



Cite this: *Environ. Sci.: Processes Impacts*, 2020, **22**, 1525

Composition and photo-reactivity of organic matter from permafrost soils and surface waters in interior Alaska†

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Yedoma permafrost soils are especially susceptible to abrupt thaw due to their exceptional thickness and high ice content. Compared to other mineral soils, yedoma has a high organic carbon content, which has shown to be particularly biolabile. The organic carbon in these deposits needs to be characterised to provide an identification toolkit for detecting and monitoring the thaw, mobilisation and mineralisation of yedoma permafrost. This study characterised organic carbon isolates from thermokarst lakes (either receiving inputs from thaw of original yedoma or refrozen-thermokarst deposits, or lacking recent thaw) during winter and summer seasons within the Goldstream Creek watershed, a discontinuous permafrost watershed in interior Alaska, to identify the extent to which thermokarst-lake environments are impacted by degradation of yedoma permafrost. Waters from lakes of varied age and thermokarst activity, as well as active layer and undisturbed yedoma permafrost soils were isolated and characterised by functional group abundance (multiCP-MAS ¹³C and SPR-W5-WATERGATE ¹H NMR), absorbance and fluorescence, and photobleaching ability. DOM isolated from winter and summer seasons revealed differing composition and photoreactivity, suggesting varied active layer and permafrost influence under differing ground water flow regimes. Water extractable organic matter isolates from permafrost leachates revealed variation in terms of photoreactivity and photolability, with the youngest sampled permafrost isolate being the most photoreactive and photolabile. As temperatures increase, release of permafrost organic matter is inevitable. Obtaining a holistic understanding of DOM composition and photoreactivity will allow for a better prediction of permafrost thaw impacts in the coming decades.

Received 29th February 2020
Accepted 27th May 2020

DOI: 10.1039/d0em00097c
rsc.li/espi

Environmental significance

Yedoma permafrost deposits are rapidly thawing in high latitudes, causing mobilisation and transformation of surface and sub-surface organic carbon pools. This study characterised organic carbon isolates from thermokarst-lakes (either receiving inputs from thaw of original yedoma or refrozen-thermokarst deposits, or lacking recent thaw) during winter and summer seasons within the Goldstream Creek watershed, a discontinuous permafrost watershed in interior Alaska, to identify the extent to which thermokarst-lake environments are impacted by degradation of yedoma permafrost.

Introduction

Northern permafrost soils hold the largest pool of terrestrial organic carbon on Earth, estimated at 1307 Gt C.¹ Of this, 450 Pg (25–35%) occurs in yedoma, yet comprises only about 7% of the permafrost area.^{1–5} This means that yedoma has a disproportionately large concentration of permafrost soil organic carbon. Yedoma deposits formed in unglaciated regions of Eurasia and North America during the late Pleistocene when syngenetic sediment, peat, and ice accumulated.^{3,6,7} Yedoma tends to be much older than non-yedoma permafrost soils, so there is arguably a more significant permafrost C feedback since the C has been locked away from the atmosphere for tens of thousands of years compared to some younger (e.g. Little Ice Age) permafrost soils.^{1,8,9}

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† Electronic supplementary information (ESI) available: Detail on chemicals utilised, elemental characterisation, NMR, and optical characterisation methods, ¹H NMR results and NMR results from IHSS isolates, as well as absorbance loss and optical index plots and representative fluorescence spectra. See DOI: 10.1039/d0em00097c



These deposits, covering 1.8 million square kilometres in North Siberia and Alaska, are especially susceptible to abrupt thaw (thermokarst) due to their exceptional thickness and high ice content.^{3,6,10-12} Ice melt leads to ground subsidence, ponding of water, and rapid talik development beneath lakes.^{6,13-15} 80% of the original late Pleistocene yedoma carbon pool has already thawed under anaerobic conditions beneath thermokarst lakes and gullies during the Holocene.^{5,8} The majority of these thermokarst deposits refroze after lakes drained, sequestering organic carbon of a different and unknown quality in less ice-rich epigenetic permafrost sediments. The remaining original yedoma is thus highly dissected across the landscape.

Thaw may be exacerbated by the infiltration of supra- or sub-permafrost ground water into unfrozen soils, further mobilizing previously held organic carbon. Release of materials from thawed permafrost soils also impacts surface water composition. For example, Toohey *et al.*¹⁶ attributed increased ions, nutrients, and observed positive monthly dissolved organic carbon (DOC) flux trends to active layer expansion and permafrost thaw in the Yukon and Tanana Rivers. In thermokarst lakes in Western Siberia, DOC concentration has been shown to be negatively correlated with lake stage development,¹⁷ with younger lakes containing more total DOC than more mature lakes, which the authors suggested was due to the progressive consumption and photodegradation of organic matter over time. However, thermokarst features on lake shores can remain active for centuries, allowing them to be a driving mechanism for the delivery of permafrost organic matter to surface waters.¹⁸

The organic carbon in original and refrozen yedoma deposits needs to be characterised in order to provide an identification toolkit for detecting and monitoring the thaw, mobilisation and mineralisation of yedoma permafrost soils. While permafrost-derived organic matter is largely comprised of smaller molecular weight, increased aliphatic moieties, and relatively less aromaticity compared to active layer carbon,¹⁹⁻²² fewer studies have characterised yedoma organic carbon.^{19,22} Organic matter in yedoma deposits in lake sediments in western Alaska were identified as largely terrestrially derived, based on C/N ratios and lipid biomarkers,²³ implying a heterogeneity in permafrost carbon composition.

Biotability of permafrost derived DOM has been observed to be dependent upon the composition of organic matter^{6,18,24,25} and lake ontogeny,²⁶ and yedoma organic matter may be especially labile.^{6,11,27} For example, Heslop *et al.*⁶ observed that increased biotability along four depths of yedoma lake sediment was due to a higher abundance of reduced and saturated organic compounds. Permafrost organic matter is less degraded, which may stimulate microbial processing of permafrost carbon upon thaw,²³ contributing significant radiative forcing from CO₂ and CH₄ emissions over the next century.²⁸

Permafrost organic carbon is also photoreactive. The extent of photo-oxidation influences its composition and thus propensity for biodegradation.^{29,30} For example, exposure to sunlight primes Alaskan Arctic soil organic carbon for microbial processing upon release and impacts its ability to produce hydroxyl radical,^{31,32} but surface waters from the

Bolshezemelskaya Tundra peatlands were found to be resistant to both bio- and photodegradation over a 1 month period.³³ Fewer studies though have focused on the photoreactivity of yedoma permafrost carbon. Stubbins *et al.*²² identified transformation but little photomineralisation of ancient yedoma-derived DOM from north-eastern Siberia near the Kolyma River. Photoreactivity can transform DOM to other products besides mineralised carbon. DOM photo-oxidation in permafrost underlain surface waters can occur *via* the formation of reactive oxygen species (ROS) such as hydroxyl radical^{30,32,34,35} or triplet excited state DOM.³⁶

DOM has been characterised in several thermokarst-impacted watersheds in Alaska, however the relationships between permafrost and surface water composition of thermokarst lakes remain poorly understood. Also, while the number of studies characterising permafrost and biogeochemical impacts upon thaw have expanded greatly over recent decades, the composition of yedoma permafrost carbon pools remains relatively unexplored. Further, even fewer studies have characterised the photoreactivity of DOM from permafrost and thermokarst-influenced waters. Given yedoma permafrost's relatively high carbon and susceptibility to form thermokarst, it is necessary to better qualify the content and behaviour of this carbon pool in order to gain understanding of its influence on the biogeochemical function of permafrost organic carbon and ultimately the ecological impacts upon its release in a warming climate.

This study characterised the composition and photoreactivity of DOM isolated from thermokarst lakes, including thaw inputs from original yedoma, thaw of refrozen-thermokarst, and lacking recent thaw, as well as surface soils, undisturbed permafrost soils, and groundwater beneath permafrost within the Goldstream Creek watershed in interior Alaska. Our objective was to identify the extent to which organic carbon pools in thermokarst-lake environments are impacted by degradation of yedoma permafrost, leaching of surface soils, and mixing with groundwater. Isolates were characterised by organic functional group abundance and optical properties, as well as susceptibility to photobleaching in simulated sunlight.

Materials and methods

Study sites

The Goldstream Creek watershed³⁷ (Fig. 1) is a sub-Arctic residential watershed underlain by discontinuous yedoma permafrost approximately 11 km northeast of Fairbanks, Alaska that drains into the Tanana River within the broader Yukon River Basin.³⁸ This study area has a mean annual temperature of -2.4°C and a mean annual precipitation of 274 mm (Fairbanks Int. Airport, 1981–2010, U.S. National Climatic Data Center) and is dominated by boreal forest vegetation. This watershed contains several thermokarst lakes, with some lakes having formed from abrupt thaw within the past 60 years.²⁸ Within this watershed, waters from multiple lakes of varying talik development and thermokarst activity (Table 1), a high carbon well, active layer soils, and permafrost soils (Table 2) were sampled. The four study lakes are located in the bottom of the watershed,



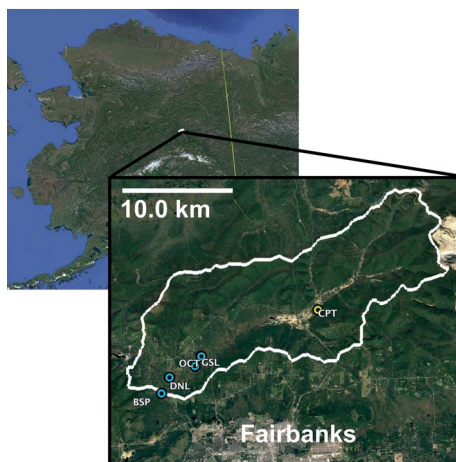


Fig. 1 Location and outline of Goldstream Watershed and sample sites. Blue locations represent surface waters. Yellow location represents CRREL Permafrost Tunnel (CPT). Active layer samples were taken at GSL site.

which is dominated by discontinuous permafrost consisting of Pleistocene organic and ice-rich deposits of the Goldstream Loess formation (e.g., yedoma), as well as its refrozen thermokarst deposits.

Goldstream Lake (informal name; GSL) has been previously described.^{38–40} The eastern margin of Goldstream Lake is expanding from active thermokarst into original yedoma.⁴⁰ Octopus Lake (informal name; OCT) appears to have minor thermokarst activity

suspected to also be into original yedoma as determined by ¹⁴C radiodating of methane and geophysical analysis.⁴¹ Blacksheep Pond (informal name; BSP) is a very small closed talik lake, less than 60 years old and resulting from abrupt thaw of refrozen thermokarst deposits. Doughnut Lake (informal name; DNL) may have also formed from thermokarst into refrozen deposits based on methane radiodating,⁴¹ but its currently insignificant thermokarst activity³⁷ allows it to serve as a negative control for thermokarst influence.

Sampling

Epilimnion surface waters were sampled for DOM isolation during summer for three lakes (DNL, GSL, OCT), and additionally in winter prior to melt for three lakes (BSP, DNL, GSL) and were transferred in a cooler to the laboratory where the waters were then analysed and processed for DOM isolation. Ground water was obtained from a high dissolved methane, high carbon residential well near Blacksheep Pond (GSRW). This water was sampled *via* an outdoor tap before filtration and softener systems in July 2016, and stored and transported as described above.

Two replicate active layer samples were obtained using a SIPRE corer during April 2018 near Goldstream Lake (GSAL1, GSAL2). The active layer depth of the cores was no more than 60 cm, however exact active layer depth was unable to be determined as supra-permafrost waters mixed with unfrozen portions of active layer while coring. Soil samples were obtained from the Cold Regions Research and Engineering Laboratory (CRREL) permafrost tunnel (CPT) with a round key hole saw attached to a power

Table 1 Physical and chemical properties of study waters

Study site ^a	Goldstream Lake (GSL)	Octopus Lake (OCT)	Doughnut Lake (DNL)	Blacksheep Pond (BSP)	Residential well (GSRW)
GPS (°N, °W)	64.916, 147.847	64.907, 147.860	64.899, 147.908	64.888, 147.920	Undisclosed
Surface area (m ²)	10 030 ^b	22 000	34 000 ^b	540	N/A
Lake max depth (m)	4.7 ^b	2	3.8 ^a	<2	N/A
Permafrost type	Original yedoma	Original yedoma	Refrozen thermokarst	Refrozen thermokarst	Unknown
Recent thermokarst	Yes	Some	No	Yes	Unknown
Est. age (years)	>120 ^b	—	1000 ^a	<60	N/A
Date(s) sampled	Jan 17	Jun 17	Aug 18	Jan 17	Aug 18
Sampled depth (m)	0	0	0	0	0
pH	6.30	8.20	8.46	5.09	8.43
Sp. conductivity (mS cm ⁻¹)	1.21	0.47	0.31	0.63	0.41
DOC (mg C L ⁻¹)	125 ± 1	556 ± 1	36.3 ± 0.7	60.2 ± 0.2	58.0 ± 0.3
TDN (mg N L ⁻¹)	5.99 ± 0.08	2.73 ± 0.03	0.16 ± <0.01	2.90 ± 0.05	0.17 ± 0.01
SUVA (L mg C ⁻¹ m ⁻¹)	2.53 ± 0.03	5.4 ± 0.1	6.3 ± 0.1	2.48 ± 0.01	3.94 ± 0.02
SUVA _{corr} (L mg C ⁻¹ m ⁻¹) ^d	1.99	5.16	5.56	2.23	3.90
S _R	0.82	0.86	0.87	0.89	1.01
E2 : E3	8.55	7.90	5.87	8.39	9.82
FI	1.70	1.62	1.54	1.56	1.52
BIX	0.74	0.70	0.60	0.70	0.72
Freshness	0.71	0.69	0.58	0.66	0.69
HIX	0.95	0.94	0.92	1.03	0.89
Dissolved Fe (μg L ⁻¹)	2960 ± 20	503 ± 2	1470 ± 30	1243 ± 2	116 ± 1
					72 ± 3
					430 ± 40

^a Unofficial names. ^b Data from Elder *et al.*³⁸ ^c TOC/TN data unavailable for this sample, reported values from a prior Aug 2016 sampling.

^d Corrections to SUVA for iron content are described in the ESI.



Table 2 Physical and chemical properties of sampled soils

Goldstream active layer (GSAL)		CRREL permafrost tunnel (CPT)		
Date(s) sampled	Apr 18		Apr 18	
GPS (°N, °W)	64.92, -147.830		64.951, -147.621	
Soil depth (m)	0–0.19	0–0.26	20	54
Soil ^{14}C age (years)	—	—	19 000 ^a	27 000 ^a
Gravimetric % water/ice	90%	88%	17%	20%
% C			3 ± 2	3.1 ± 0.3
Soil leachate properties				
pH	5.20	5.15	8.05	7.72
TOC (mg C L ⁻¹)	22.0 ± 0.4	26.1 ± 0.8	2.9 ± 0.2	11.67 ± 0.1
TDN (mg N L ⁻¹)	0.58 ± 0.03	BDL	0.49 ± 0.01	1.28 ± 0.05
SUVA (L mg C ⁻¹ m ⁻¹)	2.57 ± 0.04	1.87 ± 0.06	2.0 ± 0.1	1.72 ± 0.02
SUVA _{corr} (L mg C ⁻¹ m ⁻¹)	2.49	1.85	2.0	1.67
S_R	0.59	0.59	0.94	0.89
E2 : E3	4.69	4.66	5.24	6.26
FI	1.34	1.32	1.51	1.53
BIX	0.42	0.42	0.58	0.58
Freshness	0.40	0.39	0.57	0.57
HIX	0.88	0.90	0.95	0.93
Dissolved Fe (μg L ⁻¹)	188 ± 4	112 ± 2	13 ± 2	348 ± 3
^a Data from Mackelprang <i>et al.</i> ⁴⁷				

^a Data from Mackelprang *et al.*⁴⁷

drill at 20 m, 54 m, and 81 m from the tunnel opening. The CRREL permafrost tunnel has been well characterised in terms of its geocryology and stratigraphy,^{42–48} and is comprised of syngenetic ice-rich permafrost that has been characterised as largely containing Pleistocene aged loess and gravels of the Goldstream Formation.^{42,45,46} These depths were reported by Mackelprang *et al.*⁴⁷ as radiocarbon dated to 19 000 years, 27 000 years, and 33 000 years, respectively (Table 2). Immediately after collection, cores were wrapped in plastic and then stored at -80 °C. Soil cores were then freeze-dried to constant mass followed by storage in acid washed plastic bags in the dark.

DOM isolation and characterisation

Freeze-dried soils were leached with 18.2 MΩ water at a ratio of 1 : 200 soil : water (w/w) for a period of 7 d in the dark at 4 °C with daily agitation. Soil leachate and sampled surface waters were filtered to 0.45 μm (Pall, Port Washington, New York). Following filtering, small aliquots were sampled for total organic carbon (TOC), total nitrogen, absorbance, and fluorescence analysis. DOM was then isolated as described in Dittmar *et al.*⁴⁹ Briefly, filtered water was acidified to pH 2 with concentrated hydrochloric acid and pumped onto Bond-Elut PPL solid phase extraction cartridges (Agilent; Santa Clara, CA) at 18 mL min⁻¹. DOM was extracted from PPL cartridges with methanol, whereby eluent was rotovapped and diluted back up to 90% water before freeze-drying to constant mass. Chemicals and analytical instrumentation methods (absorbance, fluorescence, nuclear magnetic resonance; NMR) are provided in the ESI.†

Photochemical experiments

DOM isolates were reconstituted to 10 mg C L⁻¹ in aqueous solution and pH adjusted to 7.61–7.68 with NaOH or HCl and

placed in 10 mL quartz test tubes with a 1 cm inner diameter (Robson Scientific; Hertfordshire, England). Solutions were run as either pH adjusted reconstituted DOM, or with additions of 20 μM Fe(III) or 12.5 μM methanol.^{50,51} Solutions were irradiated for 24 h in an Atlas CPS+ Suntest (ATLAS; Mount Prospect, Illinois) solar simulator with a 1500 W Xe lamp set to 15 °C with duplicate samples sacrificed at each time point. Dark controls were wrapped in aluminium foil. Irradiance was monitored for consistency by a PMA2100 radiometer (Solar Light Co. Inc, Glenside, PA). Samples were stored in the dark until analysed for absorbance and fluorescence within 2 h of irradiation.

Solutions of 10 μM 2,4,6-trimethylphenol (TMP) in the presence of 10 mg C L⁻¹ DOM solutions were used to probe triplet excited DOM reactivity.³⁶ Photolysis experiments were set up as described above, and run in duplicate over 6 h with dark controls. TMP was quantified on an Agilent 1100 Series reverse-phase HPLC (Agilent Technologies, Santa Clara, CA) using a Restek Ultra C18 5 μm column (Restek Corporation, Centre County, PA) and a 60 : 40 acetonitrile : water (v/v) mobile phase at 1 mL min⁻¹, detected by fluorescence at $\lambda_{\text{ex}} = 230$ nm and $\lambda_{\text{em}} = 305$ nm.

Light screening was determined from the absorbance data by eqn (1),⁵²

$$\text{SF} = \frac{1 - 10^{-\alpha_{\lambda} l}}{2.303 \alpha_{\lambda} l} \quad (1)$$

where SF = screening factor, α_{λ} is the decadal absorption coefficient for a given wavelength λ , and l is the path length of the cell. Screening factors were integrated over 240–500 nm to determine a screening factor for the entire absorbance curve. Log-linearised pseudo-first order kinetics fits were used to determine the observed rate of absorbance loss (k_{obs}) at each wavelength or TMP photodegradation rates. Observed rates were corrected for light screening and normalised to the DOC



concentration. In order to determine if absorbance losses were due to transformation or loss of DOM by photomineralisation, DOC was monitored before and after 24 h of photolysis and corrected for light screening. Photomineralisation extent was determined by quantifying DOC loss after 24 h photoirradiation by TOC analysis (ESI†). Triplicate 10 mL samples were combined prior to TOC analysis due to volume requirements, and reported DOC losses represent an average of these, which were also corrected for light screening (eqn (1)).

Apparent quantum yields (AQY) for DOC loss were quantified using the *p*-nitroanisole (PNA) chemical actinometer, in order to compare extent of DOC loss between DOM and water extractable organic matter (WEOM) isolates, assuming all DOC was lost as CO₂.⁵³ No pyridine was added, so that PNA losses could be measured over 24 h to match the DOM photoirradiation periods. Because the solar simulator is a polychromatic light source, AQYs were calculated from 290–400 nm according to a similar method as Page *et al.*³² shown in eqn (2).

$$\phi_{\text{DOC}} = \frac{k_{\text{obs,DOC}}}{k_{\text{obs,PNA}}} \times \frac{k_{\text{abs,PNA}}}{k_{\text{abs,DOM}}} \times \phi_{\text{PNA}} \quad (2)$$

290–400 nm was chosen because of the cut-off of the daylight filter, and beyond 400 nm absorbances were very small. ϕ_{DOC} is the AQY for photochemical loss of DOC, $k_{\text{obs,DOC}}$ and $k_{\text{obs,PNA}}$ are the fitted first order rate constants of DOC and

PNA loss respectively, $k_{\text{abs,PNA}}$ and $k_{\text{abs,DOM}}$ represent the integrated light absorption over 290–400 nm for PNA and DOM respectively, and ϕ_{PNA} is the quantum yield for PNA. Uncertainty in AQY derives from standard deviations in rates, which averaged ~8% for PNA and ≤ 4% for measured DOC losses (Table 3).

Results

Surface water characterisation

For all of the sampled waters, DOC concentration was largely comparable to other boreal lakes,^{54–56} though in some cases over 100 mg C L⁻¹ in the winter months of the active thermokarst lakes (GSL, BSP). GSL's carbon content varied by over a factor of two between summer and winter, whereas DNL's carbon content was more similar (Table 1).

Specific UV absorbance at 254 nm (SUVA) values are similar to that of other lakes in interior Alaska,⁵⁴ except for GSL summer and OCT, which had SUVA > 5.0 (Table 1). Despite the similar DOC concentrations, SUVA values for GSL and DNL differ between the summer and winter. For example, GSL's SUVA values are over twice as high in the summer compared to the winter (Table 1 and Fig. S3†). Even accounting for the higher iron content in the winter (described in ESI†), compared to the summer sample, the summer sample had the higher SUVA value, indicating differences in DOM composition. Other

Table 3 Characterisation of photolability and phototransformation over 24 h photoirradiation in reconstituted DOM isolate solutions. ΔSUVA and $\Delta k_{\text{obs,254}}$ (L mg C⁻¹ h⁻¹ × 10⁻³) represent difference in SUVA after 24 h or carbon-normalised rate of absorbance loss at 254 nm upon addition of Fe(III) or methanol. TMP degradation k_{obs} are also normalised to DOC. Percent of DOC loss (% DOC_{photo}), rates of absorbance decay or TMP loss are corrected for light screening. Reported uncertainties in SUVA and % DOC_{photo} are standard deviations from analytical replicates, and photodegradation rates have uncertainties reported as standard deviation from experimental duplicates. AQY represents apparent quantum yield for DOC loss and absorbance at 254 nm loss, as determined by related rates to *p*-nitroanisole actinometer

SUVA	$\Delta\text{SUVA}_{24\text{ h}}$	% DOC _{photo}	$\text{AQY}_{\text{DOC}} \times 10^{-5}$	$k_{\text{obs,254 nm}}$ L mg C ⁻¹ h ⁻¹ × 10 ⁻³	$\Delta k_{\text{obs,254, + Fe}}$ L mg C ⁻¹ h ⁻¹ × 10 ⁻³	$\Delta k_{\text{obs,254, + MeOH}}$ L mg C ⁻¹ h ⁻¹ × 10 ⁻³	TMP k_{obs} L mg C ⁻¹ h ⁻¹	
DOM isolates								
GSL S (Jun 17)	2.64 ± 0.03	-0.63 ± 0.05	9 ± 3	32	1.15 ± <0.01	2.07 ± 0.05	-0.31 ± 0.01	0.119 ± 0.007
OCT S (Aug 18)	3.11 ± 0.02	-0.95 ± 0.03	7 ± 1	16	2.2 ± 0.4	-0.6 ± 0.4	-1.0 ± 0.4	0.129 ± 0.005
DNL S (Aug 18)	2.49 ± 0.02	-0.66 ± 0.03	Neg.	—	1.25 ± <0.01	0.18 ± <0.01	-0.11 ± 0.04	0.105 ± 0.007
GSL W (Jan 17)	2.77 ± 0.05	-0.76 ± 0.06	4 ± 3	11	1.8 ± 0.2	4.5 ± 0.2	-0.4 ± 0.3	0.11 ± 0.01
DNL W (Jan 17)	2.79 ± 0.04	-0.78 ± 0.06	6 ± 3	17	1.45 ± 0.03	2.9 ± 0.3	-0.2 ± 0.2	0.09 ± 0.02
BSP W (mar 17)	3.02 ± 0.02	-0.69 ± 0.03	7 ± 2	20	1.70 ± 0.04	7 ± 1	-0.2 ± 0.1	0.12 ± 0.03
GSRW (Jul 16)	3.82 ± 0.03	-0.69 ± 0.03	14 ± 3	29	2.0 ± 0.2	2.5 ± 0.8	-0.3 ± 0.5	0.11 ± 0.02
WEOM isolates								
GSAL1 (Apr 18)	3.73 ± 0.06	-0.68 ± 0.07	4 ± 2	8.1	1.1 ± 0.1	0.1 ± 0.1	-0.2 ± 0.2	0.050 ± 0.008
GSAL2 (Apr 18)	3.45 ± 0.06	-0.46 ± 0.07	7 ± 3	9.8	1.16 ± 0.08	0.3 ± 0.1	-0.2 ± 0.1	0.047 ± 0.003
CPT1 (Apr 18)	2.16 ± 0.02	-0.09 ± 0.04	22 ± 1	72	5.19 ± 0.04 ^a	-0.5 ± 0.3	-1.2 ± 0.6 ^a	0.14 ± 0.01
CPT2 (Apr 18)	2.73 ± 0.03	-0.80 ± 0.04	8 ± 3	23	1.3 ± 0.1	0.4 ± 0.2	-0.2 ± 0.2	0.09 ± 0.02
CPT3 (Apr 18)	2.81 ± 0.05	-0.76 ± 0.06	7 ± 3	17	1.68 ± 0.06	0.6 ± 0.1	-0.1 ± 0.1	0.09 ± 0.02
IHSS reference								
SRFA	3.86 ± 0.05	-0.50 ± 0.06	9 ± 4	15	1.1 ± 0.2 ^b	0.6 ± 0.3 ^b	-0.0 ± 0.4 ^b	0.065 ± 0.005
PLFA	2.68 ± 0.03	-0.61 ± 0.03	7 ± 3	15	0.9 ± 0.2 ^b	0.1 ± 0.3 ^b	-0.02 ± 0.3 ^b	0.066 ± 0.004

^a Rates calculated to only 6 h photolysis due minimal absorbance loss and non-first order kinetics after 6 h. ^b Errors listed are standard deviations of fitted k_{obs} from a single experiment.



optical indices are less variable across lakes and sampling events, further detailed in the ESI.†

Soil and leachate characterisation

All three permafrost soils have similar carbon content (~3% C, Table 2), consistent with previously reported measurements in the CRREL tunnel⁴⁶ and indicative of yedoma permafrost,⁵⁷ although samples were separated by over 10 000 years according to soil ¹⁴C age.⁴⁷ Ice content in permafrost soils is considerably lower than active layer moisture content (17–22% vs. 88–90%, respectively), though ice content could be underestimated due to sublimation occurring within the tunnel.⁴⁸ Soil leachate DOC differs between the sampled soils (oldest permafrost leachate had the highest DOC at $23.65 \pm 0.30 \text{ mg C L}^{-1}$), however, optical indices are relatively stable across leachates. Replicate active layer leachates from nearby GSL display near identical optical qualities (Table 2), further detailed in the ESI.†

DOM and WEOM isolate optical characterisation

In contrast to the sampled whole waters, the DOM isolates display more similar SUVA values between samples, ranging from 2.49–3.82 (Table 3). The GSL winter isolate SUVA is higher than that of the sampled filtered water (Table 1), however the summer isolate's SUVA is much lower. This could be due to the

differences in iron content, except all the isolate solutions contain less than 5 μM Fe by ferrozine (data not shown) and these differences are still present for iron corrected SUVA values (Table 1). For DNL, a similar trend occurs, where the isolate from the winter sample has a higher SUVA than the filtered water, but the summer isolate's SUVA value is lower than the sampled water, though not as dramatically as GSL. BSP and GSRW isolates display nearly the same SUVA values as their sampled waters, within the margin of error. The other optical indices are less variable between samples than their respective sampled waters and are further detailed in the ESI.†

SUVA values for WEOM isolates from soil leachates (Table 3) are higher compared to their requisite leachate solutions (Table 2), often a value of more than $1 \text{ L mg C}^{-1} \text{ m}^{-1}$ greater. Increased SUVA values have been correlated with increased aromaticity,⁵⁸ suggesting a preferential capture of aromatic groups during isolation, which may bias the carbon fraction studied relative to the entire mobilisable carbon pool. However, optical indices from soil WEOM isolates are much closer to their leachate solution optical properties than the DOM isolates are to their requisite sampled waters (Table 2 and Fig. S10†). This suggests a more uniform capture of the chromophoric and fluorescent carbon WEOM pool occurred during the isolation process compared to the sampled waters, but that non-chromophoric

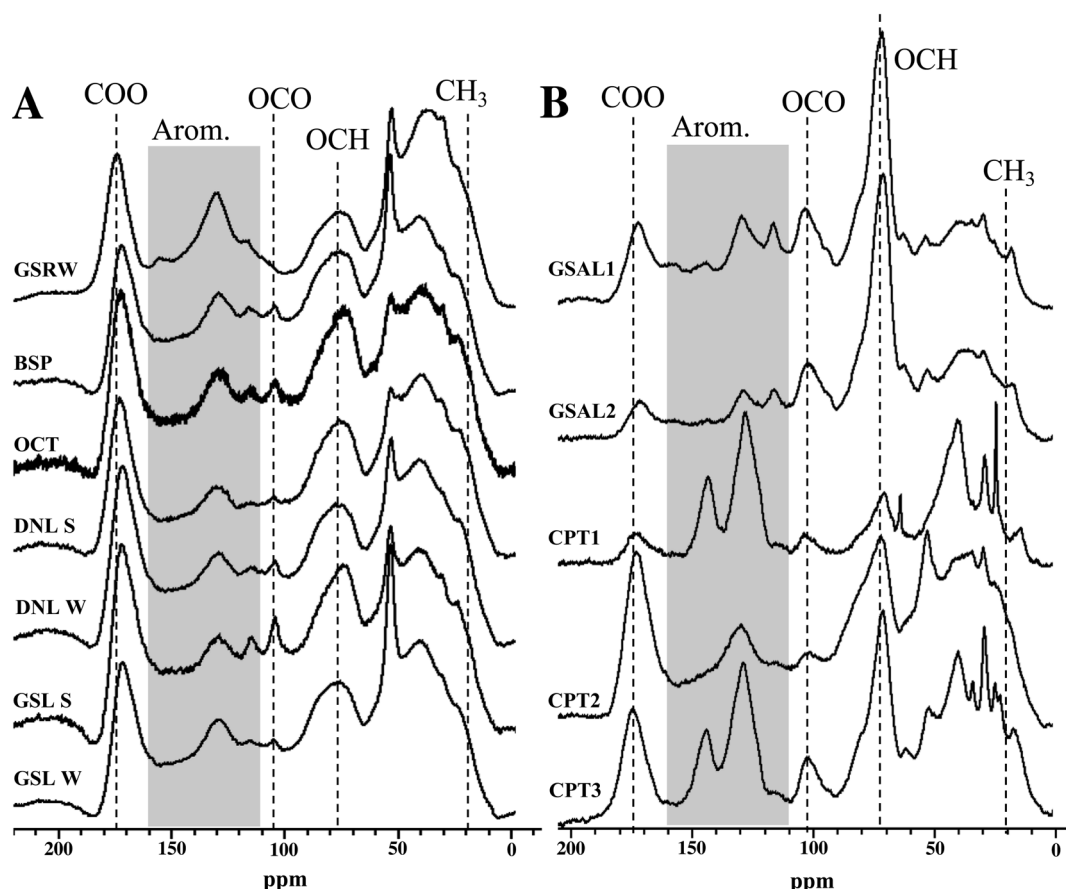


Fig. 2 ¹³C multiCP-MAS NMR spectra for PPL isolates. (A) Surface water isolates. (B) WEOM isolates.



DOM was less retained, giving rise to the higher SUVA value in the WEOM isolates compared to their leachates.

NMR spectral features

Overall, bulk functional group composition and distinct features from NMR spectra are similar across surface water DOM isolates, for both ^{13}C and ^1H spectra (Fig. 2 and S1†). More detailed results of these features are described in the ESI.† However, the WEOM isolates from sampled soils are more variable (Fig. 2). The active layer (GSAL) samples are near identical, but compared to the surface water ^{13}C spectra display a much sharper *O*-alkyl peak (70 ppm), a more defined anomeric C peak (110 ppm), a less pronounced carbonyl signal, and in some cases more defined features in the aromatic region (Fig. 2). The three permafrost tunnel isolates vary considerably, with the youngest WEOM isolate (CPT1) presenting sharper peaks in the aliphatic regions, as well as overall more defined peaks across the chemical shift range, consistent with less humification extent.⁵⁹ CPT1 contains spectral features present in some of CPT2 and CPT3 WEOM isolates. For example, CPT1 contains a less defined carbonyl peak than CPT2 or CPT3. CPT1 and CPT3 have two defined aromatic peaks at 125 and 145 ppm, but CPT2 has nearly no defined aromatic peak at 145 ppm. CPT1 also has a much less distinct *O*-alkyl peak at 75 ppm compared to CPT2 and CPT3.

Functional group distribution of DOM and WEOM isolates

Most of the DOM isolates from lake waters yield relative integrations similar to each other, with standard deviation of signal $\leq 3.5\%$ for all regions. The ability of ^{13}C multiCP-MAS to standardise signal sensitivity across chemical shift similar to DP methods⁶⁰ means that quaternary and carbonyl containing groups are likely to be elevated compared to traditional CP-MAS spectra.^{61,62} Aliphatics (0–45 ppm) and *O*-alkyl carbons (60–110 ppm) are the most abundant functional groups, averaging $26.0 \pm 3.5\%$ and $27.5 \pm 3.4\%$ of ^{13}C NMR spectra, respectively, followed by aromatic regions (110–160 ppm, $19.1 \pm 1.9\%$; Table S1†). Anomeric/acetal carbon (90–110 ppm) comprised 5.3–9.3% of the DOM isolates, with GSRW containing the smallest proportion of ^{13}C signal from this region. However, the anomeric region may actually reflect a mixture of both anomeric and aromatics found in DOM,⁶³ and are difficult to resolve from each other without other spectral editing techniques. Comparison to International Humic Substances Society (IHSS) isolates are presented in the ESI.†

Summer and winter DOM isolates from the same lakes (GSL, DNL) have similar ^{13}C NMR integrations (Table S1†), despite their differences in SUVA (Table 3). The winter DNL isolate has 3.8% more aromaticity and 3.9% less aliphatic content than its summer counterpart (Table S1†). However, in the ^1H spectra, DNL summer isolate has more aromatic signal than the winter isolate, suggesting that the increased aromaticity observed in the winter isolate ^{13}C spectrum consists of more substituted aromatics. In contrast, GSL's ^{13}C spectra from both seasons

integrate to near identical relative contributions for each region, within 2.1% for any given region.

WEOM isolates exhibit variation in both active layer and permafrost isolates, dominated by *O*-alkyl ($32.8 \pm 10.4\%$), followed by alkyl ($24.2 \pm 4.4\%$), and aromatic regions ($22.3 \pm 6.8\%$). The high average *O*-alkyl content is largely derived from the active layer and not the permafrost isolates, whose *O*-alkyl content was in some cases less than that of the DOM isolates (Table S1†). A more detailed description of WEOM isolate NMR spectra is presented in the ESI.†

The permafrost isolates differ from each other, in some cases more dramatically than compared to other isolate types. Aromaticity spread over a range from 17.7–32.9%, with no trend associated with radiocarbon age of the sampled permafrost. The isolate from the youngest permafrost sampled (CPT1) contains the highest aromaticity, followed by the isolate from the oldest (CPT3) and then the second oldest (CPT2) permafrost soil. CPT1 also contains the lowest carbonyl and *O*-alkyl content, but the highest aliphatic content of the permafrost isolates.

Photolability of DOM and WEOM isolates

Total organic carbon, absorbance and fluorescence losses upon photoirradiation were used as a probe for composition and reactivity. Several isolates saw DOC losses of over 5% from just 24 h irradiation in a solar simulator (Table 3). However, GSRW lost $13.5 \pm 2.7\%$ of initial DOC upon photoirradiation, and the WEOM isolate from the youngest permafrost soil (CPT1) lost $21.6 \pm 1.4\%$ of initial DOC. The only isolates to see DOC losses less than 5% were GSL winter, one of the active layer isolates, and negligible DOC losses were observed for only one DOM isolate, DNL summer. According to PNA actinometer degradation ($k_{\text{obs},\text{PNA}} = 0.24 \pm 0.02 \text{ h}^{-1}$), actinometry showed about $4.1 \times$ intensity compared to summer noon sunlight at 40°N , or $9.7 \times$ intensity compared to at 70°N ,⁶⁴ so these DOC losses are likely considerably higher than what would be expected to be observed in natural sunlight. It is also possible that since samples were not sterilised or sterile-filtered before experimentation, some biomineralisation could have occurred, but under 24 h of continuous photoirradiation, it is unlikely to have occurred to an extent that could account for the drastic DOC losses. Overall, significant photolability was observed.

Apparent quantum yields were calculated over 290–400 nm against PNA actinometry to account for differences in light absorption properties between the different DOM and WEOM isolates. AQYs span an order of magnitude, ranging from 8.05×10^{-5} to 7.23×10^{-4} . The highest AQYs are CPT1, followed by GSL summer, GSRW, CPT2, and BSP. The active layer WEOM isolates show the smallest AQYs of the isolates, both having an AQY < 0.0001 .

Photobleaching of DOM and WEOM isolates

Absorbance curves decay upon increased photoexposure (Fig. 3), with losses at 254 nm on average $32 \pm 10\%$ and $24 \pm 3\%$ over 24 h for DOM and WEOM isolates, respectively. Absorbance degradation follows pseudo-first order kinetics, with R^2 of



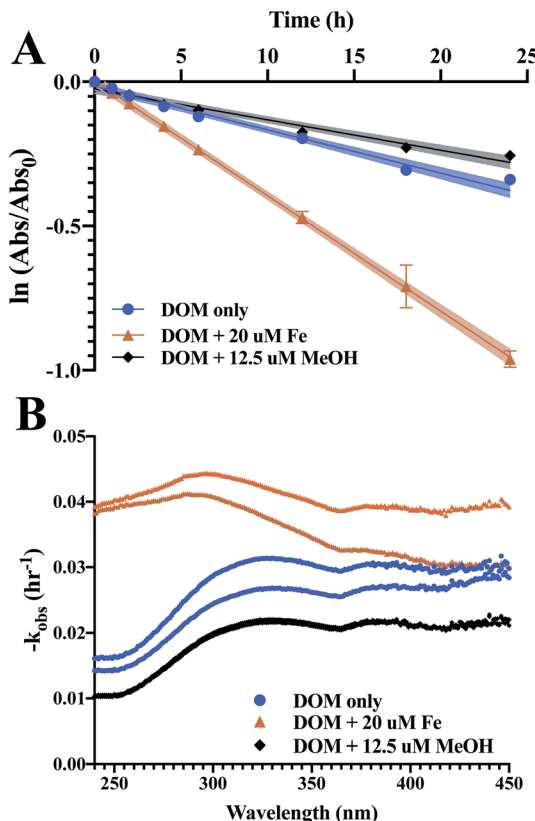


Fig. 3 Absorbance loss of 10 mg C L^{-1} GSL summer DOM isolate and amended solutions over 24 h irradiation. (A) Log linearised pseudo first-order kinetics of absorbance loss at 254 nm. (B) Fitted observed pseudo first order rates of absorbance loss across wavelengths.

first order fits >0.92 for chromophores absorbing $<400 \text{ nm}$. For several cases, fitted absorbance loss rates at higher wavelengths ($>400 \text{ nm}$) are less first order, with $R^2 < 0.90$, likely due to detection limits arising from low absorbance at those wavelengths. In general, absorbance loss rates are slowest at lower wavelengths ($<260 \text{ nm}$), and steadily increase or level off with increasing wavelength (Fig. S5 and S6†). For several DOM isolates, a local absorbance loss rate maximum was observed $\sim 320 \text{ nm}$ (BSP, DNL summer, GSL, OCT; Fig. S5†), while for the WEOM isolates a local maximum in absorbance loss appears at a slightly higher wavelength, $\sim 340 \text{ nm}$ (Fig. S6†).

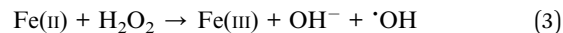
Absorbance loss rates at 254 nm range from $1.15\text{--}2.2 \times 10^{-3} \text{ L mg}^{-1} \text{ C}^{-1} \text{ h}^{-1}$ for DOM isolates, and $1.1\text{--}5.19 \times 10^{-3} \text{ L mg}^{-1} \text{ C}^{-1} \text{ h}^{-1}$ for WEOM isolates (Table 3). Of the DOM isolates, GSRW and OCT photobleach the fastest (Table 3 and Fig. 4). Though CPT1's absorbance loss could only be measured over 6 h, its absorbance loss rate is over 3–4 times faster than the other WEOM isolates (Table 3). Six-hour loss rates of the other isolates average only 1.3 ± 0.3 times faster than their 24 h loss rates (data not shown). Both DOM winter isolates photobleach faster than their summer counterparts (254 nm absorbance losses are 0.2 and $0.65 \text{ L mg}^{-1} \text{ C}^{-1} \text{ h}^{-1}$ faster, respectively, for DNL and GSL winter isolates). Only the summer DOMs, active layer WEOMs, and CPT2 photobleaching rates fall within the range of IHSS reference absorbance losses, as all

the other isolates' absorbance decay rates are faster than the IHSS references (Table 3).

Optical indices also changed during irradiation, indicating non-uniform absorbance and fluorescence losses and a transformation in overall DOM composition (Fig. S9 and S10†). SUVA values decreased on average by $0.6 \pm 0.3 \text{ L mg C}^{-1} \text{ m}^{-1}$ over 24 h photolysis, implying a loss of aromaticity and chromophoric DOM. SUVA of the OCT isolate decreased most dramatically, by $0.95 \text{ L mg C}^{-1} \text{ m}^{-1}$. While a positive relationship between SUVA and $k_{\text{obs},254}$ occurred for DOM isolates ($R^2 = 0.53$), this was not the case for the WEOM isolates and neither set of isolates had a relationship between SUVA or Al : Ar and $k_{\text{obs},254}$ that was significantly non-zero.

Influence of sensitizers and quenchers on photoreactivity

Iron(III) was added to reconstituted isolate solutions to probe for the ability of the DOM and WEOM isolates to promote photo-Fenton processes,^{65–67} whereby production of ·OH, a non-selective radical, may enhance photobleaching of DOM isolates in the presence of sunlight. Photoreduction of Fe(III) to Fe(II) reacts with photoproduced hydrogen peroxide to form ·OH (eqn (3)):



Given the high total dissolved iron present in sampled waters (Table 1), photo-Fenton processes may be extremely relevant for the phototransformation of DOM in surface waters. Most of the DOM and WEOM isolates exhibit enhanced rates of absorbance loss at 254 nm upon the addition of $20 \mu\text{M}$ Fe(III) ($1116.9 \mu\text{g L}^{-1}$), except for OCT and CPT1 (Table 3 and Fig. 4), consistent with findings of hydroxyl radical production in Arctic surface waters.³² The WEOM isolates were not especially vulnerable to photo-Fenton processes, which may indicate compositional differences between mobilised thermokarst and permafrost DOM.

Enhanced chromophore decay upon iron addition is consistent across 240–450 nm (Fig. S5†) for the DOM isolates, except for DNL summer and OCT. The winter DOM isolates and GSRW show the most enhancement of absorbance loss at 254 nm, consistent across all wavelengths. In contrast, most of the soil WEOM isolates only exhibit enhancement of absorbance loss at lower wavelengths (Fig. S6†), but at wavelengths above 275–290 nm, iron addition slows absorbance loss. Slower loss rates could be indicative of precipitation of Fe(III), as experiments were not anoxic and occurred at circumneutral pH, as observed in studied lakes (Table 1); however, no precipitate formation was observed. Differing levels of enhancement or retardation of absorbance loss could then be attributed to either efficiency of H_2O_2 photoproduction, and/or the ability to stabilise iron in solution.

Methanol was utilised to broadly scavenge radicals formed during photobleaching.⁶⁸ In most cases, the rate of absorbance loss at 254 nm upon addition of methanol ranged from $0.1\text{--}1.2 \times 10^{-3} \text{ L mg C}^{-1} \text{ h}^{-1}$ slower (Table 3 and Fig. 4), indicating that a portion of observed photobleaching was due to radical production. In general, decreased rates of absorbance loss were



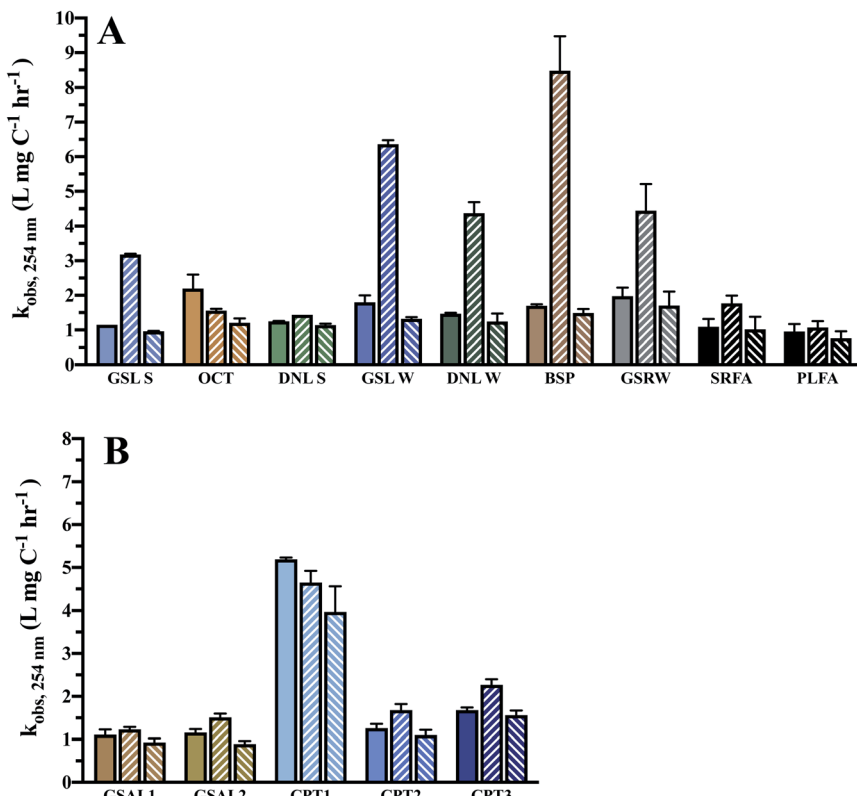


Fig. 4 Observed pseudo first order absorbance loss rates at 254 nm in 10 mg C L^{-1} reconstituted PPL and reference isolate solutions, corrected for light screening and normalised to DOC concentration. Solid fill = DOM only. Upward slanted fill = DOM + $20 \mu\text{m}$ Fe. Downward slanted fill = DOM + $12.5 \mu\text{m}$ MeOH. (A) Surface water DOM isolates. (B) WEOM isolates.

consistent across wavelengths (Fig. S5 and S6[†]). No significant difference in absorbance loss rates were observed for several isolates (Table 3), suggesting either minimal photochemical radical production or the prevalence of a quenching mechanism that did not lead to absorbance losses.

2,4,6-Trimethylphenol (TMP) degradation rates may then be indicative of either triplet excited state reactivity or singlet oxygen production.^{36,68,69} Triplet excited state DOM (${}^3\text{DOM}^*$) has been shown to react efficiently with phenolic, anilinic, and sulfidic groups.⁶⁹ TMP's one-electron oxidation potential (1.22 V) renders it susceptible to single electron transfer with several quinoid moieties, or proton coupled electron transfer through phenoxy radical formation for weaker sensitizers.⁶⁹ DOM isolates facilitated the photodegradation of TMP relatively similarly, with pseudo-first order degradation rates ranging from 0.09–0.129 $\text{L mg C}^{-1} \text{ h}^{-1}$ (Table 3). More variability was exhibited by WEOM isolates, where active layer WEOM isolates facilitated TMP photodegradation the least ($0.048 \text{ L mg C}^{-1} \text{ h}^{-1}$ on average), and CPT1 facilitating TMP photodegradation the most. Except for the active layer WEOM isolates, all isolates facilitated TMP photodegradation faster than the IHSS isolates.

Discussion

This study compared several lakes with differing thermokarst activity within a single watershed underlain by discontinuous

original yedoma or refrozen permafrost, along with WEOM isolates from permafrost and active layer soils within the watershed. Goldstream Valley is classified as yedoma permafrost. This is the same classification as yedoma permafrost in northeast Siberia, as it is defined as ice-supersaturated loess deposits that formed syngenetically in unglaciated regions of Eurasia and North America during the Late Pleistocene.^{3,5} The vast majority of yedoma permafrost occurs in northeast Siberia; however, there are pockets of it in Alaska.^{3,5} In both regions, organic carbon contents are usually less than 3%. In both regions, yedoma can be tens of meters thick.^{3,5} Unlike Siberia, where most of the yedoma occurs in the zone of continuous permafrost, the Goldstream Valley study sites are characterised by discontinuous permafrost. This means that there is a higher likelihood of surface and groundwater exchange. DOM isolates from winter sampling of active thermokarst lakes, where active layer inputs are minimal, revealed elevated photobleaching ability compared to summer isolates from the same lakes, more similar to the youngest sampled permafrost isolate than older or active layer isolates, suggesting that some permafrost thaw may enhance photo-reactivity of DOM in lakes underlain by thawing yedoma. Permafrost WEOM isolates and several DOM isolates from this watershed were found to be both photolabile and photo-reactive, even though reactivity could not necessarily be directly tied to permafrost input.



Composition of DOM and WEOM isolates

NMR spectral features of permafrost WEOM isolates appear distinct from both the active layer and surface water isolates. The carbohydrate signal (O-alkyl region, 60–90 ppm, and anomeric C peak, 110 ppm)⁶³ in the permafrost WEOM isolates is increasingly intense with older soil ¹⁴C age. Pautler *et al.*⁷⁰ associated increased carbohydrate abundance with undisturbed permafrost in the Canadian High Arctic, suggesting that depleted carbohydrate levels in exposed permafrost through active layer detachments, even recent detachments, stimulate microbial activity and degradation through release of organic material. While biolability was not measured on the WEOM isolates in this study, the permafrost WEOM isolates may have differing ability to stimulate microbial activity due to their varied carbohydrate content.

Permafrost WEOM isolates were not necessarily less aromatic than active layer isolates (Table S1†), contrasting reports elsewhere characterizing permafrost thaw waters.^{21,22,71} This elevated aromaticity may be a feature of yedoma specifically, bias of the isolation method, or a result of active layer being sampled at a different location within the watershed. The PPL isolation method likely biases toward capturing the most light-absorbing fraction of the carbon pool, preferentially extracting the chromophoric DOM and a more efficient capture of aromatic functional groups compared to natural abundance.⁴⁹ Nearly all isolation methods contain some selectivity bias, however, PPL sorbents are less biased toward aromatics than for example, XAD-8 resins.⁷² Carbon recoveries for PPL extractions are within typical ranges,⁴⁹ capturing a majority but not all of the carbon pool (>60%), which somewhat limits the representativity compared to what may be present in natural interstitial thaw waters. Other differences from the leaching process compared to a natural thaw are potentially temperature, time of water contact, and the volumes that were required in order to capture DOM by PPL isolation. However, all soil leachates were isolated according to the same conditions and method, and thus any bias would be expected to be systematic, and relative differences between samples would still be observed. A supplemental leaching study was performed on a different permafrost soil core sampled from just a few miles outside this watershed, in the Bonanza Creek Experimental Forest, to identify how representative the leached organic matter is relative to the entire carbon pool from those soils, and is detailed further in the ESI.† In general, PPL extraction from leached soils as described above captured a representative fraction of the soil organic carbon pool with the exception of aliphatics, which were largely retained on the soils even after leaching (Fig. S12†).

NMR characterisation of relative aromaticity (Al : Ar)⁷³ identified permafrost WEOM isolates as having composition intermediate of that of the DOM and active layer WEOM isolates. However, optical indices such as SUVA, S_R , and FI, which have been observed to correlate with aromaticity of DOM,^{74–77} revealed few trends (Fig. S11†). For example, CPT1's relative O-alkyl and aromatic contributions more closely resemble that of GSRW than surface water DOM isolates. GSRW was sampled

well below the active layer, so it may be possible that some of the carbon pool of this well is from deeper thawing permafrost, though more extensive sampling and characterisation is needed to determine if that is actually the case. In some cases, trends could be observed amongst various optical properties and Al : Ar, with significantly non-zero slopes for HIX, BIX, and freshness ($\beta : \alpha$), all fluorescent measurements. HIX trended negatively with Al : Ar, which is consistent with the idea that higher aromaticity indicates more humification.⁷⁷ BIX and $\beta : \alpha$ both trended positively with Al : Ar, associating less processed material with higher aliphatic content, and largely placing the permafrost WEOM isolates between the DOM isolates and the active layer WEOM isolates in terms of composition. While SUVA has been reported to correlate with aromaticity,⁷⁶ the slope in this dataset could not be determined to be non-zero.

Several of these measurements revealed a disparate pattern of trends, despite sharing commonalities in their methods. For example, FI revealed no significant trend with Al : Ar, even though BIX and freshness did, and they share similar excitation wavelength ($\lambda = 370$ or 380 nm), and are separated by as little as 10 nm in emission wavelengths. A relationship was found with Al : Ar and HIX, which shares the same excitation wavelength ($\lambda = 254$ nm) as the reported SUVA measurement, but no such trend was found with SUVA. Fundamentally the optical properties represent but a subset of organic pool (and fluorescence an even smaller subset) compared to the nearly quantitative NMR method. This could indicate a more significant presence of overlapping anomeric carbon signal such as that from carbohydrates, convoluting the determination of total aromaticity by NMR,⁶³ and that reported relative aromaticity values in this study may be higher than the actual aromatic content of the isolates. A lack of trends could also be due to the fact that the dataset is relatively small.

Permafrost WEOM is photolabile and photoreactive

Permafrost WEOM isolates are both photoreactive and photolabile. However, while the permafrost WEOM isolates were on average more photolabile than the active layer WEOM or the DOM isolates, the difference between DOC losses or AQYs was not statistically different by a student *t*-test ($p > 0.05$), likely influenced by the large variance and small sample sizes. Most strikingly, the youngest permafrost WEOM isolate, CPT1, exhibits the highest aromaticity, photolability, absorbance loss rate, and TMP photodegradation rate (Table 3). Significant photoreactivity has been observed in DOM from the Alaskan Arctic, where in headwater streams carrying DOM with less prior light exposure, sunlight was responsible for over 90% of carbon processing.²⁹ However, in other Arctic systems little photodegradability has been observed,³⁰ and limited data exists on yedoma DOM photoreactivity. Stubbins *et al.*²² found that permafrost thaw waters from near the Kolyma River, Russia was not very photolabile, observing DOC losses over 30 d in a solar simulator <5%. The increased photolability observed in this study could be due to the PPL isolation method preferentially extracting the chromophoric pool of DOM.⁴⁹ Though, even if chromophores were preferentially extracted through the



isolation process, the observed results still suggest that their sampled waters and leachates are both photoreactive and photolabile.

Integrating from 290–400 nm, AQYs are on the order of 10^{-4} to 10^{-5} , showing that these carbon pools are indeed photoreactive. As AQYs were calculated over a large range of wavelengths, it is possible that DOC photomineralisation also occurred from chromophores absorbing light outside these wavelength bounds, and thus the AQYs presented may represent a high estimate. However, these values are a similar order of magnitude as those reported by Cory *et al.* for thermokarst soils in the Arctic over a slightly narrower wavelength range (305–395 nm).³¹ Apparent quantum yields were also highest for CPT1, indicating that the high DOC losses were not simply from higher rates of light absorption for this particular isolate, which is corroborated by the lower SUVA value for CPT1 compared to the other permafrost WEOM isolates.

The small change in SUVA CPT1 suggests DOC losses included organic carbon not absorbing at 254 nm. However, there was faster absorbance loss at this wavelength for CPT1 than the other isolates. Some of the DOC loss thus could have also resulted after transformation of chromophores absorbing at 254 nm. Stubbins *et al.*²² similarly observed absorbance losses from photoirradiation of yedoma permafrost thaw waters that were considered to be associated with coloured aromatic moieties. Discrepancies between absorbance loss rates and SUVA decreases may reflect the degree of chromophore transformation, as differing chromophores will exhibit differing molar absorptivities, which may alter both absorbance loss rates and SUVA.

Both absorbance and fluorescence for all isolates decreased upon exposure to photoirradiation, however CPT1 did not stand out from the other WEOM isolates with respect to other optical indices, convoluting this observation somewhat. S_R and E2 : E3 increased with photoirradiation, corroborating a loss of aromaticity consistent with other reported observations upon photoexposure,^{77,78} while FI, BIX, and $\beta : \alpha$ decreased, indicating compositional shifts in both the chromophoric and fluorophoric pools (Fig. S10†). However, FI decreases upon photoirradiation for both DOM and WEOM isolates instead of increasing with a loss of aromaticity (Fig. S9 and S10†), contrasting with the SUVA results above, and similar to previous studies by Cory *et al.*^{31,79} While FI has been negatively correlated with aromaticity, multiple fluorescent components may give rise to detected signal in the FI region of an excitation-emission matrix (EEM).^{75,77,80} In addition, a change in fluorescence or absorbance at any single wavelength could include the formation of new chromophores or fluorophores, partly offsetting the loss of others. While net decay of fluorescence (Fig. S7 and S8†) and absorbance (Fig. S4†) occurs, optical index changes (Fig. S9 and S10†) indicate these losses are not uniform across spectra. At least some selective transformation occurs upon photoexposure, despite the fact that no significant trends (slope significantly non-zero by linear regression) between photo-bleaching rates, photolability, and AI : Ar could be identified. More studies would need to be done to ascertain which part of

the chromophoric pool is undergoing transformation *vs.* mineralisation.

All permafrost WEOM isolates facilitate the photo-degradation of TMP more dramatically than the active layer WEOM isolates (Table 3), corroborated by their lower SUVA values and despite not necessarily having a lower relative aromaticity. Differences in TMP reactivity may be due to differences in $^3\text{DOM}^*$ species or self-quenching by the higher antioxidant capacity of quinoid groups.⁸¹ For example, a variety of anilinic and phenolic compounds react with $^3\text{DOM}^*$, and are reported to have more inhibition efficiency by more aliphatic Pony Lake fulvic acid (PLFA) than Suwannee River fulvic acid (SRFA).⁸² The slower observed TMP photodegradation rates in the presence of the active layer WEOM isolates (GSAL1, GSAL2) are thus consistent with these observations. TMP degradation with isolated DOM from a North Slope river in Alaska reported TMP degradation rates ranging from 0.075 to 0.135 L mg C⁻¹ h⁻¹,³⁶ encompassing similar rates as for the DOM isolates in this study (Table 3). Quantifying the antioxidant capacity of the permafrost WEOM isolates may also help to parse out the full nature of triplet excited state reactivity of these DOM pools.

Thermokarst influence on DOM characterisation

Like the WEOM isolates, nearly all DOM isolates were photolabile, with the exception of the summer DNL isolate, spanning AQYs on the order of $1.1\text{--}2.9 \times 10^{-4}$ over 290–400 nm (Table 3). AQYs are not statistically different from that measured from the IHSS isolates. Photoreactivity was broadly similar in terms of observed absorbance loss rates at 254 nm, facilitation of TMP photodegradation, and shifts in SUVA upon photoirradiation, making it difficult to discern potential influence of thermokarst activity or original *vs.* refrozen yedoma on DOM behaviour.

In DOM isolate solutions sourced from waters with either thermokarst activity and/or minimal active layer influence (winter DOM isolates and GSRW), the addition of Fe(III) resulted in increased absorbance losses from 250–275 nm, where small aromatics absorb. This enhancement of absorbance loss upon Fe(III) addition suggests that thermokarst-influenced DOM may be more efficient at promoting photo-Fenton reactions, as iron has been correlated to aromatics in other permafrost containing systems.⁸³ However, CPT1, the most aromatic WEOM isolate by ^{13}C multiCP-MAS NMR, did not increase in photobleaching rate upon iron addition, and instead its absorbance loss slowed. This may reflect its ability for peroxide production or to associate with iron in solution, though neither of these were measured in this study. The winter BSP isolate was most responsive to iron addition, suggesting it may better stabilise iron to promote photo-Fenton pathways, potentially through better association with DOM ligands, indicating some distinct compositional differences, even if those compositional differences were not directly measured in this study.

Few ^{13}C NMR data from other Arctic or sub-Arctic lakes are reported to compare the representativity of these samples. Toolik Lake (northern Alaska, continuous non-yedoma



permafrost) had a reported aliphatic (0–60 ppm) to aromatic ratio (90–160 ppm) (Al : Ar)⁷³ of 1.94 for XAD-8 isolate,³⁴ which is slightly lower than the average Al : Ar for the PPL isolates from the lakes in this study (2.09 ± 0.32), indicating slightly more aliphatic content relative to aromatic content than Toolik Lake. However, it should be noted that the isolation method, ecosystem, permafrost type, and NMR methods reported for Toolik Lake are all slightly different from this study,³⁶ and so a direct comparison may not be completely equivalent. GSL, OCT, and BSP had the lowest Al : Ar ratios, indicating the highest proportion of aromatic content relative to aliphatic groups, and also represent surface waters with active thermokarst activity.

Winter DOM isolates present an opportunity to control for active layer inputs into lake waters, as the winter DOM is theoretically more cut off from direct active layer inputs, and thus would be receive inputs dominated by thawed sediments. However, as temperatures increase, the active layer may not completely freeze in winter, allowing lateral flow year-round.^{84–87} Thus, these relatively shallow lakes are not necessarily devoid of active layer source material in the winter, from either unfrozen active layer or recalcitrant materials remaining in the water column. The two winter isolates comprise similar relative contributions of functional groups, varying within just 2% of each other for any given region. This decrease in variability during wintertime could be due to restricted lateral flow through active layers, limiting input into the lakes. This also suggests that the similarity between GSL's summer and winter isolate composition, including aromaticity, may be from significant contributions other than the active layer, such as thermokarst activity.

DNL, in contrast, does not have active thermokarst activity,³⁷ and seasonal variation may instead reflect differences solely from surface *vs.* ground water inputs. This is supported by the fact that the DOM isolates from active thermokarst systems were more similar in composition and rates of absorbance losses to winter DOM than summer non-thermokarst DOM isolates, and also exhibited more similarity to the permafrost WEOM isolates than the active layer WEOM isolates. For example, the GSL isolates appear to have similar composition between the summer and winter samples in terms of both optical and NMR composition, despite large differences in whole water carbon and iron concentration. The GSL summer isolate was less susceptible to absorbance decay than its winter isolate, but otherwise, characterisation did not yield strong differences between DOM isolates from either winter or summer (Fig. S11†), with the exception of DNL summer standing out with a higher Al : Ar ratio.

Permafrost underlain soils and sediments around and under GSL may have the potential to enhance lateral contributions through the active layer by limiting subsurface drainage to ground waters. In contrast, winter DOM isolates may have more compositional similarities to the permafrost WEOM isolates due to being more isolated from active layer inputs, regardless of thermokarst activity. However, such a relationship cannot be conclusive, as DNL's winter isolate also had increased photobleaching ability, yet has no observed thermokarst activity.

Increased photoreactivity in winter isolates may also reflect a lack of *in situ* photodegradation during dark winter months compared to summer, where DOM may have already undergone some photobleaching at time of collection. This could be reflected in the fact that photobleaching of the winter isolates was significantly enhanced upon addition of iron, presumably through reaction with hydroxyl radical, which has been shown to correlate with photo exposure.³² However, there is no requisite observed trend in photolability that one would expect to be consistent with this hypothesis, which may be complicated by transformation (biotic and abiotic) of permafrost material prior to transport to the water column or other mechanisms. For example, several studies have reported biolability and characterised the biotransformation of permafrost material.^{6,19,70,88–90} Morgalev *et al.*⁹¹ found that the active layer/permafrost interface displayed enhanced metabolic activity in regions of discontinuous permafrost peatlands of western Siberia. Thus, any contribution of permafrost derived material to the studied thermokarst lakes may in fact be quite different in composition compared to what was characterised from the permafrost tunnel WEOM isolates. Quantifying the flux of ground waters in contact with permafrost thaw to examine the extent of thermokarst contribution to these lakes may help shed to some light on this process.

Conclusions

Overall, there were few observations that clearly separated waters with thermokarst activity from original yedoma permafrost and permafrost soils from waters of refrozen deposits or active layer. It should be noted that NMR, optical properties, and to some extent photobleaching ability are by definition bulk properties of a particular DOM pool. However, all isolates were photoreactive, and several of the permafrost influenced DOM and permafrost WEOM isolates were especially photolabile, with relatively high AQYs for DOC loss. More specific compositional analyses or broader sampling may shed light on any compositional dependence of DOM reactivity from active thermokarst inputs. Expanding the data set to develop more statistical robustness between lakes of varied thermokarst activity and exploring inter- and intra-annual variability⁹² and the impact of differing hydrological regimes will also help to resolve the extent of thermokarst influence on surface water DOM. In addition, sampling sediment pore waters in locations of active thaw rather than solely lake waters, which represent a combination of all the DOM sources to the water column, may help to identify and quantify the specific compositional influences of thermokarst activity on the surface water carbon pool. As temperatures increase, the release of permafrost organic matter is inevitable, and thus, obtaining a holistic understanding of the variability in functional group composition and photoreactivity will allow for a better understanding of how DOM's ecological role will be impacted by permafrost thaw in the coming decades.

Conflicts of interest

There are no conflicts to declare.



Acknowledgements

Funding for this project was supplied by the Alaska Space Grant Graduate Research Fellowship, National Institutes for Water Resources Graduate Student-Lead Proposal through the UAF Water & Environmental Research Center and the U.S. Department of Interior – Geological Survey, and the University of Alaska Fairbanks Undergraduate Research and Scholarly Activity program. Access to residential lakes were obtained through permissions supplied by NSF ARCSS-1500931. Research reported in this publication was supported by an Institutional Development Award (IDeA) from the National Institute of General Medical Sciences of the National Institutes of Health under grant number P20GM103395. The content is solely the responsibility of the authors and does not necessarily reflect the views of the NIH. A sincere thank you to A. Liljedahl, D. Barnes, and V. Romanovsky, for allowing access and providing intellectual support. Thank you to A. Kholodov for assistance in collecting and obtaining active layer samples and T. Douglas at Cold Regions Research and Engineering Laboratory with the U.S.A. Army for access and sampling of the CRREL Permafrost Tunnel. Thank you to A. Simpson for providing the SPR-W5-WATERGATE pulse sequence for the Bruker NMR. Further thanks to B. Eckhardt, N. Ramos, D. Knight, N. Ruckhaus, R. Noratuk, J. Sterle, A. Mutschlechner, R. Voight, R. Davey, M. Ross and T. Harms for instrumentation and sampling assistance, and G. Quesada for actinometry assistance.

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