and citations added, such as:*


Line 314: Again I find this term “abiotic inorganic” to be vague. And what do you mean by dominates? Abiotic dominates biotic, or inorganic dominates organic?

*Clarified.*

Line 351: What does “amplified inorganic carbon cycling” means?”

*Clarified.*

Line 371: The term biotic is used here, but could the DOC be degraded photochemically as well?

> Although photochemical degradation of DOC could lower $\delta^{13}$C-CO$_2$ values (references below), these streams are very turbid and therefore photodegradation is likely limited.


Line 380: Is the model intended only to bring evidence to the fact that RTS increase alkalinity export or for a possibly larger modelling/budget exercises? Could you make use of that model already in this paper for a final “wrap up” exercise?

*The model was intended to assess the influence of various landscape characteristics (including RTSs) on carbonate alkalinity yields (please see Section 2.7). We refrain from exercises on larger budgets of carbonate alkalinity, as our model reflects a snapshot of DIC dynamics within a relatively local area.*

Line 393: Dominate over what?

*Revised.*

Line 405: This “conceptual model of land-freshwater linkages” needs more elaboration. As it stands, this model seems more like a list of DIC sources and sinks across this catchment than a generalisable model. A starting point would be to determine how does it integrates with other models at lower latitudes? How do changes in DIC sources and sinks caused by RTS integrates with other water chemistry properties and C species (organic vs inorganic) that were documented in other studies?
Good point and, as noted in our reply to your comment on Figure 8 (below), we agree that the current conceptual diagram is fairly specific to the Peel Plateau. We believe the revised diagram helps to generalize our findings and, with revisions to the text, address your comments here.

Line 410: What exactly was “striking”?  
*Clarified.*

Line 414-418: How does this conclusion differ from the authors previous publication cited here?

*A key difference is the nested watershed/transect approach, which allows us to elucidate mechanisms and how they change downstream.*  
*Clarified.*

Line 432: Should your model be used to that effect? If so, it should be stated.  
*Revised.*

Line 443: “change for C cycle in the rapidly changing arctic landscape”  
*Revised.*

Authors Contribution What did D.O. do?  
*All authors contributed to manuscript writing.*  
*Clarified.*

Figures and Tables

Table 2: Should these values be presented in supplementary and only the model be presented in this table? I find the second part of the table easy to miss.

*We feel that the values in the upper panel are important to include in the main text and agree that the lower panel may be easy to miss. To address this, we revised the first sentence of the table caption to draw attention to both the upper and lower panels.*

Figure 1: The map feels quite dense, can the photos and context map be placed outside? Could the bedrock lithologies be illustrated on the map?
Thank you for the helpful suggestion. We revised the map and the individual components now have more space. Unfortunately, there are no high-resolution GIS data of bedrock lithology available.

Figure 2: The figures should be placed vertically rather than horizontally since they have the same x-axis. Also in caption, please clearly state that the top x-axis is for the dempster creek transect while the second is for the stony creek. Could the points for each transect be connected with a line for visualisation. Could HCO3 and CO2 concentration be on the same unit? Could there be a third 3 axis for the distance along the RTS runoff transect?

Good suggestion to consolidate x-axes by stacking the figures. Revised. Following your other helpful suggestion, we have also included DOC and SUVA254 in Fig. 2.

Caption revised to distinguish between the RTS FM2 runoff, Dempster Creek, and Stony Creek x-axes.

We choose to omit lines connecting the points and, as noted in the caption, instead label consecutive downstream points with their corresponding site numbers; we reserve lines to indicate significant trends.

We like and appreciate your suggestion to report CO2 in µM, which could help to facilitate a more direct comparison with the HCO3− concentrations shown in Fig. 1a. However, for Fig. 1b, we choose to report CO2 in units of partial pressure rather than µM, to make a more intuitive link with atmospheric CO2. For the readers’ reference, CO2 in µM is reported in Table 1.

We have added a third axes for RTS FM2 runoff and plotted the data (as in Table 1).

Figure 4: Can you give a reference to these end-members

Thank you for the suggestion. In the caption, we added a reference to Methods Sec. 2.4, in which we briefly elaborate on these methods and cite a publication used to derive the end-members (Zhang et al. 1995).


Figure 5: Open vs closed symbols would be clearer perhaps?

Thank you for the suggestion. This does appear to help. Figure revised.

Figure 6 and 7: Should these two figures be merged with Figure 2? This would help draw a more complete picture of simultaneous changes in water chemistry along the transects. Why isn’t there a similar figure for d13C-DIC values?
Good idea. We have included the DOC and SUVA254 plots as part of Fig. 2. We choose not to replicate this plot style for $\delta^3$C-DIC, because the downstream trends in $\delta^3$C-DIC are shown in Figures 4 and 5.

Figure 8: This is a nice schematic, but it limits the scope of the study. The schematic mostly lists the sources and transformation of DIC in this catchment. Does it only applies to this catchment, i.e. was the goal to map those processes, or can it be generalised to other catchments? I think this figure could be useful if it was to conceptualise the effect of RTS across scales, not just make a summary of all the processes identified in the data for this specific catchment. I have in mind something along the lines of Hotchkiss et al. 2015 NatGeo Figure 3.

Thank you, we agree that the figure is somewhat limiting and we appreciate your suggestion to broaden the scope of our conceptual diagram. We believe the revised conceptual diagram helps to generalize our findings.

Table A1. Why not keep the distance units the same and just add decimals for FM2 site.

Yes, good point! Revised.

Table A2: I find this could be useful in the main manuscript since it also provides a list of the DIC sources you are trying to separate.

Good suggestion. Added to main text.
Zolkos et al. present a detailed and high quality characterisation of running water chemistry in a sub-catchment of the Peel River. This work was to determine the effect of retrogressive thaw slump (RTS) on DIC sources and export. The research design, incorporating three transects at different spatial scales, is an interesting sampling approach. The dataset, including a large number of key variables, is also of very high quality. The research question is highly relevant to our understanding of the permafrost climate-feedback.

Thank you very much for the thoughtful and helpful comments.

While the design of the study is of high quality, I find that the discussion of the results needs some improvement.

I think the influence of thermokarst on fluvial inorganic carbon cycling and export is reflected in two aspects. One is the change in runoff, and the other is the change in DIC concentrations and sources. The authors have discussed the latter more clearly, but the former needs to be done further. In addition, the authors used the change in concentration and isotope of DIC to indicate the sulfuric acid carbonate weathering, but the sulfuric isotopic evidence may be the more direct one. Could they add this to further strength their conclusions?

We agree that thermokarst influences inorganic carbon cycling via changes in DIC concentrations and sources (e.g. Zolkos et al. 2018). However, discharge within thaw slumps is relatively small compared to the streams affected by slumps, so changes to runoff associated with slumping are likely to be negligible, yet we lack direct evidence for such an assertion. Also, thank you for the suggestion to consider sulfur isotopes as evidence of $\text{H}_2\text{SO}_4$ carbonate weathering. The sulfur isotopes we measured and reported in our 2018 GRL paper (Zolkos et al. 2018) do align with sulfate derived from sulfide oxidation. We now briefly consider this in our revised Discussion.

Anonymous Reviewer #3

In this study the authors investigate how permafrost thaw affects mineral weathering sources of inorganic carbon (IC), and how the fluvial IC is cycled across different scales. Specific focus is on retrogressive thaw slumps (RTS) and their major contribution to IC yields and biogeochemical processes across fluvial networks draining permafrost regions. The study is based on one synoptic summer sampling campaign of three different fluvial transects covering different scales, and where samples were taken for a comprehensive set of chemical and isotopic variables. The authors conclude that rapid weathering in the RTS runoff enhance both atmospheric CO2 emission and downstream DIC transport. They further show that the IC signal from RTS have a major downstream impact across large scales although the RTS impacted area covered less than a 1% of the total catchment area.

The manuscript focus on an important topic that is very suitable for publication in Biogeosciences. The current thaw of permafrost regions is of major concern and the response in the landscape C cycling is a central issue. Much of the literature is focusing on the mobilization of organic C stocks and the subsequent mineralization into CO2 and CH4. In comparison, relatively little focus is given to the inorganic C mobilization and to what degree mineral weathering upon permafrost act as a source or sink for atmospheric C, and how it affects biogeochemical processes in aquatic systems.

Thank you for the encouraging comments. We appreciate it!

General comments:

With this background the manuscript is an important contribution to the research field. The authors present a comprehensive and neat data set from a data scarce region, and where they disentangle different sources and processes affecting the fluvial IC in a (mostly) very convincing way. The manuscript is very well written but I have some points that need to be clarified prior to a publication. These issues are mostly to strengthen the argumentation by the authors but also to fully capitalize on their findings.

Many thanks for your helpful feedback. Please find our replies below.

Detailed comments:

Ln 15-18, a very long sentence with plenty of information. I suggest to split it.

Revised.

Ln 153-160, it is hard to grasp the uncertainty of the stream flow section. i.e. how certain the Q estimates are. On the other hand, the water or solute yields are a relatively minor part of the ms.
This is a fair point. We added a figure to the appendix which shows the relationship between our discharge measurements and the estimates from our model. As the figure shows, the 95% confidence interval is relatively larger at higher discharge levels. The strong, linear relationship provides some confidence in our estimates of discharge.

![Graph showing relationship between observed and predicted discharge](image)

Figure A2. Estimated vs. measured discharge ($Q$) ($p < 0.001, R^2 = 0.89, F_{1,18} = 150$) for 20 streams in the Stony Creek watershed. Grey band represents the 95% confidence interval around the regression. Estimates were made using measurements of stream width, $Q$, and a hydraulic geometry model (Gordon et al. 2004) (see Sec. 2.6). The model (Eq. 1) was used to estimate $Q$ in the Stony Creek tributaries.


Ln 237-239, how come these three variables were used in the MLR? Comes currently a bit out of the blue and needs to be better motivated.

Hydrology, terrain roughness, and vegetation productivity were included as covariates because they are known to be among the primary landscape controls on DIC cycling. We have clarified this and included citations.

Ln 239-245, again it is hard to judge the certainty in this modelling effort given the already above raised concern about the $Q$ estimation.

Please see our reply to your comment for Ln 153-160. We acknowledge there is some uncertainty. Yet, this approach enables us to generate a first estimate of the relevance of RTSs in
carbonate alkalinity production and export relative to other landscape conditions known to influence DIC in fluvial networks.

Ln 259-, I guess very much a question of personal taste but I feel the ms do not benefit from the mixing of results and discussion. It would be easier to keep focus by separating them in my opinion.

*We agree with you and Reviewer #1 about this. We restructured the manuscript so that the Results and Discussion are presented separately.*

Ln 278, I am not familiar with the given reference, but what is meant by “regional carbonate”? Also in this couple of sentences, I agree with the overall argumentation, but can you completely rule out a biotic source contribution? The fractionation between carbonate and CO2 (8‰ is rather theoretical. Could a mixing with geogenic and bio- genic IC be possible for generating 13C-CO2 of -11.4 to 12.1‰ You have a substantial DOC pool which is also cited by being “relatively biolabile”.

*This was meant to read “... regional carbonate bedrock”. Revised. Yes, good point about CO2 being a mix of biogenic and geogenic sources. We have added brief text clarifying that these isotopic values may also reflect some contribution of 13C-depleted CO2 from biogenic sources.*

Ln 285, how CH4 was sampled is mentioned in the methods but from what I see this is the only place where any data is presented, and then very shortly. Maybe the data is saved for another story but I believe it would further strengthen the story if it could be included for example in table 1 and with subsequent incorporation in the text.

*We tell the CH4 story in an earlier publication (Zolkos et al. 2019, e.g. Sections 3.1, 3.2, 3.3, 4.1, 4.3, 4.4). CH4 measurements in this study were done to assess for potential effects from methanogenesis on stable CO2 isotopes.*


Ln 310-313, yes it could be due to adsorption to RTS sediments, but I guess it could also be due to lower mineralization than degassing rates. Might be worth to mention.

*Yes, good point. We have clarified this in the text.*

Ln 347-349, is it really clear that biotic CO2 were the primary source of DIC in the headwaters of Stony Creek? Could not geogenic sources still be highly influential? The 13C-DIC and 13C-CO2 values (-11.6 and -13.8‰ respectively) points towards a biogenic/geogenic mixing, or?
Fair question. As noted, mixing between the stream and atmosphere was a primary CO₂ source. This is supported by the values from other measurements, such as pCO₂ at an approximately atmospheric level and also low HCO₃⁻ and pH. The former suggests relatively minor biotic CO₂ production (organic matter mineralization) and/or greater effects from degassing on CO₂ than from biotic processes. The latter suggests stronger effects on DIC speciation from variability in pH (CO₂ > HCO₃⁻) than from mineral weathering. Together, these results suggest that geogenic mixing is a less parsimonious explanation.

Ln 403-405, do the study really evaluate “across gradients of thermokarst disturbance”? I believe something like influence of RTS on IC cycling and how this signal is propagated across different fluvial scales is better describing the story.

Good point. Revised.

Ln 419-434, I somehow miss the full interpretation of the findings of the current study for the large scale picture. How do you suggest your results should be considered in large scale estimates, i.e. how does it affect the previous judgement of the area as a “modest source of CO₂”.

Fair point. We consider the broader relevance of our findings in our revised Discussion and we provide an updated conceptual model to help generalize our findings.

A general question: how common are RTS across permafrost regions worldwide? How applicable are the findings here for other areas?

Good question. We consider this in the Introduction and we believe that our edits help to clarify this.

Figure 1. For a non-north American reader, a more large-scale inset of where the area is found would be appreciated.

Good suggestion, thanks. Figure revised.
Thermokarst amplifies fluvial inorganic carbon cycling and export across watershed scales on the Peel Plateau, Canada

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Abstract. As climate warming and precipitation increase at high latitudes, permafrost terrains across the circumpolar north are poised for intensified geomorphic activity and sediment mobilization that are expected to persist for millennia. In previously glaciated permafrost terrain, ice-rich deposits are associated with large stores of reactive mineral substrate. Over geological timescales, chemical weathering moderates atmospheric CO2 levels, raising the prospect that mass wasting driven by terrain consolidation following thaw (thermokarst) may enhance weathering of permafrost sediments and thus climate feedbacks. These feedbacks depend upon the mineral composition of sediments (weathering sources) and the balance between atmospheric exchange of CO2 versus fluvial export of carbonate alkalinity (2[CO3\(^2-\) + HCO\(^3-\)]). Working in the fluvially-incised, ice-rich glacial deposits of the Peel Plateau, northwestern Canada, we determine the effects of retrogressive thaw slump (RTS) hillslope thermokarst activity on mineral weathering sources, CO2 dynamics, and carbonate alkalinity export, and how these effects integrate across watershed scales (~2 to 1000 km\(^2\)). We worked along three transects in nested watersheds with varying connectivity to RTS activity: a 550 m transect along a first-order thaw stream within a large RTS; a 14 km transect along a stream which directly received inputs from several RTSs; and a 70 km transect along a larger stream with headwaters that lay outside of RTS influence. In undisturbed headwaters, stream chemistry reflected CO2 from soil respiration processes and atmospheric exchange. Within the RTS, rapid sulfuric acid carbonate weathering, prompted by the exposure of sulfide- and carbonate-bearing tills, appeared to increase fluvial CO2 efflux to the atmosphere and propagate carbonate alkalinity across watershed scales. Despite covering less than 1% of the landscape, RTS activity drove carbonate alkalinity to increase by two orders of magnitude along the largest transect. Amplified export of carbonate alkalinity together with isotopic signals of shifting DIC and CO2 sources along the downstream transects highlight the dynamic nature of carbon cycling that may typify glaciated permafrost watersheds subject to intensification of hillslope thermokarst. The balance between CO2 drawdown in regions where carbonic acid weathering predominates and CO2 release in regions where sulfides are more prevalent will determine the biogeochemical legacy of thermokarst and enhanced weathering in northern permafrost terrains. Effects of RTSs on carbon cycling can be expected to persist for millennia and should spur their integration into predictions of weathering-carbon-climate feedbacks among thermokarst terrains.
Riverine export of carbonate alkalinity ($\Sigma$[HCO$_3^-$, CO$_3^{2-}$]), generated by the chemical weathering of silicate and carbonate minerals, is a key component of the global carbon cycle and Earth’s long-term climate (Berner, 1999; Gaillardet et al., 1999; Hilton and West, 2020; Torres et al., 2017). The degree to which carbonate alkalinity production involves CO$_2$ (as carbonic acid, H$_2$CO$_3$ = H$_2$O + CO$_2$(aq)), from atmospheric or soil-respiration sources, and liberates mineral carbon influences whether dissolved inorganic carbon (DIC = $\Sigma$(CO$_2$, carbonate alkalinity)) in fluvial networks represents a carbon sink or source. Rapid warming at northern latitudes (Serrze and Barry, 2011) is thawing permafrost (Biskaborn et al., 2019), increasing vegetation productivity ( Björkman et al., 2018), intensifying hydrologic cycles (Rawlins et al., 2010), and strengthening land-freshwater linkages (Vonk et al., 2019; Walvoord and Kurylyk, 2016). These processes are activating large amounts of mineral substrate into biogeochemical cycles, with significant implications for DIC cycling ( Lacelle et al., 2019; Wadham et al., 2019). In recent decades, increasing riverine fluxes of carbonate alkalinity and solutes across the circumpolar north reflect enhanced mineral weathering associated with active layer thickening, deepening hydrologic flowpaths into mineral soils, and greater soil acidity from increasing vegetation productivity (Drake et al., 2018a; Tank et al., 2016; Toohy et al., 2016). Glaciated permafrost terrains hosting ice-rich deposits of reactive sediments are thought to be distributed across the northern permafrost zone, raising the prospect that terrain consolidation following thaw (thermokarst) and associated carbonate alkalinity production and export may have stronger influence on climate feedbacks in such regions (Zolkos et al., 2018).

Three coupled factors primarily influence the degree to which carbonate alkalinity represents a carbon sink or source. First, the weathering source, which accounts for both the mineral composition of substrate subjected to chemical weathering and the acid(s) responsible for weathering. Silicate weathering by H$_2$CO$_3$ generates alkalinity without liberating mineral carbon and thus represents a long-term CO$_2$ sink. In contrast, H$_2$CO$_3$ carbonate weathering is a CO$_2$ sink only over $\sim$10$^{-6}$ to 10$^{-5}$ y, as half of the alkalinity produced is geogenic. HCO$_3^-$ produced during carbonate weathering in the presence of strong acids, for instance sulfuric acid (H$_2$SO$_4$) from sulfide oxidation, is a CO$_2$ source over longer timescales ($\sim$10$^{-5}$ y; Calmels et al., 2007) and can also produce CO$_2$ over shorter timescales when H$_2$SO$_4$ is present in excess (Stumm and Morgan, 1996). Second, the rate of mineral weathering and processes that further alter this rate. Rates of chemical weathering are orders of magnitude faster for carbonates and sulfides than for silicates (Stumm and Morgan, 1996). Further, weathering rates generally increase with mineral surface area, and therefore are often fast in glacial environments owing to intense physical weathering (Anderson, 2007). Indeed, hydrochemical signatures of tectonic carbonate and sulfide lithologies can dominate weathering fluxes in primarily silicate glacial environments (Anderson, 2007). The disparity is so significant that, when sediment supplies are sufficient, H$_2$CO$_3$ carbonate weathering in proglacial streams can consume dissolved CO$_2$ to below atmospheric levels (Sharp et al., 1995; St. Pierre et al., 2019). Third, the magnitude of carbonate alkalinity export, which is influenced by its production via weathering of minerals during fluvial transport (e.g. Striegl et al., 2007) and its loss via carbonate equilibrium reactions and CO$_2$ degassing along the land-freshwater-ocean continuum. From a climate perspective, the magnitude of carbonate alkalinity export is particularly relevant...
over geological timescales, because half of riverine carbonate alkalinity exported to the ocean is returned to the atmosphere as CO$_2$ via precipitation reactions within the marine carbon cycle (Calmels et al., 2007). Together, these three controls on carbonate alkalinity highlight the non-conservative nature of DIC and its susceptibility to transformation within fluvial networks. Hence, to constrain carbonate alkalinity export in rapidly changing permafrost terrains, nested-watershed sampling designs are critical for capturing DIC transformation along the land-freshwater-ocean continuum and resolving drivers and sources of carbon cycling across scales (Drake et al., 2018b).

Glaciated permafrost terrains are poised for rapid geomorphic and associated biogeochemical change as the climate warms and precipitation intensifies (Kokelj et al., 2017b). Despite glacial retreat across much of the circumpolar north, permafrost within these landscapes preserves biogeochemical legacies of glaciation across northern Canada, Alaska, and western Siberia (Kokelj et al., 2017b). In North America, the comminution of carbonate and shale bedrock during expansion of the Laurentide Ice Sheet (LIS) and the climate and vegetative protection of ice- and sediment-rich tills in the wake of its retreat endowed former glacial margins across northwestern Canada with thick inorganic tills held in ice-rich permafrost (Kokelj et al., 2017b). Today, the climate-driven renewal of post-glacial landscape change is mobilizing immense stores of minerals into modern biogeochemical cycles via hillslope thermokarst features, the largest of which include retrogressive thaw slumps (RTSs) (Kokelj et al., 2017a). On the Peel Plateau (NWT, Canada), RTSs expose carbonate- and sulfide-bearing glacialic permafrost sediments that are tens of meters thick. The chemical weathering and fluvial transport of these sediments results in increased HCO$_3^-$ immediately downstream of RTSs and greater solute and sediment loads throughout downstream systems (Kokelj et al., 2013; Malone et al., 2013; Zolkos et al., 2018). RTS activity has been suggested, but not previously proved, to be partly responsible for increasing carbonate alkalinity fluxes in the larger Peel River during recent decades (Zolkos et al., 2018). Yet, it remains unknown how hillslope thermokarst effects on mineral weathering and DIC sources and cycling integrate across watershed scales on the Peel Plateau and in relatively inorganic-rich permafrost terrains elsewhere. In this study we evaluated trends in major ions, DIC concentration, and dual 8$^\text{13}$C-DIC–8$^\text{18}$O$\text{H}_2$O isotopes along transects within three nested watersheds on the Stony Creek watershed on the Peel Plateau. Our nested watershed approach was intended to allow us to determine how RTS effects on carbon cycling integrate across scales from ~1 to 1000 km$^2$.

2 Methods

2.1 Study Area

The Stony Creek watershed is located southwest of the hamlet of Fort McPherson, in the northern, or lower Peel River watershed (Fig. 1). Stony Creek, a tributary of the Peel River, originates in the Richardson Mountains, where slopes are sparsely vegetated and mainly consist of bedrock colluvium (Duk-Rodkin and Hughes, 1992). Exposed marine shale and sandstone bedrock contain sulfide- and gypsum-bearing lithologies, but limited carbonate (Norris, 1985). As Stony Creek flows eastward, the main channel and its tributaries incise ice-rich glacial deposits and underlying Cretaceous bedrock, forming a stream network comprised of tundra flow tracks grading to incised gravel bed streams with increasing watershed size. The fluvially-incised valleys and increasing regional precipitation have

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proven conducive to thaw-driven mass wasting of ice-rich glacial deposits and formation of RTSs (Kokelj et al., 2017b). Growth of RTSs is driven by the ablation of exposed ground ice and is perpetuated by the downslope mass wasting of thawed material via fluidized earth flows, which can accumulate large volumes of debris in stream valleys (Fig. 1). Across the Dempster Creek watershed, intensifying RTS activity releases large volumes of sediment and solutes into streams relative to undisturbed headwaters (Kokelj et al., 2017b; Segal et al., 2016). This substrate is transported to streams via rill runoff channels in the scar zone and debris tongue deposits in the stream valley.

Impacts to Stony Creek are representative of numerous other major Peel River tributaries that have incised the ice-rich Peel Plateau (Kokelj et al., 2015). The ~60 km² watershed of Dempster Creek, a tributary of Stony Creek, originates in willow and open spruce taiga without RTS activity, receiving large inputs of sediments and solutes from RTSs FM2 and FM3 within several kilometers of the headwaters (Kokelj et al., 2013; Malone et al., 2013). Many small, non-RTS affected streams and several larger RTS-affected tributaries flow into Dempster Creek before its confluence with Stony Creek.

2.2 Stream Sampling

In late July 2017, we sampled along transects within three nested watersheds (Fig. 1, Table A1) to understand how the effects of RTSs integrate across watershed scales. (i) The RTS FM2 runoff transect included five sampling locations along a 550 m-long flow stream formed by a runoff channel within an active RTS. The RTS FM2 runoff received no observable hydrologic inputs during the sampling period. (ii) A 14 km transect along the mainstem of Dempster Creek, which received inputs directly from RTS FM2, was sampled at one location in undisturbed headwaters at three sites downstream of RTS FM2. Sites downstream were located on the mainstem, immediately upstream of three major tributaries. We also sampled from the tributaries near their confluence with Dempster Creek, to characterize tributary chemistry. (iii) A 70 km transect along the mainstem of Stony Creek, a 6th-order stream, was sampled at eight locations: one in undisturbed headwaters and seven on the RTS-affected reach upstream of major tributaries. We additionally sampled from one tributary of the undisturbed headwaters and from six RTS-affected tributaries near their confluence with the mainstem, Stony Creek, a major tributary of the 70000 km² Peel River watershed (Fig. 1), to determine how the effects of RTS activity on DIC integrate across broader scales.

At all sampling sites, stream temperature, specific conductance (henceforth, "conductivity"), and pH were measured using a pre-calibrated YSI Professional-Plus water quality meter. At most sites, water samples were collected for ions, DIC, CO₂, CH₄, dissolved organic carbon (DOC), UV-visible absorbance, and total suspended solids (TSS). Along the RTS FM2 runoff transect, we sampled only for DIC and CO₂ concentration, and stable isotopes of dissolved CO₂ (δ¹³C-CO₂). One day prior, additional parameters were sampled at RTS FM2 runoff site five, located near the confluence of the RTSs with Dempster Creek, for comparison with the full suite of chemistry parameters collected along the Dempster Creek transect. At the Dempster and Stony Creek sites, we additionally sampled water for stable isotopes of DIC (δ¹³C-DIC) and used dual δ¹³C-DIC, δ¹⁸O-CO₂ isotopes to characterize abiotic and biotic processes influencing DIC sources and cycling across watershed scales.
Cations and trace elements were calibrated using certified reference material (CRM). Samples for DIC were filtered using serum bottles (Sartorius) membranes. Samples for anions were filtered using 0.45 μm polyethersulfone (PES, ThermoFisher) or cellulose-acetate (CA, Sartorius) membranes. Samples for DIC were collected without headspace in airtight syringes. Samples for ions, DOC, and UV-visible absorbance were collected in acid washed (24 h, 10% v/v HCl) all-plastic syringes. Syringes were triple sample-rinsed, sealed without headspace, and stored cool and dark until processing within 10 h. For DIC was filtered (PES) into precombusted (5 h, 500°C) glass vials without headspace and sealed with a butyl rubber septum for DIC concentration or two silicone-teflon septa for δ13C-DIC. Samples for cations were filtered (CA) into acid-washed bottles and acidified with trace metal-grade HNO3, while anions were filtered (CA) but not acidified. Samples for DOC were filtered (PES) into precombusted glass vials and acidified to pH < 2 using trace metal-grade HCl (Vonk et al., 2015). Samples for UV-visible absorbance were filtered (PES) into non-acid washed 30 mL HDPE bottles. Samples were refrigerated (4°C, dark) until analysis.

Dissolved gases were collected following the headspace equilibration method (Hesslein et al., 1991) and stored in airtight syringes (for CO2 concentration) or over-pressure in pre-evacuated serum bottles sealed with pre-baked (60°C, 12 h), gas-inert butyl rubber stoppers (for δ13C-CO2-Ch4). At each site, atmospheric samples for CO2 and CH4 concentration and δ13C-CO2 were stored in the same fashion. Gas samples were stored in the dark at ~20°C prior to analysis within 10 h (CO2) or two months (δ13C-CO2, CH4). Water and air temperature, atmospheric pressure, and the volumetric ratio of sample to atmospheric headspace was recorded for correcting later calculations of CO2 partial pressure (pCO2) and δ13C-CO2 (Hamilton and Ostrom, 2007).

2.3 Hydrochemical Analyses

Upon returning from the field each day, CO2 was measured using an infrared gas analyzer (PP Systems EGM-4), which was checked monthly for drift using a commercial standard (Scotty Gases). We calculated pCO2 using Henry’s constants corrected for stream water temperature (Weiss, 1974) and accounting for the ratio of water volume to headspace during sample equilibration. DIC samples were measured by infrared CO2 detection (LiCOR 7000) following acidification within a DIC analyzer (Apollo SciTech model AS-C3). Calibration curves were made daily using certified reference material (CRM) from Scripps Institution of Oceanography. Samples with DIC concentrations < 400 μM were analyzed using solutions prepared from a 1000 ppm TIC stock (ACCUSPEC) that were calibrated with CRM. DIC species (CO2, HCO3−, CO32−) were calculated from DIC concentration and pCO2 or pH using CO2sys (v.2.3) (Pierrot et al., 2006), using field temperature and pressure at the time of sampling, and the freshwater equilibrium constants for K1 and K2 (Millero, 1979).

Cations and trace elements were measured by optical emission spectrometry (Thermo ICAP-6300) and anions by ion chromatography (Dionex DX-600) at the University of Alberta Biogeochemical Analytical Services Laboratory (BASL, ISO/EIC accreditation #17025). DOC was measured using a total organic carbon analyzer (Shimadzu TOC-V). DOC standard curves were made daily with a 1000 ppm KHP solution (ACCUSPEC) and an in-house caffeine...
standard (10 mg L\(^{-1}\)) was run every 20 samples to monitor instrument drift. Absorbance spectra were analyzed using an Ocean Optics UV-VIS instrument with a Flame spectrometer module, following Stubbins et al. (2017) and corrected for Fe interference (Poulin et al., 2014). To evaluate organic carbon reactivity, we used specific ultraviolet absorbance at 254 nm (SUVA\(_{254}\)) to infer DOC aromaticity (Weishaar et al., 2003).

\[ \delta^{13}C\text{-DIC} \]

was determined using an isotope ratio mass spectrometer (Finnigan Mat DeltaPlusXP) interfaced to a total organic carbon analyzer (OI Analytical Aurora Model 1030W) at the University of Ottawa Stable Isotope Laboratory. \(\delta^{13}C\text{-CO}_2\) and CH\(_4\) concentration were analyzed within two months using a Picarro isotope analyzer (G2201-i; < 0.2‰ precision, CH\(_4\) operational range = 1.8 - 1500 ppm) equipped with an injection module for discrete samples (SSIM). Commercial \(\delta^{13}C\text{-CO}_2\) and CH\(_4\) standards were used to check for drift during each run. We used mass balance to correct \(\delta^{13}C\text{-CO}_2\) values for the \(\delta^{13}C\) and mass of atmospheric CO\(_2\) used for equilibration (Hamilton and Ostrom, 2007). To assess \(\delta^{13}C\text{-CO}_2\) fractionation from calcite precipitation (Turner, 1982) and methanogenesis (Campeau et al., 2018) in RTS PM2 runoff, we calculated the saturation index (SI) and partial pressure of CH\(_4\) (pCH\(_4\)). SI was calculated using the hydrochemical software Aqion version 6.7.0 (http://www.aqion.de), which uses the U.S. Geological Survey software PHREEQC (Parkhurst and Appelo, 2013) as the internal numerical solver.

Samples for atmospheric and dissolved CH\(_4\) were collected in the same fashion as \(\delta^{13}C\text{-CO}_2\). pCH\(_4\) was calculated using Bunsen solubility coefficients (Wiesenburg and Guinasso, 1979) converted to the appropriate units (Sander, 2015).

TSS samples were filtered onto muffled (450°C, 4 h) and pre-weighed glass fiber filters (Whatman GF/F; 0.7 µm nominal pore size) upon returning from the field, stored frozen, and dried (60°C, 24 h) for gravimetric analysis following a modified version of U.S. Geological Survey Method I-3765.

### 2.4 Mineral Weathering and DIC Sources

We used a Piper diagram (Piper, 1944), which reflects the proportional equivalent concentrations of ions in a sample relative to mineral weathering end-members, as one method to constrain the sources of mineral weathering and HCO\(_3\)\(^{-}\). The products of Eq. 1 defined the mineral weathering end-members in the Piper diagram (Table 1). We further constrained mineral weathering and DIC sources using \(\delta^{13}C\text{-DIC}\) and pH. End-member \(\delta^{13}C\text{-DIC}\) ranges for equilibrium processes (mixing with atmospheric and/or biotic CO\(_2\)) and kinetic reactions (mineral weathering) were derived following Lehml et al. (2017) and using published isotopic fractionation factors (Zhang et al., 1995).

To evaluate potential effects on \(\delta^{13}C\text{-CO}_2\) from DIC speciation along the pH continuum (Eq. 8, Table 1) (Clark and Fritz, 1997), we compared theoretical and observed \(\delta^{13}C\text{-CO}_2\) values in the Stony Creek mainstem. Theoretical \(\delta^{13}C\text{-CO}_2\) values were calculated using mass balance to obtain \(\delta^{13}C\text{-HCO}_3\) from measurements of DIC, CO\(_2\), HCO\(_3\)\(^{-}\), \(\delta^{13}C\text{-DIC}\) and \(\delta^{13}C\text{-CO}_2\). We then used measurements of stream temperature (T) to calculate the equilibrium fractionation between CO\(_2\) and HCO\(_3\)\(^{-}\) (\(\varepsilon = -9.483 \times 10^{17} T + 23.899\); Mook et al., 1974). Finally, \(\varepsilon\) was subtracted from \(\delta^{13}C\text{-HCO}_3\) to obtain theoretical \(\delta^{13}C\text{-CO}_2\). Similarity between observed and theoretical \(\delta^{13}C\text{-CO}_2\) values was
interpreted as \( \delta^{13}C\text{-CO}_2 \) variability driven by carbonate equilibrium reactions, whereas dissimilarity was taken to reflect effects from CO\(_2\) degassing (Zhang et al., 1995) and/or biotic CO\(_2\) production (Kendall et al., 2014).

2.5 Geospatial Analyses

Stream networks and watershed areas were delineated using the ArcHydro tools in ArcGIS 10.5 from the gridded (30 m) Canadian Digital Elevation Model (CDEM). CDEM data were reconditioned using National Hydro Network stream vectors, which were first modified as needed to align with stream flow paths visible in Copernicus Sentinel-2 multispectral satellite imagery (2017; European Space Agency, https://sentinel.esa.int/). To statistically assess landscape controls on DIC cycling (Sect. 2.7), we delineated active RTSs and derived terrain roughness and vegetation productivity in the major tributary watersheds of Stony Creek. RTSs were interpreted as active where exposed sediment visibly dominated the feature surface (Cray and Pollard, 2015) in orthorectified SPOT multispectral imagery that we pan-sharpened to 1.6 m resolution using the ArcGIS Image Analysis tool. The satellite imagery was collected from September 9 to 25, 2016. Active RTSs that were connected to streams were manually delineated using ArcGIS. We used RivEx 10.25 software (Hornby, 2017) to quantify the number of active RTSs impacting streams in the Stony Creek watershed and to visualize RTS impact accumulation across the fluvial network. We defined RTS impact accumulation as the cumulative number of active RTSs impacting upstream reaches. RTSs were interpreted to impact streams based on contact or interpreted downslope flow based on slope direction and gradient from the CDEM (Supplementary Information). Where a single RTS affected multiple streams, only the upstream segment was used for the accumulation.

We used the Geomorphic and Gradients Metrics Toolbox (Evans et al., 2014) to calculate terrain roughness, which is a measure of variance across a land surface and represents topographic complexity (Riley et al., 1999). We use terrain roughness as a proxy for potential physical erosion, which is known to enhance sulfide oxidation by exposing shale regolith throughout the Peel River watershed (Calmels et al., 2007) and may therefore influence DIC. The enhanced vegetation index (EVI) was used to broadly evaluate vegetation productivity (Huete et al., 2002), which is known to influence DIC production by enhancing mineral weathering (Berner, 1999). We used the U.S. National Aeronautics and Space Administration EVI product (Didan, 2015), which is derived from gridded (250 m) moderate resolution imaging spectroradiometer (MODIS). The MODIS data were collected on July 28, 2017. The ArcGIS Zonal Statistics tool was used to calculate total RTS area, mean terrain roughness, and mean EVI in Stony Creek tributary watersheds.

2.6 Stream Flow

Water discharge \((Q)\) in Stony Creek tributaries was estimated from a hydraulic geometry model (Gordon et al., 2004) that we developed using flow measurements made in Peel Plateau streams during 2015–2017 and width \((W)\) estimated from on-site measurements or photos from 2017 with a known scale. The model reflected measurements spanning diverse stream morphologies \((W = 0.4–6.6\ m)\) and flow conditions \((Q = 0.005–0.91\ m^2\ s^{-1})\) (Fig. A1):
3 Results

3.1 pH, Ions, and Weathering Sources Across Watershed Scales

\[ Q = e^{3.061 - 0.258 / 0.661} \times 0.001, R^2 = 0.89, F_{1,18} = 150 \] (1)

Discharge values from 2015 to 2017 were calculated from measurements of stream flow (RedBack Model RB1, PVD100) and cross-sectional area made at increments equal to 10% of stream width (Gordon et al., 2004; Lurry and Kolbe, 2000), and were averaged for sites with multiple measurements.

2.7 Statistics

We tested for downstream change in HCO\textsuperscript{-3} concentration and pCO\textsubscript{2} along the Stony Creek mainstem using the non-parametric Mann-Kendall test from the R software (R Core Team, 2018) package zyp (Bronaugh and Werner, 2013), following the trend pre-whitening approach detailed by Yue et al. (2002) to account for serial autocorrelation. We developed a multiple linear regression model to evaluate the influence of RTS activity on HCO\textsuperscript{-3} export in Stony Creek tributary watersheds relative to other landscape variables known to influence DIC production, including hydrology, terrain roughness, and vegetation productivity (Berner, 1992; Drake et al., 2018a). To account for potential effects of varying tributary watershed areas on discharge (\( Q \)) and constituent concentration, we used tributary HCO\textsuperscript{-3} yields in the model. Instantaneous discharge (\( Q, \text{ m}^3\text{s}^{-1} \)) was estimated from the hydraulic geometry relationship between \( Q \) and stream width (Eq. 1). Discharge and HCO\textsuperscript{-3} flux (concentration*\( Q \)) were normalized to the respective tributary watershed area and scaled to estimate daily water yield (cm d\(^{-1}\)) and HCO\textsuperscript{-3} yield (\( \mu \text{mol m}^{-2} \text{ d}^{-1} \)). Daily HCO\textsuperscript{-3} yields in Stony Creek tributaries were modeled as:

\[ \text{HCO}_3^- \text{ yield} = \text{RTS}_s + \text{RTS}_\text{ex} + \text{Water yield} + \text{TR} + \text{EVI} \] (2)

where RTS\(_s\) is the number of active RTSs; RTS\(_\text{ex}\) is the watershed area disturbed by RTSs (%); TR is the mean terrain roughness (m); and EVI is the mean enhanced vegetation index (−1 to 1). The multiple linear regression was trimmed using the step function in the R package lmerTest (Kuznetsova et al., 2018) to eliminate covariates which did not improve the model. Highly collinear covariates were identified using a Variance Inflation Factor > 3 (Zuur et al., 2010) and removed from the trimmed models. Model fits were inspected visually with residual plots and covariates were transformed as needed to meet assumptions of independent and homoscedastic residuals (Zuur, 2009). To understand potential effects from variable rainfall on water yields prior to and during the two-day sampling window of the Stony Creek tributaries, we inspected total rainfall in 24 h increments preceding the sampling of each Stony Creek tributary. Hourly rainfall data were obtained from a Government of Northwest Territories Total meteorological station located ~1 km from the RTS FM2 (Fig. A2). Statistics were performed in the R programming environment (v.3.4; R Core Team, 2018) and significance was interpreted at \( \alpha = 0.05 \). Summary statistics are reported as mean ± standard error, unless noted.
Geochemistry of the mainstem and tributary sites are summarized in Table 2. Among sites, pH was generally circumneutral and conductivity was higher in proximity to RTS activity. pH was highest in the RTS FM2 runoff (7.69 ± 0.05, mean ± SE), intermediate in Dempster Creek (7.07 ± 0.42), and lowest in Stony Creek (6.86 ± 0.21). Along the RTS FM2 runoff transect, pH decreased from 7.72 to 7.51 between sites one and two, and increased thereafter to 7.80 at site five. pH in the Dempster Creek headwaters (5.82) was lower than in the RTS-affected reach (7.48 ± 0.1). In Stony Creek pH increased from 5.66 in headwaters to 7.70 at sites 6–8.

Similar to pH, conductivity was higher in the RTS FM2 runoff (1799 ± 111 µS cm⁻¹) than in Dempster Creek (520 ± 191) and Stony Creek (320 ± 19). Conductivity in RTS FM2 increased from 1370 to 1990 µS cm⁻¹. Along Dempster Creek, conductivity increased from 52 µS cm⁻¹ in the undisturbed headwaters to 958 µS cm⁻¹ at the first site downstream of RTS FM2, and decreased downstream thereafter. In Stony Creek, conductivity decreased between the headwaters and the fourth downstream site, and was relatively constant at ~285 µS cm⁻¹ along the lower reach of Stony Creek (sites 5–8).

Streams were characterized by Ca²⁺–Mg²⁺–SO₄²⁻–type waters (Fig. 2) with low concentrations of Cl⁻ relative to SO₄²⁻, reflecting a predominance of H₂SO₄ carbonate weathering and sulfate salt (e.g., gypsum) dissolution over other mineral weathering sources. A relatively greater proportion of SO₄²⁻ than HCO₃⁻ in the RTS FM2 runoff and along the upper reach of Stony Creek (sites 1–4) (Fig. 2a) suggests greater sulfate salt dissolution and/or carbonate weathering at these sites buffered less H₂SO₄ (Eq. 7) than in Dempster Creek headwaters and its tributaries (Eq. 3). Along the Stony Creek mainstem (sites 1–8), increasing HCO₃⁻ (Fig. 2a) reflected inputs from RTS-affected tributaries (sites 2–7) having relatively more HCO₃⁻-type waters (Fig. 2b) from H₂SO₄ and potentially H₂CO₃ carbonate weathering.

### 3.2 HCO₃⁻ Concentration and pCO₂

Carbonate alkalinity (HCO₃⁻ + CO₃²⁻) was primarily HCO₃⁻ (>99%) at all sites. HCO₃⁻ was highest in the RTS FM2 runoff (1429 ± 23 µM), intermediate in Dempster Creek (864 ± 261 µM), and lowest in Stony Creek (255 ± 59 µM). Along the RTS FM2 runoff transect, HCO₃⁻ decreased from 1510 to 1386 µM. In Dempster Creek and Stony Creek, HCO₃⁻ concentrations were relatively low in undisturbed headwaters (115 and 33 µM, respectively) and two to ten times higher at the first RTS-affected site (1321 and 69 µM, respectively). HCO₃⁻ decreased along the entire RTS-affected reach of Dempster Creek (from 1321 to 946 µM) in conjunction with inputs from dozens of tributary watersheds without active RTSs. In contrast, HCO₃⁻ increased significantly along Stony Creek (p < 0.01, Mann-Kendall test) (Fig. 3a) in conjunction with inputs from RTS-affected tributaries.

CO₂ was oversaturated at all sites (Fig. 3b) and a minor component of DIC (<10%) at most sites except the undisturbed headwaters of Dempster Creek (site 1) and upper Stony Creek (sites 1–3). pCO₂ was highest in the Dempster Creek headwaters (2467 µatm), relatively high in the RTS FM2 runoff (1023 ± 137 µatm), and consistently near atmospheric levels along Stony Creek (479 ± 12 µatm). Along the RTS FM2 runoff transect, pCO₂ increased from 1046 to 1534 µatm within the first 220 m and then decreased from 1534 to 742 µatm over the final...
Along Dempster Creek, $p_{\text{CO}_2}$ decreased from 2467 in the headwaters to 686 µatm at the first RTS-affected site, and further decreased to 600 µatm by the end of Dempster Creek. $p_{\text{CO}_2}$ in Dempster and Stony Creek tributaries were generally similar to the mainstem sites.

### 3.3 DOC Concentration and SUVA$_{254}$

DOC concentrations were highest in Dempster Creek (933 ± 83 µM), intermediate in the RTS FM2 runoff (758 ± 152 µM), and lowest in Stony Creek (303 ± 54 µM). Along the Dempster Creek transect, DOC decreased between the undisturbed headwaters (960 µM) and the first RTS-affected site (790 µM) and increased thereafter along the transect (to 1156 µM) (Fig. 3c). Along Stony Creek, DOC increased significantly (from 102 to 551 µM) ($p < 0.001$, Mann-Kendall test).

SUVA$_{254}$ values were lowest in the RTS FM2 runoff (1.85 ± 0.4 L m$^{-1}$ mgC$^{-1}$ m$^{-1}$), highest in Dempster Creek (3.10 ± 0.2 L m$^{-1}$ mgC$^{-1}$ m$^{-1}$), and intermediate in Stony Creek (2.51 ± 0.3 L m$^{-1}$ mgC$^{-1}$ m$^{-1}$). SUVA$_{254}$ values along the Dempster Creek transect followed a similar pattern to DOC and along Stony Creek SUVA$_{254}$ values doubled (Fig. 3d). DOC and SUVA$_{254}$ increased in consecutive downstream tributaries of Stony Creek, but not Dempster Creek.

### 3.4 Stable Isotopic Composition of Carbon in DIC and CO$_2$

$\delta^{13}$C-DIC values were highest in the RTS FM2 runoff (−1.0‰) and lower, on average, along the mainstem Dempster Creek (−7.5 ± 2.5‰) and Stony Creek (−8.4 ± 0.5‰). In the undisturbed headwaters of Dempster and Stony Creek, relatively negative $\delta^{13}$C-DIC values (−11.6 to −15.6‰) reflected DIC sourced from a combination of atmospheric and biogenic (soil) CO$_2$ (Fig. 4). In the RTS FM2 runoff, relatively $^{13}$C-enriched $\delta^{13}$C-DIC (−1.0‰) aligned with H$\text{SO}_4$ carbonate weathering, $\delta^{13}$C-DIC decreased from −4.2‰ at the first site downstream of the RTS FM2 runoff to −5.7‰ at the end of Dempster Creek. Along Stony Creek, $\delta^{13}$C-DIC increased from the undisturbed headwaters (−11.6‰) to the most downstream site (−7.8‰). $\delta^{13}$C-DIC signals of H$\text{SO}_4$ carbonate weathering diminished slightly downstream along the Dempster Creek transect and intensified along Stony Creek (Fig. 4).

Similar to $\delta^{13}$C-DIC, $\delta^{13}$C-CO$_2$ values were higher in the RTS FM2 runoff (−11.0 ± 0.4‰) than along the mainstem Dempster Creek (−17.9 ± 1.4‰) and Stony Creek (−16.7 ± 0.6‰) (Fig. 5). $\delta^{13}$C-CO$_2$ values were relatively low in the undisturbed headwaters of Dempster Creek (−21.6‰), and intermediate in the headwaters of Stony Creek (−13.8‰) (Fig. 5). Along the RTS FM2 runoff transect, $\delta^{13}$C-CO$_2$ values increased from sites one to four (−12.1 to −10.0‰) and decreased at site five (−11.2‰). Along the RTS-affected reach of Dempster Creek, $\delta^{13}$C-CO$_2$ values decreased from −16.0 to −18.5‰ in conjunction with inputs from non RTS-affected tributaries having relatively low $\delta^{13}$C-CO$_2$ (−18.7 ± 1.4‰) that was more similar values from soil-respired CO$_2$. Along Stony Creek, $\delta^{13}$C-CO$_2$ values decreased from −13.8 to −18.1‰, showing a trend opposite that of $\delta^{13}$C-DIC (Fig. 5). Among sites, atmospheric $\delta^{13}$C-CO$_2$ values were relatively consistent (−9.5 ± 0.4‰, mean ± SD).
Variance in δ13C of CO2 and DIC could be influenced by biotic production, CO2 conversion to HCO3−, and/or mixing with atmospheric CO2. To evaluate the relative influence of these processes, we compared measured δ13C- CO2 for Stony Creek with theoretical values reflecting DIC controlled by speciation along the pH continuum (Sec. 2.4). In the undisturbed headwaters, δ13C-CO2 indicated stronger influence from atmospheric CO2 (Fig. 6). Along the upper, RTS-affected reach of Stony Creek (sites 2–5, from ~5 to 35 km), the good agreement between measured and theoretical δ13C-CO2 values reflected equilibrium fractionation (ε = 9.7‰ at 9°C) (Mook et al., 1974) between CO2 and HCO3−, indicating greater influence from DIC speciation (Fig. 6). Along the lower RTS-affected reach of the transect (sites 6–8), δ13C-CO2 values more strongly reflected biotic CO2 production with potential effects from degassing and/or CO2 conversion to HCO3−. These trends in δ13C-CO2 values along Stony Creek show a downstream change in the processes influencing DIC source, which may be related to inputs of weathering solutes and organic matter from RTS-affected tributaries.

3.5 Stony Creek Tributary Carbonate Alkalinity Yields and Watershed Characteristics

Carbonate alkalinity yields in RTS-affected tributaries of Stony Creek (1558 ± 1135 μmol m−2 d−1, mean ± SD) were three orders of magnitude higher than in the non-RTS affected headwaters (1.8 μmol m−2 d−1) (Table 3). Consecutive downstream tributary watersheds exhibited no clear trends in the number of RTSs, the area disturbed by RTSs, terrain roughness, or EVI. In the Stony Creek headwater tributary, which had no active RTSs, terrain roughness (16.2 m) and vegetation productivity (EVI = 0.28) were higher than in the other six tributary watersheds (4.3 ± 1.3 m and 0.46 ± 0.01, mean ± SD, respectively). In the other tributary watersheds, the number of active RTSs reached 50 (15 ± 17, mean ± SD) and RTS disturbance area reached 3.5% (0.91 ± 1.29%, mean ± SD) (Table 3).

To elucidate landscape controls on carbonate alkalinity export in Stony Creek tributary watersheds, we paired geospatial data for active RTSs, terrain roughness, and vegetation productivity with estimates of carbonate alkalinity and water yields in a multiple linear regression model (Sec. 2.7). Water yield and the area of RTS disturbance were retained during automated covariate selection for the final model \( R^2 = 0.90, p < 0.001 \). In addition to the expected relationship between water yield and carbonate alkalinity yield, RTS disturbance area was a clear significant predictor of carbonate alkalinity yield and formed a stronger relationship with alkalinity than did water yield (Table 3).

4 Discussion

4.1 Rapid Carbon Cycling in Fluvial Network Headwaters

Within undisturbed headwaters and RTS runoff on the Peel Plateau, rapid carbon cycling enhanced fluvial CO2 efflux to the atmosphere. In undisturbed headwaters, δ13C-CO2 values indicate inputs of primarily biogenic CO2 from soil respiration into Dempster Creek. In the Stony Creek headwaters, we also found an influence from exchange with atmospheric CO2. In the undisturbed Dempster Creek headwaters, a 70% decrease in pCO2 within...
several kilometers downstream likely reflected degassing and diminishing inputs of respired CO$_2$ from soils to streams, relative to headwaters (Hutchins et al., 2019). These trends resemble headwater streams elsewhere, in that hydrologic inputs of respired CO$_2$ from riparian soils can drive CO$_2$ supersaturation in fluvial network headwaters (Campeau et al., 2018; Crawford et al., 2013), which is rapidly effluxed to the atmosphere over short distances downstream (Hotchkiss et al., 2015). In contrast, trends in hydrochemistry and stable isotopes within RTS FM2 runoff demonstrate that drivers of carbon cycling within RTSs are starkly different from those in undisturbed headwaters on the Peel Plateau.

Along the RTS FM2 runoff transect, the increase in conductivity corroborates experimental evidence (Zolkos and Tank, 2020) that permafrost sediments on the Peel Plateau can rapidly weather during fluvial transport within runoff. In the upper reach of the runoff transect, near RTS FM2, the decrease in HCO$_3^-$, increase in CO$_2$, and relatively enriched $\delta^{13}$C-CO$_2$ (Fig. 5) indicate rapid production of geogenic CO$_2$ via H$_2$SO$_4$ carbonate weathering (Eq. 7) and carbonate equilibrium reactions (Eq. 8) in Yedoma terrains in Siberia and Alaska, where mineral soils are relatively more organic-rich, thermokarst is associated with rapid production of biogenic CO$_2$ (Drake et al., 2018b). While respiration likely produced some CO$_2$ in RTS FM2 runoff (Littlefair et al., 2017), observed $\delta^{13}$C-CO$_2$ (~11‰) more strongly reflected H$_2$SO$_4$ weathering of regional carbonate bedrock (~0.7 to ~5.6‰) (Hitchon and Krouse, 1972) when accounting for isotopic fractionation of ~8‰ between carbonate and CO$_2$ at the temperature of FM2 runoff (18°C) (Clark and Fritz, 1997). Along the lower reach of the FM2 runoff transect, the increase in $\delta^{13}$C-CO$_2$ aligned with the preferential loss of $^{12}$C in the CO$_2$ phase via DIC fractionation and degassing (Doctor et al., 2008; Drake et al., 2018b; Kendall et al., 2014). $^{13}$C enrichment of the CO$_2$ pool by methanogenesis (Campeau et al., 2018), photosynthesis (Descolas-Gros and Fontungne, 1990), and/or calcite precipitation (Turner, 1982) was unlikely, as $\delta^{13}$C in FM2 runoff was relatively low ($\delta^{13}$C$_{\text{CH}4} = 3.6 \pm 1.9$ ‰, mean ± SD, $n = 6$), the high turbidity of FM2 runoff likely inhibited photosynthesis (Levenstein et al., 2018), and calcite was below saturation (SI = 0.79). These trends demonstrate that weathering of sediments during fluvial transport within RTS runoff can result in rapid CO$_2$ production and efflux to the atmosphere, in agreement with recent estimates of high rates of CO$_2$ efflux within RTS runoff (Zolkos et al., 2019).

High rates of weathering within RTS FM2 runoff aligns with observations of rapid solute production via the exposure and weathering of carbonate flour in glacial foreground environments (Anderson, 2007; Sharp et al., 1995; St. Pierre et al., 2019). Because minerals exposed by deeper RTSs are generally reactive, and sediment concentrations increased by three orders of magnitude between the undisturbed Dempster Creek headwaters and the first RTS-affected site, we reasoned that H$_2$CO$_2$ weathering of these sediments during fluvial transport would measurably influence pCO$_2$ along Dempster Creek (Eq. 1) (St. Pierre et al., 2019; Striegl et al., 2007). Although pCO$_2$ decreased along the RTS-affected reach of the Dempster Creek transect (sites 2–4, Fig. 2b), coincident decreases in conductivity, HCO$_3^-$, and pH (Table 2, Figs. 2a, 4) suggest that degassing and dilution associated with inputs from non RTS-affected tributaries had stronger effects on pCO$_2$ than did H$_2$CO$_2$ carbonate weathering, even at the relatively short scale of this 14 km transect. From a carbon cycling perspective, biogeochemically reactive
mineral substrate appears to be rapidly transformed in headwaters on the Peel Plateau; geogenic CO₂ production is relegated to within RTSs; and more stable weathering products, including alkalinity, are exported downstream.

4.2 RTS Activity in Headwaters Amplifies Carbonate Alkalinity Production and Accumulation Across Scales

Similar to CO₂, alkalinity production on the Peel Plateau was strongly coupled to primarily H₂SO₄ carbonate weathering mediated by RTS activity. This was reflected by a modest decrease in HCO₃⁻ along Dempster Creek in tandem with decreasing RTS disturbance area (from 3.2 to 1.2%) and some dilution by inputs from non-RTS-affected tributaries. Multiple linear regression results further indicated that RTS activity was a primary terrain control on carbonate alkalinity yields. In the Stony Creek headwaters, low carbonate alkalinity yield relative to water yield suggested that HCO₃⁻ export was limited by carbonate availability rather than by water. In RTS-affected tributaries, higher carbonate alkalinity yields relative to water yields aligned with the model results indicating that RTS activity increases carbonate weathering and alkalinity export beyond what would otherwise be expected on the Peel Plateau. HCO₃⁻ yields in RTS-affected tributaries were comparable to summertime HCO₃⁻ yields in watersheds with carbonate weathering by glacial activity (~3000 µmol m⁻² d⁻¹) (Lafrenière and Sharp, 2004; Striegl et al., 2007), emphasizing that unmodified sulfate- and carbonate-bearing sediments in regional permafrost are highly reactive (Zolkos and Tank, 2020) and primary sources of DIC production within intermediate-sized (1000 km²) fluvial networks. This aligns with stable sulfur isotopes in RTS runoff and near the Stony Creek outflow that strongly reflected sulfate oxidation (Zolkos et al., 2018). Unlike CO₂, the increase in HCO₃⁻ by orders of magnitude along Stony Creek in association with inputs from RTS-affected tributaries shows that more chemically stable (i.e., non-gaseous) weathering products accumulated across scales. This aligns with previous findings that solutes and sediments from RTSs propagate through fluvial networks (Kokelj et al., 2013; Malone et al., 2013), and suggests that future intensification of RTS activity (Segal et al., 2016) will increase HCO₃⁻ export to downstream environments.

4.3 Integration of RTS Effects on Carbon Cycling Across Watershed Scales

These findings enable us to develop a conceptual model of catchment chemical characteristics and how the effects of RTS activity on carbon cycling integrate across watershed scales on the Peel Plateau (Fig. 7). This model may be generalized to permafrost terrains elsewhere for testing hypotheses related to thermokarst effects on carbon cycling across the land-freshwater-ocean continuum (Tank et al., 2020).

In undisturbed headwaters on the Peel Plateau, DIC was primarily CO₂ and sources of CO₂ varied from relatively more atmospheric in the sparsely-vegetated and mountainous Stony Creek headwaters (Fig. 7a i), to more biogenic in the tundra-taiga headwaters of Dempster Creek (Fig. 7a ii). Downstream, CO₂ loss and mixing of streams resulted in undisturbed headwaters having relatively modest DIC comprised of a relatively large proportion of CO₂ sourced from mixing with the atmosphere and likely some inputs from soil respiration (Fig. 7a iii). Underlying the trends in CO₂ concentration, measurements of δ¹³C-CO₂ revealed shifting sources of CO₂ across scales (discussed below).

3.5 Inorganic Carbon Cycling in Thermokarst-Affected Fluvial Networks

Our study is among the first to evaluate inorganic carbon cycling across gradients of thermokarst disturbance and nested watersheds, enabling an assessment of how catchment chemical characteristics and thermokarst effects integrate across watershed scales. In this framework, our findings support a novel conceptual model of land-freshwater linkages and carbon cycling from headwaters to intermediate scales in thermokarst terrains (Fig. 8). Broadly, these findings reveal fast DIC cycling in headwaters and substantial DIC export downstream. Thermokarst activity increased CO₂ production in headwaters and fluvial HCO₃⁻ export across scales by unearthing large amounts of reactive inorganic substrate previously sequestered in permafrost (see also Kokelj et al. 2013, Malone et al. 2013). The striking carbonate weathering and DIC production that we documented in the RTS FM2 runoff indicates that rapid carbon cycling can be expected where thermokarst releases inorganic substrate with limited prior modification (Lacelle et al., 2019; Zolkos and Tank 2020). In the larger Stony Creek watershed, DIC cycling was characterized by relatively slower processes associated with HCO₃⁻ export, which reached magnitudes comparable to watersheds with carbonate denudation by glacial activity (Lafrenière and Sharp 2005, Striegl et al. 2007). Our findings directly link accelerating thermokarst activity on the Peel Plateau (Segal et al. 2016) with signals of intensifying carbon weathering across the broader Peel and Mackenzie River watersheds (Tank et al. 2016, Zolkos et al. 2018) and suggest that accelerating thermokarst activity will intensify inorganic carbon cycling across broad swaths of the circumpolar north (Zolkos et al. 2018). Permafrost terrains susceptible to hillslope thermokarst like RTSs are abundant within and outside of former glacial limits across the circumpolar north (Zolkos et al. 2018). Owing to regional variability in geology, glacial activity, climate, and ecosystem history, the mineral composition of permafrost in these regions is likely heterogeneous (Lacelle et al., 2019, Zolkos et al. 2018). The degree to which
Trends in Yedoma terrains (LaCelle et al., 2019; Zolkos and Tank, 2020) by RTS activity in Peel Plateau headwaters (see also Kokelj et al., 2013; Malone et al., 2013) promotes mineral weathering, rapidly generating CO₂ and substantial alkalinity. Alkalinity, along with large amounts of sediment (van der Sluijs et al., 2018) and organic matter (Shakil et al., 2020), are exported from RTSs into fluvial networks (Fig. 7a iv). Similar to other locations, DOC in RTS runoff on the Peel Plateau is known to be relatively biolabile (Littlefair et al., 2017), suggesting inputs from RTS FM2 to larger streams (e.g. Dempster Creek) could stimulate biotic CO₂ production.

CO₂ degassing is most pronounced within RTSs and in undisturbed headwaters that are strongly coupled with soil respiration, and active mineral weathering is less pronounced in mid-order streams (e.g. Dempster Creek). Hence, mid-order streams, which also mix with inputs from undisturbed tributaries, export HCO₃⁻ downstream at a magnitude coupled to the area of RTS disturbance (Fig. 7a v). Further, immediately downstream of the RTS FM2 inflow to Dempster Creek, the decrease in CO₂ and shift in δ¹³C-CO₂ away from a biotic source suggest that CO₂ degassing to the atmosphere was more prominent than respiration of permafrost DOC (Doctor et al., 2008; Drake et al., 2018b; Kendall et al., 2014). Thus, immediately downstream of RTSs, microbial respiration of permafrost DOC does not appear to generate substantial CO₂. This may be due to lower rates of DOC mineralization than degassing, and/or the protection of DOC from microbial oxidation via adsorption to RTS sediments (Gentsch et al., 2015). The latter aligns with the observed decrease in DOC concentration and increase in TSS (to 11800 mg L⁻¹) between Dempster Creek sites one and two (Table 2, Fig. 3c) (see also Littlefair et al., 2017). However, these effects may diminish farther downstream in mid-order streams. Along the lower reach of Dempster Creek (sites 3–4), the decrease in δ¹³C-CO₂, increase in DOC, and SUVA₂₅₄ resembling terrestrial-origin DOC from tributary streams suggest that undisturbed tributary streams may deliver biogenic CO₂ and/or stimulate organic matter respiration in RTS-affected streams. Thus, effects of RTS sediments on CO₂ are attenuated downstream as DOC inputs increase.

Up to and likely beyond scales of ~10⁴ km² (e.g. Stony Creek), the largest scale of this study, HCO₃⁻ concentrations are likely to increase significantly downstream, reflecting the export of relatively stable weathering products (see also Kokelj et al., 2013; Malone et al., 2013; Zolkos et al., 2018) and accumulation of carbonate alkalinity (Fig. 7a vi, Fig. 7b). These effects were primarily driven by inputs of HCO₃⁻ from RTS-affected tributaries, which also increased DOC significantly along Stony Creek. Potentially owing to organic matter limitation, CO₂ in the undisturbed headwaters of Stony Creek appeared to be driven by relatively faster carbonate equilibrium reactions (Eq. 8) (Stumm and Morgan, 1996). In contrast, along the lower RTS-affected reach of Stony Creek, as HCO₃⁻ and DOC increased and pH stabilized, δ¹³C-CO₂ measurements suggest that respiration of organic matter from RTS-affected tributaries contributed to CO₂ oversaturation (Fig. 6). In higher-orders streams within RTS-affected fluvial networks, biotic CO₂ production may increase together with HCO₃⁻ concentrations. This trend was not evident in δ¹³C-DIC, which primarily reflected inputs of geogenic DIC from RTS-affected tributaries. Thus, sources of CO₂ may shift across scales in RTS-affected fluvial networks, and measurements of δ¹³C-CO₂ highlight a decoupling between the drivers of CO₂ and HCO₃⁻ at larger scales (Horgby et al., 2019; Hutchins et al., 2020). A stronger signal of biogenic CO₂ production in larger streams than in permafrost thaw streams, as we observed, is opposite common trends in Yedoma terrains (Drake et al., 2018b) and may partly reflect limitation of organic substrate in Stony Creek.
headwaters that is relieved by RTS inputs farther downstream (Shakil et al., 2020). Underlying these trends, RTS disturbance area increased along the Stony Creek transect, from 0% in the undisturbed headwaters to 0.36% in the tributary watersheds (sites 4–8). Despite RTS activity occupying a small proportion of the landscape, carbonate alkalinity propagated through fluvial networks. These findings directly link intensifying RTS activity on the Peel Plateau (Segal et al., 2016) with signals of increasing weathering and carbonate alkalinity export in the broader Peel and Mackenzie River watersheds (Tank et al., 2016; Zolkos et al., 2018).

### 4.4 Implications for Carbon Cycling in Northern Permafrost Regions

Permafrost terrains susceptible to hillslope thermokarst like RTSs occur within and outside of former glacial limits across the circumpolar north (Olefeldt et al., 2016; Zolkos et al., 2018), but variability in geology, glacial activity, climate, and ecosystem history cause permafrost mineral composition to vary between regions. The degree to which carbonate weathering is coupled with sulfide oxidation will determine if mineral weathering is a CO$_2$ sink (Eq. 1) or source (Eqs. 3, 7) over the coming millennia (Zolkos et al., 2018). Where thermokarst releases inorganic substrate with limited prior modification – as in deeper RTSs on the Peel Plateau (Lacelle et al., 2019; Zolkos and Tank, 2020) – carbon cycling can be expected to be rapid and driven by inorganic processes, and strengthen abiotic components of the permafrost carbon-climate feedback (Schuur et al., 2015). Current dynamic-numerical biogeochemical models for the Mackenzie River basin suggest the ubiquity of sulfide minerals reduces weathering consumption of atmospheric CO$_2$ by half (Beaulieu et al., 2012). These models do not account for enhanced H$_2$SO$_4$ carbonate weathering associated with RTS activity, which our results show is significantly and positively correlated with alkalinity production and export across watershed scales. Further, climate feedbacks associated with RTS activity appear to be scale-dependent. RTSs rapidly generate CO$_2$, but its outgassing occurs mostly within runoff and comprises a small proportion of watershed-scale fluvial CO$_2$ efflux (Zolkos et al., 2019). Carbonate alkalinity generated within RTSs represents a much larger positive feedback to climate change, albeit over geological timescales, via carbonate precipitation reactions within the marine carbon cycle (Calmels et al., 2007). Future intensification of RTS activity (Segal et al., 2016) can thus be expected to increase geogenic CO$_2$ production within headwaters (see also Zolkos et al., 2019) and carbonate alkalinity export across scales (Fig. 7b) (Tank et al., 2016). Cross-scale watershed investigations will help to understand these effects across terrains with varying lithologies and permafrost composition, and the implications of hillslope thermokarst for climate feedbacks.

### 5. Conclusions

Climate-driven renewal of deglaciation-stage geomorphic activity and associated carbonate weathering in the western Canadian Arctic is amplifying aquatic carbon export across scales, despite RTSs disturbing only a fractional proportion of the landscape. Primary consequences include geogenic CO$_2$ production that is rapid, and localized to RTSs, and augments soil-respired CO$_2$ efflux from undisturbed headwater streams. Significant carbonate alkalinity production within and export from RTSs project through fluvial networks and likely to Arctic coastal marine environments, forecasting stronger land-freshwater-ocean linkages (Tank et al., 2016) as RTS activity intensifies in...
glacial margin landscapes across northwestern Canada (Kokelj et al., 2017a). Legacy effects of RTSs on carbon cycling can be expected to persist for millennia and should spur the integration of dynamic-numerical biogeochemical models (Beaulieu et al., 2012) into predictions of weathering-carbon-climate feedbacks (Zolkos et al., 2018) among northern thermokarst terrains (Turetsky et al., 2020).

Data availability. All data used in this study are available in the supplement.

Supplement. The supplement for this article is available online.

Author contributions. SZ and SET designed the study with contribution from RGS and SVK. SZ led the field research, laboratory analyses, and manuscript writing. JK contributed to geospatial analyses. CEA contributed to laboratory analyses. All authors (SZ, SET, RGS, SVK, JK, CEA, DO) contributed to manuscript writing.

Competing interests. The authors declare no conflicts of interest.

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References


### Table 1. Mineral weathering equations used to create Piper diagram end-members. $\text{H}_2\text{CO}_3 = \text{carbonic acid}, \text{H}_2\text{SO}_4 = \text{sulfuric acid}. \text{H}_2\text{O}$ includes dissolved $\text{CO}_2\text{(aq)}$.

<table>
<thead>
<tr>
<th>Eq.</th>
<th>Reaction</th>
<th>Equation</th>
<th>Reference</th>
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<tbody>
<tr>
<td>1</td>
<td>H$_2$CO$_3$ carbonate weathering (CACW)</td>
<td>$\text{H}_2\text{CO}_3 + (\text{Ca},\text{Mg})\text{CO}_3 \rightarrow (\text{Ca}^{2+},\text{Mg}^{2+}) + 2\text{HCO}_3^-$</td>
<td>Lehn et al. (2017)</td>
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<td>2</td>
<td>H$_2$CO$_3$ silicate weathering (CASW)</td>
<td>$2\text{H}_2\text{CO}<em>3 + 3\text{H}<em>2\text{O} + (\text{Ca},\text{Mg})\text{AlSi}</em>{2}\text{O}</em>{8} \rightarrow (\text{Ca}^{2+},\text{Mg}^{2+}) + 2\text{HCO}_3^- + 2\text{AlSiO}_4\text{(OH)}_4$</td>
<td>Lehn et al. (2017)</td>
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<td>3</td>
<td>H$_2$SO$_4$ carbonate weathering (SACW)</td>
<td>$\text{H}_2\text{SO}_4 + 2(\text{Ca},\text{Mg})\text{CO}_3 \rightarrow 2(\text{Ca}^{2+},\text{Mg}^{2+}) + \text{SO}_4^{2-} + 2\text{HCO}_3^-$</td>
<td>Lehn et al. (2017)</td>
</tr>
<tr>
<td>4</td>
<td>H$_2$SO$_4$ silicate weathering (SASW)</td>
<td>$\text{H}_2\text{SO}<em>4 + \text{H}<em>2\text{O} + (\text{Ca},\text{Mg})\text{AlSi}</em>{2}\text{O}</em>{8} \rightarrow (\text{Ca}^{2+},\text{Mg}^{2+}) + \text{SO}_4^{2-} + \text{AlSiO}_4\text{(OH)}_4$</td>
<td>Lehn et al. (2017)</td>
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<td>5</td>
<td>Sulfate salt dissolution (SSD)</td>
<td>$(\text{Ca},\text{Mg})\text{SO}_4 \rightarrow (\text{Ca}^{2+},\text{Mg}^{2+}) + \text{SO}_4^{2-}$</td>
<td>Lehn et al. (2017)</td>
</tr>
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<td>6</td>
<td>Sulfide oxidation</td>
<td>$\text{FeS} + 15/4\text{O}_2 + 7/2\text{H}_2\text{O} \rightarrow \text{Fe(OH)}_3 + 2\text{H}^+ + \text{SO}_4^{2-}$</td>
<td>Calmes et al. (2007)</td>
</tr>
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<td>7</td>
<td>Carbonate weathering by H$_2$SO$<em>4$ in excess (SA$</em>\text{ex}$CW)</td>
<td>$2\text{H}_2\text{SO}_4 + (\text{Ca},\text{Mg})\text{CO}_3 \rightarrow \text{Ca}^{2+} + \text{Mg}^{2+} + 2\text{SO}_4^{2-} + 2\text{H}_2\text{CO}_3$</td>
<td>Stallard and Edmond (1983)</td>
</tr>
<tr>
<td>8</td>
<td>Carbonate equilibrium / DIC speciation</td>
<td>$\text{H}_2\text{O} + \text{CO}_2\text{(aq)} + \text{H}_2\text{CO}_3 \leftrightarrow 2\text{H}^+ + \text{HCO}_3^- \leftrightarrow 2\text{H}^+ + \text{CO}_3^{2-}$</td>
<td>Stumm and Morgan (1996)</td>
</tr>
<tr>
<td>Type</td>
<td>Site</td>
<td>pH</td>
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- tributary and
- DIC

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- were

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### Updated:
- Table 2: Geochemistry of mainstem and tributary sites along Dempster and Stony Creeks. Retrogressive thaw slump (RTS) FM2 runoff samples collected on July 31, 2017, except where noted (July 30, 2017). RTS FM2 runoff site #5 was nearest the confluence with Dempster Creek (Fig. 1). Area = watershed area, SE = standard error. *Not RTS*

### Deleted:
- Sampled on July 30, 2017, while other

### Table 2:

#### Dempster (Mainstem)

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<th>DOC (µM)</th>
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### Updated:
- Table 2: Geochemistry of mainstem and tributary sites along Dempster and Stony Creeks. Retrogressive thaw slump (RTS) FM2 runoff samples collected on July 31, 2017, except where noted (July 30, 2017). RTS FM2 runoff site #5 was nearest the confluence with Dempster Creek (Fig. 1). Area = watershed area, SE = standard error. *Not RTS*

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- cal characteristics
- tributary and
- DIC

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### Updated:
- Table 2: Geochemistry of mainstem and tributary sites along Dempster and Stony Creeks. Retrogressive thaw slump (RTS) FM2 runoff samples collected on July 31, 2017, except where noted (July 30, 2017). RTS FM2 runoff site #5 was nearest the confluence with Dempster Creek (Fig. 1). Area = watershed area, SE = standard error. *Not RTS*

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- cal characteristics
- tributary and
- DIC

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Table 3. Characteristics of Stony Creek tributary watersheds (upper panel) and results from the multiple linear regression model (lower panel). The Stony Creek watershed contained 109 retrogressive thaw slumps (RTS), 92 of which were in the major tributaries of the Stony Creek mainstem. TR = terrain roughness. EVI = enhanced vegetation index. Values were used in the multiple linear regression model to determine the drivers of HCO$_3$– yields in Stony Creek tributary watersheds. Model results are shown in the lower panel. Covariates eliminated during model selection (RTS, TR, EVI) are not reflected in the lower panel or final model: lnHCO$_3$– yield = 1.04 lnWater yield + 0.35 lnRTS$_{area}$ + 8.76.

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--- [3]
Figure 1. Sampling sites on the Peel Plateau (NWT, Canada). Water samples were collected along the mainstem Dempster and Stony Creeks (n = 12) and major tributaries (n = 10), and from the rill runoff at retrogressive thaw slump (RTS) FM2. Numbers within symbols are sampling sites (Tables 1 and A1). RTS impact accumulation represents the number of active RTSs affecting upstream reaches (n = 109) (see Methods Sect. 2.6). (a) Aerial photograph of Stony Creek where it was first impacted by RTS activity. (b) RTS FM2 runoff transect sampling scheme. RTS FM2 spans ~40 ha, its headwall (e) reaches ~25 m in height, and the debris tongue contains 2 × 10^6 m^3 of sediment (van der Sluijs et al., 2018). (c) Aerial photograph of the Stony Creek (lower left) flowing into the Peel River. (d) Satellite image of RTS FM2 in September 2017 (j) obtained from Copernicus Sentinel data (European Space Agency, https://sentinel.esa.int/). Basemap: Esri ArcGIS Online © OpenStreetMap contributors, GIS User Community.
Figure 2. Piper diagrams (modified to show the upper half of the diamond plot) showing stream chemistry of the (a) mainstem sites and (b) tributary and retrogressive thaw slump (RTS) FM2 runoff sites. Axes and corresponding text in gray and black reflect the proportions of cations and anions, respectively. Mineral weathering end-members were derived from the proportional concentration (meq L⁻¹) of solutes generated by H₂CO₃ carbonate weathering (CACW, Eq. 1), H₂SO₄ carbonate weathering (SACW, Eq. 2), H₂SO₄ silicate weathering (SASW, Eq. 4), sulfate salt (e.g. gypsum) dissolution (SSD, Eq. 5), and carbonate weathering by H₂SO₄ in excess (SAexCW, Eq. 7). Site numbers given within symbols (Table A1). *Site was not affected by RTSs.
The retrogressive thaw slump (RTS) FM2 runoff is a tributary to Dempster Creek.

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Figure 3. (a) HCO₃⁻, (b) pCO₂, (c) dissolved organic carbon (DOC), and (d) SUVA₂₅₄ along the RTS FM2 runoff transect and the mainstem Stony and Dempster Creeks (see locations in Fig. 1). For the RTS FM2 runoff, DOC and SUVA₂₅₄ were sampled only at 0.55 km. Note different x-axis scales for the FM2 runoff transect (0–0.55 km, upper x-axis), Dempster Creek (0–14 km, below RTS FM2 x-axis) and Stony Creek (0–70 km, lower x-axis). Regression lines in (a) and (b) are from Mann-Kendall test (details in Sect. 2.7). Bars on x-axis indicate where RTS FM2 runoff enters the Dempster Creek transect (5.3 km) and where Dempster Creek enters Stony Creek (55 km). Site numbers are given within symbols (Table A1). *Site was not affected by RTSs.
Figure 4. The pH and composition of dissolved inorganic carbon stable isotopes ($\delta^{13}$C-DIC) in streams. The upper and lower reference lines depict theoretical end-members for equilibrium reactions (mixing with atmospheric and biotic CO$_2$, respectively). Gray boxes span theoretical end-member values for kinetically controlled mineral weathering reactions (SACW = H$_2$SO$_4$ carbonate weathering, CACW = H$_2$CO$_3$ carbonate weathering, CASW = H$_2$CO$_3$ silicate weathering) (see Sect. 2.4 for derivation of end-members). The vertical line corresponds to the pH at which $\geq$ 90% of DIC is HCO$_3^-$, for the mean observed stream water temperature (11.7°C). At pH < 7.4, $\delta^{13}$C-DIC values primarily reflect equilibrium (rather than kinetic) controls on DIC cycling. Arrows reflect increasing downstream distance from the headwaters in Stony Creek and from the first retrogressive thaw slump (RTS) affected site in Dempster Creek. Site numbers given within symbols (Table A1). *Site was not affected by RTSs.
Figure 5. The composition of DIC and CO₂ stable isotopes at varying DIC concentrations along the Dempster and Stony Creek mainstems, and in the rill runoff of retrogressive thaw slump (RTS) FM2. Arrows reflect increasing downstream distance from headwaters in Stony Creek, from the first RTS-affected site in Dempster Creek, and from the start of the FM2 runoff transect. Site numbers given within symbols (Table A1). *Site in headwaters and not affected by RTSs.
Figure 6. Observed and expected $\delta^{13}C$ CO$_2$ values along the Stony Creek mainstem. Theoretical $\delta^{13}C$ CO$_2$ values were calculated as detailed in Sect. 2.6 and reflect changes in CO$_2$ due to DIC speciation (i.e. $HCO_3^- \rightarrow H^+ + HCO_3^-$, Eq. 8). Deviation from theoretical $\delta^{13}C$ CO$_2$ values by observed values thus indicates isotopic effects from degassing and/or microbial oxidation of organic matter (OM), as indicated by the arrows. Site numbers given within symbols (Table A1). *Site was not affected by retrogressive thaw slumps.

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Figure 2. (a) Conceptual model of retrogressive thaw slump (RTS) activity and mineral weathering effects on carbon cycling in glaciated thermokarst terrains like the Peel Plateau. Source abbreviations: bio = biogenic, atm = atmospheric, geo = geogenic. (b) RTS effects on CO2 and DIC (CO3, carbonate alkalinity) observed in this study (dark solid line), projected across broader scales in the modern-day (dark dotted line), and under hypothetical future scenarios of increasing RTS activity (medium- and light-gray dashed lines). Shaded regions along x-axis depict relative RTS area approximated for modern-day (black) and for hypothetical future increases in RTS area (medium- and light-gray).

Notes:
- Deleted: Evidence for processes in the diagram are shown as numbers and discussed in the main text:
  1. Low CO2; intermediate δ13C; high SO4; and δ13C in DIC;
  2. Low δ18O in DIC; decreasing CO2; high DIC, pH, and δ18O in DIC;
  3. Relatively δ13C-enriched δ18O; and increasing δ13C in DIC, high but decreasing DIC and conductivity;
  4. Increasing DIC, pH, and δ18O in DIC, and δ13C in DIC reflect microbial activity during thermokarst formation.
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- Deleted: fluvial inorganic
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- Deleted:
- Deleted: NWT.
- Deleted: Richardson Mountains
- Deleted: Dempster Creek
- Deleted: 
- Deleted: (RTS FM2)
- Deleted: *Non RTS-affected regions along x-axis depict relative RTS area approximated for modern-day (black) and for hypothetical future increases in RTS area (medium- and light-gray).
Table A1. Sampling site characteristics. Retrogressive thaw slump (RTS) FM2 runoff was a tributary to Dempster Creek (confluence upstream of site 2) and Dempster Creek was a tributary to Stony Creek (confluence upstream of site 8). Coordinates reported in decimal degrees. Site was not affected by RTSs.

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Deleted: 'FM2 runoff distances are in m.'
Figure A1. Estimated vs. measured discharge ($p < 0.001$, $R^2 = 0.89$, $F_{1,18} = 150$) for 20 streams in the Stony Creek watershed. Grey band represents the 95% confidence interval shown around the regression. Estimates were made using measurements of stream width, $Q$, and a hydraulic geometry model (Gordon et al., 2004) (see Sect. 2.6). The model (Eq. 1) was used to estimate $Q$ in the Stony Creek tributaries.

$y = 0.003x + 1.013$
Figure A2. Total rainfall in 24 h increments preceding the sampling of each Stony Creek tributary. Rainfall data were obtained from a Government of Northwest Territories weather station on the Peel Plateau located near the RTS FM2. Locations of tributary sampling sites and the weather station are shown in Fig. 1. *Indicates no rainfall in the 24 h window.