# Soil profile connectivity can impact microbial substrate use, affecting how soil CO<sub>2</sub> effluxes are controlled by temperature

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**Abstract:** Determining controls on the temperature sensitivity of heterotrophic soil respiration remains critical to incorporating soil-climate feedbacks into climate models. Most information on soil respiratory responses to temperature come from laboratory incubations of isolated soils, and typically subsamples of individual horizons. Inconsistencies between field and laboratory results may be explained by microbial priming supported by crosshorizon exchange of labile C or N. Such exchange is feasible in intact soil profiles, but is absent when soils are isolated from surrounding depths. Here we assess the role of soil horizon connectivity, by which we mean the degree to which horizons remain layered and associated with each another as they are in situ, on microbial C and N substrate use and its relationship to the temperature sensitivity of respiration. We accomplished this by exploring changes in C:N, soil organic matter composition (via C:N, amino acid composition and concentration, and nuclear magnetic resonance spectroscopy), and the  $\delta^{13}$ C of respiratory CO<sub>2</sub> during incubations of organic horizons collected across boreal forests in different climate regions where soil C and N composition differ. The experiments consisted of two treatments: soil incubated (1) with each organic horizon separately, and (2) as a whole organic profile, permitting cross-horizon exchange of substrates during the incubation. The soils were incubated at 5°C and 15°C for over 430 days. Enhanced microbial use of labile C-rich, but not N-rich, substrates were responsible for enhanced, whole-horizon respiratory responses to temperature relative to individual soil horizons. This impact of a labile C priming mechanism was most emergent in soils from the warmer region, consistent with these soils' lower C bioreactivity relative to soils from the colder region. Specifically, cross-horizon exchange within whole soil profiles prompted increases in mineralization of carbohydrates and more <sup>13</sup>C-enriched substrates and increased soil respiratory responses to warming relative to soil horizons incubated in isolation. These findings highlight that soil horizon connectivity can impact microbial substrate use in ways that affect how soil effluxes of CO<sub>2</sub> are controlled by temperature. The degree to which this mechanism exerts itself in other soils remains unknown, but these results highlight the importance of understanding mechanisms that operate in intact soil profiles – only rarely studied – in regulating a key soil-climate feedback.

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#### 1 Introduction

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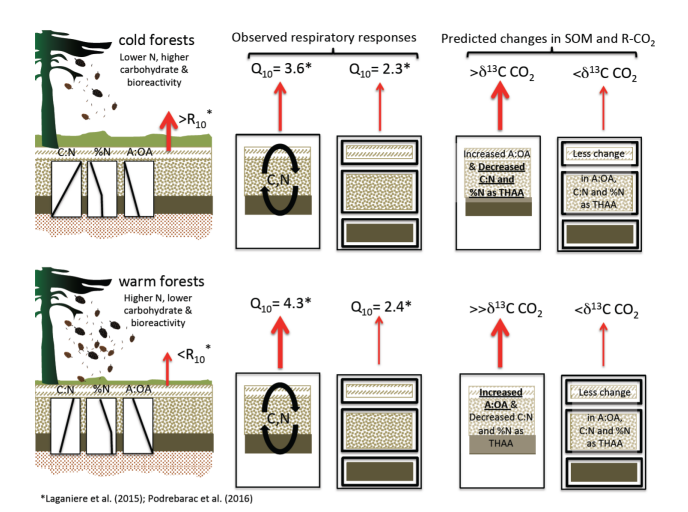
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Increased understanding of the controls on soil respiration, a globally significant flux of  $CO_2$  (Bond-Lamberty and Thomson, 2010; Stocker et al., 2013), and its response to temperature is required in developing Earth System Models. Global scale surveys suggest that temperature sensitivity of soil respiration is largely attributed to responses occurring at the level of the whole microbial community, with the greatest temperature sensitivities occurring in high C:N ratio, C-rich soils of high-latitude boreal and arctic ecosystems (Karhu et al., 2014). Congruent with these laboratory studies, temperature sensitivity of soil respiration from field experimental warming studies indicates that the greatest enhancement occurs in high latitude soils (Carey et al., 2016). Microbial mechanisms for these high latitude soil responses as well as differences between field and laboratory studies may lie within differences in what soil horizon or collection of horizons are assessed. For example, temperature sensitivity of soil respiration can increase with depth in association with a reduction in soil organic matter bioreactivity, consistent with the idea that increased temperature sensitivity is associated with more slow-turnover, and perhaps higher  $E_a$ , substrates (Conant et al., 2008; Lefevre et al., 2014; Leifeld and Fuhrer, 2005). In boreal forest soils warming appears to enhance bacterial use of labile surface soil C sources and fungal use of deeper slower-turnover soil C pools (Ziegler et al., 2013), with lower bacterial to fungal ratios associated with increases in the temperature sensitivity of soil respiration (Briones et al., 2014).

Association between soil depth or bioreactivity and temperature sensitivity of soil respiration are not ubiquitous (Fang et al., 2005; Liski et al., 1999), nor have these laboratory findings always been supported by *in situ* whole-profile investigations of respiration that reveal consistent heterotrophic respiration of relatively young soil C and elevated Q<sub>10</sub> of soil respiration to 100 cm (Hicks Pries et al., 2017). In fact, enhanced temperature responses of soil respiration observed within whole soil profiles suggests deeper soil profiles contribute significantly to the temperature response of soil respiration (Hicks Pries et al., 2017). This raises questions regarding soil profile attributes, such as root and dissolved organic matter inputs or microbial substrate and nutrient exchange, that may control respiratory responses not revealed in commonly used laboratory experiments where horizons are isolated.

Interactions among soil horizons may be important features driving temperature responses of the C rich organic layers common in boreal forests. For example, the temperature sensitivity of soil respiration was up to 30% higher in whole boreal forest organic profiles from a warmer versus colder climate despite the fact that the temperature sensitivity of soil respiration from the individual organic horizons isolated from those same organic profiles did not differ by climate (Laganière et al., 2015; Podrebarac et al. 2016). Differences in SOC and SON composition between these soil's climate regions are consistent with the differences in the bioreactivity of these soils (left Fig. 1; Laganière et al., 2015) but the differences in temperature responses of respiration consistent with bioreactivity was only realized within the whole soil profiles (center Fig. 1; Podrebarac et al., 2016). Here the temperature response of the whole organic profile respiration was over 50% greater than the respiration from the same soils incubated as isolated horizons (middle Fig. 1). This suggests microbial access to the different C or N substrates available among horizons when soils were incubated as a whole organic profile can regulate soil respiratory responses to temperature perhaps in analogous ways observed for root exudates (Zhu and Cheng, 2011). Specifically, labile substrates from the less degraded L horizon (same as Oi in U.S. soil classification)



**Figure 1.** Conceptual figure depicting differences in relatively cold and warmer boreal forest organic horizon composition, including bioreactivity ( $R_{10}$ ) given as C-normalized soil respiration at  $10^{\circ}$ C (left column), previously reported temperature sensitivities of heterotrophic respiration ( $Q_{10}$ ) in two incubation designs ("whole" vs. "isolated", left to right in center column; Podrebarac et al. 2016), and hypothesized responses of incubated soils with and without interlayer exchange of microbial C and N substrates (right column). The carbon to nitrogen (C:N), nitrogen content as %N and the alkyl to O-alkyl ratio (A:OA) reflect decreasing relative content of carbohydrate with depth and in the warm relative to colder climate forest sites and decreasing N content with depth and in the colder sites (left side). Enhanced temperature sensitivities in the "whole" incubations relative to the "isolated" prompted hypotheses that (1) soils in the "whole" incubation, especially from the warmer forests, would experience greater relative losses of carbohydrates signified by more  $^{13}$ C enriched respired CO<sub>2</sub>, increased ratio of alkyl-C to O-alkyl-C and reductions in C:N relative to the isolated soil incubations signifying a greater use of labile C in support of the enhanced temperature sensitivity of respiration; (2) soils in the "whole" incubation, especially from the colder forests, would experience a reduction in %N as total hydrolyzable amino acids signifying a greater use of soil organic N in support of the enhanced temperature response of soil respiration.

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may enhance the use of more complex, high  $E_a$  substrates found in the lower F or H horizons (Oa and Oe in U.S. soil classification) via microbial priming (Cheng et al., 2014; Finzi et al., 2015; Fontaine et al., 2007; 2011).

Understanding the presence or absence of cross-horizon substrate exchange and use will help determine the mechanisms driving SOM compositional changes with temperature, as well as those governing the temperature sensitivity of soil respiration. Differences in microbial access to distinct substrates (e.g. labile C or N-rich compounds) may affect respiratory responses to temperature (Billings and Ballantyne, 2013) given that labile inputs such as rhizosphere C can provide a significant control on soil C and N cycling (Cheng et al., 2014; Finzi et al.,

2015). For example, labile substrates mixed into soils can enhance decomposition of extant soil organic matter and the temperature response of soil respiration (Di Lonardo et al., 2019; Wang et al., 2016; Wild et al., 2016). More labile substrates are typically found in surficial soil horizons, particularly in boreal forest podzols where thick organic horizons are characterized by a surface litter horizon. The transport of these labile substrates to deeper horizons, where slower-turnover organic matter is present, occurs via mobilization of dissolved organic matter (Kaiser and Kalbitz, 2012; Kalbitz and Kaiser, 2008), or microbial use of neighboring horizons' substrates via hyphae or mycelia (Dijkstra et al., 2013; Fontaine et al., 2011). More fungal dominated communities in surface horizons may access N-rich, higher  $E_a$  substrates from deeper soil horizons, a mechanism found to support priming effects in some soils (Li et al., 2017). In forest soils, increased N availability can enhance substrate use by bacteria relative to fungi and Actinobacteria, and can suppress soil respiration rates (Butnor et al., 2003; Ziegler and Billings 2011; Maier and Kress, 2000). Given these N-driven alterations in microbial strategies and the relatively low bacterial to fungal ratios in boreal forest soils (Hogberg and Hogberg 2002), often associated with increased temperature sensitivity of microbial activity (Briones et al., 2014), we may expect increased N to enhance temperature sensitivity of soil respiration in boreal forest soils. However, most studies exploring relevant issues leverage isolated soil layers to address the question via incubations, and when horizons are separated the exchange and availability of labile C and N substrates across horizons is inhibited, potentially altering microbial decomposition processes and their response to temperature.

By following soil C and N use in soils from a boreal forest transect where SOC and SON composition differ both by depth and climate region (left side Fig. 1), we addressed two hypotheses describing how whole soil profile connectivity, or interlayering, affects the temperature response of soil respiration (center Fig. 1). Firstly, we hypothesized that priming of soil respiration and C loss from slower turnover F and H horizon soils is induced by microbial use of more labile C from the overlying L horizon and greater N availability from deeper F and H horizons, which combine to enhance respiratory responses to increased temperature. Secondly, we hypothesize that the evidence for a labile C priming mechanism is most emergent within the warmer region soils given these soils' lower SOC bioreactivity relative to soils from the cold region (right side Fig. 1). However, we also anticipate that if the priming mechanism is supported by cross-horizon N availability it would be most emergent within the colder region soils, where SON availability is lower relative to the warmer region. By assessing changes in soil organic matter composition via C:N, nuclear magnetic resonance spectroscopy (NMR), and amino acid profiling, as well as  $\delta^{13}$ C of respiratory CO<sub>2</sub>, we investigated whether the elevated Q<sub>10</sub> of soil respiration within the whole organic profiles, as observed in Podrebarac et al. (2016), was associated with increased use of more labile soil C or N relative to horizons incubated individually. Support for these hypotheses would suggest a potentially important means by which the temperature sensitivity of soil microbes' CO<sub>2</sub> release may be governed that is rarely explored.

#### 2 Methods

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### 2.1 Study Area

This study was conducted using soils from the two end-member climate regions of the Newfoundland and Labrador Boreal Ecosystem Latitudinal Transect (NL-BELT) where mean annual temperature differs by approximately 5.2°C and mean annual precipitation is 1074 and 1505 mm, for the highest latitude region (hereafter referred to as the cold region) and lowest latitude region (hereafter referred to as the warm region), respectively (Cartwright and Doyles, NL weather station climate normals between 1981-2010; Environment-Canada, 2014; Table S1). Within these two climate regions, three mesic forest sites dominated by mature balsam fir stands (*Abies balsamea* L.) and underlain by humo-ferric podzols were established. The forest transect sites used here, established in 2011, provide a unique opportunity to determine the impact of climate history of soil organic matter cycling and the fate of *in-situ* reservoirs (Ziegler et al. 2017).

#### 2.2 Soil sampling

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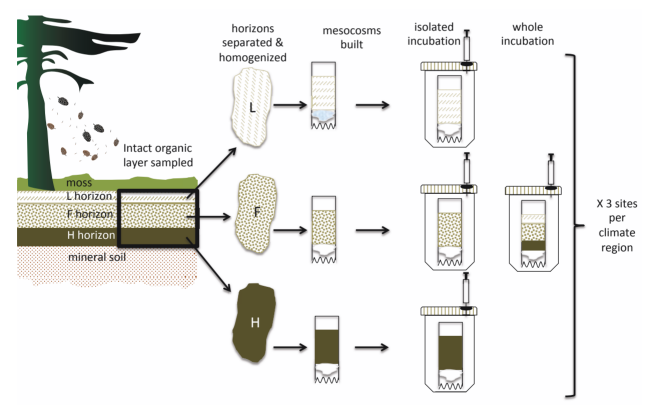
Soil was collected in October 2011 as described in Laganière et al. (2015). Briefly, three soil sampling plots each with a diameter of 10 m were established within each site. A 20 x 20 cm intact 'cake' of the whole profile including the L, F, and H horizon was collected using a sharp knife and a trowel within each soil sampling plot. The Canadian Soil Classification of L, F, and H horizons is synonymous with O<sub>i</sub>, O<sub>e</sub> and O<sub>a</sub> sub-horizons, respectively, in the U.S. Soils Classification and collectively hereafter will be referred to as the 'organic profile' or the LFH. Although they are technically horizons of the O layer in the Canadian Soils Classification, the L, F and H will be referred to as sub-horizons from here on in order to make it easier to associate with the more commonly used U.S. Soils Classification. Mean LFH depth is  $8.1 \pm 0.3$  cm (L  $1.0 \pm 0.0$  cm, F  $6.1 \pm 0.3$  cm, H  $1.0 \pm 0.0$  cm) and  $8.4 \pm 0.4$ cm  $(1.0 \pm 0.0 \text{ cm}, 6.3 \pm 0.4 \text{ cm}, 1.1 \pm 0.1 \text{ cm})$  for the L, F and H, respectively) for the cold and warm regions, respectively. On site, half of the intact organic profile was separated by hand into the L, F, and H horizons and placed in a cooler while field sampling. Samples were transported to the laboratory and stored at 5°C until analysis and experimental set-up prior to the incubation as described in Laganière et al. (2015) and Podrebarac et al. (2016). Briefly in preparation for the incubation, the L was homogenized by cutting the large soil pieces into 1 cm lengths; whereas the F and H were homogenized separately by soil sampling plot through the use of a 6mm sieve. If present, large roots (>6 mm) were removed. For the incubation, the homogenized soils were pooled by site to yield three site composite samples per region.

### 2.3 Soil incubations

As described in Laganière et al. (2015) and Podrebarac et al. (2016), the microcosms consist of a plastic tube (5 cm diameter x 15 cm height) with an acid-washed glass wool plug and V-notches on one end to enable aeration and drainage of the soil samples (Fig. 2). The selected soil horizon(s) are placed on the glass wool horizon inside the plastic tube and the tube placed in a 1 L mason jar with these microcosms allowed to equilibrate for 1 week at 5 °C to reduce handling effects (Robertson et al., 1999). Replicate microcosms were then incubated at 5 °C and 15 °C, the average range in temperatures across the study sites in April-August (growing season). To maintain gravimetric water holding capacity at approximately 70% for between 438 and 482 days, depending on the experiment, the soil moisture was adjusted weekly by adding water to the top of the soil core, based on mass loss measured for each

microcosm. Laganière et al. (2015) incubated the L, F, and H horizons separately to determine the soil bioreactivity and Q<sub>10</sub> of soil respiration specific to each separate horizon, an approach hereafter referred to as the 'isolated' experiment. In Podrebarac et al. (2016), the organic profile was reconstructed in the same proportions by mass as found in-situ using the same soils used in the isolated experimental treatment; an approach hereafter referred to as 'whole' experiment (Fig. 2). Total soil mass in grams dry weight equivalent was the same across all microcosms (~11 gdw) and between the two experimental treatments. Actual mass of the individual L, F and H horizons in the whole treatment totaling ~11 gdw was based upon the average proportion of each horizon measured from across sites in all regions (Laganiere et al. 2015). In a previous study, intact soil cores of the whole organic horizon were collected at the same time as those used to obtain the homogenized separated horizons. These were incubated to compare respiratory responses among intact profiles, the reconstructed whole profiles and those incubated as isolated horizons. This confirmed similar soil respiratory responses to temperature between the intact and reconstructed whole profiles indicating that the homogenized soils used in this study behaved in a similar way as intact soil profiles from these forest sites (Podrebarac et al. 2016). Furthermore, the overall climate region differences in temperature sensitivity of forest soil respiration observed in the laboratory incubations of the intact and reconstructed whole profiles were found to be in line with field observations of total soil respiration in these forest sites (Podrebarac et al. 2016)

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**Figure 2.** Conceptual figure depicting the experimental design. Organic horizon soils were collected as an intact 20X20 cm layer from each of three field sites in each climate region and separated into the separate L, F and H (equivalent to the Oi, Oe, and Oa in U.S. soil classification) horizons. After homogenization, soil was used as illustrated to set up two sets of incubations. The treatment labeled "isolated" incubated each soil horizon in isolation. The treatment labeled "whole" incubated the L, F and H horizons together, with their masses reflecting the average proportional masses observed at the sites.

The temperature sensitivity of the soil respiration rates were taken directly from Laganière et al. (2015) and Podrebarac et al. (2016) and reported as Q<sub>10</sub> simply calculated as the ratio of cumulative respiration over the course of the entire incubation at 15°C over that measured in the 5°C incubations. This simple approach was chosen because it avoids possible bias introduced when fitting data to a least-squares regression line (Sierra, 2012) enabling us to illustrate a direct comparison in the temperature sensitivity of respiration among treatments with which to place the soil compositional results from this study.

The whole experiment was incubated in triplicate by site over the 438+ day incubation and compared with results of Laganière et al. (2015). The summation of the results of the isolated experiment using the same proportions of L, F and H horizons as in the whole experimental treatment were expected to represent the organic profile without the cross-horizon exchange that exists naturally *in-situ* in whole horizons. This summed approach is hereafter referred to as the 'predicted whole' experiment. The predicted whole values for a given soil metric, such as respiration or %C was derived using Eq. (1),

$$X_{predicted\ whole} = X_L \frac{(M_L)}{(M_{whole})} + X_F \frac{(M_F)}{(M_{whole})} + X_H \frac{(M_H)}{(M_{whole})}$$
(1)

where  $X_{predicted\ whole}$  represents the value of X (e.g. %C, respiration rate, C:N) for the whole experiment soil treatment as predicted from measurements of X from the isolated horizons reported in Laganière et al. (2015) or this current study. The "M" refers to the total mass of dry soil with subscripts designating the isolated sub-horizon (L, F, or H) incubated.  $M_{whole}$  represents the total soil dry mass for the incubated microcosm of the whole experiment. Using this equation we generated a predicted measurement of each soil metric (e.g. soil %N, C:N, total hydrolyzable amino acid content, ratio of alkyl-C to O-alkyl-C) for the whole profile without cross-horizon exchange with which to make direct comparisons with values observed for the actual or measured whole LFH profiles. As opposed to the instantaneous respiration rates, sampling for soil chemical composition was destructive and soil C and N composition was not expected to change enough to warrant sampling for soil composition throughout the incubation experiment. Therefore we limited sampling for soil composition to the end of the incubation experiment. However, we do acknowledge that the most significant differences in respiratory responses to soil temperature among treatments and sites were observed before the end of the incubation period (Fig. S3). Thus, the experimental and site effects observed in soil composition may be conservative.

### 2.4 Soil respiration and the $\delta^{13}$ C of respiratory CO<sub>2</sub>

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The CO<sub>2</sub> production rate and  $\delta^{13}$ C of respiratory CO<sub>2</sub> ( $\delta^{13}$ C-CO<sub>2</sub>) measured at 6 and 3 time points, respectively, occurred over the course of the 438+ day incubation according to Laganière et al., (2015) and Podrebarac et al. (2016). Briefly, each microcosm was flushed with ambient air before being sealed with an airtight lid with a rubber septum. Prior to gas sampling a volume of N<sub>2</sub> was injected into the sealed 1 L microcosm to avoid the generation of a vacuum upon sampling. A gas tight syringe was then used to sample the initial gas sample at the same volume as the injected N<sub>2</sub>. The final gas sample was collected following the same method after 16 hours and 4 hours, respectively, for 5°C and 15°C treatments. The samples collected were injected into evacuated gas tight vials (Labco Limited, Lampeter, UK) and stored (less than 2 weeks) alongside standards until analysis for CO<sub>2</sub>

concentration using an Agilent 6890A gas chromatograph with a thermal conductivity detector (Agilent Technologies, Santa Clara, CA, USA). The  $CO_2$  production rate was calculated as the difference in the headspace  $CO_2$  concentration between final and initial gas samples per gram of initial soil C per unit of time (mg C-CO<sub>2</sub> g<sup>-1</sup> initial C h<sup>-1</sup>). In the case of the whole experiment,  $CO_2$  production was measured on day 0, 7, 42, 96, 149, 243, and 438. The  $CO_2$  production rate for the isolated experiment was measured on day 0, 7, 42, 91, 156, 245, and 482 as reported in Laganière et al. (2015). The  $\delta^{13}$ C-CO<sub>2</sub> measured on days 0, 96, and 438 and day 0, 91, and 482, respectively, for the whole and isolated experiments was determined on an Agilent 6890 gas chromatograph (Agilent Technologies, Santa Clara, CA, USA) using a Carboxen 1010 PLOT column (30m X 0.32mm X 15 mm; Sigma Aldrich) interfaced to a Delta V+ isotope ratio mass spectrometer (ThermoFinnegan). To determine the  $\delta^{13}$ C of the  $CO_2$  produced via respiration over the entire incubation a linear extrapolation through measured values over the incubation was used to obtain  $\delta^{13}$ C of respired  $CO_2$  on days not measured. These values were used to estimate the  $\delta^{13}$ C of the total cumulative  $CO_2$  over the entire incubation period using Eq. (2).

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$$\delta^{13}$$
C of total cumulative respired CO<sub>2</sub> =  $\sum_{n=1}^{6} \left( \frac{X_n - X_{n-1}}{X_6} \right) \times Y_n$  (2)

where X is the cumulative respiration (mg C-CO<sub>2</sub> g<sup>-1</sup> initial C) measured for a given sampling time point n. Y is the  $\delta^{13}$ C of respired CO<sub>2</sub> at each sampling time point n.

#### 2.5 Soil chemistry

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All initial and final soil samples were analyzed for %C, %N,  $\delta^{13}$ C,  $\delta^{15}$ N, total hydrolyzable amino acids (THAA), and relative differences in the proportion of main C functional groups via nuclear magnetic resonance spectroscopy (NMR). The O-alkyl and di-O-alkyl C proportions were tracked to assess relative changes in carbohydrate content of the soils, a labile source of C. The alkyl-C and more specifically its ratio to O-alkyl C (A:O-A) was used to assess the degradative state of soil C. The A:O-A increases with soil depth and degradative state of plant tissues or soil (Preston et al., 2009; 2000). To track the degree of SON use we followed changes in the %N as THAA and mol% glycine, given that amino acids make up the bulk of SON and exhibit significant losses relative to total soil N (Philben et al. 2016). Soil samples were air-dried and ground prior to analyses. The %C, %N,  $\delta^{13}$ C, and  $\delta^{15}$ N were analyzed with a Carlo Erba NA1500 Series II elemental analyzer (Milan, Italy) coupled to a DeltaV Plus isotope ratio mass spectrometer via a Conflo III interface (Thermo Scientific). Solid-state cross polarization magic-angle spinning (CP-MAS) experiments were performed using a Bruker AVANCE II 600 MHz with a magic-angle-spinning probe for H, C, N, and <sup>2</sup>H (MASHCCND). Samples were run at 150.96 MHz (<sup>13</sup>C) and spun at 20kHz at 298K. Experiments run for each replicate sample (n=3) were each deconvoluted using a 19component model within the 'DM fit' base software (Massiot et al., 2002). Chemical shift regions assigned to the following functional groups: alkyl-C (50-0 ppm), amine+methoxy-C (65-45 ppm), O-alkyl-C (90-65 ppm), di-Oalkyl-C (110-90 ppm), aromatic-C (145-110 ppm), carbonyl-C+amide (190-165 ppm), were expressed as % of total area resolved (Preston et al., 2009; Wilson et al., 1987).

For the THAA analyses, ground samples (5-10 mg) were added to glass hydrolysis tubes followed by an addition of 1 ml of 6 M HCl. The hydrolysis tubes were sparged with N<sub>2</sub>, sealed with Teflon-lined caps and heated to 110°C for 20 h. Each hydrolysis tube was opened, and an aliquot of the hydrolysate was dried under a stream of N<sub>2</sub> gas. Samples were then redissolved in 0.01 M HCl and norvaline was added as an internal standard. Amino acids were recovered from the hydrolysate using solid phase extraction and derivatized using the EZ:Faast kit for amino acid analysis (Phenomenex, USA). The derivatized samples were separated by gas chromatography using a Phenomenex ZB-AAA column (110–320°C at 30° min<sup>-1</sup>) and quantified with a flame ionization detector using an HP6850 gas chromatograph (Agilent Technologies, Santa Clara, CA, USA). This resulting in 15 quantified amino acids (AA) including alanine, glycine, valine, leucine, isoleucine, threonine, serine, proline, aspartic acid, hydroxyproline, glutamic acid, phenylalanine, lysine, histidine, and tyrosine. Glutamine and asparagine are converted to glutamic acid and aspartic acid, respectively, during hydrolysis and are included in the measurement of these amino acids. The THAA yield was expressed as a percentage of total soil C or N based on the total of all 15 AA according to Eq. (3),

$$THAA (\%C \ or \ N) = \sum \left(\frac{Yield_{AA}}{(C \ or \ N)}\right) \times [Wt\% \ (C \ or \ N)_{AA}]$$

$$\tag{3}$$

where  $\frac{Yield_{AA}}{C \ or \ N}$  is the C- or N- normalized yield of each AA given in mg amino acid per 100 mg C or N and Wt% (C or N)<sub>AA</sub> is the weight %C or N in the AA.

#### 2.6 Statistical Analyses

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The initial soil measures (C:N, %N of THAA, mol% glycine, %alkyl-C, alkyl-C:-O-alkyl-C ratio (A:OA), and %di-O-alkyl-C,  $\delta^{13}$ C,  $\delta^{15}$ N) were analyzed using a two-way ANOVA to test the effects of climate region, horizon, and their interaction term to quantify meaningful differences in soil properties prior to these experiments. These are expected to support previous observations of lower N, elevated carbohydrate (lower A:OA and %alkyl, higher %di-O-alkyl-C) and higher bioreactivity in the colder relative to warmer region forest soils (left side Fig. 1). Previous work investigated the impact of climate region on the soil respiratory responses to temperature (Laganière et al., 2015; Podrebarac et al., 2016). This work indicated no real difference in respiratory responses to temperature between the colder versus warmer forest soils when horizons were incubated in isolation. In contrast, and consistent with the differences in bioreactivity and soil chemical properties, a larger temperature response was observed in the warmer relative to colder forest soils when incubated as a whole organic layer (center Fig. 1). This study focuses on the mechanisms controlling those different responses, which appear dependent upon the layering of the soil horizons. As such, we focus on the impact of the intact nature of whole soil horizons on the temperature responses of soil respiration and its relationship to C and N substrate use. To investigate how microbial use of C and N components of the SOM pools relate to the previously reported Q<sub>10</sub> of soil respiration (Laganière et al., 2015; Podrebarac et al., 2016), the soil mass loss, loss of C and N, change in soil composition ( $\Delta$  of C:N,  $\delta^{13}$ C,  $\delta^{15}$ N, THAA, NMR resolved C chemistry), and the  $\delta^{13}$ C-CO<sub>2</sub> were tracked over the incubation within the (1) isolated experiment, (2) whole experiment, and (3) predicted wholse experiment based on the isolated experimental results. Given our two hypotheses, we expected to observe a more <sup>13</sup>C enriched respired CO<sub>2</sub>, greater relative losses of

carbohydrates signified by increased %alkyl and/or A:OA and decreased %di-O-Alkyl, and reductions in C:N in the whole-soil relative to the isolated soil incubations, signifying microbial use of labile C resulting in enhancement of observed temperature sensitivity of CO<sub>2</sub> release. We also expected to observe greater reduction in %N as THAA and increase in mol% glycine in the whole relative to the isolated soil incubations signifying microbial use of SON that promoted the enhanced temperature sensitivity. Finally, we anticipated that the enhanced use of labile C and SON would be most emergent within the warm and cold forests, respectively, as a result of the climate region differences in soil C and N (right side Fig. 1). The difference between the initial and final soil composition metric (given as final - initial =  $\Delta$ ) was calculated, with negative values indicating a decline in metric value over the incubation (e.g., a  $\Delta$  of -10 for soil C:N indicates that C:N decreased by 10 units). Due to large standard errors associated with some of the changes in composition, a Student's T-test was used to determine if the initial and final soil values were significantly different from each other (i.e., if  $\Delta$  was different from zero). Because of a significant effect of region on multiple response variables (see Results), we performed region-specific tests to assess mechanisms responsible for respiratory responses. For the isolated experiment, the effect of incubation temperature, horizon, and their interaction was assessed using a two-way ANOVA where levels of temperature were 5 °C and 15 °C, and horizons defined as L, F, and H within each climate region. Using both the isolated and whole horizon treatments we explored the effect of incubation temperature, experiment type (i.e., whole, isolated), and their interaction using a two-way ANOVA within climate region. Here, a significant effect of experimental type denotes some impact of cross-horizon exchange on SOM processing.

For tests in which residuals did not meet the assumptions of normality and homoscedasticity, data were log<sub>10</sub>-transformed prior to testing (Zar 1999). The Tukey's Honestly Significance Differences test was used to determine which combination of treatments or effects were significantly different. All statistical analyses were performed using R with a significance threshold set at 0.05 (R-Core-Team, 2017, 2014).

#### 3 Results

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### 3.1 Initial soil organic matter chemistry and nitrogen differ between the cold and warm regions

The initial soil N content in the present study was greater in the warm relative to the cold region (p<0.0001), consistent with soil sampled in earlier studies from the same climate regions (Philben et al. 2016). The initial soil C:N ranged from 32-54. These ratios did not exhibit any region by horizon effects (p=0.1272), but were lowest in the warm region (p<0.0001) and decreased with depth (p<0.0001; Fig. 3a). The initial %N as THAA ranged from 38-43% and was similar in soils for the two climate regions, and decreased with depth (p=0.0011; Fig. 3b) in a manner consistent with increasing soil organic N degradation with depth (Philben et al. 2016). Consistent with this feature, mol% glycine increased with depth (p<0.0001) and was elevated in the warmer region soils (p=0.0035) and to a greater extent within the H horizon (p=0.0126), consistent with less soil organic N degradation in the cold relative to the warm region (Fig. 3c; Philben et al., 2016). The initial %alkyl-C ranged from 27-33% and exhibited a region by horizon interaction (p=0.0443), revealing higher values in F relative to L, and H relative to F horizons for the warm region (Fig. 3d). Higher %alkyl-C was observed in the L and H of the warmer region

(p=0.0301) that increased with depth (p=0.0137). The initial %di-O-alkyl-C, ranging from 7-9%, was elevated in the cold region soils (p<0.0001; Fig. 3e) but exhibited no other trends. The alkyl-C to O-alkyl-C ratio exhibited a regional (p=0.0014) effect only with an elevated ratio in warm relative to cold region soils (Fig. 3f).

# 3.2 Losses of soil mass and declines in soil C and N concentrations were enhanced by increased temperature and to a greater extent in the cold region soils

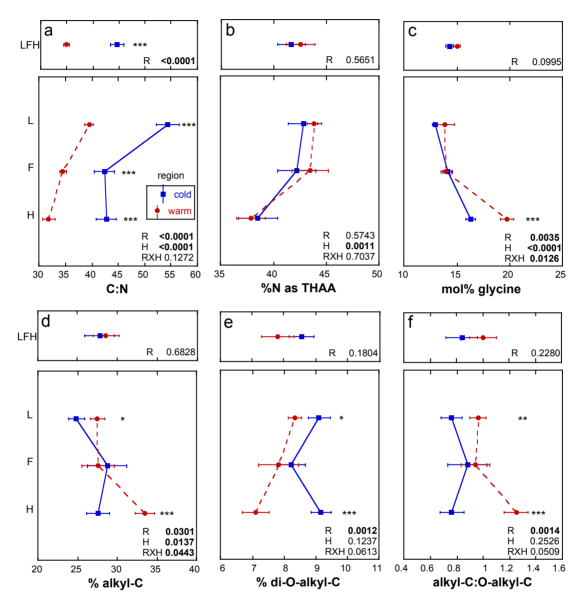
Over the 438+ day incubation, mass loss, %C loss, and %N loss were greatest in the 15°C incubation and for the cold region soils (Table 1). In the isolated experiment, mass loss ranged from  $16 \pm 1$  to  $39 \pm 5\%$  and  $16 \pm 3$  to  $30 \pm 2\%$  (mean  $\pm$  standard error), respectively, for the cold and warm regions. The % C loss ranged from  $21 \pm 3$  to  $82 \pm 4\%$  in the cold region soils with 82% loss at 15°C in the F sub-horizon. The warm region soils exhibited a % C loss that ranged from  $24 \pm 5$  to  $46 \pm 3\%$ . The % N loss in the cold region soils ranged from  $11 \pm 9$  to  $45 \pm 5\%$  and was greatest in the 15°C treatment. In the warm region % N loss ranged from  $13 \pm 3$  to  $52 \pm 23\%$  and did not exhibit a temperature effect (p = 0.227). In the whole organic profile (whole experiment) and the predicted whole experiment soils, %mass loss was greatest at 15°C and ranged from  $16 \pm 2$  to  $37 \pm 1\%$  and  $12 \pm 2$  to  $28 \pm 2\%$ , respectively, for the cold and warm regions. In the warm region, %mass loss was greater in the predicted experiment relative to the whole profile experiment (p = 0.011). Effect of temperature and experiment was observed for %C loss in the cold and warm regions with greatest %C losses having occurred at 15 °C and in the predicted whole experiment. The %N loss in these whole profile experiments ranged from  $11 \pm 4$  to  $38 \pm 5\%$  in the cold climate soils where greater loss was observed at 15 °C yet no experimental effect was observed (p = 0.119). The warm climate soils exhibited a similar range in % N loss from  $13 \pm 3$  to  $40 \pm 13\%$  with no temperature effect (p = 0.064).

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**Table 1.** The mean (standard error) of % mass loss, % C loss and % N loss within each experiment where individual horizons were incubated in isolation from each other (isolated experiment), then calculated as a whole profile values based upon those isolated horizon results (predicted), and given also as the actual measured incubation results for whole organic profiles (measured). The effect of temperature (T), horizon (H), and interaction term (T x H) are given for the isolated experimental results (Top). The effect of T, experiment (E), and interaction term (T x E) are given for the tests conducted across both the predicted and measured whole profile experimental treatments (Bottom). Significance ( $\alpha = 0.05$ ) is denoted in bold.

| % Mass                |           |             |        | s loss      |        | % C loss    |         |             |        | % N loss    |         |             |         |
|-----------------------|-----------|-------------|--------|-------------|--------|-------------|---------|-------------|--------|-------------|---------|-------------|---------|
| -                     |           | Cold region |        | Warm region |        | Cold region |         | Warm region |        | Cold region |         | Warm region |         |
| Experiment Horizon(s) |           | 5°C         | 15°C   | 5°C         | 15°C   | 5°C         | 15°C    | 5°C         | 15°C   | 5°C         | 15°C    | 5°C         | 15°C    |
| Isolated              | L         | 22.62       |        | 18.27       |        | 34.22       | 65.80   |             |        | _           | 23.36   | 12.56       | 9.16    |
|                       | _         | (3.46)      | (4.68) | (1.63)      | (3.52) | (6.90)      | (10.13) | (4.09)      | (4.54) | (9.57)      | (14.75) | (3.31)      | (4.92)  |
|                       | F         | 23.54       |        | _           | 29.73  | 34.53       | 81.52   | _           | 46.15  |             |         | _           |         |
|                       |           | (1.55)      |        |             | (2.36) | (2.74)      | •       |             | (2.79) |             |         |             | (22.83) |
|                       | Н         | 15.81       | 27.91  | 15.75       | 22.12  | 21.05       | 56.70   |             | 42.92  |             |         |             | 31.86   |
|                       |           | (1.13)      | (2.02) | (2.48)      | (1.77) | (3.23)      | (12.17) | (4.66)      | (3.23) | (8.49)      | (5.22)  | (7.46)      | (6.44)  |
|                       | Effects   | F           | р      | F           | p      | F           | Р       | F           | р      | F           | р       | F           | p       |
|                       | Т         | 27.43       | 0.0002 | 15.16       | 0.0021 | 39.09       | <0.0001 | 24.42       | 0.0003 | 5.41        | 0.0383  | 1.63        | 0.2265  |
|                       | Н         | 4.84        | 0.0288 | 3.67        | 0.0572 | 3.33        | 0.0709  | 0.85        | 0.4535 | 2.16        | 0.1582  | 2.98        | 0.0887  |
|                       | ΤxΗ       | 0.14        | 0.8709 | 0.14        | 0.8723 | 0.57        | 0.5783  | 0.12        | 0.8886 | 0.24        | 0.7891  | 1.02        | 0.3904  |
|                       |           |             |        | 1           |        | 1           |         | 1           |        | •           |         | 1           |         |
| Whole                 | predicted |             |        |             | 27.87  | 32.02       |         |             | 44.98  |             |         |             |         |
| Profile               |           | (0.20)      |        |             | (2.26) | (0.79)      |         |             | (2.33) |             |         |             |         |
|                       | measured  | 15.46       | 36.82  | 12.32       | 21.81  | 21.07       | 67.62   | 21.31       | 38.04  | 10.61       | 31.70   | 13.14       | 15.78   |
|                       |           | (1.34)      | (2.24) | (2.44)      | (1.18) | (1.29)      | (4.73)  | (3.35)      | (4.19) | (3.68)      | (4.30)  | (3.30)      | (0.98)  |
|                       | Effects   | F           | р      | F           | р      | F           | Р       | F           | р      | F           | р       | F           | р       |
|                       |           |             | -      |             | -      |             | <0.0001 |             | •      |             | •       |             | 0.1946  |
|                       | E         | 3.83        |        |             |        |             | 0.0082  |             | 0.0372 |             |         |             | 0.0643  |
|                       | ΤxΕ       | 4.70        | 0.0621 | 0.13        | 0.7312 | 0.78        | 0.4026  | 0.02        | 0.8829 | 0.05        | 0.8213  | 1.15        | 0.3151  |



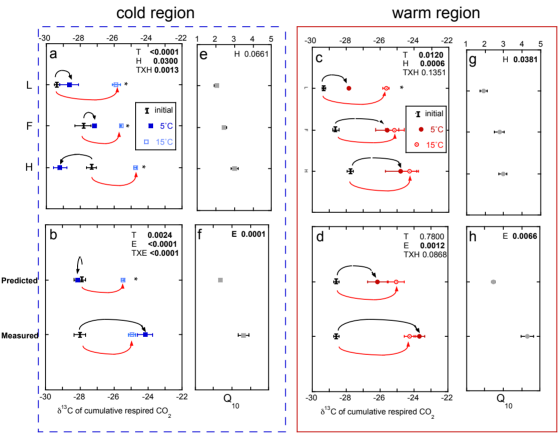
**Figure 3.** Initial mean of the three sites  $\pm$  standard error of soil metrics for the whole organic profile (LFH; upper panel) and individual horizons (L, F, H) from the isolated experiment (lower panel). Within the organic profile, an analysis of variance was utilized for the effect of region (R). Significance is denoted by asterisks:  $p \le 0.0001$  \*\*\*,  $p \le 0.005$  \*, and p > 0.05 n.s.. The effect of R within horizon is denoted with the asterisks.

# 3.3 The $\delta^{13}C$ of cumulative respired $CO_2$ increased in whole-profile incubations, and with incubation temperature

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The  $\delta^{13}$ C of respired CO<sub>2</sub> increased throughout the incubation period. However, an exception to this includes all the soil horizons from the cold region incubated within the isolated experiment and, as a result, the predicted whole experimental treatment (Fig. 4). The  $\delta^{13}$ C of respired CO<sub>2</sub> over the course of the incubation ranged from -29 to -24‰, with more  $^{13}$ C-enriched respired CO<sub>2</sub> released from the deeper soil horizons and soils exposed to the higher incubation temperature. Overall the effect of temperature (p=0.0024), experiment (p<0.0001) and their interaction (p<0.0001) was observed in the cold region soils while only the effect of experiment (p = 0.0010) was observed in the warm region soils. Regardless of region or temperature, the whole experiment exhibited more  $^{13}$ C-enriched respired CO<sub>2</sub> akin to the impact



**Figure 4.** The  $\delta^{13}$ C of the total cumulative respired CO<sub>2</sub> comparing individual horizons incubated in isolation (isolated experiment; L, F and H; a,c) and whole-profile values (b,d) predicted from those isolated horizons (predicted) to those measured directly as a whole-profile (measured) with the corresponding temperature sensitivity (Q<sub>10</sub>; e,f,g,h) of the total cumulative respiration for the entire incubation period for the soils from both the cold and warm regions. Values are given as the mean of three sites  $\pm$  standard error with the initial bulk soil  $\delta^{13}$ C included for reference. The effect of temperature (T), horizon (H) or experiment (E) and their interaction term (TxH or TxE) are given for all three effects with significance ( $\alpha = 0.05$ ) denoted in bold. Within horizon effect of T is denoted with an asterisk (\*; a,c). The significant effect of T within experiment is denote with an asterisk (\*; b,d).

of increased temperature across all soils and experiments. For example, the whole experiment incubations at both temperatures exhibited a 3-4% increase in  $\delta^{13}$ C-CO<sub>2</sub> relative to both the initial and the isolated experiment at 5°C (Fig. 4b). In contrast, the cold region soils incubated as individual L, F and H horizons exhibited a temperature difference with a ~3% increase in  $\delta^{13}$ C-CO<sub>2</sub> in the 15°C relative to both the initial values and the 5°C incubations (Fig. 4a). These results indicate that the whole profile structure enhanced the respiration of more <sup>13</sup>C-enriched substrates, to an extent similar to that observed in the warmer incubations. Furthermore, the respiration of more <sup>13</sup>C-enriched substrates was associated with higher Q<sub>10</sub> of soil respiration regardless of climate region.

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Though initial soil organic matter  $\delta^{13}C$  and  $\delta^{15}N$  differed by climate region, the changes in soil  $\delta^{13}C$  and  $\delta^{15}N$  over the course of the incubation were relatively small (typically less than 1‰) across all horizons, and with both temperatures and experimental types (Table S2). No temperature, horizon or experiment effect was observed in the changes in bulk soil of  $\delta^{13}C$  and  $\delta^{15}N$  in these soils.

# 3.4 Degradation of soil nitrogen occurred to a greater extent in the cold relative to the warm region soils with no evidence of an effect of whole profile structure.

The magnitude of change in C:N (ΔC:N) over the course of the experiment decreased with soil depth regardless of region (Fig. 5abcd). Increased temperature resulted in a greater decrease in soil C:N which exhibited decreases of 3-14 and 0-9 in the cold and warm regions, respectively, over the course of the entire incubation (Fig. 5 abcd). In the cold region, where the soil C bioreactivity is generally greater (Laganiere et al. 2015), this trend was similar across both the predicted and measured whole experiments, suggesting that soil profile structure was not an important factor for this variable. However, in the warm region soils, the decrease in C:N was evident in the individual L and H horizons as well as the whole profile experiment but not when calculated as a whole profile from the individual horizons. Although a temperature effect was noted within the isolated and both whole treatments of the cold and warm region soils, no effect of soil profile treatment was detected for the ΔC:N within either region. Changes in %C as THAA were only noted in the F sub-horizon of the warm region soils incubated at 15°C (Fig. S1). No effects of temperature, horizon or experiment were observed for changes in %C as THAA. Changes in %N as THAA were variable, but exhibited temperature effects consistent with the  $\Delta$ C:N in the cold climate soils in the whole profile soils whether incubated as a whole profile or predicted from the isolated horizons (Fig. 6abcd). Changes in mol\% glycine were also consistent with these observations of soil organic N processing in the cold region soil (Fig. 6efgh). The mol% glycine, an indicator of greater microbial reworking of soil organic N including in these soils (Dauwe and Middelburg, 1998; Hedges et al., 1994; Philben et al., 2016), increased with temperature in the L and H sub-horizons as well as the whole organic profiles from the cold region. In the warm region soils, temperature effects were only noted in the isolated experiment where decreases in mol% glycine were observed in the 5°C relative to  $15^{\circ}$ C incubation temperatures (p = 0.0028). This temperature effect was observed within the H sub-

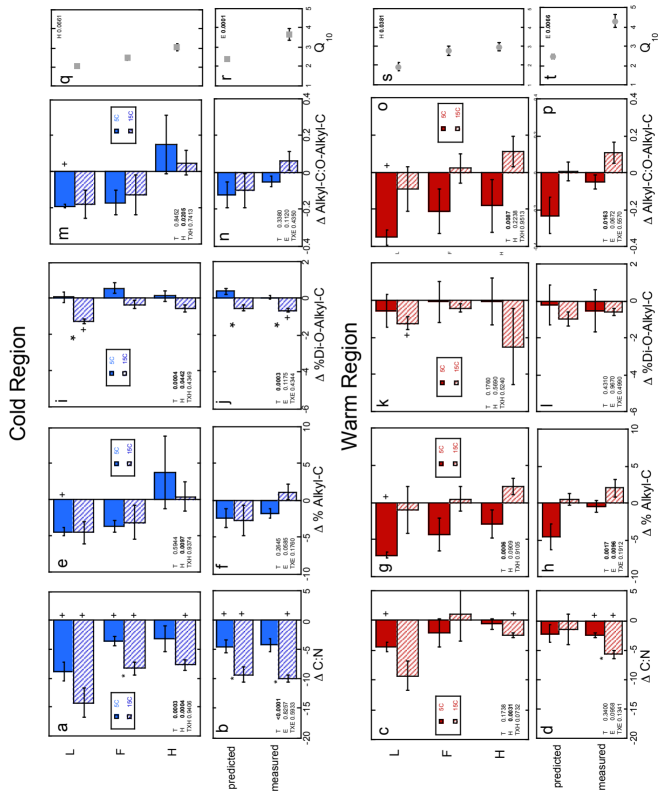


Figure 5

Figure 5. The change in soil C:N<sub>molar</sub> ( $\Delta$  C:N), percent of soil carbon as alkyl-C ( $\Delta$ % Alkyl-C) and as di-O-alkyl-C ( $\Delta$ % Di-O-Alkyl-C), and the ratio of alkyl-C to O-alkyl-C ( $\Delta$  Alkyl-C:O-Alkyl-C) given as the final minus initial absolute values comparing the experiment where individual horizons were incubated in isolation from each other (upper panels in each labeled by horizon; L, F and H) to both the calculated whole profile values based upon those isolated horizon results (predicted) and the actual measured incubation results for whole organic profiles (measured). These results are given for both the cold (a,b,e,f,i,j,m,n) and warm regions (c,d,g,h,k,l,o,p). The corresponding temperature sensitivity of the total cumulative respiration for the entire incubation period (Q<sub>10</sub>; q and r for cold region; s and t for warm region) taken from Podrebarac et al. (2016) are provided for reference. All values provided are the mean of the three sites ± standard error with a significant change from 0 denoted by symbol "+". For the effect of temperature (T), horizon (H) or experiment (E) and their interaction term (TxH or TxE) significance (α ≤ 0.05) is denoted in bold. Significance of the effect of temperature within horizon or within experiment is denoted by an asterisk.

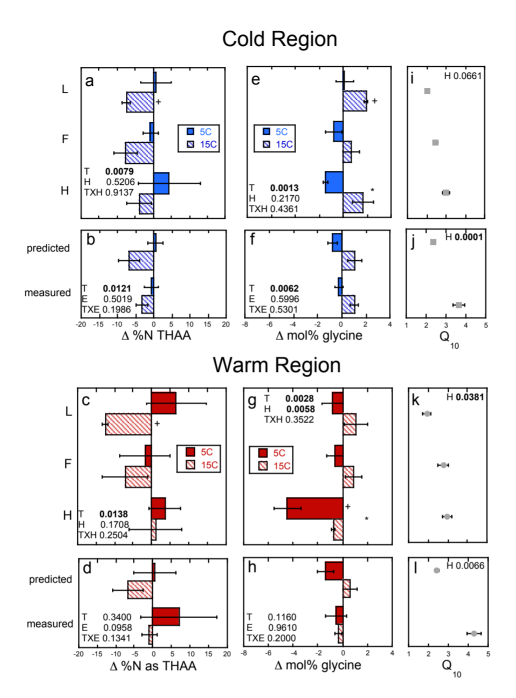


Figure 6. The change in soil %N as total hydrolyzable amino acid (Δ %N as THAA) and mole percent glycine (Δ mol% glycine) given as the final minus initial absolute values comparing the experiment where individual horizons were incubated in isolation from each other (upper panels in each labeled by horizon; L, F and H) to both the calculated whole profile values based upon those isolated horizon results (predicted) and the actual measured incubation results for whole organic profiles (measured). These results are given for both the cold (a,b,e,f) and warm regions (c,d,g,h). The corresponding temperature sensitivity of the total cumulative respiration for the total incubation period (Q<sub>10</sub>; i and j for cold region; k and l for warm region) taken from Podrebarac et al. (2016) are provided for reference as they were in Fig.3. All values provided are the mean of the three sites ± standard error with a significant change from 0 denoted by symbol "+". For the effect of temperature (T), horizon (H) or experiment (E) and their interaction term (TxH or TxE) significance (α ≤ 0.05) is denoted in bold. Significance of the effect of temperature within horizon or within experiment is denoted by an asterisk.

horizon (p = 0.0058), indicating a relative increase in final mol% glycine following incubation at the higher relative to lower incubation temperature.

# 3.5 Change in soil carbon chemistry during incubation was affected by both temperature and whole soil profile structure

Changes in the alkyl-C to O-alkyl-C ratio ( $\Delta$ A:O-A) were observed within the L horizons from both climate regions. However, temperature effects on the  $\Delta A$ :O-A (p = 0.0087) were noted only in the warm region soils and horizon effects (p = 0.021) only in the cold climate soils (Fig 5 mnop). Unexpectedly, %alkyl-C (Fig 5 efgh) decreased while %O-alkyl-C increased in the warm region soils over the incubation at 5 °C resulting in the decreased  $\Delta A$ :O-A observed (Fig. 5 mnop). Decomposition of vascular plant tissues typically results in increases in A:O-A as a result of losses in carbohydrates (rich in o-alkyl-C) and relative retention of plant aliphatics (rich in alkyl-C) (Preston et al., 2009; 2000). This unexpected finding is perhaps a result of the significant surface inputs of moss tissues rich in structural carbohydrates that can be resistant to decomposition (Turetsky et al., 2008; Hájek et al., 2011; Philben et al., 2018). The cold region soils exhibited a horizon effect on the change in %O-alkyl-C (p=0.026) and  $\Delta$ A:O-A (p = 0.021) such that %O-alkyl-C generally increased and A:O-A decreased in the L subhorizon while %O-alkyl-C decreased and A:O-A increased in the H sub-horizon over the incubation (Fig 5 e). In the warm region soils, increased incubation temperature appeared to have eliminated these unexpected changes in %alkyl-C, %O-alkyl-C and the A:O-A. No change in %alkyl-C, %O-alkyl or A:O-A was noted over the course of the 15°C incubation. However, the unexpected trend of decreasing A:O-A and %alkyl-C was not observed in the warm climate soils incubated as whole soil profiles. An effect of both temperature (p=0.0017) and experiment type (i.e., whole vs isolated) (p=0.0096) was noted in the change in %alkyl-C results. This is observed as an increase in %alkyl-C within the whole profile experiment only when incubated at 15°C as opposed to the reduction in %alkyl-C at 5°C and no change at 15°C in the predicted whole profile (Fig. 5 efgh). No experimental effects were noted for either the A:O-A (p = 0.067) or %O-alkyl-C (p = 0.143) changes in the whole soil profiles from the warm region. The %di-O-alkyl-C exhibited a decrease only following the 15°C incubation of the L horizons from both climate regions (Fig 5 ik). However, the isolated horizons from the cold region exhibited both a temperature (p = 0.0004) and horizon (p = 0.044) effect, indicating decreases in %di-O-alkyl-C with warming and primarily within the L subhorizon. Further, a temperature effect (p = 0.0003), observed as a decrease in %di-O-alkyl-C, was also observed in the whole soil profiles from the cold region regardless of experiment type (Fig 5 j). The relative changes in the other carbon types as detected via <sup>13</sup>C-NMR were generally below detection. However, we did note increases in %aromatic-C over the course of the 5°C incubation of the warm region soils, consistent with the proportional decreases as observed in %alkyl-C (data not shown).

### 4 Discussion

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Prior work at these same sites demonstrated how the temperature response of respiratory C loss from mesic boreal forest organic soils can be enhanced in warmer relative to colder regions, but only when horizons were

incubated together and layered as a whole soil profile; this is also congruent with field observations of soil respiration in the same forest sites (Fig. 1 center; Podrebarac et al., 2016). Isolated horizons of the same soils, under the same conditions, exhibited lower respiratory responses to increased temperature, and no regional climate differences despite clear differences in soil bioreactivity between the two climate regions (Laganière et al., 2015; Podrebarac et al., 2016). These previous studies suggest that some mechanism facilitated by the connectivity among soil horizons can enhance the temperature sensitivity of soil respiration over weekly to monthly time scales. Here, we explore the potential role of soil priming as a mechanism supporting the enhanced temperature responses of respiration in these whole soil profiles. We found evidence for labile C but not labile N priming of microbial activities, supported by soil horizon connectivity that indirectly stimulates respiratory responses to increased temperature (Fig. 1 right). Additionally, the data indicate that the labile C priming mechanism is most emergent in the warmer region soils, consistent with lower SOC bioreactivity in the warmer region relative to soil from the colder region.

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# 4.1 Enhancement of microbial labile carbon use supports an indirect mechanism for increased temperature sensitivity of soil respiration in whole organic soil profiles

The inclusion of relatively faster-turnover soil substrates from the surface L horizons with the slower turnover pools within the lower F and H horizons promoted microbial use of more bioreactive substrates and an enhanced soil respiratory response to temperature in these whole soil profiles. Enhanced temperature sensitivity of microbial activity supported indirectly through increased availability of labile substrates, or a priming effect, has been observed with root exudates and other fresh plant inputs (Curiel Yuste et al., 2004; Zhu and Cheng, 2011) but, to the best of our knowledge, has not yet been linked to soil profile connectivity. These whole soil profiles exhibited increases in the  $\delta^{13}$ C of respired CO<sub>2</sub> and relative enhancement in the decomposition of carbohydrates compared to soil horizons incubated in isolation. These increases were associated with the enhanced temperature response of respiration in the whole organic profiles not predicted by the response of the same individual horizons incubated in isolation. Most strongly observed in the warm region forest soils, where composition metrics indicates initially lower bioreactivity, the enhanced temperature sensitivity of respiration with the whole soil profile structure appears to be largely supported by labile C priming as we had hypothesized. These respiratory responses to cross horizon exchange were observed throughout the incubation (Podrebarac et al. 2016) and not just at the end when we were able to observe soil chemical composition. This highlights the challenges for in situ profile studies where a combination of short and longer term investigations are needed to capture labile C priming influences on soil substrate use and respiratory responses to temperature.

Enhanced carbohydrate use suggests that catabolism of more bioreactive,  $^{13}$ C-enriched substrates supported the use of lower  $E_a$  substrates, helping to explain the elevated  $Q_{10}$  of soil respiration within the whole profile soils relative to the sum of the individual horizons from the same profiles. Soil substrates are not uniformly available to microbes, and both temperature and changes in the suite of available substrates can alter the catabolism or incorporation of soil substrates by microbes (Bölscher et al., 2017; Fontaine et al., 2007; Frey et al., 2013; Streit et al., 2014; Zogg et al., 1997). During incubations, catabolism of more recent, fast turnover soil inputs relative to the

bulk soil can occur along with an increased proportion of older, slower-turnover soil substrates incorporated into microbial biomass (Blagodatskaya et al., 2011). The increases in the  $\delta^{13}$ C of respired CO<sub>2</sub> observed with the whole profile structure, independent of region or incubation temperature, were congruent with increases in  $\delta^{13}$ C of respired CO<sub>2</sub> observed with incubation temperature in the isolated horizons (Fig. 4). These increases exceed those attributed to isotopic discrimination associated with respiration (<1‰; (Breecker et al., 2015; Czimczik and Trumbore, 2007) and likely resulted from enhanced use of more <sup>13</sup>C-enriched substrates, increased reuse of bioreactive C pools incorporated into microbial biomass, and/or shifts in microbial composition of the active community (Blagodatskaya et al., 2011).

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Congruent with the increased δ<sup>13</sup>C of respired CO<sub>2</sub>, the relative retention of alkyl-C observed in the warmer region soils was likely a consequence of enhanced use of carbohydrates which are relatively <sup>13</sup>C enriched components of soil organic matter (Benner et al., 1987; Hobbie and Werner, 2004). When the soil horizons were incubated at 5°C as isolated horizons we observed the *opposite* of the typical relative loss of carbohydrates (reduced O-alkyl-C) and retention of plant waxes (retained alkyl-C) associated with the decomposition of vascular plant tissues (Preston et al., 2000; 2009). The unexpected trend of decreasing alkyl-C and A:O-A, observed in the 5°C incubation of the cold and warm region L horizons, was notably absent in the 15°C incubations of warm region soils (Fig. 5). The experimental effect on the change in alkyl-C indicated that the whole horizon structure enhanced decomposition more typical of increased carbohydrate use which was further enhanced with increased temperature.

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The lack of change in %alkyl-C or %O-alkyl-C within the cold region soils likely resulted from the larger relative concentration of carbohydrates (Fig. 3 ef), a consequence of the greater moss contributions to these soils relative to the warm forest soils (Kohl et al., 2018). The cold region profiles lack the increase in the A:O-A with depth observed in the warm region profiles and typically observed in vascular plant dominated soils including boreal forests (Kane et al., 2010). Therefore, the increases in  $\delta^{13}$ C-CO<sub>2</sub> associated with temperature and the proximity of horizons within the whole profile may still have been due to enhanced mineralization of carbohydrates in the cold region soils not clearly detected in bulk changes in the chemistry of the SOM.

Microbial use or catabolism of lower  $E_a$  compounds (e.g. carbohydrates) in support of microbial responses

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to increasing temperature likely enhanced substrate assimilation and use efficiency in the whole profile soils. The enhanced labile substrate use and temperature response of soil respiration in the whole profile incubations coincided with lower soil C losses (Table 1) as compared with the isolated horizons. This could be explained by a priming effect within these soil profiles enhancing the use of more complex higher  $E_a$  substrates consistent with respiratory temperature responses closer to intrinsic values relative to those of isolated horizons lacking this priming effect (Davidson and Janssens, 2006). Root exudates have been found to increase the availability or use of complex, high  $E_a$  substrates via a priming effect (Bingeman et al., 1953; Cheng et al., 2014), by accelerating SOM decomposition via co-metabolism or the increased production of polymer degrading enzymes breaking down macromolecules and generating more soluble molecules (Schimel and Weintraub, 2003; Wallenstein and Weintraub, 2008; Zhu and Cheng, 2011). Similarly, the whole-soil profiles likely promoted availability of a diversity of substrates and activity of more diverse microbes than in isolated horizons, supporting co-metabolism or increased polymer degrading enzyme activity (Basler et al., 2015) as has been noted with litter additions (Malik et al., 2016).

The whole profile structure studied here results in the contact between communities with high fungal to bacterial ratios (F:B) within carbohydrate rich L horizons with communities exhibiting low F:B within less carbohydrate rich F and H horizons (F:B L>F>H in these forest soils; Kohl et al. 2015). Relative to bacteria, fungi can exhibit greater substrate use efficiencies (Bölscher et al., 2016; Kallenbach et al., 2016) and their hydrolytic activities may support the cross horizon enhancement of substrate use including higher  $E_a$  substrates in these organic horizons, congruent with the reduced respiration rates and enhanced carbohydrate decomposition observed in the whole profile soils. By initiating key steps in decomposition of more complex soil organic matter (Paterson et al., 2008), fungi can enhance the decomposition of higher  $E_a$  substrates found within the F and H layers, and likely to a greater extent when incubated in contact with the fungal rich L layer. Enhanced fungal enzyme activity is not exclusive of enhanced bacterial respiration and use of carbohydrates in these soil profiles; rather enhanced bacterial relative to fungal respiration could explain the increases in  $\delta^{13}$ C of respired CO<sub>2</sub> observed in the whole profiles (Dijkstra et al., 2006; Glaser and Amelung, 2002). This suggests that enhancement in bacterial relative to fungal respiration, and specifically bacterial catabolism of carbohydrates, occurred in the whole soil profiles, support the idea of a labile C priming effect.

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## 4.2 Enhanced temperature sensitivity of respiration is not associated with enhanced N use in whole soil profiles

Soil N availability can impact the soil microbial community, its substrate use and growth efficiency (Blagodatskaya et al., 2014; Mooshammer et al., 2014), and priming effects supported by fungi (Dijkstra et al., 2013). Therefore, we additionally hypothesized that enhanced soil N availability supported by the whole soil profile structure, has the potential to enhance the temperature response of soil respiration given differences in soil N content and microbial community composition by horizon. Soil N content also differed by climate region in these forests, providing us with an opportunity to assess the role of soil N exchange and use across a climate relevant range of soil N availability in this boreal forest region (Philben et al. 2016). Substrate use could therefore be influenced by exchange across horizons as both fungal abundance and soil organic N concentration and composition vary with depth in most soil profiles and particularly in the boreal forest organic horizons explored here (Fig. 1 left side; Kohl et al., 2015; Philben et al., 2016).

Despite the observed temperature effects on soil N losses consistent with enhanced N<sub>2</sub>O production with warming observed in these organic horizons (Buckeridge et al. 2020), and clear differences in availability of soil N represented by the two climate region soils, we observed no difference in soil N use between soils incubated as a whole profile versus as isolated horizons. In particular the lower N availability of the cold region soils, indicated by lower initial %N and higher C:N (Fig. 3), suggests that if we were to see an enhancement in soil organic N use it would likely have been observed in the cold region soils as noted. However, the enhanced use of soil organic N in the colder region with increased incubation temperature was not impacted by whether soil horizons were incubated in isolation or connected as a whole profile. The observed changes in %N, C:N, and indicators of amino acid degradation (%N as THAA, mol% glycine) were similar in both soil profile treatments of the cold region soils (Fig 6). Congruent with the greater bioreactivity and soil C respiratory losses observed in the cold relative to warm

region soils (Laganière et al., 2015), decreases in soil C:N were primarily observed in the cold region soils where soil C:N was initially higher and temperature effects were noted regardless of soil incubation type (isolated horizons or as a whole profile). This contrasted with the warm region soils where the change in C:N only differed with incubation temperature in the whole profile treatment consistent with the carbohydrate losses.

We observed some increased degradation of amino acids in the isolated horizons during the incubation. For example, decreases in %N as THAA were detected in the isolated L horizons from both the warm and cold region indicating we were able to detect soil N use over the incubation period used, and found that the greater availability of N in the surface L horizons supports enhanced N use with increased temperature but without necessarily impacting overall N use in whole profiles. These results suggest that the enhanced temperature sensitivity of soil respiration observed in the whole profile soils relative to the sum of their isolated horizons is not due to changes in N substrate use but rather C substrate use facilitated by the whole profile structure.

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# 4.3 Cross horizon exchange supports labile C priming as a whole soil profile mechanism enhancing soil respiratory responses to temperature

We demonstrate how inter-layering of soil horizons as a whole profile, and as found in situ, supports crosshorizon exchange that can enhance the use of labile C and respiratory responses to temperature (Fig. 1). Thus we reveal an additional mechanism that can control in situ respiratory responses to changing temperature, an important soil-climate feedback. This result further explains discrepancies between laboratory and in situ observations of the temperature sensitivity of soil respiration highlighting the role of in situ processes that must be captured to better predict this feedback within Earth System Models.. Given that recent photosynthates, e.g. root exudates, can contribute similarly to soil respiration across surface to deep horizons (Pumpanen et al., 2009) the role of a priming mechanism suggested by our study is worthy of investigating in deeper mineral soil profiles where enhanced temperature sensitivity of soil respiration is supported by soil C of recent origin (Hicks Pries et al., 2017). Deeper soils are certainly key to uncovering the full soil response to climate change, and understanding controls on those responses appears to require an increased understanding of the cross-horizon exchange processes suggested by this study. For example, root inputs and hydrologic regimes transferring dissolved organic matter and nutrients as well as regulating redox conditions represent relevant factors likely controlling interactive effects of cross-horizon exchange on soil C use. The degree to which this mechanism exerts itself in other soils remains unknown, but these results highlight the importance of understanding priming mechanisms that operate within whole soil profiles – only rarely studied – in regulating respiratory responses to changing temperature.

Appendices. Supplemental material related to this article available online.

Data availability. All data are included in the paper tables and Supplement.

### **Author contribution**

Authors JL, SB, and SZ contributed to the general conceptions of the study. SB, KE, JL, and SZ designed the sampling. KE, JL, and SZ contributed to field sample collections while incubation experiment set up and sampling was conducted by FP and JL. FP, JL and MN contributed to sample analyses including soil CO2 fluxes, sample extractions and preparations for isotope and elemental analyses, NMR and amino acids. FP conducted data and statistical analyses. FP and SZ jointly wrote the manuscript which received edits from all co-authors.

### **Competing interests**

The authors declare that they have no conflict of interest.

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### Conflicts of interest.

The authors declare that they have no conflicts of interest or competing interests.

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