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1 Journal: Biogeochemistry

2 *Title:* Using radon to quantify groundwater discharge and methane fluxes to a shallow,

3 tundra lake on the Yukon-Kuskokwim Delta, Alaska

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20 Abstract: Northern lakes are a source of greenhouse gases to the atmosphere and contribute substantially to

- 21 the global carbon budget. However, the sources of methane (CH₄) to northern lakes are poorly constrained
- 22 limiting our ability to the assess impacts of future Arctic change. Here we present measurements of the natural

23 groundwater tracer, radon, and CH₄ in a shallow lake on the Yukon-Kuskokwim Delta, AK and quantify

- 24 groundwater discharge rates and fluxes of groundwater-derived CH₄. We found that groundwater was
- 25 significantly enriched (2000%) in radon and CH₄ relative to lake water. Using a mass balance approach, we

26 calculated average groundwater fluxes of 1.2 ± 0.6 and 4.3 ± 2.0 cm d⁻¹, respectively as conservative and upper

- 27 limit estimates. Groundwater CH₄ fluxes were 7 24 mmol m⁻² d⁻¹ and significantly exceeded diffusive air-
- 28 water CH_4 fluxes $(1.3 2.3 \text{ mmol m}^{-2} \text{ d}^{-1})$ from the lake to the atmosphere, suggesting that groundwater is an
- 29 important source of CH₄ to Arctic lakes and may drive observed CH₄ emissions. Isotopic signatures of CH₄
- 30 were depleted in groundwaters, consistent with microbial production. Higher methane concentrations in
- 31 groundwater compared to other high latitude lakes were likely the source of the comparatively higher CH₄
- 32 diffusive fluxes, as compared to those reported previously in high latitude lakes. These findings indicate that
- deltaic lakes across warmer permafrost regions may act as important hotspots for CH₄ release across Arctic
- 34 landscapes.
- 35 *Keywords:* radon-222; methane; tundra; groundwater; wetland; subarctic;

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- 39 samples (B2, U1, U3, B3) (Ludwig et al. 2017a) and to Jordan Jimmie for stream discharge data and sampling.
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43 1. Introduction

Perennially frozen ground, also known as permafrost, underlies up to 25% of the land in the Northern Hemisphere (Brown et al. 2002). On average, 16% of the terrestrial permafrost landscape is covered by water (Lehner and Döll 2004), and in some areas, like on the Yukon-Kuskokwim Delta in Alaska, it exceeds 30% (US Fish & Wildlife Service 2002). These aquatic systems are closely linked to the terrestrial environment through hydrology. Intense Arctic warming and permafrost thaw may alter the tight connection between terrestrial and aquatic ecosystems. For example, permafrost thaw is causing changes in aquatic systems by changing transit times and shifting flow paths between organic and mineral-rich soils (Vonk et al. 2015).

51 Groundwater is a source of water and solutes to marine and freshwater systems. In temperate and tropical 52 environments, groundwater discharge has been well-documented as a source of nutrients (Charette and 53 Buesseler 2004; Paytan et al. 2006) and carbon (Beck et al. 2007; Richardson et al. 2017; Kim and Kim 2017) 54 to surface waters. In Arctic environments, there are few studies on groundwater discharge, many of which lack 55 information on quantified fluxes of solutes like carbon and nitrogen (see (Lecher 2017) for a review). Permafrost 56 limits most groundwater flow to the shallow, thawed active layer (Williams 1970; Woo 2012). Potential 57 groundwater supply through sediment beds also depends on the presence or absence of continuous permafrost. 58 Taliks-or perennially unfrozen sediments often found beneath lakes and streams-allow for groundwater 59 exchange between a lake and underlying sediments (Woo 2012). Expanding taliks in a warming climate are 60 expected to enhance exchange between lakes, rivers and underlying aquifers via groundwater supply (Walvoord 61 and Kurylyk 2016).

62 Many lakes in polar regions are known to be substantial sources of carbon to the atmosphere ((Wik et al. 63 2016) and references therein), which may be influenced by groundwater-surface water interactions. In addition 64 to delivering dissolved organic carbon that can be mineralized to CO₂ and CH₄, groundwater may directly 65 transport carbon dioxide (CO_2) and methane (CH_4) that was produced in active layer soils to lakes (Kling et al. 66 1992) where CH_4 can be oxidized or released to the atmosphere. Paytan et al. quantified CH_4 transport to a lake 67 in the Arctic suggesting that carbon-rich soils in the northern latitudes, and the release of carbon from permafrost 68 thaw, provide fuel for CH_4 production (Schuur et al. 2008; Natali et al. 2015; Paytan et al. 2015). With the 69 expected shift to greater subsurface flow due to warming combined with future permafrost thaw (Walvoord and 70 Striegl 2007; Bring et al. 2016; Walvoord and Kurylyk 2016), groundwater may become an increasingly 71 important source of CH₄ to lakes in permafrost environments. This is important in the context of the global

carbon cycle because lakes in the Arctic constitute a substantial portion of Arctic CH₄ sources and represent
6% of global natural CH₄ emissions (Wik et al. 2016).

74 In this study, we investigated the importance of groundwater as a source of CH_4 to a shallow tundra lake. 75 Radon (²²²Rn) was used as a natural geochemical tracer of groundwater discharge (Charette et al. 2008; Dimova 76 and Burnett 2011; Dimova et al. 2013), an approach that is advantageous in regions like northern wetlands 77 because it captures groundwater flow despite their low landscape gradients and microtopographic features that 78 inhibit the use of traditional hydrologic methods such as seepage meters and water table elevation measurements 79 (Morison et al. 2017b). In contrast, ²²²Rn allows for the integration of these heterogeneities. As radon is 80 produced naturally from decay of uranium-series radionuclides in sediments and soils, it is an ideal tracer of all 81 groundwater sources including those present above the permafrost in the seasonally-thawed active layer, in 82 permafrost, and in subpermafrost aquifers (Woo 2012). We used a mass-balance approach (Charette et al. 2008) 83 to quantify groundwater discharge rates and estimate groundwater-derived CH4 fluxes to the lake and compared 84 them to measured air-water diffusive fluxes and stable isotopes.

85 2. Materials and Methods

86 2.1. Study site

87 The study site (Fig. 1; 61.264 °N, 163.246 °W) is located 93 km NW of Bethel, AK in the Yukon Delta 88 National Wildlife Refuge (YDNWR). Fieldwork was conducted over two field seasons from July 1 - 13, 201789 and June 30 - July 10, 2018. The majority of groundwater and lake sampling was conducted in 2017. Gas 90 exchange coefficients and CH_4 air-water fluxes were measured in 2018. Average air temperatures in this region 91 (1981 – 2010 average for Bethel; US National Weather Service) are -0.8 °C annually, -14.4 °C in January, 13.4 92 °C in July, with above freezing average monthly air temperatures from April to October. Annual precipitation 93 is ~470 mm, with 60 mm occurring in July on average. The average temperatures in July 2017 and 2018, 94 respectively, were 14.4 °C and 13.9 °C. The recorded precipitation in July 2017 was 92 mm and in July 2018 95 was 38 mm (US National Weather Service). The study site is located in a zone of continuous to discontinuous 96 permafrost (Brown et al. 2002) that is moderate in thickness (~180 m) (Ferrians Jr. 1965) with taliks underlying 97 most wetlands and water bodies. Thaw depth was 30 - 40 cm in July 2017 in areas without taliks. The sediments 98 beneath the thick organic layer in this region were deposited in the early Pleistocene (Wilson et al. 2015). This 99 region is characterized by polygonal peat plateaus beside low-lying wetlands. The maximum elevation in this 100 region is approximately 15 meters above sea level and the minimum elevation is approximately 8 m. The 101 elevation of the lake surface and neighboring peat plateaus are 13 and 15 m, respectively.

102 Lakes and ponds occupy about one third of the YDNWR in surface area (US Fish & Wildlife Service 103 2002). Most lakes in this region have a maximum depth of <1-3 m (Bartlett et al. 1992) and range widely in 104 surface area from several square meters to several square kilometers. The lake in this study, colloquially termed 105 "Landing Lake," has an average depth of 0.53 ± 0.03 m and a surface area of approximately 0.36 km² and is 106 therefore representative of the numerous small, high latitude lakes of the YDNWR. Much of the lake's 107 watershed is in a region of the YDNWR that experienced a wildfire in 2015, as visible by satellite imagery and 108 evident in the field by a lack of vegetation and the presence of leftover charred materials (Fig. 1). Fire frequency 109 has been found to increase with warming in northern Alaska (Higuera et al. 2011) and on the Yukon-Kuskokwim 110 Delta (Sae-Lim et al. 2019), and can cause permafrost thawing, vegetation shifts, and carbon release (Loranty 111 et al. 2016). Although fire effects were not the focus of this study, statistical tests were performed when enough 112 data was available, and potential implications are discussed (Section 4.1.3). Only one surface water channel 113 was connected to Landing Lake at its southeast corner; it was ~0.33 m wide and ~0.15 m deep, and discharge 114 flowed away from the lake at 0.003 m³ s⁻¹.

115 2.2. Sample collection

116 Surface water and groundwater samples for all analyses were collected on July 1 - 12, 2017 and June 30 117 - July 10, 2018 (Fig. 1, Table 1). Samples from active layer soils and lake and pond bottom sediments were 118 collected in 2017 for analysis and incubation experiments in the laboratory. A lake sediment sample 119 (groundwater symbol next to the weather station, Fig. 1) was collected from the top 5 cm using gloved hands, 120 stored in a clean plastic bag, and frozen until analysis (~4 months). Active layer soils (n = 4, 0 - 30 cm) were 121 cored using a sharpened steel coring barrel, sample tube and hand drill, and then frozen within 48 hours of 122 collection. Samples were thawed for biogeochemical analyses (available online: (Ludwig et al. 2017a)) ~2 123 weeks after sample collection and refrozen for ~4 months before radionuclide analyses. Air temperature, wind 124 speed and direction, and rainfall rates were collected every 12 minutes using a weather station (AcuRite 5-in-1 125 Weather Sensor) placed ~5 m above the lake surface on a peninsula (Fig. 1). At each surface water and 126 groundwater sampling event, we measured temperature, dissolved oxygen, and electrical conductivity (YSI 6-127 Series Sonde (2017), YSI ProPlus multiparameter probe (2018)). Instruments were calibrated immediately prior 128 to fieldwork and in the field.

129Lake water samples (2017, n = 18, Table 1) for 222 Rn were collected in two ways. A RAD AQUA system130(Durridge Inc.; (Schubert et al. 2012)) was used for 222 Rn collection for 17 of the samples. One sample, WP4,131was collected in a calibrated 2-L plastic bottle with no headspace that was analyzed within four hours. One 100-

L surface water sample (5 20-L "cubitainers") was collected to estimate ²²²Rn supported by its parent, ²²⁶Ra. At the sampling sites in both years, dissolved CH₄ was collected by vigorously shaking 30-mL of the water sample with 30-mL of ambient air for 60 seconds. The headspace was then transferred into pre-evacuated 12-mL Exetainer vials until slightly over-pressurized. Two separate gas samples were collected for separate analyses of CH₄ concentration and δ^{13} CH₄, respectively. Samples for water isotope (δ^{2} H and δ^{18} O) analysis were also collected in 2017 in 4.5-mL glass vials with no headspace.

Groundwater samples (2017, n = 7, Table 2) were collected from the active layer at 20-40 cm depth below the soil surface using a push-point piezometer (MHE Products, Inc.) and peristaltic pump with gas impermeable tubing. Groundwater samples were limited by the maximum thaw depth of ~40 cm. Samples for ²²²Rn were collected in 250-mL glass bottles (RAD H2O, Durridge) that were flushed by at least three volumes of sample water and then sealed with no headspace. The same sampling procedures described above for dissolved CH₄ and δ^{2} H and δ^{18} O isotopes were used for groundwater samples. One set of water samples was also collected from the southeastern stream discharging Landing Lake for CH₄ and water isotopes (Fig. 1, Table 1).

145 Spatial and temporal variation in CH₄ flux was examined across Landing Lake in 2018 to provide context 146 for groundwater fluxes of CH₄. Seven chambers were deployed for a 24-hour measurement period around the 147 lake. Gas samples from chamber headspace and dissolved surface water were collected upon chamber 148 deployment and after 12-24 hours (Bastviken et al. 2004). Flux rates were calculated from the difference in 149 initial and final concentrations of CH_4 in the chamber, assuming the flux decreased over time in response to a 150 decreasing concentration gradient between the lake water and chamber headspace (Bastviken et al. 2004). To 151 compare the impact of different flux estimate approaches, instantaneous CH₄ fluxes (averaged triplicate 152 measures, each 5 min duration) were measured during the same period. The CH₄ concentrations in the chamber 153 headspace were measured instantaneously using a Los Gatos Research Ultraportable Greenhouse Gas analyzer, 154 and the increase in concentration over the sampling period was used to calculate chamber fluxes by fitting a 155 linear slope to the data.

- 156 2.3. Sample analysis
- 157 2.3.1. Radioisotopes

Surface water measurements of ²²²Rn were conducted in two ways. At all lake sampling locations (except WP4) ²²²Rn was measured using a radon-in-air monitoring system (RAD7, Durridge) connected to a drying unit, spray chamber (RAD AQUA, Durridge, Inc.) and bilge pump. The temperature in the spray chamber was recorded using a stainless-steel temperature probe and data logger (HOBO U12-008, ONSET). At each station, the detector was run for 45 - 75 minutes, including 30 minutes of equilibration. Uncertainties (standard errors) were $\sim 3 - 5\%$ for each sample for the integrated measurement periods. The amount of ²²²Rn in water was calculated using the measured temperature in the spray chamber and its solubility (Dimova and Burnett 2011). At station WP4, ²²²Rn was measured in a 2-L sample at the field site using the Big Bottle accessory (Durridge) for the RAD7. The uncertainty or standard error for this method was ~16%.

167 Groundwater ²²²Rn activities were measured using two different techniques. In the field, groundwater 168 samples (n = 7) were analyzed using the RAD H2O accessory (Durridge, Inc.) within 24 hours of collection. 169 Activities were corrected for decay between collection and measurement times. Uncertainties were 9-45 % 170 $(1\sigma, \text{standard error})$. To determine equilibrium ²²²Rn activities in groundwater as additional endmembers in the 171 model, soils (n = 4) and lake sediments (n = 1) were incubated in the laboratory (Corbett et al. 1997; Chanyotha 172 et al. 2014). One soil sample (B2, Table 2) was collected >5 km away from the lake, but was included as an 173 endmember due to its similar bulk density to the average bulk density of all other burned soils (Table 3). Radon 174 activities were measured using a radon emanation approach (Key et al. 1979). Efficiencies were determined 175 using a set of radium-fiber standards containing 20 dpm ²²⁶Ra (NIST-certified SRM#4967A). Uncertainties 176 were 3 - 15% (1 σ , standard error). The ²²²Rn activities were converted into groundwater endmember activities 177 using porosity and bulk density (Section 2.3.4, Table 3) (Chanyotha et al. 2014).

178 Experiments in the laboratory were carried out with lake bottom sediments to determine the diffusive flux 179 of ²²²Rn to the lake. Wet sediments were incubated in gas tight flasks with air stones and radium-free water and 180 connected in a closed loop with two charcoal columns as described by Chanyotha et al. (2016). Radon activities 181 were monitored for 10-20 hours. The exponential ingrowth of ²²²Rn activity was linearly approximated (errors 182 of 4 - 10% at 10 - 25 hours) (Chanyotha et al. 2016). This slope was used to calculate the diffusive flux of 183 ²²²Rn. Leakage of the system over 20 hours was corrected for using a radium-fiber standard containing 20 dpm 184 ²²⁶Ra. A second method was used in which lake bottom sediments were incubated and analyzed with the radon 185 emanation approach described above (Key et al. 1979; Corbett et al. 1997; Chanyotha et al. 2014). The total 186 equilibrium ²²²Rn activity was multiplied by the decay constant and normalized to the area of the flask to obtain 187 an estimate of the diffusive flux of ²²²Rn to overlying water. The standard error of 3 trials was reported as the 188 uncertainty. Blanks were run using the same experimental setups and subtracted from any reported values. To determine the amount of ²²⁶Ra dissolved in the Landing Lake that was supporting ²²²Rn in the water 189

190 column, the ~100-L sample was filtered onsite at <1 L min⁻¹ through a Mn-impregnated acrylic fiber to extract

191 the radium (Moore and Reid 1973). The fiber was analyzed for the activity of ²²²Rn supported by ²²⁶Ra. The

- 192 fiber was ashed, packed in a polystyrene vial, and sealed with epoxy to prevent ²²²Rn loss (Charette et al. 2001).
- 193 The activity of ²²⁶Ra was measured by gamma spectrometry in a well-type germanium gamma detector
- 194 (Canberra). The detector was calibrated using a ²²⁶Ra standard (NIST-certified SRM#4967A) in the same
- 195 geometry as the sample. The standard error (1σ) was reported as the uncertainty in this measurement.
- 196 Table 1. ²²²Rn activities and dissolved CH₄ concentrations measured at Landing Lake. Latitude (Lat) and
- 197 longitude (Lon) are in decimal degrees. Depth represents depth of water sample collection. Cond. =
- 198 conductivity; δ^{13} C of methane are presented relative to Pee Dee Belemnite (PDB). H₂O stable isotopes
- 199 reported relative to Vienna Standard Mean Ocean Water (VSMOW). Average value ± standard error is

Station	Туре	Lat.	Lon.	Depth cm	Cond mS cm ⁻¹	O_2 mg L^{-1}	Water Temp <i>°C</i>	²²² Rn <i>dpm</i> <i>m</i> ⁻³	$\begin{array}{c} \mathrm{CH}_4 \\ \mu mol \\ L^{-l} \end{array}$	${}^{\delta^{13}}C_{CH4} \\ {}^{\%}_{\%}$	$\overset{\delta^2 H_{H2O}}{\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \$	$\overset{\delta^{18}O}{\overset{_{H2O}}{_{\%}}}$
WP4	Lake	61.264	-163.246	10	ND^{a}	ND	ND	2700	ND	ND	ND	ND
WP6	Lake	61.265	-163.244	45	0.133	14.7	15.7	2640	ND	ND	ND	ND
WP7	Lake	61.263	-163.244	30	0.066	13.7	17.3	1660	ND	ND	ND	ND
WP8	Lake	61.264	-163.243	30	0.051	14.9	15.6	1140	0.1	-45.7	-67.3	-7.5
WP17	Lake	61.268	-163.240	50	0.072	13.9	16.5	1350	1.4	-47.2	-67.1	-7.1
WP18	Lake	61.267	-163.238	50	0.067	10.0	17.4	1370	1.3	-47.1	-67	-7.3
WP19	Lake	61.268	-163.237	43	0.067	10.7	17.7	1051	1.3	-47.0	-67	-7.3
WP20	Lake	61.270	-163.237	40	0.065	9.5	17.9	1390	1.5	-47.3	-67.1	-7.4
WP21	Lake	61.269	-163.241	44	0.062	12.8	18.1	1080	1.5	-46.3	-67.1	-7.3
WP22	Lake	61.268	-163.243	45	0.062	9.2	18.3	1020	1.3	-41.3	-66.9	-7.4
WP31	Lake	61.263	-163.240	35	0.089	10.0	18.3	1620	1.5	-48.0	-65.7	-7.2
WP32	Lake	61.264	-163.239	35	0.087	10.3	18.9	1394	1.8	-44.6	-65.8	-7.2
WP33	Lake	61.265	-163.238	35	0.080	10.4	19.9	1008	1.6	-46.0	-65.7	-7.2
WP34	Lake	61.266	-163.240	50	0.077	11.1	19.4	570	1.5	-48.7	-65.5	-6.5
WP35	Lake	61.267	-163.243	45	0.072	11.1	19.7	1040	1	-49.2	-65.6	-7.3
WP42	Lake	61.267	-163.246	50	0.076	10.5	15.9	1550	1.7	-47.8	-67.5	-7.5
WP46	Lake	61.265	-163.247	40	0.058	10.6	18.9	1640	6.1	-51.4	-68.8	-8.5
WP47	Lake	61.264	-163.248	50	0.059	10.7	18.7	1650	2.9	-48.5	-67.9	-7
Stream	Stream	61.260	-163.241	10	0.020	6.4	8.5	ND	5.5	-47.0	-87.2	-11.5
Avg.					0.073	11.4	17.9	1400	1.8	-47.1	-66.8	-7.3
±					0.004	0.4	0.3	300	0.5	0.6	0.2	0.1

200 reported. Stream sample not included in average.

 a ND = no data.

202 Table 2. ²²²Rn activities and dissolved CH₄ concentrations measured in groundwater and incubated soils.

203 Evidence of the 2015 wildfire is noted for each sample. Depth is below the soil surface. Cond. =

- 204 conductivity. δ^{13} C of dissolved CH₄ are presented relative to Pee Dee Belemnite (PDB). H₂O stable
- 205 isotopes reported relative to Vienna Standard Mean Ocean Water (VSMOW). Average value ± standard
- error is reported.

Station	Туре	Fire ^a	Lat.	Lon.	Depth cm	Cond mS cm ⁻¹	O_2 mg L^{-1}	Temp °C	²²² Rn <i>dpm</i> <i>m</i> ⁻³	CH_4 μmol L^{-l}	${}^{\delta^{13}}C_{CH4} \\ {}^{\%0}$	δ ² H ‰	δ ¹⁸ O ‰
WP5	GW^b	Ν	61.263	-163.245	30	ND ^c	ND	ND	48000	ND	ND	ND	ND
WP29	GW	Y/N	61.270	-163.237	30	0.085	2.8	5.1	15000	550.8	-73.6	-94.8	-13.2
WT7-3	GW	Y/N	61.270	-163.237	30	0.030	3.3	3.7	35000	7.8	-50.5	-90.9	-13.1
WP43	GW	Y	61.267	-163.247	30	0.186	5.1	16.8	36000	563.2	-58.0	-95.7	-13.2
WP45	GW	Y	61.265	-163.238	40	0.215	5.7	10.4	29000	612.2	-65.2	-106	-14.6
WP30	GW	Ν	61.270	-163.239	25	0.118	3.2	10.8	ND	456.2	-50.0	-94.0	-14.0
WT8-2	GW	Y	61.270	-163.236	30	0.043	0.9	12.6	ND	25.1	-73.9	-92.1	-12.9
Bottom	Inc ^d	Y/N	61.264	-163.246	0-5	ND	ND	ND	38000	ND	ND	ND	ND
B2-T1	Inc	Y	61.321	-163.243	0-30	ND	ND	ND	5000	ND	ND	ND	ND
U1-T3	Inc	Ν	61.258	-163.247	0-30	ND	ND	ND	2000	ND	ND	ND	ND
U3-T1	Inc	Ν	61.270	-163.237	0-30	ND	ND	ND	1000	ND	ND	ND	ND
B3-T2	Inc	Y	61.271	-163. 235	0-30	ND	ND	ND	32000	ND	ND	ND	ND
Avg.						0.113	3.5	9.9	24000	370	-61.9	-95.6	-13.5
<u>+</u>						0.031	0.7	2.0	5000	110	4.4	2.2	0.3

 a Y/N = yes or no for samples collected within the 2015 fire. b GW = groundwater. c ND = no data. d Inc =

Incubated soil or sediment. See ref. (Ludwig et al. 2017a) for more details on soil samples B2, U1, U3 and B3.

209 2.3.2. Methane

210 Methane concentrations were analyzed using a greenhouse gas chromatograph (Shimadzu GC-2014) at 211 the Woods Hole Research Center, and stable carbon isotopic composition of CH₄ was measured at Northumbria 212 University using a Delta V Plus IRMS interfaced to a Trace Gas Pre-Concentrator and Gas Bench (Thermo 213 Scientific). Each isotope measurement run contained three standards (Liso1, Tiso1, Hiso1; Isometric 214 Instruments), run in full at the beginning and end, with individual standards interleaved throughout (precision 215 <0.5‰). Both CH₄ concentration and isotopic signatures were blank corrected for atmospheric contamination 216 assuming the global mean surface atmospheric CH₄ concentration of 1.8 ppm and δ^{13} C-CH₄ of -47.2% 217 (Warwick et al. 2016) and reported relative to Pee Dee Belemnite (PDB).

In 2018, air-water diffusive fluxes (F_{atm}) of CH₄ from the lake were measured directly via the instantaneous and 24-hr measurement period methods described above. From these data, we calculated the gas transfer coefficient (k_x) from the following equation:

$$k_{x} (m d^{-1}) = F_{atm} (mol m^{-1} d^{-1}) / ([X]_{water} (mol m^{-3}) - [X]_{air}),$$
(1)

where $[X]_{water}$ is the measured concentration of dissolved CH₄ in the lake, and $[X]_{air}$ is the concentration of CH₄ expected in the lake when in equilibrium with the ambient air (Emerson and Hedges 2008). The equilibrium concentration of CH₄ was calculated using lake temperature, ambient air CH₄ concentration, and Bunsen solubility constants (Wiesenburg and Guinasso 1979). Two models of gas exchange coefficiencients (k_x) (Crusius and Wanninkhof 2003; Holgerson and Raymond 2016) for the lake was used to derive air-water diffusive fluxes of CH₄ concentrations for Landing Lake in 2017 given similar average wind speed observations for the two years.

228 2.3.3. $\delta^{18}O$ and $\delta^{2}H$

To examine hydrologic processes and sources of water into the lake, $\delta^{18}O$ and $\delta^{2}H$ stable isotope values of lake water, stream, and groundwater samples were measured at Northumbria University using a Water Isotope Analyzer (LGR LWIA-24d, San Jose, USA). Ratios were measured to a precision of 0.2‰ for $\delta^{2}H$ and 0.03‰ for $\delta^{18}O$ and reported relative to Vienna Standard Mean Ocean Water (VSMOW).

233 2.3.4. Soil characterization

234 Porosity and bulk density were measured in order to calculate equilibrium groundwater radon (22Rn) 235 activities (Table 3) (Chanyotha et al. 2014). Soil and sediment were sampled volumetrically, dried at 60° C 236 (organic soils) or 100° C (sediments) for 48 hours, and bulk densities (B_D) calculated as dry mass/volume. 237 Landing Lake bottom sediment characteristics were averaged for the top 5 cm (measured in 0.5 cm intervals, 238 (Ludwig et al. 2017c)). For porosity measurements, soils and sediments were dried in an oven at 50 °C. Dry 239 sediment/soil was gently packed into a pre-weighed, volume-calibrated test tube. Deionized water was added 240 to the test tube until it just covered the soil surface. The mass of the dry soil and test tube was subtracted from 241 the new mass of the test tube, soil and water. Porosity (φ) was then calculated as follows:

$$\varphi =$$
[Water added (g) / Density of water (g cm⁻³)] / Volume of soil (cm³). (2)

After measuring the equilibrium 222 Rn activities (A_{222. TOTAL}) via radon emanation (see section 2.3.1), the

243 following equation was used to calculate groundwater (GW) ²²²Rn activities (Chanyotha et al. 2014) :

GW ²²²Rn (dpm m⁻³) =
$$[A_{222, \text{TOTAL}} / \text{wet mass of soil (g)}] \cdot B_D (g \text{ cm}^{-3}) \cdot (1 \text{ cm}^3 / 1 \text{ x } 10^{-6} \text{ m}^3) / \phi.$$
 (3)

Other soil and sediment characteristics were measured (C, N, moisture, etc.) and can be found online (Ludwiget al. 2017a, c).

246 2.4. Statistical analyses

Linear regressions were fit to CH_4 and water stable isotope data with a 99% confidence interval. ANOVAs were used to report *p*-values indicating the significance of the relationship. These analyses were performed 249 across all samples and with the two groups of surface waters and groundwaters, but only statistically significant 250 relationships (p < 0.05) were reported.

251 Although the effects of wildfires on groundwater hydrology and CH_4 are beyond the scope of this study, 252 statistical tests (t-test, two-sample, unequal variances) were performed with sample data to test the potential 253 impacts of the 2015 wildfire. First, the relationship between fire and activities of ²²²Rn in groundwater samples 254 was examined across all groundwater samples taken during the field campaign, including those not adjacent to 255 Landing Lake (Table 4). The same statistical test was performed for CH₄ in burned and unburned groundwaters. 256 The impact of fire on soil bulk density was also tested using a two-sample t-test, assuming unequal variances 257 for soils collected in 2017 (see data online: (Ludwig et al. 2017a)). Only soils from peat plateaus in 2017 were 258 included to eliminate other environmental variables.

259 Table 3. Measured soil and sediment characteristics used in the incubation experiments. Visible evidence

260 of the 2015 wildfire is noted for each sample. Sediments were collected from the top 5 cm. Soils were

261 collected from the 0 - 30 cm. Sample names started with B represent burned soils; U, unburned.

Sample name	Туре	Fire in 2015?	Porosity	Dry bulk density $g \ cm^{-3}$	GW ²²² Rn dpm m ⁻³
Bottom Sediment	Landing Lake Sediment	Y/N	0.82	0.45	38000
B2-T1	Soil	Y	0.86	0.19	4800
U1-T3	Soil	Ν	0.93	0.09	1600
U3-T1	Soil	Ν	0.90	0.13	1000
B3-T2	Soil	Y	0.74	0.37	32000

262 *3. Results*

263 Water quality data from Landing Lake and nearby groundwaters are presented in Tables 1 and 2. The 264 conductivity of surface water and groundwater was on average 0.073 ± 0.004 and 0.113 ± 0.031 mS cm⁻¹, 265 respectively. All measurements in Landing Lake indicated that it was well oxygenated and thermally well 266 mixed. The average dissolved oxygen concentration was 11.4 ± 0.4 mg L⁻¹ (115% saturation). Water 267 temperatures were 15.6 to 19.9 °C with an average of 17.9 °C. Groundwater had a lower average dissolved 268 oxygen concentration of 3.5 ± 0.7 mg L⁻¹ and a lower average temperature of 9.9 ± 2.0 °C. The stream outlet 269 of Landing Lake had an intermediate dissolved oxygen concentration of 6.4 mg L⁻¹ and a temperature of 8.5 °C, 270 which was similar to that of groundwater.

271 3.1. Radioisotopes

272	Radon activities were ~20 times more enriched in groundwater than in surface water samples (Tables 1
273	and 2). Groundwater samples in burned soils did not significantly differ with respect to ²²² Rn compared to other
274	soils ($p = 0.84$, Table 4). However, soils collected in 2017 (see data online: [30]), did significantly differ ($p < 1000$
275	0.01) in bulk density between recently burned (mean = 0.170 g cm ⁻³ , $\sigma^2 = 0.024$ g cm ⁻³) and unburned peat
276	plateaus soils (mean = 0.087 g cm ⁻³ , σ^2 = 0.005 g cm ⁻³). In the lake, ²²² Rn activities were on average 1,400 ±
277	300 dpm m ⁻³ (range = $570 - 2,700$ dpm m ⁻³) while groundwater activities were 24,000 ± 5,000 dpm m ⁻³ (range
278	= $1,000 - 48,000$ dpm m ⁻³ , Tables 1 and 2). The highest surface water activities were near the southern and
279	western edges of the lake, and the lowest activities were in the center of the lake (Fig. 2a). The lowest radon
280	activities in groundwater were for the three soil samples incubated in the laboratory (Table 2). The measured
281	activity of 226 Ra in lake water was 24 ± 2 dpm m ⁻³ (standard error) and was a minor contributor to the 222 Rn
282	inventory in the lake. The diffusive flux of 222 Rn from bottom sediments was 850 ± 90 dpm m ⁻² d ⁻¹ and 640 ±
283	90 dpm m ⁻² d ⁻¹ as found using the hourly flux method (Chanyotha et al. 2016) and equilibration method (Corbett
284	et al. 1997), respectively. The average of the two techniques was 740 ± 140 dpm m ⁻² d ⁻¹ .

Table 4. Groundwater samples collected in 2017 (including those near Landing Lake and other lakes) and the
 associated ²²²Rn activities and methane concentrations. Evidence of fire in 2015 is indicated by Y/N.

Sample name	Туре	Fire in 2015?	Lat.	Lon.	Depth cm	²²² Rn <i>dpm m</i> - ³	CH4 µmol L ⁻¹
WP43	GW^1	Y	61.267	-163.247	30	36000	563.2
WP45	GW	Y	61.265	-163.238	40	29000	612.2
WT8-2	GW	Y	61.270	-163.236	22	ND^2	25.1
B1-WP27	GW	Y	61.284	-163.247	36	18000	520
B2-WP28	GW	Y	61.273	-163.230	55.5	36000	418.6
B3-WP37	GW	Y	61.284	-163.259	37	19000	ND
B4-WP39	GW	Y	61.284	-163.259	52	32000	628.6
B5-WP40	GW	Y	61.288	-163.262	52	26000	2.9
B2-T1	Inc ³	Y	61.321	-163.243	0-30	4800	ND
B3-T2	Inc	Y	61.271	-163.235	0-30	32000	ND
WP5	GW	Ν	61.263	-163.245	30	48000	ND
UB1-WP10	GW	Ν	61.258	-163.246	45	30000	635.3
UB1-WP15	GW	Ν	61.258	-163.246	36	25000	517.3
UB2-WP25	GW	Ν	61.321	-163.238	35	65000	98.4
WP30	GW	Ν	61.270	-163.239	25	ND	456.2
U1-T3	Inc	Ν	61.258	-163.247	0-30	1600	ND
U3-T1	Inc	Ν	61.270	-163.237	0-30	1000	ND
Fire, average						26000	395.8
σ^2						$1.07 \ge 10^8$	7.3 x 10 ⁴
No fire, average						28000	426.8
σ^2						6.39 x 10 ⁸	$5.4 \ge 10^4$

287 288

 1 GW = groundwater. 2 ND = no data. 3 Inc = incubation (See section 2.3.1) for description of incubation methods.

289 3.2. Methane

290 Like radon, dissolved CH₄ was more enriched in groundwater (~200x) than in lake water (Tables 1 and 291 2). Groundwater samples in burned soils did not significantly differ in dissolved CH₄ compared to unburned 292 soils (p = 0.85, Table 4). In the lake, CH₄ varied from 0.1 to 6.1 µmol L⁻¹ (Fig. 2b) with an average concentration 293 of $1.8 \pm 0.5 \mu$ mol L⁻¹ (Table 1). The highest concentrations were at stations WP46 and WP47 at the southwestern 294 edge of the lake (Fig. 2b). The lowest concentrations were in the center of the lake. Dissolved CH₄ 295 concentrations in groundwater varied over a larger range from 8 to 612 µmol L⁻¹, and the average groundwater 296 concentration of CH₄ was $370 \pm 110 \mu$ mol L⁻¹ (Table 1). Dissolved CH₄ in the stream was ~5.5 μ mol L⁻¹, 297 intermediate between average lake waters and groundwaters.

298 Dissolved CH₄ in groundwater was on average more depleted in 13 C than surface water (-61.9 ± 4.4‰ and 299 -47.1±0.6‰, respectively; Tables 1 and 2). The most depleted δ^{13} C value of -51.4‰ in surface water was found 300 at station WP46, coinciding with the highest concentration of CH_4 observed in the lake (Fig. 2b). The stream 301 outlet had a δ^{13} C value of -47.0%, similar to lake waters. There was a significant negative relationship between 302 δ^{13} C and logged CH₄ concentrations in all samples (δ^{13} C = -5.98 log [CH₄, µmol L⁻¹] – 46.9‰., R² = 0.729, p 303 < 0.01, Fig. 3); however, this was largely driven by differences between lake and groundwater samples, and 304

there was no detected relationship between δ^{13} C and dissolved CH₄ within each group (p > 0.01).

305 In 2018, the average CH₄ concentration in Landing Lake was $1.1 \pm 0.4 \mu$ mol L⁻¹ (Table 6), similar to the 306 average in 2017 of 1.8 \pm 0.5 μ mol L⁻¹. Air-water CH₄ fluxes measured using the instantaneous and 24-hr 307 measurement period methods were 13.5 \pm 3.3 mmol m⁻² d⁻¹ and 2.7 \pm 1.0 mmol m⁻² d⁻¹, respectively. The 308 calculated gas exchange coefficients, k₆₀₀, using the instantaneous and 24-hr measurement period flux methods 309 were 1.32 ± 0.50 m d⁻¹ and 0.251 ± 0.014 m d⁻¹, respectively (Table 6).

3.3. $\delta^{18}O$ and δ^2H 310

311 Stable isotopes of H and O in groundwater were more depleted than lake water (Tables 1 and 2, Fig. 4). 312 Lake water $\delta^2 H$ and $\delta^{18} O$ values were -66.8 \pm 0.2‰ and -7.3 \pm 0.1‰, respectively, and groundwater $\delta^2 H$ and 313 δ^{18} O values were -95.6 ± 2.2‰ and -13.5 ± 0.3‰, respectively. The stream draining Landing Lake had 314 intermediate $\delta^2 H$ and $\delta^{18}O$ values, respectively, of -87.2‰ and -11.5‰. When $\delta^2 H$ values were plotted as a 315 function of δ^{18} O values (Fig. 4), groundwater samples (Table 2) fell close to the Global Meteoric Water Line 316 (Craig 1961), and were represented by following best-fit line: $\delta^2 H_{H2O} = 6.87 (\delta^{18}O) - 2.90\%$ (R² = 0.70, p =

317 0.04). Stable isotope values for all lake and pond samples collected in 2017 (see data online: (Ludwig et al. 318 2017b)) were represented by the following relationship: $\delta^2 H_{H2O} = 4.31(\delta^{18}O) - 36.55\%$ (R² = 0.96, *p* << 0.01). 319 Landing Lake surface water samples fell below the GMWL line, but within the range of all lake samples. The 320 stream sample was more depleted than Landing Lake surface waters and was on the line represented by all lakes 321 and ponds.

322 4. Discussion

323 4.1. Radon sources and sinks

324 Consistent with previous studies, ²²²Rn was much more enriched in groundwater than in surface water (Dimova and Burnett 2011; Dimova et al. 2013; Paytan et al. 2015). Groundwater ²²²Rn activities (1.000 – 325 326 48,000 dpm m⁻³) were less than those observed in sandy, Floridian soils (~170,000 dpm m⁻³) (Dimova et al. 327 2013) and in silty soils near Toolik Lake, Alaska (~490,000 dpm m⁻³) (Paytan et al. 2015). The lower activity 328 of ²²²Rn in soils near Landing Lake was likely due to the organic-rich soils that are low in mineral content (by 329 weight) than most sandy or silty soils and therefore lower in its parent isotope ²³⁸U that produces ²²²Rn. The 330 surface water activities (Fig. 2a, 570 - 2,710 dpm m⁻³) were similar to those reported in a small lake in Florida 331 $(1200 - 4800 \text{ dpm m}^{-3})$ (Dimova and Burnett 2011) and Toolik Lake in Alaska (2900 - 5700 dpm m}^{-3}) (Paytan 332 et al. 2015).

To quantify groundwater discharge to Landing Lake using ²²²Rn as a tracer, we constructed a mass balance model that includes all sources and sinks of radon to the lake (Fig. 5). Similar models have been used to study groundwater discharge in both marine and lacustrine environments (Corbett et al. 1997; Dulaiova et al. 2010; Dimova and Burnett 2011; Dimova et al. 2013). The spatial and temporal heterogeneity of groundwater discharge precludes direct quantification; therefore, we use a "flux-by-difference" approach (Charette et al. 2008). Assuming steady state over a few weeks, the change in ²²²Rn over time should be equal to zero, and the sources must be balanced by the sinks:

$$0 = d^{222} Rn/dt \ (dpm \ m^{-2} \ d^{-1}) = F_{222,GW} + F_{226} + F_{benthic} - F_{atm} - \lambda \cdot I_{222} - F_{stream} - F_{recharge}.$$
 (2)

The sources in this equation other than groundwater ($F_{222,GW}$) of ²²²Rn include alpha-decay of ²²⁶Ra in the water column (F_{226}) and diffusive inputs from lake bottom sediments ($F_{benthic}$). We found no surface water streams entering the lake. The sinks in this model include loss to the atmosphere via gas exchange (F_{atm}), decay ($t_{1/2}$ = 3.82 days), which is equivalent to the inventory of ²²²Rn (I_{222}) multiplied by its decay constant ($\lambda = 0.181$ days⁻ 1), and loss via the stream draining Landing Lake (F_{stream}). Recharge of lake water into downgradient soils and sediments ($F_{recharge}$) was not measured, although its potential impact on the mass balance is discussed below. Sources of uncertainty for each mass balance model term are described in Table 7. Generally, the largest sources
 of uncertainty in ²²²Rn mass balances are natural variability in endmember ²²²Rn activities and atmospheric
 evasion, as well as mixing with offshore waters for coastal zones (Burnett et al. 2007).

349 4.1.1. Sinks of ²²²Rn: gas exchange, decay, streams, and recharge

350 To determine the loss of radon via gas exchange, two empirical models were compared to field 351 measurements of the gas exchange coefficient at Landing Lake. The air-water flux of radon was calculated 352 using Equation 1 (Emerson and Hedges 2008). In this case, [X]_{water} and [X]_{air} are the activities of radon 353 measured in the lake and the activity expected when the lake is in equilibrium with the atmosphere, respectively. 354 We assumed that atmospheric ²²²Rn was negligible relative to the lake ²²²Rn ([X]_{air}=0). The gas exchange 355 coefficient, k_{Rn}, was first estimated based on relationships to temperature (Wanninkhof 1992) and wind speed 356 (Crusius and Wanninkhof 2003). For this mass balance, we used the linear relationship for the SF_6 gas exchange 357 coefficient as a function of wind speed $(0 - 5 \text{ m s}^{-1}, 20 \text{ °C})$ for a lake similar in surface area (0.13 km²) to 358 Landing Lake (0.36 km²) (Crusius and Wanninkhof 2003). Then, k_{SF6} (n = 14, (Crusius and Wanninkhof 2003)) 359 was converted to k_{Rn} for the average water temperature in this study (17.9 ± 0.3 °C) using the appropriate 360 Schmidt numbers (Sc(SF₆, 20 °C) = 956, Sc(Rn, 20 °C) = 883, Sc(Rn, 17.9 °C) = 991) (Wanninkhof 1992; 361 Crusius and Wanninkhof 2003). This resulted in the following best-fit linear relationship as a function of wind 362 speed, u: $k_{Rn}(17.9 \text{ °C}, \text{ m d}^{-1}) = 0.28 \cdot u(\text{m s}^{-1}) - 0.13$, which had a slope error of 19%, similar to the 20% error 363 that is typical for empirical wind-speed relationships (Dimova and Burnett 2011). We used the average water 364 temperature (17.9 \pm 0.3 °C, n = 18) and wind speed (3.83 \pm 0.05 m s⁻¹, n > 1000) over the 12-day study period, 365 which resulted in an average gas exchange coefficient of $k_{Rn} = 1.1 \pm 0.2$ m d⁻¹ and atmospheric flux (F_{atm}) of 366 $1,600 \pm 300$ dpm m⁻² d⁻¹ (upper limit of gas exchange, Fig. 6).

In another study, an empirical relationship based on surface area, rather than wind speed, across 309 small lakes and ponds over a range of latitudes was used to estimate gas exchange (Holgerson and Raymond 2016). To apply this to Landing Lake, we used the gas exchange coefficient for surface areas of $0.1 - 1 \text{ km}^2$ ($k_{600} = 0.80 \text{ m d}^{-1}$) (Holgerson and Raymond 2016) and the Schmidt number for radon at the average lake temperature of $17.9 \pm 0.3 \text{ °C}$ (Sc = 991) (Wanninkhof 1992) to obtain a second estimate for the gas exchange coefficient of $k_{Rn} = 0.62 \text{ m d}^{-1}$. This produced a lower estimated atmospheric flux (F_{atm}) of 900 ± 200 dpm m⁻² d⁻¹ (lower limit, Fig. 6).

We compared these literature-derived estimates of the gas transfer coefficient with those obtained from direct measurements of gas exchange in 2018 on Landing Lake via 2-min (instantaneous) and 24-hr 376 measurement floating chambers (Sections 2.2, 2.3.2, Table 6). The coefficients (k_{600} , 12.5 °C) were 1.3 ± 0.5 377 and 0.25 ± 0.01 m d⁻¹, respectively, according to each method. When the coefficients were converted for radon 378 at the average lake temperature in 2017, it resulted in values of 1.0 ± 0.4 and 0.20 ± 0.01 m d⁻¹, respectively for 379 k_{Rn} (17.9 °C). CH₄ concentrations and weather conditions were similar in 2017 and 2018, so we expect these 380 gas exchange coefficients to apply to both years. The instantaneous method resulted in gas transfer coefficients 381 similar to the wind speed model, but was likely influenced by ebullition, resulting in overestimates of the 382 diffusive flux, and thus the gas transfer coefficient. The 24-hr measurement period fluxes were less than both 383 the surface area model and wind speed model, which may have been due to the lower temperature of Landing 384 Lake in 2018 compared to 2017 and the shielding of surface water from wind due to the chamber. To encompass 385 uncertainty due to gas exchange in the ²²²Rn mass balance, we used the surface area model as a conservative 386 estimate and the wind speed model as an upper limit estimate of groundwater fluxes.

To calculate radon loss from the lake due to decay, we first estimated the inventory of radon in the lake by multiplying the average depth (0.53 ± 0.03 m) by the average activity of ²²²Rn in the lake (1400 ± 300 dpm m⁻ 3, Table 1). The flux due to decay is the product of this inventory and the decay constant (λ ·I₂₂₂), and was equal to 130 ± 10 dpm m⁻² d⁻¹ (Fig. 6). Of the combined sinks for ²²²Rn, decay accounted for $8 \pm 1\%$ and $13 \pm 1\%$, while atmospheric exchange was $92 \pm 18\%$ and $87 \pm 17\%$ of total losses of ²²²Rn, for the upper limit and conservative gas exchange estimates, respectively.

393 We were not able to directly measure the loss of ²²²Rn due to recharge or the single stream outlet. However, 394 if we assume negligible evaporation and negligible stream outflow to determine the maximum impact of 395 recharge on the mass balance, we expect that lake water would recharge into adjacent wetland areas at the same 396 rate as groundwater influx $(\sim 1 - 4 \text{ cm d}^{-1})$ with a ²²²Rn activity equal to average lake water (1400 dpm m⁻³). The 397 222 Rn loss rate for this process would be 20 – 60 dpm m⁻² d⁻¹, or only 2 – 3% of the combined losses due to 398 decay and gas exchange. In the case of the stream outlet, discharge was ~ 0.003 m³ s⁻¹, which is equivalent to 399 0.07 cm d⁻¹ when integrated over the lake's area, as with the other mass balance terms. If the ²²²Rn activity of 400 the stream is assumed to be that of average lake water (1400 dpm m³), then the ²²²Rn loss would be 1.0 dpm m⁻ 401 2 d⁻¹, or 0.06 – 0.10% of the combined sinks of decay and gas exchange. Therefore, both recharge and the stream 402 outlet are considered negligible sinks in the ²²²Rn mass balance, well within the uncertainty of most of the model 403 terms (Table 7).

404 *4.1.2. Sources of*²²²*Rn: dissolved*²²⁶*Ra, sediments, groundwater*

405 Potential sources of ²²²Rn in this system other than groundwater are production via decay of dissolved 406 226 Ra and diffusive inputs from bottom sediments (Fig. 5). We first calculated the dissolved inventory of 226 Ra 407 by multiplying the measured activity of ²²⁶Ra in the lake (24 dpm m⁻³) by the average depth (0.53 m). The inventory of 222 Rn supported by 226 Ra is equivalent to the dissolved inventory of 226 Ra (12 ± 1 dpm m⁻²) 408 409 multiplied by the decay constant of 222 Rn. This results in a flux (F₂₂₆) of 30 ± 2 dpm m⁻² d⁻¹ (Fig. 6). In our 410 steady state model where we assume that sources are equal to sinks, the input of ²²²Rn from ²²⁶Ra can only 411 account for 2 - 3% of the radon inputs to the lake, consistent with other lake ²²²Rn budgets (Corbett et al. 1997; 412 Dimova et al. 2013).

413 The diffusive input of ²²²Rn, which was measured in the laboratory using Landing Lake sediments, agreed 414 well between the two methods. The short-term measurement over 10 - 20 hours resulted in a greater flux than 415 the equilibration method, likely due to the larger concentration gradient between sediment and overlying water 416 for shorter incubation periods. Because the short-term measurement approximates the decay as a linear function, 417 up to 10% error is expected in addition to any experimental error. In the mass balance, we used the average of 418 the two techniques $(740 \pm 140 \text{ dpm m}^{-2} \text{ d}^{-1})$ for the sediment-water diffusive flux (Fig. 6). The flux was less than 419 that of freshwater lake sediments from Cambodia (2040 dpm $m^{-2}d^{-1}$) (Chanyotha et al. 2016), although this is 420 expected because radon is derived from natural uranium in minerals (Charette et al. 2008), and the lake 421 sediments in the YDNWR have a low mineral content. The ²²²Rn diffusive flux accounted for 42 and 72% of 422 sources in the radon budget for the upper limit and conservative estimates, respectively (Fig. 6). This 423 contribution from diffusion is higher than most lake budgets (Dimova et al. 2013, 2015); since Landing Lake is 424 only ~0.5 m deep, the ratio of bottom sediment area to lake volume is relatively large, which likely explains 425 why diffusion is estimated to be a major contributor to the Landing Lake ²²²Rn inventory.

Together, diffusive inputs and dissolved ²²⁶Ra decay account for 44 to 73% of the sources in the mass balance. Assuming negligible transport of ²²²Rn out of Landing Lake via recharge and streams (Section 4.1.1), groundwater must be the missing source that contributes 27 to 56% of radon to the lake inventory (Fig. 6).

429 4.1.3. Quantifying groundwater fluxes

With measurements of groundwater endmembers, one can convert the ²²²Rn fluxes into groundwater fluxes and volumetric discharge estimates. The remaining 25 ± 10 to $58 \pm 24\%$ of the ²²²Rn inventory was 300 ± 100 to 1000 ± 400 dpm m⁻²d⁻¹, for the conservative and upper limit estimates, respectively (Fig. 6). In the following equation (Charette et al. 2008),

$$F_{GW} (m d^{-1}) = F_{222,GW} (dpm m^{-2} d^{-1}) / A_{GW} (dpm m^{-3}),$$
(3)

434 F_{222.GW} is the flux of ²²²Rn via groundwater and A_{GW} is the activity of ²²²Rn in groundwater. There is a 435 significant amount of variability in the groundwater samples when considering both field samples and 436 incubations.²²²Rn activities in groundwater at Landing Lake are likely controlled by the mineral content of soils, 437 which is known to increase with depth in peatlands (Morison et al. 2017a). Using the average endmember (Table 438 2, 24,000 \pm 5,000 dpm m⁻³), the ²²²Rn flux via groundwater (300 \pm 100 to 1000 \pm 400 dpm m⁻²d⁻¹) and Equation 439 3, we calculated groundwater fluxes of 0.012 ± 0.006 and 0.043 ± 0.020 m d⁻¹ (1.2 ± 0.6 , 4.3 ± 2.0 cm d⁻¹, Table 440 5), respectively, for conservative and upper limit estimates. If we use the highest activity endmember (48,000 441 dpm m⁻³), the groundwater flux is 0.6 ± 0.3 to 2.1 ± 0.9 cm d⁻¹ (Table 5), for conservative and upper limit 442 estimates, respectively. Since these groundwater fluxes were calculated using the average ²²²Rn inventory for 443 the whole lake surface, they represent inflow averaged over the lake's area. We only have one sample for lake 444 bottom sediment porewater (222 Rn = 38,000 dpm m⁻³) that may be representative of possible subpermafrost 445 groundwater, which is higher in activity that the average groundwater endmember. If subpermafrost 446 groundwater were a significant source of water to this lake, it would likely have a ²²²Rn activity similar to that 447 of our porewater sample, which is greater than our average groundwater endmember but less than the highest 448 activity endmember; therefore, it would not impact our estimate of ²²²Rn-based groundwater discharge fluxes. 449 Another factor that could influence these groundwater fluxes is the impact of the 2015 wildfire. Fire did not 450 seem to have a significant impact on ²²²Rn activities in groundwater, but it did result in significantly higher bulk 451 densities (Table 3). Higher soil density usually lowers hydraulic conductivity, which could cause the 452 groundwater fluxes to be lower in fire-affected areas of the watershed. A dedicated process study would be 453 needed to truly determine the environmental impacts of fire on groundwater hydrology.

The volumetric input of groundwater to the lake of $4,000 \pm 2,000$ to $15,000 \pm 7000$ m³ d⁻¹ was estimated by multiplying the groundwater flux $(1.2 \pm 0.6 \text{ to } 4.3 \pm 2.0 \text{ cm } \text{d}^{-1})$ by the lake area $(3.6 \times 10^5 \text{ m}^2)$. Such a discharge rate would flush the lake about 3 - 7% by volume per day, equivalent to a residence time of 15 - 53days (Table 2). For lakes in the US with depths <2 m, residence times on average are 30 - 300 days (Brooks et al. 2014), which agrees well with the residence times calculated here.

459 Table 5. Estimates of groundwater fluxes, residence times, and methane fluxes for Landing Lake

460 compared to other studies. Average and high activity endmembers refer to the concentrations of radon in

461 groundwater.

Lake name	GW ^a flux cm d ⁻¹	Residence time days	$\begin{array}{c} {\rm GW} \ [{\rm CH}_4]^{\rm b} \\ \mu mol \ L^{-l} \end{array}$	GW CH ₄ flux mmol m ⁻² d ⁻¹	Lake [CH ₄] $\mu mol L^{-1}$	Air-water CH ₄ flux ^c mmol m ⁻² d ⁻¹
Landing Lake average endmember	1.2 ± 0.6 to 4.3 ± 2.0	12 – 44	370 (8 - 612)	4 ± 2 to 16 ± 7	1.8 ± 0.3	1.3 – 2.3 (1.3 – 5.7)
Landing Lake high activity endmember	0.6 ± 0.3 to 2.1 ± 0.9	25 - 88	370 (8 - 612)	2 ± 1 to 8 ± 3	1.8 ± 0.3	1.3 – 2.3 (1.3 – 5.7)
Toolik Lake (Paytan et al. 2015; Garcia-Tigreros Kodovska et al. 2016)	1.4 ± 0.9	ND ^d	8 – 35 (0.01 – 150)	0.1 – 0.7	0.02 - 0.8	0.06 - 0.2
Northern peatland ponds (n = 38) (Wik et al. 2016)	ND	ND	ND	ND	ND	7 (2 – 10)

462 ^a GW = groundwater. ^b Average listed along with minimum and maximum in parentheses. Other values listed

with entire range of estimates or as average \pm standard deviation. ^c Estimated air-water fluxes calculated using Equation 1 and measured air-water fluxes via 24-hr measurement period flux chambers in 2018 listed in parentheses. See Table 6 in Appendix for details. ^d ND = no data.

466 Unless the lake volumes were increasing over the study period, any groundwater inputs to the lake must 467 be lost to surface water flow, wetland recharge, or evaporation. Surface water flow was estimated to drain only 468 0.5% of Landing Lake's volume per day, and we had no means to quantify recharge from the lake to the 469 subsurface. If the talik beneath the lake does not penetrate the permafrost completely, the main recharge pathway 470 for water flow would be through the wetland areas near the lake, visible as a darker green color just north and 471 west of the lake (Fig. 1), or through outlet streams. The elevation difference between the plateaus and low-lying 472 areas, such as the lake surface and wetlands, was approximately 2 meters, likely enough to support some level 473 of hydrologic outflow. 474 Stable isotopes (δ^{18} O and δ^{2} H of H₂O) provide quantitative evidence for evaporation at Landing Lake 475 (Fig. 4). All lakes and ponds sampled in 2017 (Ludwig et al. 2017b) fall on the following best-fit line: $\delta^2 H_{H2O}$ 476 $= 4.31(\delta^{18}O) - 36.55\%$ (R² = 0.96), which we define as the Local Evaporation Line (LEL). A slope of 4.31 is

within modeled slopes of 4 - 6 for lakes at 60°N (Gibson et al. 2008) and measured slopes of 4.1 - 7.1 in Canadian lakes and wetlands (Gibson et al. 2005). Landing Lake surface waters fell on the LEL and seem to be

479 more impacted by evaporation than the majority of the lakes and ponds sampled, which is expected since 480 Landing Lake had the highest surface area and a similar depth compared to the other sampling sites. The 481 intersection of this evaporation line with the meteoric water line indicates the source of water to the lake (Fontes 482 1980) was locally sampled active layer groundwaters. Stable isotopes in groundwaters were close to the GMWL

- +62 1960) was locally sampled active rayer ground waters. Stable isotopes in ground waters were close to the GM WE
- 483 and therefore were similar to precipitation. Another study of water stable isotopes in also found that summer
- 484 precipitation was the major source of water to the active layer on the Alaskan tundra (Throckmorton et al. 2016).

485 These data show that evaporation was a significant loss of water during the study period, although the exact 486 percentage is not quantifiable with the available data.

487 *4.2. Methane in Landing Lake*

488 Using the radon-derived groundwater fluxes ($F_{GW} = 1.2 \pm 0.6$ to 4.3 ± 2.0 and 0.6 ± 0.3 to 2.1 ± 0.9 cm d⁻ 489 ¹) and dissolved CH₄ concentration measurements, we estimated groundwater fluxes of CH₄ to Landing Lake 490 from the following equation,

$$F_{CH4,GW} \text{ (mmol } m^{-2} d^{-1}\text{)} = F_{GW} \text{ (m } d^{-1}\text{)} \cdot [CH_4]_{GW} \text{ (mmol } m^{-3}\text{)},$$
(4)

491 in which $F_{CH4,GW}$ is the flux of CH₄ to Landing Lake via groundwater, and [CH₄]_{GW} is the concentration of 492 CH₄ in groundwater (average = 370 μ mol L⁻¹). The groundwater flux of CH₄ to Landing Lake (F_{CH4,GW}) for 493 July 2017 was 4 ± 2 to 16 ± 7 mmol m⁻² d⁻¹ (High ²²²Rn endmember: 2 ± 1 to 8 ± 3 mmol m⁻² d⁻¹, Table 5). A 494 study at Toolik Lake, AK conducted during July in 2011 and 2012, the same time of year as this study, included 495 similar methods to determine radon-derived groundwater fluxes (Paytan et al. 2015). The groundwater flux of 496 CH₄ to Landing Lake is an order of magnitude greater than to Toolik Lake (Table 5, 0.1 - 0.7 mmol m⁻² d⁻¹), 497 despite having similar groundwater fluxes (Table 5, 1.2 ± 0.6 to 4.3 ± 2.0 cm d⁻¹ at Landing Lake; 0.5 - 2.3 cm 498 d^{-1} at Toolik Lake). This is largely due to the greater Landing Lake groundwater CH₄ concentrations (370 µmol 499 L^{-1}) compared to Toolik (21 µmol L^{-1}). These higher fluxes may lead to the observed higher surface water 500 dissolved CH₄ in Landing Lake than at Toolik (Table 5, $1.8 \pm 0.3 \mu$ mol L⁻¹ and $0.02 - 0.8 \mu$ mol L⁻¹, 501 respectively). A fraction of CH₄ measured in groundwaters may be oxidized before reaching lake surface 502 waters, and other sources of CH₄, such as methanogenesis in lake sediments may drive the observed differences. 503 Further investigation is recommended to confirm the role that groundwater plays in CH₄ lake budgets.

504 The depleted carbon-isotopic signature of CH₄ in groundwater (-61.9 \pm 4.4‰, Table 2) is consistent with 505 microbial production (Hornibrook et al. 1997; Whiticar 1999), and the large range in isotopic values suggests 506 both methanogenesis and oxidation may be occurring. If oxidation is a dominant process removing CH_4 , it is 507 expected that δ^{13} C will increase logarithmically as CH₄ decreases because lighter CH₄ is preferred in the 508 reaction (Whiticar and Faber 1986; Whiticar 1999), a pattern which was observed in Landing Lake between 509 groundwater and lake water samples (Fig. 3). We assume that the highest concentration of CH_4 observed in 510 groundwater was the starting concentration and stable isotopic composition before any oxidation ($[CH_4]_{GW}$ = 511 612 μ mol L⁻¹, $\delta^{13}C_{CH4, GW}$ = -65.2‰, Table 2). The final composition after oxidation was assumed to be the average concentration and stable isotope value in Landing Lake ([CH₄]_{LAKE} = 1.8 μ mol L⁻¹, $\delta^{13}C_{CH4, LAKE}$ = -512 513 47.1‰, Table 1). Following the equation below (Whiticar and Faber 1986):

$$\delta^{13}C_{CH4, LAKE} = [\delta^{13}C_{CH4, GW} + 1000([CH_4]_{LAKE} / [CH_4]_{GW})^{1/\alpha - 1}] - 1000,$$
(5)

514 The fractionation factor (α) between starting groundwater CH₄ and average lake CH₄ was 1.003, in good 515 agreement, considering the margin of error, with the expected α of 1.005 – 1.030 for bacterial CH₄ oxidation 516 (Whiticar 1999), which supports the idea that CH₄ in the lake was produced in the active layer and then 517 transported by groundwater movement, as has been qualitatively observed in other lakes and streams (Kling et 518 al. 1992; Crawford et al. 2013).

519 Additionally, CH₄ produced in bottom sediments may also be transported into the lake by diffusion and 520 ebullition. Additional measurements of CH₄ concentrations and $\delta^{13}C_{CH4}$ in sediment porewater profiles and 521 floating chambers would be necessary to completely quantify sediment-water diffusive fluxes and ebullitive 522 fluxes, respectively, and their contribution to the lake's CH₄ budget. Diffusion, ebullition and advection may 523 collectively contribute to the CH₄ budget, and each may be impacted by environmental changes. As 524 precipitation increases in the Arctic (Rawlins et al. 2010; Wrona et al. 2016), groundwater flow is expected to 525 increase, impacting advective transport of CH₄ (Walvoord and Kurylyk 2016). Recent work has also revealed 526 that abrupt thaw beneath Arctic lakes can accelerate carbon emissions from lakes (Walter Anthony et al. 2018), 527 potentially increasing future diffusive and ebullitive CH₄ fluxes from sediments.

528 Once CH₄ enters a lake, it may be lost in the water column via oxidation, to the atmosphere by gas 529 exchange, to groundwater recharge, or surface transport. We calculated diffusive air-water CH₄ (Section 2.3.2), 530 using the observed $(1.8 \pm 0.3 \text{ µmol } \text{L}^{-1})$ and saturated concentrations of CH₄ in the lake (0.004 µmol $\text{L}^{-1})$ and 531 two modeled gas exchange coefficients ($k_{CH4} = 1.36$ m d⁻¹ and 0.79 m d⁻¹, at 17.9 °C). The flux from Landing 532 Lake to the atmosphere for July 2017 was 1.3 - 2.3 mmol m⁻² d⁻¹, approximately 3 - 18 times less than lake 533 input of CH₄ via groundwater (Table 5). The 24-hr measurement period CH₄ fluxes in 2018 were 1.3 - 5.7534 mmol $m^{-2} d^{-1}$ (Tables 5 and 6), which agreed well with the calculated diffusive air-water fluxes. This suggests 535 that groundwater sources of CH_4 can support all observed diffusion of CH_4 from the lake surface and that they 536 may be a driver of observed diffusive CH₄ emissions.

That the groundwater fluxes of CH₄ were higher than air-water diffusive losses is likely due to the additional removal of CH₄ via oxidation in the water column (Whiticar 1999; Bastviken et al. 2002), a determination supported by $\delta^{13}C_{CH4}$ (Fig. 3). Oxidation of CH₄ in the water column of freshwater lakes is expected by CH₄ oxidizing bacteria (Whiticar 1999) and is typically 30 – 99% of CH₄ produced in sediments or anoxic waters (e.g. (Bastviken et al. 2002, 2008)). Typical oxidation rates can therefore account for this "missing" CH₄ in Landing Lake. Climate warming will increase both methanogenesis and CH₄ oxidation, but 543 oxidation rates are typically less temperature dependent than production rates, and lower solubility of CH_4 in 544 warmer warmers may cause CH_4 release via bubbles that escape oxidation (Dean et al. 2018).

545 The air-water diffusive flux in this study was similar to the diffusive methane flux of $2 - 10 \text{ mmol m}^2 d^{-1}$ 546 for 38 peatland ponds across the Arctic and subarctic (Table 5) (Wik et al. 2016). Another study of 40 lakes in 547 Alaska (~65°N) with similar surface areas found average air-water CH₄ fluxes in summer of 0.6 mmol m⁻² d⁻¹ 548 (Sepulveda-Jauregui et al. 2015). It is important to note that this study was done in the summer season, so these 549 fluxes are likely to change with better temporal coverage. Polar regions are expected to become warmer (Schuur 550 et al. 2008, 2015; Vihma et al. 2016) and wetter (Rawlins et al. 2010; Wrona et al. 2016) over the following 551 decades, so higher CH₄ production in soils is expected if increasing precipitation increases soil moisture (Natali 552 et al. 2015) which can then be transported to aquatic systems by groundwater flow.

In this study, we used naturally occurring ²²²Rn to quantify groundwater discharge and dissolved CH₄ 553 554 fluxes to a lake in a subarctic terrestrial wetland. Groundwater fluxes were similar to those at another lake in 555 Alaska measured with the same radon-budget method (Paytan et al. 2015). We found that groundwater is a 556 source of CH_4 to the lake as suggested by the fact that groundwater CH_4 fluxes substantially exceeded diffusive 557 fluxes from the lake surface. The concentrations of CH₄ and diffusive fluxes were higher than the well-studied 558 Toolik Lake. Increased CH₄ production with warming and wetting of the Arctic may lead to higher rates of 559 delivery of CH₄ to aquatic environments due to the combined increase in CH₄ production (Natali et al. 2015) 560 and the shift to greater subsurface flow as permafrost thaws (Walvoord and Kurylyk 2016).

561 Appendix 1: Methane fluxes

562 Table 6. The methane concentrations, measured fluxes, and measured gas exchange coefficients for

- 563 Landing Lake, July 2018. Each method is described in Section 2.2. Average wind speed over the 3 days
- 564 was 4.6 ± 1.1 m s⁻¹. All measurements were made at the same location (latitude, longitude): 61.26583, -
- 565 163.24199.

Sample	Method	Length of deployment	Temp. °C	Lake CH ₄ µmol L ⁻¹	CH ₄ flux mmol m ⁻² d ⁻¹	k ₆₀₀ m d ⁻¹
7_8	24-hr period	3.6 hr	13.3	2.35	5.7	0.24
7_8	Instantaneous	15 min	13.3	2.35	21.8	0.93
7_9	24-hr period	28.7 hr	12.1	0.93	2.0	0.23
7_9	Instantaneous	15 min	12.1	0.93	6.3	0.71
7_10A	24-hr period	16.6 hr	11.6	0.65	1.8	0.29
7_10A	Instantaneous	15 min	11.6	0.65	17.1	2.32
7_10B	24-hr period	21.4 hr	12.7	0.58	1.3	0.24
7_10B	Instantaneous	15 min	12.7	0.58	17.0	ND^{a}
7_10C	Instantaneous	15 min	13.7	ND	5.1	ND

24-hr measurement period average	12.4	1.1	2.7	0.251	
±	0.7	0.8	1.0	0.014	
Instantaneous average	12.7	1.1	13.5	1.32	
±	0.9	0.8	7.4	0.50	
^a ND = no data.					

567 *Appendix 2:* Uncertainty estimates in the mass balance model

- 568 Table 7. The parameters in the mass balance and the methods for estimating the uncertainty in each
- 569 parameter.

566

Parameter	Estimation of uncertainty			
Gas exchange, Wind Speed Model	Slope error (19%); standard deviation of measured wind speeds (<1%, n > 1000)			
Gas exchange, Size Class Model	Estimated at 20% (std error = 7-25% in Holgerson & Raymond, 2016)			
Gas exchange, direct measurement	Standard error of measurements (24-37%)			
Gas exchange in mass balance	Two estimates: Conservative = Size Class, Upper Limit = Wind Speed; 30th errors ~20%			
Decay	Standard error of lake ²²² Rn inventories (11%, n = 18)			
Recharge	Impact on mass balance discussed in Section 4.1.1			
Stream discharge (out of lake)	Impact on mass balance discussed in Section 4.1.1			
Diffusion from bottom sediments	Propagated measurement error of the two methods (19%)			
Dissolved ²²⁶ Ra	Measurement error of ²²⁶ Ra by gamma spectrometry (8%)			
Groundwater ²²² Rn flux	Propagated uncertainty of all model terms (41%)			
Groundwater flux (cm d ⁻¹)	Reported range for each estimate propagated from: 1) uncertainty of the ²²² Rn flux (41%); 2) standard error in average groundwater endmember (21%, n = 10) OR measurement error in high activity groundwater endmember (8%)			

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576 S.M.N, J.D.S..

578 study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision

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580 References

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Title: Using radon to quantify groundwater discharge and methane fluxes to a shallow, tundra lake on the Yukon-Kuskokwim Delta, Alaska

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Fig. 1 "Landing Lake" sampling locations and the study site location within Alaska, USA shown in inset (star symbol). One groundwater sample (B2) is not shown because it was 5 km north of Landing Lake (Figure was made using ArcMap 10.5.1).



Fig. 2 Concentrations of (a) dissolved 222 Rn and (b) dissolved methane in Landing Lake. Sizes of symbols represent relative concentrations. CH₄ error = 30% for all samples; 222 Rn error = 16% for WP4, 0.1-6% for all other samples. (Figure was made using ArcMap 10.5.1).



Fig. 3 Stable carbon isotopes of dissolved CH₄ as a function of CH₄ concentration in groundwater (light blue squares) and surface water samples (dark blue circles) at Landing Lake in 2017. Notice the logarithmic scale on the x-axis. The regression equation is $\delta^{13}C = -5.98 \log [CH_4, \mu mol L^{-1}] - 46.9\%$ and includes both the lake waters and groundwaters. PDB = Pee Dee Belemnite standard. (Figure made using Microsoft Excel).



Fig. 4 The stable isotope values for δ^2 H and δ^{18} O in water for samples collected in 2017. The stream sample (triangle) drains Landing Lake. The dashed line is the Global Meteoric Water Line (Craig 1961). Diamonds represent all lake and pond samples collected in 2017 (see data online: (Ludwig et al. 2017b)) which were fit with a Local Evaporation Line (LEL): δ^2 H_{H2O} = 4.31(δ^{18} O) – 36.55‰ (R² = 0.96, *p* << 0.01). The dotted black line is the best-fit line for Landing Lake groundwaters: δ^2 H_{H2O} = 6.87(δ^{18} O) – 2.90‰ (R² = 0.70, *p* = 0.04). (Figure made using Microsoft Excel).



Fig. 5 A conceptual model showing the sources and sinks of ²²²Rn in Landing Lake. Sources (dark blue arrows) include decay of dissolved ²²⁶Ra in lake water, diffusion from lake bottom sediments and groundwater. Sinks (light blue arrows) include ²²²Rn decay, loss to the atmosphere via gas exchange, recharge into soils, and the stream outlet. (Figure made using Microsoft Powerpoint).



Fig. 6 The fluxes of ²²²Rn for each source (dark blue) and sink (light blue) in the mass balance model for Landing Lake. The radon flux due to groundwater is highlighted with a black outline because it is the difference between the sinks and the other two sources. Error bars are propagated errors. Lower limits are the conservative estimate discussed in the text. (Figure made using Microsoft Excel).