Abstract

The thawing and subsidence of frozen peat mounds (palsas) in permafrost landscapes results in the formation of organic-rich thermokarst lakes. We examined the effects of palsa degradation on CH$_4$ and CO$_2$ emissions by comparing thermokarst lakes at two peatland locations in subarctic Québec, Canada: in the northern discontinuous permafrost region, and in southern sporadic permafrost where palsas are more rapidly degrading. The lakes were shallow (<3 m) but stratified at both sites, and most had anoxic bottom waters. The surface waters at both sites were supersaturated in CH$_4$ and CO$_2$, and to a greater extent in the southern lakes, where the surface CH$_4$ concentrations were up to 3 orders of magnitude above air equilibrium. Concentrations of CH$_4$ and CO$_2$ increased by orders of magnitude with depth in the southern lakes, however these gradients were less marked or absent in the North. Strong CH$_4$ and CO$_2$ emissions were associated with gas ebullition, but these were greatly exceeded by diffusive fluxes, in contrast to thermokarst lakes studied elsewhere. Also unusual relative to other studies to date, the surface concentrations of both gases increased as a linear function of water column depth, with highest values over the central, deepest portion of the lakes. Radiocarbon dating of ebullition gas samples showed that the CH$_4$ had $^{14}$C-ages from 760 yr to 2005 yr before present, while the CO$_2$ was consistently younger. Peatland thermokarst lakes may be an increasingly important source of greenhouse gases as the southern permafrost limit continues to shift northwards.

Global levels of atmospheric methane (CH$_4$) continue to rise, with values close to 2 ppm observed locally in the Arctic (NOAA 2014). The origins of this CH$_4$ are uncertain, and one source of uncertainty is the net flux to the atmosphere from thermokarst lakes and ponds (Negandhi et al. 2013; Tan et al. 2015). These waterbodies are formed by the thawing and collapse of ice-rich permafrost (Hopkins 1949), and they have been recognized as sites of intense biogeochemical activity in northern landscapes (Vincent et al. 2013, and references therein). Permafrost soils have the potential to release up to 508 Pg of carbon by 2100, depending on different response scenarios (MacDougall et al. 2012); however the current estimates of greenhouse gas (GHG) emissions from thermokarst lakes range from major (Schuur et al. 2013) to insignificant (Gao et al. 2013). This large variability in estimates is in part related to the diversity of thermokarst lake types, and the sparseness of observations across the circumpolar North.

Much of the emphasis to date on measuring gas fluxes from thermokarst lakes has focused on water bodies that are located on yedoma, wind-blown surface deposits that cover vast areas of Siberia and Alaska. Yedoma permafrost soils contain 1–5% organic carbon (Tarnocai et al. 2009), and the thermokarst lakes in these landscapes have elevated rates of CH$_4$ release, dominated by ebullition. For example, in a comparison of 40 Alaskan lakes along a north-south transect, the CH$_4$ emissions were sixfold higher than from non-yedoma lakes, the latter including waters unaffected by permafrost; ebullition accounted for 86% and 65% of the annual CH$_4$ emissions from yedoma and non-yedoma lakes, respectively (Sepulveda-Jauregui et al. 2015). In general, however, there is a sparseness of information about the biogeochemistry of non-yedoma thermokarst lakes, despite an estimated storage of 75% of the global carbon permafrost pool in non-yedoma permafrost (Vonk et al. 2015).

Non-yedoma thermokarst lakes include water bodies associated with the collapse of palsas, ice-rich permafrost mounds and plateaus that are found in northern peatlands,
particularly in subarctic regions of discontinuous permafrost (e.g., Karlsson et al. 2010; Saemundsson et al. 2012). These mounds form during periods of climatic cooling and permafrost aggradation, and they rise up to 10 m above the initial ground level as a result of interstitial water and groundwater that is aspirated towards growing ice lenses in fine sediments at the base of the mound (“cryosuction”); this uplifts the overlying frozen peat, which in turn reduces snow accumulation and favors the cooling, freezing and ongoing mound development (An and Allard 1995). These periglacial features are sensitive to subsequent increases in temperature and precipitation, and their thawing and collapse result in thermokarst basins that fill with water, derived in part from the melting ice (Luoto and Seppälä 2003). Palsas contain a large organic carbon pool, with concentrations around 50% dry weight (Krüger et al. 2014). This is an order of magnitude higher than yedoma soils (Tarnocai et al. 2009), and although the lability of this moss-derived material is uncertain, observations in northern Norway point to degrading palsa soils as sites of abundant methanogens and high rates of methanogenesis, particularly during the thermokarst pond stage, prior to eventual infilling by sediment and plants (Liebner et al. 2015).

Our primary objective in the present study was to quantify the CH$_4$ and CO$_2$ emissions from thermokarst lakes formed by subsiding palsas in subarctic peatlands. We undertook this research in northern Québec (Nunavik), where mean annual air temperatures have increased over the last three decades, and where permafrost thawing and degradation are proceeding rapidly (Bhiy et al. 2011; Bouchard et al. 2015). We aimed to determine the age of the source carbon of these emissions for comparison with the stratigraphic history of these sites, and to evaluate the relative importance of gas fluxes to the atmosphere via ebullition and diffusion. We hypothesized that climate-induced thawing and expansion of palsa-associated thermokarst lakes would mobilize ancient carbon reserves and would stimulate CH$_4$ and CO$_2$ emissions as a result of the aquatic microbial degradation of these materials. To address this hypothesis, we measured the concentrations, surface fluxes and $^{14}$C age of CH$_4$ and CO$_2$ along with profiles of limnological variables in thermokarst lakes at two contrasting peatland sites: in a valley of rapidly degrading palsas near the southern permafrost limit, and in a colder palsa valley located further to the North where permafrost degradation is less advanced. We also examined the questions of how gas emissions in these waters vary as a function of water depth and within a 24-h period of repeated sampling.

**Methods and materials**

**Study sites**

The sampled lakes and ponds (hereafter referred to as thermokarst lakes) lie in two peatland valleys, at southern and northern locations in subarctic Québec, Canada (Fig. 1; Supporting Information Table S1). Sampling took place during mid- to late summer (July–August) over the course of three field seasons (2012, 2013, 2014). The vegetation in both valleys includes mixed shrubs and semi-aquatic plants, notably Carex aquatilis, Carex rariflora, and Eriophorum angustifolium. Both valleys also contain small stands and isolated individuals of black spruce (Picea mariana).

The southern thermokarst lakes are in the valley of the Sasapimakananiskw River (hereafter SAS valley), 8 km southwest of the village of Whapmagoostui-Kuujjuarapik. Sampling in the SAS valley was at two locations: thermokarst lakes formed alongside palsas north of the river at 55°13′N, 77°41′W (SAS2 lakes), and similar palsa-associated lakes located south of the river at 55°13′N, 77°42′W (SASI lakes). The SAS lakes are located in the sporadic permafrost zone, near the southern limit of subarctic permafrost that underlies <2% of the land surface, and now mostly present in the remaining palsa mounds. Stratigraphic and macrofossil analyses of this valley (Arlén-Pouliot and Bhiy 2005; Fillion et al. 2014) have shown that the organic soil deposits date back to 5800 yr BP and are underlain by marine clays derived from the Tyrell Sea, which flooded the area immediately after ice retreat around 8000 yr BP. The soil stratigraphy shows periods of marsh, bog, and fen development with the most recent moss peatland bog forming between approximately 2100 yr BP to 400 yr BP. Permafrost development took place during the Little Ice Age (LIA), from 400 yr BP to 200 yr BP, with the formation of palsas that likely fused to form an extensive frozen peatland plateau. The valley currently has more than 100 palsas, mostly 3–5 m in height. Permafrost thawing in this region dates from the end of the LIA onwards (ca. 150 yr BP) and resulted in the production of thermokarst lakes and ponds. Some of these have now filled in completely, while others are newly formed or are continuing to expand.

The northern thermokarst lakes are located 280 km north of the SAS valley, 136 km north of the village of Umiujaq and 10 km south of the treeline, in the palsa-rich floodplain of the Boniface River at 57°45′N, 77°20′W (BON lakes). This site is in the subarctic discontinuous permafrost zone, where the permafrost extends over more than 50% of the landscape (Allard and Seguin 1987). Stratigraphic and macrofossil analyses at this site (Bhiy et al. 2007) have shown that peat formed over marine silt from around 6800 yr BP onwards, with subsequent periods of accumulation and decomposition. In contrast to the SAS site where the palsas formed only during the LIA, three phases of permafrost aggradation and palsa development have been identified at BON: 3200 yr, 2000 yr, and 400 yr BP, but with most of the palsas formed during the LIA. These are elongate mounds mostly around 6.5 m high, separated by linear moist depressions including thermokarst lakes. The lakes likely began to form subsequent to the LIA, with expansion over the 20th century.
A comparison of aerial photographs of this region showed that the total surface area of palsa decreased by 23% between 1957 and 2001, while the area of thermokarst lakes and ponds increased by 76%; these changes were much less extensive, however, than at southern, warmer sites in subarctic Québec (Vallée and Payette 2007), and palsa subsidence appears to be proceeding more slowly than in the SAS valley at this northern cooler site.

**Air and ground temperatures**

Air and ground temperatures were measured at minute intervals with YSI 44033 thermistors (Yellow Springs Instruments, Yellow Springs, Ohio, U.S.A.; accuracy ± 0.1°C in the observed range; response time 0.12 s to 10 s), and the averages were recorded each hour in a Campbell CR10 data logger (Campbell Scientific, Edmonton, AB, Canada; accuracy ± 0.02%). The air temperature probes were housed in radiation screens (RM Young 6 plate radiation shield; Campbell Scientific, Edmonton, Alberta, Canada) and positioned 2 m above the ground. The ground temperatures were measured in boreholes drilled into a palsa mound at SAS2 to a depth of 5 m, and in a hole dug into gravel soils to a depth of 1.0 m at BON (Allard et al. 2015).

**Limnological profiling**

Vertical profiles of temperature (± 0.15°C), conductivity (± 0.001 mS cm⁻¹), oxygen (± 0.2 mg L⁻¹), and pH (± 0.2 units) were recorded using a YSI 6000 profiler (Yellow Springs Instruments, Yellow Springs, Ohio, U.S.A.) with all sensors calibrated and verified according to manufacturer specifications. Vertical profiles of photosynthetically available radiation (PAR; ± 1%) were obtained with a LI-192 Underwater PAR Quantum Sensor (LI-COR Biosciences, Lincoln, Nebraska, U.S.A.).

Near-surface lake water samples were collected in 125 mL borosilicate amber glass bottles (Wheaton, Millville, New Jersey, U.S.A.) for total and soluble reactive phosphorus (TP and SRP) and total nitrogen (TN), and analyzed with a Lachat Autoanalyzer according to the methods of Stainton et al. (1977). The detection limits were 0.001 mg TP L⁻¹, 0.5
Dissolved organic carbon (DOC) was analyzed in water samples that were filtered through pre-rinsed 0.2-μm cellulose acetate filters (4°C) and measured by high temperature catalytic combustion in a Total Organic Carbon analyzer (Shimadzu VCPH, TOC-5000A) calibrated with potassium biphthalate (detection limit of 0.05 mg DOC L⁻¹). Phytoplankton biomass was estimated from the chlorophyll a concentration (Chl-a) quantified by high performance liquid chromatography using the methods as in Bonilla et al. (2005).

**Gas concentrations and fluxes**

Water for gas concentrations was sampled with a thinline sampler based on the design shown in Fig. 2.4 of Goltstein and Clymo (1971). This was composed of two 140-mm diameter circular thermoplastic plates that were set 63.5 mm apart. The upper plate was drilled with a 6.4 mm diameter central hole, which was fitted with a tube that extended to a peristaltic pump onboard the boat. The pump was adjusted to a constant speed to allow laminar flow (Reynolds number = 5.6 × 10⁻³) and the sampler was lowered through the water column to collect samples at 0.1 m depth intervals. The water was pumped into 2-liter LDPE bottles that were overfilled three times. Immediately after collection (on site), 20 mL of lakewater was replaced by ambient air using a syringe connected to the bottom of bottle to create a headspace; the bottle was then vigorously shaken for 3 min to extract the dissolved gases, and the headspace was subsampled (Hesslein et al. 1990). Ambient air samples were also taken directly in situ. All samples (10 mL) were stored in 5.9 mL Labco Exetainer vials (Labco Limited, UK) that were sealed with butyl rubber septa. These had been previously flushed with helium and evacuated, and the vials were kept at cool temperature until laboratory analysis.

Concentrations of \( \text{CH}_4 \) and \( \text{CO}_2 \) in the samples were determined by gas chromatography with flame ionization detection (Varian 3800, COMBI PAL head space injection system, CP-Poraplot Q 25m with flame ionization detector). The diffusive flux (\( F \)) at the air–water interface was calculated as Fickian diffusion corrected for low solubility gases (Vachon et al. 2010):

\[
F = k \cdot K_{H} \cdot \Delta P
\]

where \( k \) is the gas transfer velocity of either \( \text{CO}_2 \) or \( \text{CH}_4 \), \( K_{H} \) is the Henry’s constant, and \( \Delta P \) is the air–water gradient of the partial gas pressure. The \( k \) values were calculated using Cole and Caraco (1998) parameterization, and the Schmidt numbers were computed according to the algorithm given in Crusius and Wanninkhof (2003). A correction for near-surface wind mixing was made using the wind model of MacIntyre et al. (2010) adjusted for the energy loss by surface cooling. The horizontal component of wind speed was measured at 2 m above the lake surface with a portable Kestrel weather probe (Kestrel 4500, Nielsen-Kellerman Co., Boothwyn, PA; range 0.4–60.0 m s⁻¹, resolution 0.1, accuracy ± 3% of least significant digit) and averaged over 10 min. These values were then corrected to provide wind speed at 10 m; these extrapolations were made as in Vachon et al. (2010) assuming a logarithmic wind profile.

\( \text{CH}_4 \), \( \text{CO}_2 \) and \( \text{O}_2 \) concentrations were also measured in the water column using an automated continuous GHG monitoring system. This instrument measured gas concentrations with three independent sensors in a gas stream equilibrated with the source water; further details are provided in Bastien et al. (2009) and Laurion et al. (2010).

Gas emissions by ebullition were measured using opaque submerged gas traps. They consisted of a gas–tight inversed funnel with a 0.5 m² opening at the bottom, and a 140 mL plastic syringe (Kendall Monoject™ piston syringe) mounted on the top and closed with a one-way luer-lock valve. Each trap was installed with the opening submerged at a depth of 0.5 m, for 12 h to 36 h (depending on ebullition rates). Two to five traps were deployed per lake on each sampling date, along transects from inshore to offshore sites. At the end of each deployment, the accumulated gas was transferred to 5.9 mL Labco Exetainer vials as above for subsequent analysis.

**Radiocarbon dating**

A gas volume of 100 mL was collected from some of the gas traps, for \( ^{14} \text{C} \)-dating of \( \text{CO}_2 \) and \( \text{CH}_4 \). These samples were transferred to pre-vacuumed 50 mL serum bottles (Wheaton, U.S.A.) closed with a thick butyl rubber stopper (Chemglass Life Science, U.S.A.) and were subsequently analyzed by accelerator mass spectrometry (AMS) at the Keck Carbon Cycle AMS Facility of the University of California, Irvine, California, U.S.A., as in Bouchard et al. (2015). Firstly, \( \text{CH}_4 \) and \( \text{CO}_2 \) were separated by a continuous flow line consisting of purification and combustion traps: \( \text{CO}_2 \) was frozen in liquid nitrogen, then carbon monoxide (CO) was oxidized to \( \text{CO}_2 \) in a 300°C CuO furnace and frozen in a second liquid nitrogen trap, and finally noncondensable \( \text{CH}_4 \) was oxidized to \( \text{CO}_2 \) in a CuO furnace at 975°C (Linberg/Blue M Tube Furnace, Thermo Scientific). The resulting \( \text{CO}_2 \) and \( \text{H}_2\text{O} \) from \( \text{CH}_4 \) combustion were further separated cryogenically on the vacuum line, and the purified \( \text{CO}_2 \) was graphitized using the sealed tube zinc reduction method. This laboratory reports a day-to-day analysis relative error of 2.5%, to 3.1% based on secondary standards, and including extraction, graphitization and AMS measurement. A sediment core was also obtained from Lake SAS1B at the deepest point in the lake with a mini-Glew gravity corer, and the black, floucculent organic over layer (top 0–3 cm) along with three cohesive, deeper layers of organic sediment (3–6, 6–10, and 10–12 cm) were dated. These were also analyzed at the Keck Carbon Cycle AMS Facility. The \( ^{14} \text{C} \) results were expressed as fractions of the modern standard, \( \Delta^{14} \text{C} \), and conventional
radiocarbon age following Stuiver and Polach (1977). All results were corrected for isotopic fractionation.

Results

Air and ground temperatures

The mean annual air temperature measured at the southern SAS site (−3.42°C) was ca. 2.4°C higher than at the northern BON site (Fig. 2), and the number of thawing degree-days was 26% higher (1437 vs. 1144°C-days). The ground temperature thermistors showed that both sites exhibited large seasonal temperature variations (Fig. 2a), and that the palsa permafrost core remained at temperatures below freezing even at the southern (SAS) site (Fig. 2b). The palsa at SAS2 had an active layer that extended down to 0.6 m depth (Fig. 3). Permafrost temperatures varied seasonally below that depth, down to 3.5 m, where the changes were minimal throughout the year (Fig. 3, inset), with the zero annual amplitude depth at 3.55 m. The permafrost temperature at that site was only slightly below freezing (−0.48°C).

Limnological profiling

All of the palsa lakes were thermally stratified on all dates of observation. However, the southern (SAS) lakes showed stronger and more stable stratification, especially towards the end of the open water season. This tendency corresponded to the greater light attenuation in the SAS lakes, with 1% irradiance found around 0.25 m depth vs. almost 1 m in the northern (BON) palsa lakes (Figs. 4a,b,c, 5a,b,c).

Lakes at both northern and southern locations demonstrated a decrease in pH with depth. The BON lakes (Fig. 4d,e,f) ranged from close to neutral pH = 6.8 at the surface (Fig. 4d) to acidic pH < 5.0 at the bottom (Fig. 4f). The southern SAS lakes had more acidic waters throughout the entire water column (Fig. 5d–f), with a maximum value of pH = 6.2 at the surface of the lake SAS2A and a minimum of pH = 4.5 just below the oxycline in that lake (Fig. 5e). There were also large differences in specific conductivity particularly at the bottom of these lakes. While of similar magnitude at the surface (BON = 2.9 ± 0.7 mS m⁻¹; SAS = 4.6 ± 0.3 mS m⁻¹), specific conductivity increased sharply with depth in the SAS lakes (up to 21 mS m⁻¹ in SAS2A lake, Fig. 5e), but stayed around the same order of magnitude (around 1.2 mS m⁻¹) throughout the entire water column towards the bottom of BON lakes (Fig. 4d–f).

Surface waters at both locations were undersaturated in O₂. Despite their shallow depths, all palsa lakes exhibited oxygen stratification during the entire period of observation, with strongest stratification in August in the southern SAS lakes, where hypoxic conditions prevailed from 0.4 m down to the bottom of the water column (Fig. 5g,h,k). Most of the lakes had anoxic bottom waters; an exception was lake BON2A where there was an unusual increase to 5 mg O₂ L⁻¹.

Fig. 2. Air and ground temperatures at northern BON (a) and southern SAS (b) study sites during the study period. The ground temperature records are from thermistors installed at a depth of 1 m near BON palsas or inside a palsa mound at SAS.

Fig. 3. Ground temperature profiles and annual variations in the palsa at site SAS2A. The vertical dashed line marked by the asterisk (*) denotes the palsa permafrost core temperature of −0.48°C. The dashed horizontal lines mark the depth of the permafrost table (0.65 m) and the zero annual amplitude depth (3.55 m). Days of the months label the corresponding temperature profiles in 2013. Inset: Ground temperature at 0.5 m and 3 m depth at site SAS2A in 2012–2014.
near the bottom (Fig. 4k), possibly related to groundwater inflow.

CH$_4$ and CO$_2$ concentrations and fluxes

All of the palsa lakes were organic-rich (> 5 mg DOC L$^{-1}$) and had high concentrations of CH$_4$ and CO$_2$ (Table 1). The surface concentrations of both gases were at least one order of magnitude greater in the southern sites compared to those of northern sites in the less degraded palsa fields, and both exceeded the air equilibrium values on average by one order of magnitude for CO$_2$ and by three orders of magnitude for CH$_4$ (Table 2).

The gas concentrations increased with depth, especially in the southern SAS lakes (Tables 1, 2). The vertical gradients of CH$_4$ and CO$_2$ concentrations down the water column were steeper in the southern palsa lakes than in the northern lakes (Figs. 4, 5). Gas concentrations increased towards the bottom of SAS lakes by 3 (CH$_4$) and 2 (CO$_2$) orders of magnitude respectively (Fig. 5g,h,k). In two of the northern lakes, BON1A and BON2A, CH$_4$ concentrations increased in deeper waters ($Z>2$ m) by ca. 2 orders of magnitude in the anoxic hypolimnion (Fig. 4g,k), while the shallow (1.3 m) lake BON1B exhibited a small decrease in CH$_4$ concentrations with depth (Fig. 4h). CO$_2$ concentrations varied little throughout the water column of lake BON1A (Fig. 4g), but increased by an order of magnitude in lakes BON1B (Fig. 4h) and BON2A (Fig. 4k).

Such gradients at the air-water interface generated large diffusive fluxes of both CH$_4$ and CO$_2$ (Table 3). Large ebullition fluxes were also recorded in the lakes, but the diffusive
fluxes were higher by 1–3 (CH₄) and 3–6 (CO₂) orders of magnitude, in striking contrast to thermokarst lakes studied elsewhere (Table 3). The molar ratios of CH₄ to CO₂ fluxes were highly variable among sites, including within the same lake and valley. For the SAS lakes, the diffusive flux ratios averaged 0.010 (± 0.007, n = 50), while the ebullition flux ratio averaged 1.40 × 10⁻⁶ (± 1.97 × 10⁻⁶, n = 24); for the BON lakes, these values were 0.013 (± 0.005, n = 14) and 0.80 × 10⁻⁶ (± 0.71 × 10⁻⁶, n = 11).

There was a consistent spatial pattern in the gas fluxes, with greater emissions of both CH₄ and CO₂ measured at the deeper, middle part of the lake, and lower values inshore, near the edge of the lake. This is seen in the overall data set of all sampled lakes, where there was a strong relationship between the gas partial pressures at the lake surface and the depth of the water column underneath: surface concentrations of both gases increased as a linear function of water column depth (Fig. 6; for CO₂ vs. lake depth, R² = 0.32, df = 38, p = 0.013; for CH₄ vs. lake depth, R² = 0.41, df = 38, p < 0.0001). This relationship was stronger when the depths for each sample were normalized to the maximum depth of the lake (data not shown): for CO₂ vs. relative lake depth, R² = 0.44, df = 39, p = 0.004; for CH₄ vs. lake depth, R² = 0.71, df = 39, p < 0.0001). There was no correlation between mid-lake diffusive fluxes and lake area; for the seven lakes in Supporting Information Table S1, CH₄ flux vs. area, r = −0.37, df = 5, p = 0.41; CO₂ flux vs. area, r = −0.62, df = 5, p = 0.14.
Sources of variation

Our estimates of CO₂ and CH₄ concentrations and fluxes are subject to multiple sources of variation, including analytical error, temporal variability, spatial variability (within and among lakes), and errors associated with model calculations. As an estimate of analytical and manipulation error, we examined the coefficients of variation (CV) for duplicate samples taken on 5 dates from the mid-lake surface waters of SAS2A. For CO₂, the CVs averaged 6.2% (range = 1.6–13.1%) and for CH₄ they averaged 9.7% (range = 2.1–23.1%). These variations are small relative to the observed inshore-offshore gradients (> 50%; Fig. 6) in gas concentrations.

As an indication of the potential error introduced in sampling at different periods of the day, a further measure of variability is provided by the measurements of gas concentrations in the near surface waters of SAS2A over 24 h, with duplicate samples for headspace determinations taken every 6 h from 18:00 h August 2 to 12:00 h August 3, 2013. The diurnal variation in CO₂ concentrations was small, with an average value for the four sampling times of 405 μmol L⁻¹ (CV = 11%). An ANOVA analysis of all data (duplicate measurements at each of four times) showed a significant effect of time (F₃,₄ = 200, p = 0.007); subsequent Holm-Sidak pair-wise tests showed that only the 0:00 value was significantly different from those at 12:00 h, 18:00 h, and 24:00 h (t = 6.2, 7.0, and 5.4, respectively; df = 2, p ≤ 0.022), and was 16% above the overall mean. The near-surface CH₄ concentrations showed a greater degree of variability among sample times, with a mean of 2.74 μmol L⁻¹ (CV = 25%). The ANOVA analysis for all the CH₄ data also showed a significant effect of time (F₃,₄ = 92, p = 0.029). The subsequent pair-wise tests showed only one significant difference: the 18:00 h sampling was significantly below that at 24:00 h (t = 5.0, df = 2, p = 0.045), and was 25% below the overall mean.

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As measure of variability associated with model parameter estimations, we calculated the effects of a 20% error on gas concentrations and wind speed for estimates of diffusive CH₄ fluxes using Eq. 1. For an increase in CH₄ concentration from 3.60 μmol L⁻¹ to 4.32 μmol L⁻¹, the estimated CH₄
flux would increase by 21% from 4.8 mmol m\(^{-2}\) d\(^{-1}\) to 5.8 mmol m\(^{-2}\) d\(^{-1}\). Changes in wind speed have a slightly greater effect: for a 20% increase from 3.37 m s\(^{-1}\) to 4.04 m s\(^{-1}\), the estimated CH\(_4\) flux would increase by 25% from 4.8 mmol m\(^{-2}\) d\(^{-1}\) to 6.0 mmol m\(^{-2}\) d\(^{-1}\).

**Radiocarbon dates**

Gas concentrations in the bubbles were highly variable, and the two gases differed markedly in their radiocarbon ages, with the CH\(_4\) containing much older carbon (Table 4). There were also large differences between SAS and BON. The \(\Delta^{14}C\) values ranged from 83 to \(-277\) ppt for CH\(_4\) and from 24 to \(-517\) ppt for CO\(_2\), which corresponded to ages up to 2005 yr before present (BP) for CH\(_4\) and 415 yr BP for CO\(_2\) in the southernmost sites, and up to 760 yr BP for CH\(_4\) and 120 yr BP for CO\(_2\) in the northernmost sites (Table 4).

**Dating of surficial lake sediments at SAS1B**

Dating of surficial lake sediments at SAS1B gave values of 2735 yr BP for the top 3 cm of flocculent material, 2775 yr BP for the layer 3 cm to 6 cm, 3000 yr BP for the layer 6 cm to 10 cm, and 3285 yr BP for the bottom layer of the sediment core, from 10 cm to 12 cm.

**Discussion**

Palsa landscapes are common throughout the discontinuous permafrost zone, and descriptions of their geomorphology...
often include mention or photographic evidence of their associated thermokarst lakes; for example, in eastern Canada (Kuhry 2008), western Canada (Turner et al. 2014), western Siberia (Pokrovsky et al. 2014), northern Sweden (Christensen et al. 2012), Iceland (Saemundsson et al. 2012), and interior Alaska (Kanevskiy et al. 2014). However, despite the circum-polar abundance of such lakes, their limnology has been little explored. The results of the present study show that palsa-associated thermokarst lakes in two contrasting regions of permafrost degradation shared a number of distinctive limnological characteristics, including high DOC, darkly colored waters, small surface areas, shallow depths, and strong physical and chemical gradients down their water columns. Concentrations of CO₂ and especially CH₄ were well above air-cal and chemical gradients down their water columns. Concentrations of CO₂ and especially CH₄ were well above air-equilibrium values in the surface waters of all of these lakes, and both gases increased in concentration by orders of magnitude with depth, generally in tandem with a drop in oxygen tensions to hypoxic or anoxic conditions near the bottom sediments.

The highly stratified nature of certain types of thermokarst lakes has been attributed to their short wind fetch for mixing, the strong near-surface absorption of solar radiation, and the accumulation of solutes in the bottom waters due to mineralization processes in the sediments and freeze-concentration effects during winter (Deshpande et al. 2015). Palsa-associated thermokarst lakes represent an extreme set of conditions for these effects, with wind fetches of the order of tens of m or less, and the attenuation of solar irradiance to 1% of surface values within the upper 2 m of the water column (Figs. 4, 5). Additionally, the lakes are partially sheltered from winds coming from the direction of their associated palsa (Fig. 2), which may be 10 or more m wide and several m in height, thereby dampening the extent of wind-induced mixing. The resultant stratification combined with strong respiratory demand for oxygen in these DOC-rich waters would favor the loss of oxygen near the sediments, and anoxic conditions that are conducive to planktonic as well as benthic methanogenesis.

The diffusive fluxes for both gases were at or above the high end of ranges reported for lakes elsewhere, including other types of thermokarst waters (Table 1). This was particularly striking for the CH₄ fluxes, which had maximum values at and above 10 mmol m⁻² d⁻¹, compared with <1 mmol m⁻² d⁻¹ in other types of thermokarst lakes and ponds. In a comparison of lakes in the Canadian boreal zone, south of the present study, diffusive fluxes for CH₄ averaged around 1 mmol m⁻² d⁻¹, but maximum values extended up to 20 mmol m⁻² d⁻¹ in some lakes (Rasio et al. 2015). In that study, the CH₄ fluxes increased with decreasing lake area, which the authors attributed to shallower water columns, the increased tendency towards bottom anoxia and the higher ratio of sediments to water volume, which are all also features of the small thermokarst lakes sampled in the present study.

Although the diffusive CH₄ fluxes measured in the present study are large relative to other types of thermokarst lakes, they are not unusually high relative to peatland lakes and ponds further to the south. In a recent set of global estimates for northern waters, Wik et al. (2016) report that peatland lakes (outside the permafrost zone) have diffusive fluxes that are on average 2.6 times those measured in thermokarst lakes, and their compiled diffusive rates for this peat-based class of waterbodies (mean of 5.4, range of 1.6 mmol CH₄ m⁻² d⁻¹ to 7.6 mmol CH₄ m⁻² d⁻¹) overlap with our data from the SAS valley peatland thermokarst lakes: mean of 4.8, range of 1.4 mmol CH₄ m⁻² d⁻¹ to 22.5 mmol CH₄ m⁻² d⁻¹. In part this may reflect the order of magnitude greater soil carbon concentrations found in peatlands, including permafrost peatlands, relative to yedoma soils. The small size of peatland water bodies in general is also likely to be a factor contributing to their high rates of methanogenesis and CH₄ emission. All of the lakes in the present study were below the remote-sensing cut-off of 0.002 km² in the analyses by Wik et al. (2016), and there is increasing attention to the importance of small ponds that can have exceptionally high rates of greenhouse gas emissions (Holgerson and Raymond 2016).

One factor that may contribute to greater CH₄ fluxes from smaller lakes is the close physical coupling between the central regions of the lake and the littoral zone (Rasio et al. 2015), where fluxes can be much higher than offshore. These high littoral fluxes have been attributed to warmer sediment temperatures inshore and the effects of wave action on sediment resuspension (Hoffmann et al. 2010, 2013). In thermokarst lakes, the erosion of the surrounding permafrost may also play a role. For example, in a series of Siberian and Alaskan thermokarst lakes, a major source of CH₄ was from point source ebullition fueled by carbon released from permafrost degradation at the lake margins; offshore sources of CH₄ were also substantial, and derived from older sediment strata (Walter et al. 2006). In the present study, however, ebullition fluxes of CH₄ and CO₂ were likely limited to background ebullition (sensu Walter et al. 2006), while the diffusive emissions increased with overall water column depth, and maximum emission occurred in the central, deepest waters of the lake (Fig. 6). Several features of palsa-associated thermokarst lakes may give rise to this reverse pattern. Firstly, the lakes typically have steep sides, and they lack extensive shallows that could differentially heat up to provide warmer, inshore habitats for methanogens. Secondly, the short fetch of the lakes means that little wave production occurs, and wave-induced sediment stirring is likely to be minimal. Finally, the transfer of eroded permafrost soils to the deeper part of the lake by sediment focusing, and the low oxygen conditions at the bottom of this central region, would likely favor the most rapid rates of methanogenesis, resulting in bubble production and transfer of gases to the surface, ultimately the atmosphere.
The production of quasi-stable microbubbles in supersaturated water has been identified as a potentially important pathway of CH$_4$ transfer from lakes to the atmosphere (Prairie and del Giorgio 2013; McGinnis et al. 2014), and could operate in the studied palsa lakes, where CH$_4$ concentrations are orders of magnitude above air equilibrium. Such microbubbles would rise at minimal rates through the water column, and their transfer to the lake surface would be largely dependent on turbulent mixing. However, if the bubbles formed through nucleation in the highly supersaturated bottom waters and then continued to grow (Boudreau et al. 2001), they could eventually achieve sufficient flotation velocities that would allow them to float up into the surface waters. The shaking of water samples for head space analysis would extract the gas from such microbubbles (Prairie and del Giorgio 2013), contributing to the higher gas concentrations, hence diffusive flux estimates, over the deeper sites of the lakes.

Although ebullition is recognized to be a major mechanism for gas transfer from thermokarst lakes to the atmosphere (Walter et al. 2006), it is difficult to accurately quantify. Bubble production is highly variable in space and time, with bubble size being one of several important factors influencing this variability (DelSontro et al. 2015). The ebullition fluxes that we measured here were unexpectedly lower in magnitude (Table 1). Our highest measured ebullition rate (0.8 L m$^{-2}$ d$^{-1}$) lies near the low end of the ebullition range for thermokarst lakes as compiled in Wik et al. (2016): 0.4–8.2 mmol CH$_4$ m$^{-2}$ d$^{-1}$. The bubble CH$_4$ concentrations (Table 4) overlapped with other studies: in percentage terms, the mean CH$_4$ concentration in all our ebullition data was 16%, with a maximum up to 83%; this compares with the range of 4–107% (all lakes) and 34 to >80% (Arctic lakes) as compiled in Walter Anthony et al. (2010). However, the volumetric gas fluxes were low. The mean flux for all of our ebullition measurements was 0.046 L m$^{-2}$ d$^{-1}$, with large site-to-site variability (range = 0.006 L m$^{-2}$ d$^{-1}$ to 0.118 L m$^{-2}$ d$^{-1}$). The mean ebullition flux for thermokarst lakes reported in Wik et al. (2016) was 87.5 mg CH$_4$ m$^{-2}$ d$^{-1}$, which for an average CH$_4$ concentration in the bubbles of 40% would convert to 0.306 L m$^{-2}$ d$^{-1}$, more than a factor of 6 higher than in the present study. Our measurements covered a wide range of lakes and within-lake locations, and were made during the warmest time of year when CH$_4$ production and ebullition rates might be expected to be high. It is possible that our sampling funnels failed to capture sparsely distributed active point sources such as seeps and hotspots (Walter Anthony et al. 2010) within the lakes, although these were never observed despite many sampling visits, and are not apparent in automated camera images of the lakes during freeze-up (Pienitz et al. 2016). The traps were deployed for only a short period of time, and possibly missed important periods of bubble production. For these reasons, the ebullition fluxes (Table 3) could be substantially underestimated. It is also possible that palsa-associated thermokarst lakes, underlain by a continuous layer of organic-rich, ancient peat, have more homogeneous sediments than yedoma systems, and that ebullition is dominated by low background rates rather than by point sources.

The dominance by diffusive fluxes (which could also include microbubble fluxes; Prairie and del Giorgio 2013) was especially due to the markedly high concentrations of CH$_4$ and CO$_2$ in the surface waters. The high CH$_4$ concentrations are consistent with microbiological studies in these lakes, which show an unusually high percentage of aerobic methanotrophs among the bacterial RNA sequences (Crevecoeur et al. 2015). These organisms would be favored by the high CH$_4$ availability, however their activity along with that of anaerobic CH$_4$ oxidizers, appears to be insufficient to deplete the continuous supply of CH$_4$.

The present study was conducted during warm, open-water conditions over three summer periods, but other times of year may also be important for GHG production, or their release to the atmosphere. These lakes are covered by ice for more than 6 months of the year, and the liquid water remaining under the ice becomes anoxic soon after freeze-up (Deshpande et al. 2015). Such conditions would be favorable for ongoing methanogenesis over winter, leading to large accumulations of CH$_4$ as well as CO$_2$ that could be subsequently released to the atmosphere after ice-out, as found at more southern latitudes (e.g., Ducharme-Riel et al. 2015). However, heating and stratification occur in early spring in these waters, and it is not until convective mixing in fall that the water column is fully mixed (Deshpande et al. 2015); the winter as well as summer hypolimnetic accumulation may only then be fully vented as a storage flux to the atmosphere. Such emissions may occur as short but intense bursts that would be difficult to capture by punctual sampling visits (Ducharme-Riel et al. 2015). This release of stored gases (Table 4) is likely to be considerable relative to the daily fluxes; for example the integrated water column values at SAS2A (2.8 m) were 0.4 mol CH$_4$ m$^{-2}$ and 8 mol CO$_2$ m$^{-2}$. Expressed as potential storage fluxes, these are equivalent to 40 d and 8 d, respectively, of the maximum measured diffusive fluxes from this lake.

The thermokarst lakes in the southern part (SAS) of the study region had orders of magnitude greater CH$_4$ and CO$_2$ concentrations compared to the northern sites (BON), and the calculated gas fluxes differed accordingly. Several features may contribute to these large differences, including the organic-rich permafrost deposits at the SAS site that are now degrading rapidly and collapsing into the lakes. Northern Québec has experienced rapid warming over the last few decades, and lakes in the SAS valley have become more numerous and larger (Bhiry et al. 2011; Fillion et al. 2014). As shown by the thermal profile data (Fig. 3), the SAS2A palsa still retains a permafrost core, but its temperature is only
slightly below freezing and it is vulnerable to further warming. The BON site also lies on thick peat deposits (Bhiry et al. 2007), however it remains within the northern discontinuous permafrost zone where we measured cooler air temperatures and a lesser number of thawing degree days. Ongoing warming could lead to the BON lakes following a trajectory towards SAS conditions, with lake expansion, permafrost carbon release into the waters and increased microbial activity leading to increased GHG emissions, amplified also by higher autochthonous production of organic matter under warmer catchment and lake conditions. Increased DOC concentrations would lead to the greater absorption of solar irradiance by surface waters, as observed in the SAS lakes, and this would exacerbate thermal stratification and anoxia. Climate warming is also likely to have a direct effect on microbial heterotrophs and CH₄ producers, especially given that methanogenesis has been observed to respond strongly to increased temperatures, across multiple scales, from microbial cells to ecosystems (Yvon-Durocher et al. 2014). However, if climate warming leads to colder hypolimnia and stronger water column stability persisting for longer periods each year, this effect could be slowed down.

The ¹⁴C dating analysis was based on a limited number of samples, and will require further sampling and corroboration in the future. Nevertheless, results indicated a striking difference in radiocarbon age between CH₄ and CO₂ in the ebullition samples from the palsas. This implies that in these thermokarst systems, CH₄ oxidation is not the primary source of CO₂, and that the emitted CO₂ and CH₄ are at least partly derived from separate carbon pathways, as reported for lakes in the Canadian High Arctic (Bouchard et al. 2015). The highly variable ratio of CH₄ to CO₂ fluxes (CV ≥ 40%) also indicates that these two processes are largely uncoupled, as does the difference in the timing of maximum concentrations during the diurnal sampling. At BON, and to a greater extent at SAS, the CH₄ was at least partially derived from old carbon, likely associated with materials that were only recently mobilized by permafrost thawing.

The differences in CH₄ dates between SAS and BON are consistent with differences in the extent of permafrost degradation at the two sites. In both regions, peat bogs developed soon after the marine recession, at around 5800 yr BP at SAS and 4000 yr BP at BON (Bhiry et al. 2011). At BON, three phases of palsa formation have been identified, at 3200 yr BP, 2000 yr BP, and 400 yr BP (Bhiry et al. 2007), while at the lower latitudes of SAS and the James Bay region, palsas did not form until towards the end of the Little Ice Age, around 200 yr BP (Tremblay et al. 2014). Palsa degradation likely started earlier in the warmer conditions at SAS relative to BON, and is currently progressing rapidly (Bhiry et al. 2011; Fillion et al. 2014), mobilizing older peat bog carbon. The oldest date for the CH₄ in SAS approaches that of the surficial sediments of SAS lakes (2735 yr BP), implying recent mobilization of ancient organic carbon delivered to the bottom of the lake. The younger age at BON is consistent with a lesser extent of thawing, and mobilization of carbon in younger permafrost strata than at SAS. Dilution with carbon from modern sources such as macrophytes, phytoplankton and recent terrestrial vegetation will have reduced the average carbon age in the CH₄ samples. At both locations, the CO₂ carbon had a younger age, implying that more recent photosynthesis produced the carbon sources for aerobic bacterial heterotrophy in these waters, in contrast to the old carbon used in anaerobic methanogenesis.

In summary, our results show that peatland thermokarst lakes share a number of distinct limnological features including high DOC concentrations and strong emissions of CH₄ and CO₂. These freshwater ecosystems are widely distributed throughout discontinuous permafrost peatlands, and they differ from the yedoma class of thermokarst lakes, which lie in less organic soils. The measurements obtained in the present study are too limited in time and space to provide a robust estimate of the total greenhouse gas flux from all thermokarst lakes in northern permafrost peatlands, but given the vast area of this landscape type (1.4 million km²; Tarnocai et al. 2009) and the high diffuse emission rates that we measured, there is a need to measure flux rates throughout the subarctic for this type of non-yedoma system. The much greater gas fluxes from SAS relative to BON waters implies that old permafrost carbon stocks will be increasingly thawed and converted to CH₄ in peatland thermokarst lakes as the southern limit of permafrost continues to move northwards with ongoing climate change.

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