# Effect of water table drawdown on northern peatland methane dynamics: Implications for climate change

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[1] As natural sources of methane (CH<sub>4</sub>), peatlands play an important role in the global carbon cycle. Climate models predict that evapotranspiration will increase under a 2  $\times$ CO<sub>2</sub> scenario due to increased temperatures leading to lowered water tables at many northern latitudes. Given that the position of the water table within a peatland can have a large effect on CH<sub>4</sub> emissions, climate change may alter the CH<sub>4</sub> emissions from peatlands in this area. Research was conducted during 2001–2003 on natural and drained (8 years prior) sites within a poor fen in central Quebec. Flux measurements were made for each site at different microtopographical features that varied in depth to water table and vegetation cover. The quantity of CH<sub>4</sub> dissolved in the pore water was measured in the field and the potential of the peat for  $CH_4$  production and consumption was determined in the laboratory. Methane emissions and storage were lower in the drained fen. Growing season  $CH_4$  emissions at the drained site were 55% lower than the control site, primarily due to significantly reduced fluxes from topographic highs (up to 97% reduction), while the flux from topographically low areas remained high. The maintenance of high fluxes at these hollow sites was related to hydrological and ecological effects of the water table drawdown. The removal of standing water removed a potential zone of  $CH_4$  oxidation. It also enabled plant colonization at these locations, leading to an increase in gross ecosystem photosynthesis (GEP). At the hollow sites, seasonal CH<sub>4</sub> emissions were significantly correlated to seasonal GEP ( $R^2 = 0.85$ ). These results suggest that the response of northern peatland  $CH_4$  dynamics to climate change depends on the antecedent moisture conditions of the site. Moreover, ecological succession can play an important role for determining future CH<sub>4</sub> emissions, particularly from wetter sites. INDEX TERMS: 0315 Atmospheric Composition and Structure: Biosphere/ atmosphere interactions; 1615 Global Change: Biogeochemical processes (4805); 1851 Hydrology: Plant ecology; 1890 Hydrology: Wetlands; KEYWORDS: climate change, methane, peatland

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# 1. Introduction

[2] Northern peatlands play an important role in the global carbon cycle storing an estimated 500 Pg C [*Gorham*, 1991]. They are also large sources of atmospheric methane (CH<sub>4</sub>) [*Fung et al.*, 1991], with estimated annual releases of 46 Tg C-CH<sub>4</sub> [*Gorham*, 1991]. Large uncertainty exists regarding the ecohydrological response of peatland ecosystems to climate change [*Moore et al.*, 1998] making accurate predictions about the magnitude of future peatland CH<sub>4</sub> fluxes difficult. This research investigates CH<sub>4</sub> emissions and storage in a natural and drained poor fen in central Quebec

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in order to better predict the response of these systems to the water table drawdown expected to occur with climate change.

[3] Methane is produced under the saturated conditions present in peatlands by anaerobic decomposition of organic material by methanogenic bacteria [Zinder, 1993]. Once produced, CH<sub>4</sub> can be released to the atmosphere by diffusion, ebullition, and transport via vascular plants. While transport by diffusion is slow and often results in the majority of CH<sub>4</sub> being lost to oxidation in the unsaturated, aerobic zone of the peat profile [Oremland and Culbertson, 1992], transport through the aerenchyma of wetland vascular plants bypasses the oxic zone, serving as an important pathway for CH<sub>4</sub> emission at many sites [Schutz et al., 1991; Greenup et al., 2000]. Vascular vegetation can also affect CH<sub>4</sub> emission by supplying fresh substrate for CH<sub>4</sub> production to anoxic layers through root exudation and decay [Bellisario et al., 1999] and by increasing CH<sub>4</sub> oxidation in the rhizosphere [Bellisario et al., 1999; Popp et al., 2000]. Overall, the presence of

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Microtopography	Average Growing Season Water Table (cm relative to peat surface)	Dominant Vegetation
		Control Site
Hummock	-21 to $-14$	Sphagnum rubellum, Chamaedaphne calyculata, Carex oligosperma
Lawn	-14 to $-6$	Sphagnum papillosum, Carex limosa
Hollow	0 to 20	Sphagnum cuspidatum, Rhynchospora alba, Scirpus subterminalis
		Drained Site
Hummock	-42 to $-26$	Polytrichum strictum, Chamaedaphne calyculata
Lawn	-19 to $-14$	Sphagnum magellanicum, Carex oligosperma
Hollow	-10 to $-6$	Sphagnum papillosum, Sphagnum cuspidatum, Rhynchospora alba

Table 1. Site Descriptions Along the Microtopographic Gradient at the Control and Drained Sites

vascular vegetation leads to enhanced CH<sub>4</sub> emissions relative to unvegetated sites [*Shannon et al.*, 1996; *Waddington et al.*, 1996; *Greenup et al.*, 2000; *Tuittila et al.*, 2000].

[4] Owing to these mechanisms of production, oxidation, and emission, overall CH<sub>4</sub> flux from a peatland is dependent on the water table position [Granberg et al., 1997], peat temperature [Daulat and Clymo, 1998], substrate quality [Kelly and Chynoweth, 1981], and primary productivity [Whiting and Chanton, 1993]. Global climate change will likely alter all of these variables, thus affecting CH<sub>4</sub> emissions. Using a  $2 \times CO_2$  scenario in which temperature and precipitation in northern Canada increased 3°C and 1 mm day<sup>-1</sup>, respectively [Mitchell, 1989], Roulet et al. [1992] used a simple hydrological model and estimated that water tables would decline by an average of 14 cm in northern Canadian peatlands. Incorporating this value into contemporary CH<sub>4</sub>-water table-temperature relationships resulted in the conclusion that  $CH_4$  emissions from northern peatlands would decline by 74-81%. This is supported by several studies that have reported lower CH<sub>4</sub> fluxes from drained peatlands relative to undrained controls [Roulet et al., 1993; Nykänen et al., 1995, 1998]. Waddington et al. [1998] investigated the effect of the 14-cm decline in water table position predicted by Roulet et al. [1992] on net ecosystem CO<sub>2</sub> exchange (NEE) at a fen by considering the potential for ecological succession. When the response of the vegetation community to the water table shift was considered, the resulting carbon balance at the site depended on its initial moisture conditions. It was determined that wet sites (initial mean water table position above 0.1 m below the surface) would become larger sinks for atmospheric CO<sub>2</sub>, while drier sites would become sources. Thus, under climate change, ecological succession caused by water table drawdown may lead to increased carbon uptake at wet (hollow) sites, providing substrate for CH<sub>4</sub> production. If ecological succession leads to an increased coverage of vascular plants with aerenchymatic tissues, an increase in plant-mediated transport of CH<sub>4</sub> may impact overall emissions. The extent of this ecological succession and its impact on CH<sub>4</sub> emissions requires further investigation.

[5] The objectives of this study are (1) to determine differences in  $CH_4$  emissions between a natural poor fen and one in which the water table has been reduced by approximately 20 cm, (2) to determine differences in belowground pools of dissolved methane between the two sites, and (3) to investigate both hydrological and ecological

explanations for these changes in order to better predict the effect of climate change on peatland CH<sub>4</sub> fluxes.

# 2. Materials and Methods

#### 2.1. Study Area

[6] The study was carried out in a poor fen fragment (46°40'N 71°10'W) near St. Charles-de-Bellechasse, Quebec, Canada. Within the fragment are several pool-ridge complexes, one of which had the water table lowered approximately 20 cm 8 years prior to the study (drained). This pool was compared to a natural pool-ridge complex (control). Peat depth at the control site is approximately 120 cm, while at the drained site it is 80 cm. Nine sampling sites were arranged along the microtopographic gradient at each pool. These locations represented a moisture gradient, with three sites at each of hummock, lawn, and hollow. The general characteristics and site description of sampling locations, including mean water table position and dominant vegetation, for both the control and drained sites are shown in Table 1.

# 2.2. CH<sub>4</sub> Flux

[7] CH<sub>4</sub> flux was determined using the static chamber method [*Tuittila et al.*, 2000] at the sampling locations described above. Samples were analyzed within 48 hours of collection on a Varian 3800 gas chromatograph (GC) equipped with flame ionization detector at 250°C and Porapak N column at 50°C with helium as the carrier gas and a flow rate of 30 mL min<sup>-1</sup>. CH<sub>4</sub> emissions were measured weekly from July to September 2001 and May to September 2002 and monthly from October 2001 to March 2002 and November 2002 to January 2003.

#### 2.3. Subsurface CH<sub>4</sub>

[8] Pore water samplers, consisting of a 20 cm length of 2.5 cm inner diameter (i.d.) PVC pipe, closed at both ends, slotted at the middle 10 cm, and covered in mesh to prevent clogging, were installed at the lawn areas of each pool. Tygon tubing was inserted at the top and extended above the surface of the peat to allow for collection of water with a syringe. Samplers were placed at 25, 40, and 60 cm and 25, 40, 60, 85, and 100 cm below the surface at the drained and control site, respectively. Pore water was collected weekly from May to September in 2002 and 2003, and CH<sub>4</sub> concentration was determined using headspace analysis

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[*Ioffe and Vitenberg*, 1982] after equilibration with nitrogen  $(N_2)$ .

# 2.4. Potential Methane Production (PMP) and Consumption (PMC)

[9] Cores were hand cut and removed from hollow, lawn, and hummock locations at both sites and returned immediately to the lab, where they were frozen until analyzed. Peat samples for incubation were taken at 10-cm-depth intervals from all cores, resulting in samples centered at 5, 15, 25, 35, and 45 cm below the surface for all sites.

[10] PMP and PMC were determined under anaerobic and aerobic conditions, respectively. For the aerobic experiment, approximately 15 to 25 g of wet peat was placed in 250-mL incubation jars and spiked with 20 mL of a 53 ppmv CH<sub>4</sub> standard, resulting in an initial mean CH<sub>4</sub> concentration of 5.1 ppm. For the anaerobic experiment, peat slurries of approximately 15 to 25 g of wet peat were made in 250-mL incubation jars by the addition of enough distilled water to saturate samples without allowing for standing water. These slurries were then flushed with N<sub>2</sub> for 15 min and sealed. Aerobic and anaerobic samples were incubated at 4° and 20°C in the dark and were sampled initially and every 12 hours over a 48-hour period. Anaerobic samples were mechanically agitated for 20 min prior to sampling to mix the gases within the peat pore spaces and the jar headspace.

#### 2.5. Environmental Variables

[11] Temperature profiles in the peat (surface to 30 cm depth) were measured at each collar on each sampling date using a thermocouple thermometer. Water table position was determined at each collar on each sampling date using 1.3-1.9 cm i.d. PVC wells installed adjacent to the collar.

[12] Leaf area index (LAI) was determined by counting all vascular plant leaves within five  $7.5 \times 7.5$  cm subplots distributed throughout each collar each month. Total leaf numbers within the collar were extrapolated from these values. Individuals of the dominant species were selected outside of the collar at each site, and their leaves were measured biweekly. An average biweekly surface area of leaves was computed and multiplied by leaf numbers to determine LAI.

[13] Gross ecosystem photosynthesis (GEP), at a variety of radiation (PAR) levels, and ecosystem respiration were determined weekly at each collar using a clear Plexiglas chamber and a PP systems EGM-1 portable infrared gas analyzer. Light levels were controlled with shades, and the chamber was covered with an opaque shroud for respiration measurements. The temperature inside the chamber was maintained within 5°C of ambient temperatures using a radiator cooling system. Results were fitted to PAR values using Photosyn Assistant 1.1 software (Dundee Scientific, UK), and these relationships were combined with continuously recorded PAR data to estimate seasonal  $CO_2$  uptake at each collar.

# 3. Results

### 3.1. Site Characteristics

[14] As shown in Table 1, water table position was lower at the drained site, ranging from 6 to 42 cm below

the surface along the microtopographical gradient, compared to the control site, which had water table positions in the range of 26 cm below the surface to 20 cm above the surface. The differences in water table position have led to the dominance of different vegetation communities (Table 1).

[15] Peat temperature at 30 cm depth was warmer at the control site for all microtopographic units, with lawns and hollows being significantly warmer (t-test, lawns, p = 0.01; hollows, p < 0.001). Average 30-cm peat temperatures from May 20 to September 23 for 2002 at hummocks, lawns, and hollows were 16.7°, 17.5°, and 18.8°C at the control site and 16.0°, 15.8°, and 16.5°C at the drained site.

[16] In addition to differences in vegetation community (Table 1), there were differences in LAI between sites. Maximum LAI between May and September 2002 at hummocks, lawns, and hollows was 0.8, 1.0, and 1.4 m<sup>2</sup> m<sup>-2</sup> at the control site, and 0.3, 4.7, and 1.0 m<sup>2</sup> m<sup>-2</sup> at the drained site. Despite having a high maximum LAI, the control hollows only had a substantial cover of vascular vegetation after July and a sparse cover of Sphagnum (0–3%). In contrast, drained hollows had vascular vegetation present throughout the sampling period and a *Sphagnum* cover of 80–97%.

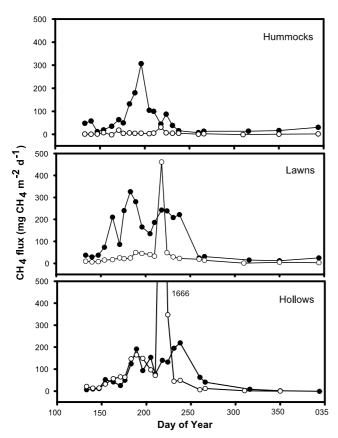
# 3.2. CH<sub>4</sub> Fluxes

[17] The average growing season CH<sub>4</sub> fluxes were 143.4 and 64.4 mg  $CH_4$  m<sup>-2</sup> d<sup>-1</sup> at the control and drained sites, respectively. Flux values varied greatly both spatially and temporally (Figure 1) at both sites. The standard deviation on a particular date at control hummocks, lawns, and hollows was 1.0-330.7, 2.7-196.5, and 0.8-206.6 mg  $CH_4 m^{-2} d^{-1}$ . At the drained sites the standard deviation was 0.5–12.5, 1.4–243.4, and 0.9–245.9 mg  $CH_4 m^{-2} d^{-1}$ at hummocks, lawns, and hollows, respectively. Spatial variability is to be expected as the sampling locations were placed along a topographic gradient to reveal the effects of environment on CH<sub>4</sub> flux. Average winter fluxes for each microtope (Table 2) were less variable and substantially lower than growing season fluxes. They are relatively unimportant for the calculation of annual CH<sub>4</sub> emissions from low-lying areas, but may make up about 20% of total emissions from hummocks.

[18] CH<sub>4</sub> emissions were significantly higher at the control site than the drained site for hummock and lawn collars (t-test, p < 0.01). However, no significant difference was observed for hollows between the two sites. Average CH<sub>4</sub> emissions were weakly related to average water table position at both sites (Figure 2), showing a unimodal ( $R^2 = 0.25$ ) and exponential relationship ( $R^2 = 0.76$ ) at the control and drained sites, respectively. No significant relationship was found between daily water table position and individual flux measurements. There was also no correlation between average peat temperature and average CH<sub>4</sub> fluxes.

# 3.3. Subsurface CH<sub>4</sub>

[19] The concentration of  $CH_4$  in the pore water at lawn areas was significantly higher (p < 0.01) at all depths at the control site when compared to the drained site for both 2002



**Figure 1.** Measured  $CH_4$  fluxes for control (solid symbols) and drained (open symbols) sites along a microtopographic gradient. Each point is the average of the three collars representing a particular microtopographic unit (hummock, lawn, hollow). Data collected between May 2002 and January 2003 are presented. Values outside of the scale of the figure are given as text on the plot.

and 2003. At the control site, average seasonal values of dissolved CH<sub>4</sub> were 1.4–5.8 mg L<sup>-1</sup> in 2002 and 1.3– 5.3 mg L<sup>-1</sup> in 2003. At the drained site values were 0.3–0.7 and 0.6–1.7 mg L<sup>-1</sup> in 2002 and 2003, respectively. The profile of average dissolved CH<sub>4</sub> was similar for both years (Figure 3); however, temporal shifts in concentration generally occurred simultaneously for all depths and sites in 2002, but were more erratic in 2003 (data not shown). Profiles of CH<sub>4</sub> collected as bubbles within the peat had a

**Figure 2.** Average growing season CH<sub>4</sub> flux at each collar plotted against average growing season water table at each collar (negative values are below the surface) based on data collected from July 2001 to September 2002. Water table explains some of the variability at the control site (solid symbols, solid line,  $R^2 = 0.25$ ) but is more significant at the drained site (open symbols, dashed line,  $R^2 = 0.76$ ).

similar pattern to pore water concentrations in both years (data not shown).

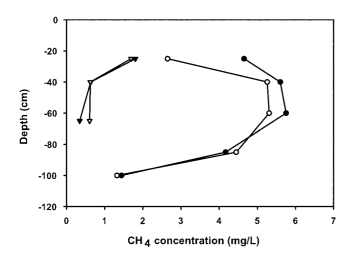
# 3.4. Potential CH<sub>4</sub> Production and Consumption

[20] In most cases the rate of CH<sub>4</sub> production and consumption by the peat was not notably different at the two sites. At lawns sites, PMP and PMC were in the range of 2.5–44.7 and 2.8–32.3 ng CH<sub>4</sub> cm<sup>-3</sup> (fresh peat) d<sup>-1</sup> at the control site and 4.2–10.7 and 10.2–18.8 ng CH<sub>4</sub> cm<sup>-3</sup> d<sup>-1</sup> at the drained site. At hummocks, control and drained sites had PMP of 1.8–5.7 and 2.9–6.0 ng CH<sub>4</sub> cm<sup>-3</sup> d<sup>-1</sup> and PMC of 6.3–18.0 and 10.6–26.1 ng CH<sub>4</sub> cm<sup>-3</sup> d<sup>-1</sup>, respectively. The only notable differences between the sites occurred at 20–40 cm for lawns and 10–20 cm for hummocks for which the drained site had higher consumption rates than the control site lawn, which had higher production rates than the drained peat.

[21]  $Q_{10}$  values between 4° and 20°C were less than 2 for all cases except for production in the top 10 cm ( $Q_{10} = 2.1$ ) and consumption in the upper 20 cm ( $Q_{10} = 2.6$  from 0– 10 cm and  $Q_{10} = 2.8$  from 10–20 cm) of the control lawn peat sample. This suggests that the reaction rate was limited

Table 2. Winter and Growing Season CH<sub>4</sub> Fluxes

Location		Total CH <sub>4</sub> Emitted mg CH <sub>4</sub> m <sup><math>-2</math></sup> d <sup><math>-1</math></sup> (Mean (Std Dev))		Winter Flux as
	Site	Winter (Nov. 1 to April 15)	Growing Season (April 15 to Oct. 31)	Percentage of Total
Hummocks	control	3.2 (1.5)	9.7 (1.5)	23.8
	drained	0.2 (0.2)	1.2 (0.9)	17.6
Lawns	control	2.8 (2.7)	16.3 (6.8)	14.4
	drained	1.5 (0.9)	7.6 (1.5)	15.7
Hollows	control	0.1 (0.2)	14.5 (14.3)	1.7
	drained	0.2 (0.2)	22.0 (4.9)	0.8



**Figure 3.** Average dissolved CH<sub>4</sub> concentration at depth in 2002 (solid symbols) and 2003 (open). Circles and triangles represent the control and drained sites, respectively.

by a factor other than temperature for both  $CH_4$  production and consumption at both the control and drained sites.

# 4. Discussion

### 4.1. Changes in CH<sub>4</sub> Dynamics

[22] When averages of flux measurements for all collars at each site were compared, growing season  $CH_4$  fluxes were 55% lower at the drained site than the control site. The postwater table drawdown decline was the result of large changes in  $CH_4$  emissions at hummocks (97% decline) and lawns (71% decline), while hollow fluxes were unchanged. This reveals that the antecedent moisture conditions of the site play a significant role in its response to water table changes and thus potentially in its response to climatic change.

[23] Since PMP and PMC are relatively similar at hummocks at the two sites, the reduction in flux is due primarily to the decrease in the size of the anoxic zone and subsequent increase in the size of the oxic zone as opposed to a shift in the peat properties (peat quality, microbial biomass, microbial community). At the lawns the large declines in CH<sub>4</sub> emissions are mirrored by significantly smaller pools of subsurface CH<sub>4</sub> at the drained site. These smaller subsurface pools suggest a reduction in CH<sub>4</sub> production and increase in CH<sub>4</sub> oxidation at the drained site lawns compared to the control site. This is in accordance with the reduced size of the anoxic zone and increased size of the oxic zone. The maintenance of the drained hollow CH<sub>4</sub> fluxes at the same level as the control site is related to both the hydrological and ecological impacts of the water table drawdown.

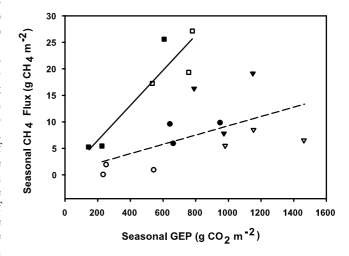
# 4.2. Hydrological Effects of Water Table Drawdown on Hollows

[24] The relationship between water table and  $CH_4$  flux at the control site is unimodal, with highest fluxes occurring at intermediate water table positions while sites with large depths of standing water have lower fluxes (Figure 2). Thus a water table drawdown of 20 cm shifts the hollow sites

from the inundated low fluxes toward an "optimal" water table position, suggesting a maintenance or increase in  $CH_4$  emissions. This concept is supported by the exponential water table- $CH_4$  flux relationship at the drained site, which results from the lack of sites with standing water (the cause of the reduced fluxes at the control site).

[25] Reduced fluxes at sites with standing water are likely the result of differences in substrate availability, temperature regime and the potential for oxidation between flooded and nonflooded sites. A lower availability of fresh substrate at flooded areas is suggested by the lower productivity of these sites. In Figure 4 the two sites with standing water are present in the lower left-hand corner of the figure and thus have the lowest GEP of all the sites studied. Analysis of the peat temperature profile measured on each sampling day reveals that peat temperatures were significantly higher (p <0.001) at the control site hollows  $(18.8^{\circ}C)$  than drained hollows (16.5°C) at 30 cm depth. The dark color of the water and sediment likely increases the absorption of radiation leading to preferential heating of the pools. A similar observation was made by Weyhenmeyer [1999], who found high sediment temperatures in beaver ponds and suggested the dark color of the sediment as the cause of this warming. Thus temperature actually declines at the hollows due to drainage, which should promote a decline in CH<sub>4</sub> emissions; however, this was not observed.

[26] Wind mixing of open water may introduce dissolved oxygen (DO) into the water column, creating a partially aerobic zone in which  $CH_4$  can be oxidized as it is transported to the surface by diffusion. The deep control hollows have standing water year round, whereas the hollows at the drained site are only periodically flooded at snowmelt and after heavy summer rains. At this time, standing water is



**Figure 4.** Cumulative growing season CH<sub>4</sub> emissions as a function of cumulative growing season gross ecosystem photosynthesis (GEP) for 2002. Solid symbols represent control site plots, while open symbols are drained sites. Circles, triangles, and squares represent hummocks, lawns, and hollows, respectively. Relationship between GEP and CH<sub>4</sub> emissions for hollows shown as solid line, (p < 0.01,  $R^2 = 0.85$ ), and for hummock and lawns sites, dashed line, (p = 0.07,  $R^2 = 0.29$ ).

Site	Initial CH <sub>4</sub> Flux, mg CH <sub>4</sub> m <sup><math>-2</math></sup> d <sup><math>-1</math></sup>	Perturbed CH <sub>4</sub> Flux, mg CH <sub>4</sub> m <sup><math>-2</math></sup> d <sup><math>-1</math></sup>	Percent Change
Overall	143	64	-55
Hummocks	125	3	-97
Lawns	160	46	-71
Hollows	152	144	-0.05
Roulet et al. [1992] floating	116	30	-74
Roulet et al. [1992] nonfloating	74	14	-81

**Table 3.** Comparison of Average Growing Season  $CH_4$  Emissions Between Initial Conditions (Control Site) and Perturbed Conditions (Drained)<sup>a</sup>

<sup>a</sup>Calculations of changes in CH<sub>4</sub> emissions with water table drawdown from *Roulet et al.* [1992] model predictions are also shown for comparison.

only present for several days. Dissolved oxygen content in standing water at the control hollows was 6.2 mg/L at the surface, declining to 2.3 mg/L at a depth of 25 cm. This indicates that the water column may be a zone of oxidation. The slow rate of diffusion in water ( $10^4$  times slower than air) allows substantial time for CH<sub>4</sub> oxidation as CH<sub>4</sub> moves through the water column, leading to reduced CH<sub>4</sub> fluxes from these sites.

[27] The maintenance of CH<sub>4</sub> emissions at the drained hollows in a range similar to the control site results from the fact that at these locations the 20-cm water table reduction removes standing water, but maintains water tables relatively close to the surface. Therefore the oxic zone within the peat profile of drained hollows remains small, limiting CH<sub>4</sub> oxidation and maintaining high CH<sub>4</sub> fluxes. This effect is enhanced by the fact that the removal of standing water (which contains some dissolved oxygen) may actually eliminate a source of CH<sub>4</sub> oxidation. As well, peat in hollow areas may be "floating," having a surface that oscillates with water table changes. Thus, when the water table drops, the peat surface also drops, maintaining relatively stable moisture conditions within the peat column and limiting the development of an oxic zone. Dise et al. [1993] found no significant relationship between CH<sub>4</sub> flux and water table at floating sites because of surface level oscillations. A similar response is likely at play at drained hollow sites in this study.

# 4.3. Ecological Effects of Water Table Drawdown on Hollows

[28] The removal of standing water from the drained hollows has also enabled ecological succession, which may in turn have affected CH<sub>4</sub> fluxes. Of the hollows at the control site, those with standing water have no Sphagnum cover and contain only Scirpus subterminalis, which does not have a substantial leaf area until July. This results in low levels of GEP at these sites in May and June (data not shown). Drained hollow sites have a high surface cover of Sphagnum (>80% cover) and a diverse vascular community. These differences in vegetation community between the control and drained hummocks result in large differences in seasonal gross ecosystem production (GEP), with average seasonal GEP of 325 and 690 g  $CO_2 m^{-2} d^{-1}$  at control and drained hollows, respectively. At the hollows of both the control and drained sites, a good relationship between seasonal GEP and seasonal CH4 emissions was found (Figure 4, p < 0.01,  $R^2 = 0.85$ ). Several other studies have

also found a linear relationship between site productivity and CH<sub>4</sub> emissions [Whiting and Chanton, 1993; Waddington et al., 1996; Bellisario et al., 1999]. The correlation between productivity and CH<sub>4</sub> emissions is the result of the provision by vascular vegetation of fresh substrate for CH<sub>4</sub> production and a conduit for CH<sub>4</sub> release. A much weaker relationship ( $p = 0.07, R^2 = 0.29$ ) was present between seasonal GEP and seasonal CH<sub>4</sub> emissions for hummock and lawn sites (Figure 4) where vascular plants with aerenchymatic tissues were less abundant (Table 1). Waddington et al. [1996] found a similar pattern in which the role of vascular vegetation was more pronounced when water tables were near the surface. Since lawn and hummock sites are much drier than the hollows, it is likely that moisture conditions are exerting an overriding influence on CH<sub>4</sub> production and oxidation, making the impact of ecosystem productivity secondary. Thus at sites with relatively dry initial conditions and vascular plant community whose roots are located in the permanently oxic layer, the impact of ecological succession caused by water table drawdown plays a less significant role in the maintenance of CH<sub>4</sub> flux as these sites become drier.

# 5. Climate Change Implications

[29] The reduction in CH<sub>4</sub> flux by 55% observed in this study is much lower than the 74-81% reduction predicted by Roulet et al. [1992] (Table 3). The reduction observed at the lawn sites (71%) is similar to the Roulet et al. [1992] model prediction. This similarity may result from the fact that the lawn sites in this study had a similar water table position to the average used for calculations by Roulet et al. [1992]. Our findings reveal the importance of antecedent moisture conditions and peatland ecotope in determining the response of a site to water table drawdown and thus potentially to climate change. Sites situated at the low-lying extreme of the microtopographic gradient did not experience a reduction in CH<sub>4</sub> fluxes with water table drawdown because at these sites this resulted in the removal of standing water and allowed for ecological succession, which has the potential to enhance CH<sub>4</sub> flux. This study has shown that ecological succession can have a significant impact on the response of peatland CH<sub>4</sub> fluxes to long-term shifts in water table, and this feedback should be included in future predictions of peatland climate change response. As well, further research is needed to quantify the evolving effects of vascular vegetation on CH4 fluxes between natural and drained sites, as well as the time frame for ecological succession following water table drawdown.

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