Low methane emissions from a boreal wetland constructed on oil sand mine tailings

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Abstract. A 58 hectare mixed upland and lowland boreal plains watershed called the Sandhill Fen Watershed was constructed between 2008 and 2012. In the years following wetting in 2013, methane emissions were measured using manual chambers. The presence of vegetation with aerenchymous tissues and saturated soils were important factors influencing the spatial variability of methane emissions across the constructed watershed. Nevertheless, median methane emissions were equal to or less than $0.51 \text{ mg CH}_4 \text{ m}^2 \text{ h}^{-1}$ even from the saturated organic soils in the lowlands. Although overall methane emissions remained low, observations of methane ebullition increased over the three study years. Ebullition events occured in 10 % of measurements in 2013, increasing to 21 % and 27 % of measurements in 2014 and 2015, respectively at the plots with saturated soils. Increasing metal ion availability and decreasing sulphur availability was measured using buried ion exchange resins at both seasonal and annual timescales potentially as a result of microbial reduction of these ions. Using principle component analysis, methane fluxes had a significant positive correlation to the leading principle component which was associated with increasing ammonium, iron, and manganese and decreasing sulphur availability ($r = 0.31$, $p < 0.001$). These results suggest that an abundance of alternative inorganic electron acceptors may be limiting methanogenesis at this time.

1 Introduction

The boreal biome stores large quantities of soil carbon (C) due to the high density of peatlands with some estimates as high as 165 kg C m^2 (Beilman et al., 2008). A legacy of open pit oil sands mining in northern Alberta, Canada will be that almost 5000 km² of the boreal landscape (Alberta Government, 2017) will require reclamation to restore it to

- 5 the "equivalent capability" of the pre-mining ecosystems (Environmental Protection and Enhancement Act 2017). These engineered landscapes will be important in determining the long-term C footprint of the region, potentially helping to offset ecological losses and industrial emissions. Rooney et al. (2012) used land cover classification and reclamation plans to estimate a potential net loss of $11.4 - 47.3$ Tg of the stored soil C from the pre-mining landscape and a reduction in CO₂ sequestration capacity of the new constructed landscape by $5.7 - 7.2$ Gg C year⁻¹. This loss is
- 10 expected to occur as a result of both the decomposition of peat when it is used in the reclamation of uplands and from the diminished C sequestration capacity of engineered landscapes. In particular, Rooney et al. (2012) predicted that an insufficient proportion of the disturbed area will be replaced with the C sequestration capacity of the original boreal wetlands. However, wetland reclamation in the region is still quite novel, and little is known about the surfaceatmosphere exchange of C in these newly engineered wetlands (Nwaishi et al. 2015; Clark et al. 2019).
- 15 Undisturbed peatlands are long-term CO₂ sinks, accumulating C as peat over millennia (Rydin and Jeglum, 2006; Vasander and Kettunen, 2006). Yet, on annual timescales the C cycling within peatlands is highly variable and only partially understood, particularly with regards to methane (CH4) emissions (Vasander and Kettunen, 2006). Wetland CH4 emissions are thought to be the single greatest driver of inter-annual variability in atmospheric CH4 concentration (Bousquet et al., 2006) and globally wetlands are the largest non-anthropogenic source of CH4 to the atmosphere
- 20 (Kirschke et al., 2013). Atmospheric CH₄ has 28-34 times the global warming potential of CO_2 over a 100 year time period but with a perturbation lifetime of 12.4 years in the atmosphere, its radiative forcing is much greater on decadal timescales (IPCC, 2013). Therefore, CH4 production and release is an important land-atmosphere climate change feedback process (Kirschke et al. 2013). Since methanogenesis is only favoured at the lowest reduction/oxidation (REDOX) potentials known to support life, the high rates of CH4 emission and low rates of C turnover in peatlands
- 25 are linked to a scarcity of inorganic electron acceptors in anoxic, saturated soils (Han-Schofl et al., 2011; Peters and Conrad, 1995; Thomas et al., 2009). Around the time that Rooney et al. (2012) estimated the C balance for the postmining landscapes of Alberta, the first two large-scale wetlands were being constructed in the Fort McMurray region. These wetlands were designed and constructed to test theories on how to return the peat forming and C capturing capacities to the mined landscape (Wytrykush et al., 2012). A research gap exists, however, between the modelled
- 30 predictions of C cycling by Rooney et al. (2012) and the actual C cycling of constructed wetlands. Clark et al. (2019) discussed the increasing CO_2 sink strength over the first three years of one of these wetlands but did not address emissions of CH4, which can contribute substantially towards the long-term C balance of some ecosystems (Vasander and Kettunen, 2006). The aim of this study is to evaluate CH4 emissions from this constructed ecosystem and identify the factors which influence their temporal and spatial variability.
- 35 Methane emissions from wetlands are highly variable in space and time (Moore et al., 1998). Four key wetland characteristics are typically linked to the temporal and spatial variability of CH4 flux in peatlands: water table position,

REDOX, vegetation composition, and temperature(Kim et al., 2012; Limpens et al., 2008; Turetsky et al., 2014; Vasander and Kettunen, 2006). Approximately 60 to 90 % of the CH4 produced in peatlands is re-oxidized before it reaches the atmosphere (Le Mer and Roger, 2001). Emissions tend to be higher in areas with a high water table as

- 40 this limits the potential for CH4 oxidation (Moore et al., 2011) while deep rooting vegetation with aerenchymatous tissues (Couwenberg, 2009; Kludze et al., 1993) allow CH4 to bypass the aerated surface peat where aerobic methanotrophic communities reside. Aerenchymatous tissues can also transport oxygen into the rhizosphere which can increase REDOX potentials and create unfavourable niches for anaerobic microbial communities (Blossfeld et al. 2011). Rhizospheric microbial communities readily metabolize the labile organic compounds exuded by roots such
- 45 that plant net primary productivity has positively correlated with CH4 production (Dacey et al., 1994; Megonigal and Schlesinger, 1997; Vann and Megonigal, 2003; e.g. Whiting and Chanton, 1993; Ziska et al., 1998). Numerous studies have also shown CH₄ emissions increase in warmer soils (Bubier et al., 2005; Bubier and Moore, 1994; Roulet et al., 1992; Whalen, 2005; Whiting and Chanton, 1993) although this effect may be masked when water table or other environmental factors co-vary (Moore et al., 2011).
- 50 Although constructed peatlands are relatively novel, restoration of peatlands has occurred for decades by rewetting drained peat, often by blocking the ditches originally used to drain them (Rochefort and Lode, 2006). In a comprehensive review of the literature, the IPCC suggested that CH4 emission rates from rewetted boreal sites with rich organic soils can be estimated as 2.1 mg CH₄ m⁻² h⁻¹ but encourage site specific study because the range in the published literature is very large and not normally distributed (Blain, et al., 2014). A follow up review of the literature
- 55 by Wilson et al. (2016) provided additional evidence that CH4 emissions were 1.7 to 22.6 times greater for rewetted vs. drained boreal organic soils. The data examined by Wilson et al. (2016) was extrapolated using models of average area (to include multiple surface types) and annual rates, to conform with IPCC reporting standards. Our review of paired studies which directly compare rewetted and drained environments show that rewetted boreal and temperate organic soils have an average of 23 and 64 times the CH4 emissions of their drained counterparts, respectively
- 60 (supplementary Table S1). However these differences in CH4 emissions are highly variable and some studies show a decrease in CH4 emissions after wetting (Christen et al., 2016; Juottonen et al., 2012; Urbanová et al., 2012; Waddington and Day, 2007). The few studies which compare emissions from rewetted and undisturbed wetlands (supplementary Table S1) show a wide range of results with rewetted wetland emissions <1% (Juottonen et al., 2012), 19% (Beetz et al., 2013), 43% (Urbanová et al., 2012), and 127% (Christen et al., 2016) of the emissions observed in
- 65 undisturbed wetlands.

To determine the CH4 emissions of a closure watershed in the oil sands mining region of northern Alberta and how these emissions compare to those from the restored peatlands listed above, this paper presents three years (2013-2015) of CH4 emission measurements from the Sandhill Fen Watershed (SFW), one of the first wetland complexes constructed in the Athabasca oil sands region (AOSR). Once water was added to the SFW, approximately 17 ha of the

70 58 ha construction had permanently saturated peat soils with little horizontal or vertical water flow (Nicholls et al., 2016) and thriving plant communities including species with aerenchyma (e.g. *Carex* spp.) (Vitt et al., 2016). We hypothesized that these areas would be a significant source of CH4 and these CH4 emissions would be an important component of the C balance of the SFW. Results of this research will provide important information on the greenhouse

gas and C budget for reclamation ecosystems in the AOSR and help guide future strategies for ecosystem design and 75 C management.

2 Study Site

The SFW (57.0403 N, 111.5890 W) was designed to reclaim sand-capped soft tailings with the conditions needed to promote the long-term development of a peat forming boreal plains ecosystem (Wytrykush et al., 2012). Details of the SFW construction were described by Wytrykush et al. (2012) and others (Biagi et al., 2019; Nicholls et al., 2016; 80 Oswald and Carey, 2016; Vitt et al., 2016). Clark et al. (2019) described the three main topographic features of the

- SFW as a lowland region (the wetland), a midland region (drained but moist organic soils), and upland region lying 2 to 6 m above the wetland with well drained sandy soils and two experimental perched wetland sites with moist organic soils (Figure 1). The sand of the lowland, midland and perched wetland regions were covered with 0.5 to 1 m peat coarsely mixed with some underlying mineral soil via excavation of nearby peatlands before mining. A goal of the
- 85 SFW design was to limit vertical transport of tailings and process-affected waters (i.e. water with some component of industrial wastewater known to contain salts and naphthenic acids) to the surface (Wytrykush et al., 2012). To achieve this, fine sediments with 26.4 – 46.4 % clay (based on texture analysis) were placed in the lowland regions of the SFW to help minimize vertical transport of process affected waters from the tailings below.
- Initially, fresh water was pumped into the containment pond in the west end of the SFW in 2013 (Figure 1). This 90 water flowed through a leaky gravel dam into the SFW to initially saturate the lowland region. A downstream pump was used to remove water from an outlet V-notched weir (Figure 1). Except for a few hours of operation, the pumps were not used in 2014 or 2015. Four underdrains, which run along the central area of the lowland region, were placed to limit the upwelling of the salt-rich process water. The underdrains were not operated after the beginning of the 2014 growing season.
- 95 After placement and before wetting, the top 0.25 m of the donor peat/mineral material in the midland and lowland had a mean $(\pm SD)$ C to nitrogen (N) ratio of 22.7 (\pm 3.4), N content was 0.98 (\pm 0.36) % by dry weight, Ca and Na concentrations were 786.1 (\pm 409.4) and 184.2 (\pm 79.0) mg kg⁻¹, respectively and electrical conductivity was 1980 (\pm 600) μ S cm⁻¹ (n = 107). The concentration of Fe was 8028 (\pm 2712, n=13) mg kg⁻¹ and SO₄² concentration was high, with a mean of 914.0 (\pm 425.5, n=107) mg of S kg⁻¹. Total S concentration by dry weight was also high at 1.00 % $(\pm 0.50 \text{ %}, \text{n=12})$ determined by
- 100 oxidation in an induction furnace then quantified through infrared mass spectroscopy with a LECO IR (LECO Corporation, Saint Joseph, MI).

In the third growing season (2015) since the ecosystem was seeded/planted (see Vitt et al., 2016 for more details), there were four distinct plant communities in the lowland roughly distributed along a soil moisture and water depth gradient (Vitt et al., 2016). The areas with standing water (Figure 1) were dominated by *Carex aquatilis* Wahlenb.*, Typha latifolia* L. and to a

105 lesser extent *Carex utriculata* Stokes, while areas with the water table near the surface contained the most species associated with peatlands including sedges and bryophytes (Vitt et al., 2016). The drier areas contained the largest percentage of weedy species, or species not associated with peatlands (Vitt et al., 2016). The midland and organic soil regions of the upland was populated by planted trees (*Populus tremulodies* Michx., *Betula papyearifera* Marshall, *Picea glauca* Monech, *Picea mariana* Mill., *Larix laricina* Du Roi, and *Pinus banksiana* Lamb.) and local grasses (e.g. *Hordeum jubatum* L.) that had 110 naturally colonized the region. Vegetation cover throughout the whole watershed increased over time; peak season leaf area index increased from 1.5 ± 0.6 to 2.2 ± 1.1 over the three years as measured at 15 plots (5 midland, 10 lowland) using a plant canopy analyzer (LAI-2200, LI-COR Inc., Lincoln, NE).

3 Materials and methods

- This study used plot-scale measurements of CH4 fluxes over three years to assess the temporal and spatial variations 115 in CH4 emissions and the factors that influence their variability. Fluxes were measured using non-steady state static chambers following methods described by Wilson and Humphreys (2010). Plots were established over the three years of study to sample the dominant landforms and built features of the SFW. In 2013, 15 plots were established across the lowland and midland of the SFW. In 2014, the number of plots was increased to 21 by including plots on the organic soils in the perched wetlands (upland region) and in 2015 was increased to 29 by adding more plots in the
- 120 lowland (Figure 1). The 2013 plots were chosen before the pumps were activated and were selected to capture the surface moisture and vegetation heterogeneity while maintaining multiple plots on similar landforms. The midland plots were chosen to capture differences in the vegetative cover. Two plots were placed on sandy soils, one directly on exposed tailings sand and the other on the salvaged mineral soil mixture. In 2014, the new collars were placed on the experimental wetland sites that were depressions built into the upland hills. Note that these upland plots remained
- 125 moist but well drained for the duration of the measurements. The eight new collars in the lowland added in 2015 were placed along a moisture gradient transect also used for REDOX monitoring (described below). The midland plots had water table depths over 0.5 m below the surface and all upland plots had water table depths exceeding 1 m. For the remainder of this paper, midland and upland plots were combined into one group called 'Upland' due to their similar soil, water table depth and vegetation characteristics. In the lowland, the 'Saturated'
- 130 plots were those with volumetric soil water content \geq 86% (i.e. fully saturated) and all remaining plots were part of the 'Unsaturated' group. Only 5 lowland plots switched categories in 2013 and early spring 2014 and 2015, all periods when CH4 emissions were uniformly low with no discernible differences among the groups. Water table depth was not monitored at each plot but the soil moisture measurement method is described below.
- Each plot included a pair of 0.19 m tall collars with a surface area of 0.07 m^2 that were inserted until nearly flush with 135 the soil surface to minimize changes to the microclimate as a result of the collar (Parkin and Venterea, 2010). Each pair consisted of one collar that was maintained free of vegetation by clipping and the other was left undisturbed (Figure 2). A limitation of the undisturbed collar was that any vegetation larger than the flux chambers (0.4 m tall, 0.03 m³) was trimmed to fit within the chamber. This trimming had the largest effect in the lowland, where some *Typha* died once trimmed. The collars were made of SDR35 12" PVC sewer pipe with a groove cut into the top edge
- 140 where an acrylic chamber was placed during measurement. The chambers were constructed of acrylic and were covered in opaque black plastic to reduce heating within the chamber and eliminate photosynthetic uptake of CO₂ for a respiration analysis discussed by Clark et al. (2019). A seal between the chamber and collar was made by filling the groove with water. A small coiled vent tube on the top of the chamber maintained equal pressure with the surroundings (Hutchinson and Livingston, 2001). During a measurement, the air inside the chamber was mixed by pumping a 60
- 145 mL syringe connected to the chamber sampling line and then pulling 24 mL of air from the chamber volume at 0, 5,

10, 15, and 20 minute intervals. The sampled air was injected into a 12 mL evacuated vial containing a small amount of magnesium perchlorate to remove any water vapour from the air sample. Flux measurements were made between late May and early August in all three years, referred to hereafter as the growing season. The air samples were transported to Carleton University where CO₂ and CH₄ concentrations were measured on a gas chromatograph (CP

- 150 3800, Varian, CA) within a few months of sampling. The operational details of the gas chromatograph (GC) are described in Wilson and Humphreys (2010). During the initial collar installation, five thermocouples were buried to a depth of 2, 5, 10, 20, and 50 cm at each plot between the two collars. At the time chamber flux measurements were made, soil temperatures were recorded from the buried thermocouples. A portable soil water sensor that integrated over the upper 0.2 m of soil (Hydrosense,
- 155 Campbell Scientific Inc., Utah, USA) was used to measure volumetric water content of the soil at three locations surrounding each of the collars. When there was standing water over 0.05 m deep, no manual soil water measurements were made. In those cases, the soil moisture was estimated at 87 % (an estimate of saturated conditions based on an average soil bulk density of 0.28 g cm⁻³).

Fluxes in units of mg C-CH₄ m⁻² h⁻¹ were calculated from the linear rate of change in CH₄ mixing ratios (nmol CH₄

- 160 mol⁻¹ air), the molar density of the air in the chamber (mol air m⁻³), the chamber volume (m³) and area (m²) and the molecular weight of carbon (mg C nmol⁻¹ CH₄). To determine air density, barometric pressure was recorded at a nearby micrometeorological station (Figure 1) and chamber temperature was estimated using the 0.02 m thermocouple at each plot. The chamber volume was adjusted for the different collar heights and depth of standing water. All calculations and statistical analyses were carried out with MATLAB 2015 (Mathworks Inc., Massachusetts, USA).
- 165 The R^2 coefficient from the linear regression of CH₄ concentration over time is not suitable as a quality control metric by itself (Lai et al., 2012). For example, when the fluxes approach zero, the slope also approaches zero and small variability in gas concentrations results in low \mathbb{R}^2 values. In total, 16 % of the fluxes had an \mathbb{R}^2 over 0.9 and 28 % over 0.8. Instead of the typical \mathbb{R}^2 filtering, quality control of the data was done by a visual inspection of each time series used to calculate the fluxes. Any flux measurement which had a distinctly non-monotonic or non-linear trend
- 170 due to individual data point anomalies were removed from further analysis (6.4 %, 12.3 %, 11.2 % of the flux measurements in 2013, 2014 and 2014, respectively). These anomalies included isolated large decreases in concentration or a return to ambient concentration or isolated or unsustained large increases in concentration and represented fluxes from a leaking chamber or an ebullition event (Tokida et al., 2007). In 2013 REDOX potential was measured every 15 min with nine HYPNOS III rods (Vorenhout et al., 2011) inserted into the soil every 2 m alongside
- 175 the 20 m transect of new plots to evaluate the impacts of a moisture gradient on CH4 emissions (Figure 1). The HYPNOS rods were equipped with four platinum probes and thermistor temperature sensors, of which the sensors 0.2 and 0.4 m below the surface were used in this study. The reference was a pH probe with a 0.1 M KCl standard buried below the water table near the middle of the transect. The probes were connected to two HYPNOS dataloggers and the standardised REDOX potentials ($n = 36$) were calculated as the sum of the measured potential and the reference
- 180 potential. No pH correction was applied to probe measurements because pore water remained circumneutral and stable throughout the season. During summer 2015, the mean pH was 7.1 \pm 0.4 (\pm SD) as determined using weekly measurements at a nearby pore water sampling well that integrates the water from ~1 m depth to the surface. Early

spring (May 20th) had the largest discrepancy between the well and surface water measurement locations, with the surface water at a pH of 7.6 and the well water at 6.2. By early July the two sampling locations had almost converged 185 at neutral, with a slightly higher pH in the surface water (~7.4 vs 6.9).

- Within each plot, three replicate sets of Plant Root Simulator (PRS) probes (Western Ag., Saskatoon, Canada) were buried at a depth of 0.1 m (shallow) and 0.2 m (deep) outside the collars. PRS probes are ion exchange membranes designed to mimic in-situ soil-root exchange of nutrients and other ions in a non-destructive manner (Qian et al., 2008; Qian and Schoenau, 2005). PRS probes provide a time-integrated representation of the soil nutrient availability/net
- 190 adsorption rates in units of ion mass per membrane area per burial time. These values differ from the more common point-in-time soil extraction measurements (typically element mass per mass of dry soil) although studies show good correspondence between methods for N, P, K, and S (Harrison and Maynard, 2014; Qian et al., 1992). The probes were buried for 28 days, (\sim one month) with three consecutive burial periods monitored each year starting on day 149, 146 and 146 for 2013, 2014, and 2015, respectively. For simplicity, since the midpoint of each of these three one-
- 195 month burial periods corresponds roughly to the start of June, July and August, results from each period were referred to as the result from those months (i.e. the July ion adsorption rate refers to the moles of ions absorbed by the PRS probes period between 174th and 202nd day of the year). An example plot with collars, PRS probes and thermocouples is shown in Figure 2.
- Temporal trends in the PRS probe ion data were assessed using Spearman rank correlations using three different 200 temporal groupings of the data. First, the data were binned into categories 1 to 9, representing each of the three months across three years of measurements (for example, June of the third year was category 7). Second, the data were binned into three categories corresponding to the three months of measurements each year, regardless of year, to ignore the inter-annual trends. Finally, the data were binned by year, regardless of the collection month, to ignore the monthly/seasonal trends. This allowed some quantification of the seasonal trends relative to the inter-annual trends in 205 all three groups (upland and the saturated and unsaturated lowland groups).
- To relate CH4 fluxes, REDOX and PRS probe results, a principal component analysis (PCA) was performed on the standardized measurements (z-scores) of net ion adsorption rates from the PRS probes buried at 0.2 m depth in the nine plots with REDOX probes. The availability of alternative electron acceptors including NH_4^+ , Mn^{4+} , Fe^{3+} , and SO₄² are linked to varying soil REDOX potential through their participation in microbially mediated REDOX
- 210 reactions. It should be noted that although PRS probes adsorb all mobile forms of S, most are expected to be SO_4^2 (Li et al., 2001). Using Pearson correlation, the relationship between the leading two principal components and 0.2 m REDOX measurements were assessed. Pearson correlation was also used to assess the relationship between the leading two principal components and the log-transformed CH4 flux (transformed to account for skew) averaged over the burial period.. Methane fluxes were increased by a common absolute value of the minimum flux observed in the
- 215 study (-0.10 mg C-CH₄ m⁻² h⁻¹) to permit the log transformation. Pearson correlation was also used to assess any linear relationships between the transformed CH4 flux and soil moisture and 0.02 m soil temperature. Individual Pearson correlations were calculated between the mean transformed CH4 flux and the net ion adsorption rates at the two burial depths for a burial period.

The effects of vegetation and standing water on CH4 fluxes were evaluated using a linear mixed effect 220 (LME) model:

Methane Flux_{ij} = α_{ij} + ζ_{0i} + β_1 Vegetated_{ij} + β_2 Saturated_{ij} + ϵ_{ij} [1]

where *Vegetated* and *Saturated* were binary vectors indicating if the measurement came from a vegetated plot, or a saturated plot, respectively. *i* and *j* subscripts represented measurements from each sampling day and site respectively. β was the slope of the fixed effect, ε was the residual, α was the intercept, and ζ was 225 the random effect from the repeated measures occurring at each site.

An independent but similar LME model was constructed to detect significant effects of soil depth and standing water on PRS net ion adsorption rates and REDOX potentials; the *Vegetated* covariate was replaced with the binary vector *deep*, which represented the relative depth of the REDOX or PRS measurement (i.e. deep or shallow):

230 *PRS ion/REDOX potential*_{ij} =
$$
\alpha_{ij} + \zeta_{0j} + \beta_1 Deep_{ij} + \beta_2 Saturday_{ij} + \epsilon_{ij}
$$
 [2]

4 Results

The three growing seasons (1 May through 31 September) were warmer and wetter than the long-term average. The average air temperatures recorded at 3 m above the lowland were 16.4, 15.3, and 15.3ºC and total rainfall was 375.1, 299.1, and 231.3 mm for the 2013, 2014, and 2015 growing seasons, respectively. The 1981-2010 climate normal for

235 this period was 13.3ºC and 211.1 mm at a nearby weather station (Fort McMurray Airport; 48 km from study area; Environment and Climate Change Canada, 2016). Over the three-year study period, the upland plot soils were drier and slightly warmer than the lowland plots with an average volumetric soil moisture of 34.7 % compared to the 57.5 % in the lowland (Table 1). There were only small differences in growing season 0.02 m soil temperatures at the plotlevel (Table 1). Near the climate monitoring station in the centre of the lowland the EC was measured to be 1171 \pm 240 269 , 2109 \pm 306, and 2163 \pm 248 μ S cm⁻¹ in 2013, 2014 and 2015, respectively where the typical electrical conductivity of boreal wetland pore waters ranges from 400 to 2770 μ S cm⁻¹ (Trites and Bayley, 2009).

4.1 Spatial and temporal CH4 and ion relationships in the SFW

During each growing season, CH₄ emissions were generally very low, with median values less than or equal to 0.04 mg C-CH₄ m⁻² h⁻¹ for all plot groups (Table 1). The greatest median CH₄ emissions were from the saturated plots (all 245 located in the lowland) in July 2015 with median emissions reaching 0.51 mg C-CH₄ m⁻² h⁻¹ (Figure 3).

The proportion of measurements where substantive CH_4 emissions were detected increased for the saturated plots over the study period (Table 1). This included occasions with ebullition or when fluxes were greater than 0.5 mg C-CH₄ m⁻ $2 h^{-1}$, which is equivalent to ten times the maximum CH₄ uptake rate observed at this site and is in the upper range of average uptake rates observed in grassland and forest soils around the world (Yu et al. 2017). Among all three groups

250 and years, the 2015 saturated plots had the highest proportion of CH₄ emissions exceeding 0.5 mg C-CH₄ m⁻² h⁻¹ (6.5)

%) and highest number of ebullition events (26.6 %). Vegetated collars had a higher median CH4 flux than in collars with vegetation excluded, particularly in the saturated plots (Figure 3). There appeared to be an increase in $CH₄$ emissions over time in the vegetated collars within the saturated plots both within the growing season and interannually.

255 The rates of Mn^{2+} , Fe²⁺ and NH₄⁺ ion adsorption on the PRS probes were greatest in the saturated lowland plots and lowest in the upland plots. By 2015, net adsorption of S was lower in the saturated lowland plots than in the upland plots (Figure 4). The ion with the strongest absolute temporal trend in adsorption was S (Table 2, Figure 4), with most of this trend likely a result of changes in SO_4^2 availability (Geer and Schoenau, 1994; Li et al., 2001). In the saturated plots, mobile S decreased seasonally, annually and over the duration of the study (the greatest Spearman rank 260 coefficient was for 0.25 m deep S ion adsorption rates and chronological time represented as increasing values 1 through 9, Table 2). Declines in net S adsorption were also observed in the other plot groups but the greatest decline was in the saturated lowland plots, where relative to 2013 values, 2014 values were 40 % lower and 2015 values 50 % lower (Table 1 and Figure 4). Correspondingly, net adsorption rates for Mn^{2+} , Fe²⁺ and NH₄⁺ ions significantly

265 **4.2 Spatial and temporal relationships along a moisture gradient in the lowland**

increased over time at the saturated plots (Table 2).

Along the moisture gradient established by a slightly sloping surface in the north-eastern part of the lowland (Figure 1), CH4 emissions were higher in the 4 saturated plots than in the 5 unsaturated plots (Figure 5). Methane emissions were also higher in vegetated collars than in collars with vegetation excluded. Using a LME model (Eq. 1), and controlling for sampling location as a random effect, CH₄ emissions were 0.34 ± 0.11 mg m⁻² h⁻¹ greater in saturated 270 plots than unsaturated and 0.17 ± 0.06 mg m⁻² h⁻¹ greater in vegetated collars than collars without vegetation over the August period (Table 3), when CH4 fluxes were greatest, both in magnitude and proportion of emissions exceeding

- 0.5 mg C-CH₄ m⁻² h⁻¹ (Table 1). Temperature and soil moisture were also included in an earlier version of the LME model, but they did not have a significant effect on CH4 emissions so were removed from the model. When CH₄ emissions were greatest in the saturated plots, both REDOX potential and the PRS ion adsorption rates
- 275 indicated reduced soil conditions (higher Mn²⁺ and Fe²⁺ and lower S net adsorption; Figure 5). Using just the August data from all lowland plots the LME model indicated that the average net adsorption of Mn^{2+} was 16.0 ± 3.7 µg 10 cm⁻² month⁻¹ higher in saturated conditions, and $7.1 \pm 2.3 \,\mu$ g 10 m⁻² month⁻¹ higher for the deeper (0.2 m) probes. The net adsorption of Fe²⁺ was 188 ± 36 µg 10 m⁻² month⁻¹ higher in saturated conditions but there was no significant difference at depth ($\beta_2 = 42 \pm 24$). The net adsorption of S was $588 \pm 92 \,\mu$ g 10 m⁻² month⁻¹ lower in the saturated plots,
- 280 but there was no significant difference between the two depths ($\beta_2 = -74 \pm 63$). The median REDOX potentials were -127 and -168 mV in the saturated plots at 0.2 and 0.4 m below the surface, respectively. The LME model (Eq. 2) indicated REDOX potentials were 116 mV \pm 47 (hydrogen standard) lower in saturated conditions and 85 mV \pm 33 lower at a 0.4 m depth (Table 3).

The PCA's leading component related increasing net adsorption rates of NH₄⁺ and the metals with decreasing S. 285 Together, the leading two components of the PCA explained 89.8 % of the variability in net ion adsorption rates for the deep probes (with 79.1 % explained by the first component) for the nine plots with REDOX potential measurements. All ions loaded relatively evenly in magnitude on the leading component (PC1; Table 4), but S was inverse to the others. Although the negative sign of the correlation coefficient between PC1 and the REDOX potential $(R = -0.24)$ intuitively matched what would be expected for a REDOX gradient, where more negative potentials were

- 290 associated with less S availability and more Mn^{2+} and Fe²⁺, it was not significantly different from zero (p = 0.243). The correlation coefficient between PC1 and log-transformed CH₄ fluxes ($R = 0.378$) was also not significant ($p =$ 0.06), perhaps due to relatively few observations at the REDOX monitoring transect in 2015 (n=25). However, the same analysis conducted for all plots' PRS data ($n=163$) found similar weighting on PC1 and a significant ($p < 0.001$) correlation between PC1 and log-transformed CH₄ fluxes ($R = 0.31$). This suggests a link between increasing CH₄
- 295 emissions to decreasing S and increasing Mn^{2+} , Fe²⁺ and NH₄⁺ availability. Some of the individual ion adsorption rates were correlated to log-transformed CH4 fluxes for the 9 plots along the moisture gradient in the lowland and for the entire dataset (Table 5). At the REDOX monitoring transect in 2015, only S adsorption was negatively and significantly correlated to CH_4 emissions, but when the full data set was included, CH_4 emissions were significantly correlated with the following ion adsorption rates in the following order, by correlation strength: $Mn^{2+} = Fe^{2+} > NH_4$ ⁺
-

 300 > NO₃⁻ (Table 5). Overall, the correlations were similar for shallow and deep probes (Table 5). A significant correlation was also found between CH₄ fluxes and 0-20 cm integrated volumetric soil moisture ($R = 0.24$), whereas no significant relationship was found for 0.02 m soil temperature ($R = -0.04$).

5 Discussion

Methane emissions typically increase with increasing saturation/rising water tables within (Moore et al., 2011) and 305 among peatlands (Turetsky et al., 2014), and also tend to increase when drained peatlands are restored by wetting (supplementary Table S1). However, CH_4 emissions in the newly constructed SFW unexpectedly remained very low over the first three years since wetting, despite abundant vegetation with aerenchymatous tissues, peat soils, and high water tables. Over the three-year study period, the only plots that showed small but increasing trends in CH₄ emissions over time were vegetated plots with standing water above the soil surface. The vegetation at these plots was primarily

310 *Carex aquatilis,* and *Typha latifolia*. These species have aerenchymatous tissues that enable plant-mediated transport of $CH₄$ to the surface and have been reported in the peatland restoration literature to promote $CH₄$ emissions (Mahmood and Strack, 2011; Wilson et al., 2009).

Median CH4 fluxes from this system are 0.2 to 50 % of the values published from other studies on rewetted peatlands (Table 1; supplementary Table S1). Instead, the fluxes in this study are similar to those reported from the other

- 315 constructed wetland in the AOSR (median CH₄ emissions were below 0.08 mg m⁻² h⁻¹) and an undisturbed saline fen (Murray et al., 2017). Murray et al. (2017) accredited the small CH4 emissions and low CH4 pore water concentrations at the constructed wetland to the supply rate of mobile S. Methanogens, CH4-producing microorganisms, are obligate anaerobes and an abundance of alternative electron acceptors such as $SO₄²$ can support microbial communities that can outcompete methanogens. This effect has been described using the conceptual framework of the REDOX 'ladder'
- 320 (for a detailed definition see Bethke et al., 2011). Simply, a conceptual pristine aquifer will contain zones with distinct electron acceptors for metabolic REDOX reactions as reduction potentials decrease (Lovley et al., 1994). The sequence starts with the reduction of oxygen until it is consumed, then oxidized nitrogen (NO₃⁻), oxidized metals (non-

mobile MnO₂ and Fe(OH)₃) and S (e.g. SO_4^2), finally with the production of CH₄ through the reduction of CO₂ or acetate. In incubation studies, methanogens can be outcompeted by both metal reducing bacteria (MRB) (Achtnich et

- 325 al., 1995; Miller et al., 2015) and by sulphur reducing bacteria (SRB) (Achtnich et al., 1995; Akunna et al., 1998; Gauci and Chapman, 2006; Granberg et al., 2001; Han-Schofl et al., 2011; Kang et al., 1998; Kuivila et al., 1989; Lovley and Klug, 1983; Peters and Conrad, 1995; Watson and Nedwell, 1998). Although the community dynamics between MRB, SRB and methanogens is complex (Bethke et al., 2011), increased interactions among these organisms often leads to the suppression of methanogenic activity since microbial communities are competing for H_2 and acetate,
- 330 which methanogenic microbes require exclusively for their metabolism. In some fen peatlands, drought conditions have been shown to increase alternative electron acceptor abundance and suppress CH₄ production after rewetting (Estop-Aragonés et al., 2013). Two lines of evidence suggest that methanogenesis was inhibited in the SFW through this mechanism. Within the plots with REDOX probes, the largest (albeit still very small) CH4 fluxes were observed where reduction potentials at both 0.2 and 0.4 m below the surface were close to hydrogen standards of -200 mV;
- 335 potentials known to be favourable for methanogenesis (Akunna et al., 1998). At these potentials, the net adsorption of reduced metals (Fe²⁺ and Mn²⁺) were greatest and mobile S, which would be largely SO_4^2 ⁻, was lowest. In those nine plots where REDOX was observed, only S had a significant relationship to CH₄, but Fe²⁺, Mn²⁺ were also found to have a significant correlation to CH₄ with the full data set. In addition, CH₄ emissions and ebullition events increased over time in the saturated plots (those with standing water) throughout the lowland while mobile S appeared to be
- 340 declining in abundance as the presence of mobile metals (Fe^{2+} and Mn^{2+}) increased. This suggests the SFW soil became more reduced with a decrease in SRB abundance, which eased the competitive exclusion of methanogenic organisms. This conclusion follows Christiansen et al. (2016) who also found that Fe^{2+} measured with PRS probes was highest in conditions which promoted greater CH₄ emissions.

Wetter soils are expected to increase the probability of mobile ions diffusing toward the PRS probes. For example,

- Wood et al. (2015) found that the variability of net adsorption rates of Mn^{2+} , Fe²⁺ and S flux was greatest when the soil volumetric moisture content was highest in three undisturbed wetland sites from the same region as this study. Wood et al. (2015) interpreted these results as an increase in ion availability as well as increased mobility. However, if either a change in availability or mobility was the driving force of the observed variability in this study, all abundant ions should increase (or decrease) as they did in the Wood et al. (2015) study. Here, mobile S fluxes decreased in the
- 350 saturated soils where reduced metal ions were increasing. The loading of net adsorption rates of S on PC1 was almost equal to the other ions, but was negative (Table 4), suggesting that the leading mode of variability in ion adsorption rates had an opposite relationship between the reduced and oxidized ions. This suggests that microbially mediated REDOX reactions such as those by SRB and MRB are important in the changing biogeochemistry of the SFW peat soils. Negative correlations between time and mobile S and $NO₃$, while positive correlations between time and
- 355 reduced metals and NH₄⁺ provide additional evidence that alternative electron acceptors are being consumed over time (Table 2). The results presented here are comparable to a study by Kreiling et al. (2015) who found an increase in PRS-adsorbed Mn^{2+} and Fe²⁺ along with a decrease in mobile S with increasing flood frequency within the Mississippi River floodplain. They too attributed these changes in ion adsorption to decreasing REDOX potentials.

Although there is evidence that the soil REDOX conditions and alternative electron acceptor abundance is varying in 360 time and space at the SFW as described by the REDOX 'ladder', the variability in CH4 emissions were not strongly explained by these factors. This may be due in part to the complexity associated with electron acceptor abundance and CH4 production. In one study, methanogen communities were documented to become better competitors, relative to MRB, for scarce resources in Arctic tundra soils with increasing temperatures at higher REDOX potentials (Herndon et al., 2015). Granberg et al. (2001) demonstrated that vegetation, rapid temperature shifts, and N and S deposition all

- 365 had significant dependent effects on CH4 fluxes. In that study, the effects of the interaction terms were as large as, and sometimes inverse to the effects of each variable correlated to CH4 fluxes alone. Other factors might include the role of humic substances, soil heterogeneity and microsites, salinity, pH, and a breakdown of the REDOX ladder conceptual framework. For example, there is evidence that humic substances can suppress CH_4 production, by becoming anaerobic election acceptors (Blodau and Deppe, 2012), and more recently particulate organic matter itself
- 370 has been shown to be an important electron acceptor in peat soils (Gao et al. 2019). Microsites $\langle 10 \mu m \rangle$ with limited gas and water exchange with surrounding soil pore space also may affect overall soil REDOX potential and CH4 production by permitting localized electron acceptor depletion or abundance (Sey et al., 2008). Soil salinity may also have impacted the CH4 emissions from this wetland constructed on top of mine tailings; in 2013, the electrical conductivity was $792 \pm 616 \,\mu\text{S cm}^{-1}$ which increased to $2163 \pm 248 \,\mu\text{S cm}^{-1}$ in 2015 (Biagi et al., 2019). Although the
- 375 effects of salinity on CH4 production are not well understood, Poffenbarger et al. (2011) reviewed the literature and found CH₄ emissions were suppressed only in polyhaline wetlands (18 g L⁻¹ or \sim 24 mS cm⁻¹) where salinity far exceeded that of the SFW. In this constructed wetland the dominant form of cation is Ca^{2+} not Na+ and thus, it is unknown what effect this may have on the microbes at this time. However, $Na⁺$ concentrations have been increasing from 56 ± 52 mg L⁻¹ in 2013 to 130 \pm 109 mg L⁻¹ in 2015 (Biagi et al., 2019), suggesting a trend towards more common
- 380 Na⁺ dominated saline environments. The assemblage of anaerobic microbes that are thermodynamically favoured in a soil also varies with pH since the Gibbs energies of some alternative electron accepting metabolic process, but not others, vary with hydrogen ion concentration (Bethke et al., 2011; Flynn et al., 2014). Bethke et al. (2011) concluded that at neutral pH, the Gibbs energies of the major metabolic pathways of Fe^{3+} reduction, SO_4^2 reduction, and methanogenesis all converge. This contradicts the competitive exclusion concept of the REDOX 'ladder' at neutral
- 385 conditions. Other reactions can also have a cascading effect on the thermodynamics of the system. For example, Kreiling et al. (2015) found that precipitation of Fe^{2+} and H_2PO_4 lead to non-linear trends in Fe^{2+} ion adsorption to ion exchange resins despite increasing time in anoxic conditions. Kreiling et al. (2015) demonstrated that the precipitate removed waste products and maintained the system's relative abundance of oxidized iron ($Fe³⁺$), thereby maintaining forward reactions within Fe reducing metabolic pathways. Currently, microsite conditions are very

390 difficult to assess at the level Bethke et al. (2011) argue are needed to explain anaerobic microbial community dynamics on a thermodynamic basis.

At the SFW, it is reasonable to assume that N and S deposition near an oil sand processing plant could influence soil biogeochemistry and the results described here. Even before wetting, high concentrations of total S and available S in the peat used to construct the SFW were similar to an undisturbed fen in Alberta as reported by Chagué-Goff et al. (1996).

395 Such high concentrations of S occur when the groundwater that supplies the fen passes though coal and dark shale

deposits. Because Alberta is rich in both types of deposits, fens classified as "moderate rich" or "extreme rich" in terms of ion abundance, such as the four boreal fens described by Hartsock et al. (2016), are not uncommon in the region. High concentrations of S are also found in natural tidal wetlands where CH4 emissions are typically low (Poffenbarger et al., 2011). However, due to the abrupt change in environmental conditions affecting the salvaged peat with placement and

400 flooding in the SFW, the mobile S appears to be declining in abundance in the anoxic regions of this system. In the future, surface soil SO_4^2 may be replenished by diffusion from the underlying tailings which has high concentrations of gypsum added post-processing to promote aggregation of soft tailings (Oil Sands Wetland Working Group, 2014). However, there is no indication that the upward vertical transport of salts from the tailing sands is occurring at a rate to offset the current decline in mobile S (Biagi et al., 2019). Therefore, without any other processes liming production, CH4 405 emissions may increase in the future at the SFW.

6 Conclusions

Carbon cycling processes in constructed boreal plains lowlands are not yet well understood as these ecosystems have only existed for a few years. This study shows that CH4 emissions are very low in the SFW, one of the first of two boreal plains lowlands constructed in boreal northern Alberta. Methane emissions were low compared to rewetted 410 and restored peatlands but similar to another newly constructed wetland in the Alberta oil sand region. Changes in ion adsorption rates on buried ion exchange resins (PRS probes) were related to decreasing REDOX potentials and were used to support the argument that methanogen activity may be competitively suppressed by an abundance of alternative electron acceptors. Seasonal and interannual variations in net ion adsorption rates suggest microbially mediated changes in soil chemistry. In particular, decreases in PRS S adsorption occurs concurrently with modest increases in CH4 emissions in the

415 lowlands of the SFWsuggestingCH4 emissions are likely to increase in the future, if the trends in the abundance of electron acceptors continue. The findings of this research indicate that the design of the SFW promotes, at least initially, highly reduced soils in the lowlands with limited CH₄ production. This is significant as the REDOX conditions necessary for long term C accumulation appear to be achieved while limiting the emissions of a potent greenhouse gas.

7 Author Contributions

420 Funding was obtained by Humphreys and Carey who also supervised Clark during this work. The experiments were designed by Clark and Humphreys, with Clark conducting them in the field. This manuscript was written by Clark, who also performed the primary investigation, with contributions from all co-authors.

8 Data Availability

Data will be provided on request.

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11 Tables and Figures

Table 1: Soil and CH4 flux characteristics of the three location groups. The standard deviation from the mean is reported in brackets. Dominant vegetation is the most common plant type among the vegetated collars (by % cover).

***As the water table declined, plots were reclassified as unsaturated in the lowland at the time of sampling. Therefore, the sum of the groups each year exceeds the number of chambers reported in the text for that year.**

1When standing water was observed, soil moisture was not measured, and the volumetric soil moisture was set to 87 %. The depth of standing water varied from 0.03 m to 0.33 m.

Table 2: Spearman rank correlations to assess temporal trends in net ion adsorption rates for the three plot groups. Plant Root Simulator (PRS) data were categorized into chronological order (1-9 for the three burial periods over three years, 'Combined'), seasonal order regardless of year (1-3, 'Seasonal'), and annual order regardless of season (1-3, 'Annual'). Bold values indicate a significant trend at $a = 0.01$. The greatest temporal trend across all plot groups for a given ion and depth is shaded in **black.**

		Lowland saturated			Lowland unsaturated			Upland		
	Ion	Combined	Seasonal	Annual	Combined	Seasonal	Annual	Combined	Seasonal	Annual
Shallow	Mn^{2+}	0.26	0.22	0.21	-0.21	-0.29	-0.11	-0.35	-0.32	-0.23
Deep	Mn^{2+}	0.54	0.17	0.52	0.49	0.07	0.50	-0.27	-0.29	-0.17
Shallow	$Fe2+$	0.47	0.48	0.37	0.26	0.06	0.29	-0.01	-0.26	0.08
Deep	$Fe2+$	0.64	0.34	0.60	0.54	0.26	0.52	0.07	-0.16	0.10
Shallow	S	-0.64	-0.45	-0.54	0.03	-0.31	0.15	-0.02	-0.08	0.05
Deep	S	-0.74	-0.32	-0.71	-0.43	-0.48	-0.25	-0.08	-0.20	0.03
Shallow	TN	-0.19	-0.10	-0.19	-0.33	-0.19	-0.22	-0.17	0.04	-0.19
Deep	TN	-0.05	-0.09	-0.03	-0.28	-0.02	-0.28	-0.25	-0.01	-0.25
Shallow	NO ₃	-0.36	-0.36	-0.26	-0.30	-0.35	-0.10	-0.08	-0.06	-0.04
Deep	NO ₃	-0.14	-0.26	-0.05	-0.35	-0.27	-0.22	-0.15	-0.04	-0.13
Shallow	NH_4 ⁺	0.19	0.18	0.15	-0.14	0.17	-0.16	-0.33	0.43	-0.54
Deep	NH_4 ⁺	0.21	0.25	0.16	0.21	0.38	0.08	-0.37	0.31	-0.50

Table 3: Results of the linear mixed effects models for CH4 emissions, PRS ions and REDOX potential (Equations 1 and 2) measured August 2015 at the nine plots along the moisture gradient in the lowland of the Sandhill Fen Watershed. Values are the mean effect \pm the standard deviation. Bold values indicate parameters significantly **different from zero at the α = 0.05 level. Vegetation effects are only for the CH⁴ emission model (Equation 1). Plot effect is the standard deviation of the random effects from each plot.**

Response		β_1	β_2	α	$\text{Var}(\zeta)^{1/2}$
Variable	F-value	(Saturated)	(Depth/Vegetation)	(Intercept)	(Plot Effect)
CH ₄	0.84	0.34 ± 0.11 mg m ⁻² h ⁻¹	0.17 ± 0.06 mg m ⁻² h ⁻¹	-0.04 ± 0.07 mg m ⁻² h ⁻¹	$0.18 \,\mathrm{mg}\,\mathrm{m}^2\,\mathrm{h}^1$
Mn^{2+}	3.12	16.0 ± 3.7 µg 10 cm ⁻² month ⁻¹	$7.1 \pm 2.3 \,\mathrm{\mu g}$ 10 cm ⁻² month ⁻¹	$1.7 \pm 2.5 \,\mu g \, 10 \,\text{cm}^{-2} \,\text{month}^{-1}$	$5.5 \,\mu g \, 10 \, \text{cm}^2 \, \text{month}^{-1}$
$Fe2+$	2.99	$188 \pm 36 \,\mu g$ 10 cm ⁻² month ⁻¹	$42 \pm 24 \,\mu g \, 10 \,\text{m}^2 \, \text{cm}$ onth ⁻¹	$57 \pm 25 \,\mu g$ 10 cm ⁻² month ⁻¹	$51 \mu g 10 \text{ cm}^2$ month ⁻¹
S	3.8	$-588 \pm 92 \,\mu g \, 10 \,\text{cm}^{-2} \,\text{month}^{-1}$	$-74 \pm 62 \,\mu g \, 10 \,\text{m}^2 \,\text{cm}$ onth $^{-1}$	$1145 \pm 64 \,\mu g \, 10 \,\text{cm}^{-2} \,\text{month}^{-1}$	$126 \,\mu g \, 10 \,\text{cm}^2 \,\text{month}^{-1}$
REDOX	0.42	-116 ± 47 mV	-85 ± 33 mV	13 ± 32 mV	44 mV

Table 4: PCA loadings and the Pearson correlation for principle components (PC) 1 and 2 and REDOX potential and log transformed CH4 flux (p-values given in parentheses). Methane fluxes were averaged over the same one-month periods the PRS probes were buried. Bold numbers indicate significant correlations at $\alpha = 0.05$ **,** $n = 25$ **.**

PC loadings:			Pearson correlations:		
Ion	PC ₁	PC ₂	Predictor	PC ₁	PC ₂
NH_4 ⁺	0.41	0.76	$REDOX$ 0.2 m	$-0.275(0.166)$	0.254(0.201)
Mn^{2+}	0.54	0.1	$REDOX$ 0.4 m	$-0.06(0.767)$	$-0.025(0.901)$
Fe^{2+}	0.58	-0.15	$ln(CH_4 flux)$	0.295 (≤ 0.001)	0.187(0.016)
S	-0.45	-0.62			

Table 5: Pearson correlation coefficients between natural logarithm transformed CH4 fluxes and Plant Root Simulator (PRS) ion exchange resin measurements. Methane fluxes were averaged over the same one-month periods the PRS probes were buried, n = 25 for transect data n= 163 for entire study. Deep probes were buried outside the collars at 0.2 m, shallow probes were buried at 0.1 m outside the collars. Significant correlations (α **= 0.05) are in bold.**

		2015 Transect Only	All Data				
		PRS Probe Location		PRS Probe Location			
Ion	Deep	Shallow	Deep	Shallow			
NO ₃	-0.21	-0.37	-0.14	-0.16			
NH_4 ⁺	0.36	0.13	0.23	0.15			
Mn^{2+}	0.34	0.17	0.31	0.37			
$Fe2+$	0.31	0.40	0.31	0.38			
S	-0.47	-0.36	-0.06	0.12			

Figure 1: Map of the Sandhill Fen Watershed. Standing water was limited to the lowland region only.

Figure 2: Example of a plot with a collar pair in the wetland area illustrating a collar excluding vegetation (on the left) and a vegetated collar (right), surface PRS probes (orange and purple tabs with orange flagging tape, locations outside the collar are marked with circles). The permanent soil thermocouple profile post is marked by an 'X'. This plot is from the lowland unsaturated category (bottom of the eastern most boardwalk, Figure 1) and the photograph was taken July 3rd 2015.

Figure 3: Water table, precipitation and median CH4 fluxes by sampling date. Water table depth and precipitation (top panel) were measured at the central meteorological station (Figure 1). Circles indicate median CH₄ flux on the day of sampling and lines indicate the $25th$ to 75th percentiles for collars with vegetation (middle panel) and **collars maintained free of vegetation (bottom panel). An offset between vegetation groups has been added to the x-axis to reduce overlapping points. All groups were sampled on the same day, roughly once a week. Tick marks on the x-axis indicate first day of the labelled month. If the upper 75th percentile exceeded the range of the y axis, the value is printed on the plot.**

Figure 4: Ion availability as measured using Plant Root Simulator ion exchange resins over the three monthly burial periods for three years expressed in units of µg 10 cm-2 month-1 at two depths (0.1 m and 0.2 m). Units are a function of time the **probes were buried for. Metal ions were measured in their reduced forms and mobile S is the oxidized form of sulphur. An offset has been added between groups on the x-axis to reduce overlapping points.**

Figure 5: CH4 fluxes (top panel), net rates of ion adsorption measured with PRS ion exchange resins (middle panels) and the REDOX potential (bottom panel) measured every 15 min at 0.2 m and 0.4 m over the burial period in the lowland moisture gradient plots August 2015. Boxplots represent the interquartile range, with whiskers as 95 $%$ and lines representing the median for the nine plots (n = 4 for the saturated category where there was standing water, n = 5 for the unsaturated category, where **mean 0-0.20 m volumetric water content was 57 %)**