Magnitude and seasonality of wetland methane emissions from the Hudson Bay Lowlands (Canada)

C. A. Pickett-Heaps¹*, D. J. Jacob¹, K. J. Wecht¹, E. A. Kort¹, S. C. Wofsy¹, G. S. Diskin², D. E. J. Worthy³, J. O. Kaplan⁴, I. Bey⁵, and J. Drevet⁶

¹School of Engineering and Applied Sciences & Department of Earth and Planetary Sciences, Harvard University, Cambridge MA 02138, USA
²NASA Langley Research Center, Hampton, Virginia, USA
³Environment Canada, Toronto, Ontario, Canada
⁴Environmental Engineering Institute, École Polytechnique Fédérale de Lausanne, Lausanne, Switzerland
⁵Center for Climate Systems Modeling (C2SM), ETH Zürich, Universitätstrasse 16, Zürich, Switzerland
⁶Laboratoire de Modélisation de Chimie Atmosphérique, École Polytechnique Fédérale de Lausanne, Lausanne, Switzerland

*current address: CSIRO – Marine and Atmospheric Research, GPO Box 3023, Canberra, ACT, 2601, Australia

Abstract. The Hudson Bay Lowlands (HBL) is the second largest boreal wetland ecosystem in the world and an important natural source of global atmospheric methane. We quantify the HBL methane emissions by using the GEOS-Chem chemical transport model to simulate aircraft measurements over the HBL from the ARCTAS and pre-HIPPO campaigns in May–July 2008, together with continuous 2004–2008 surface observations at Fraserdale (southern edge of HBL) and Alert (Arctic background). The difference in methane concentrations between Fraserdale and Alert is shown to be a good indicator of HBL emissions, and implies a sharp seasonal onset of emissions in late May (consistent with the aircraft data), a peak in July–August, and a seasonal shut-off in September. The model, in which seasonal variation of emissions is mainly driven by surface temperature, reproduces well the observations in summer but its seasonal shoulders are too broad. We suggest that this reflects the suppression of emissions by snow cover and greatly improve the model simulation by accounting for this effect. Our resulting best estimate for HBL methane emissions is 2.3 Tg a⁻¹, several-fold higher than previous estimates (Roulet et al., 1994; Worthy et al., 2000).

1 Introduction

Methane is the second most important anthropogenic greenhouse gas after carbon dioxide (IPCC, 2007). Methane concentrations have increased from 700 ppbv in the pre-industrial atmosphere to 1700 ppbv by the early 1990s (Etheridge et al., 1998). This increase is presumably driven by direct emissions from industry and agriculture (IPCC, 2007), but could also reflect changes in the chemical sink (reaction with the OH radical) and the effects of climate change on natural emissions (Worthy et al., 2000). Wetlands are the largest natural source of methane and are highly sensitive to changes in climate (Kaplan et al., 2006), especially in the boreal zone (Zhuang et al., 2006; Sitch et al., 2007). Here we use aircraft observations over the Hudson Bay Lowlands in northern Ontario as well as surface observations at Fraserdale and Alert to better quantify this boreal wetland source.

Methane is produced in boreal wetlands by heterotrophic respiration of partially decomposed organic material under anoxic conditions. As it rises to the surface, it may encounter oxic conditions resulting in oxidation. The remaining methane escapes to the atmosphere (Walter et al., 2001). Production of methane primarily depends on soil temperature, and the distribution of anoxic and oxic zones depends primarily on the water table level (Pelletier et al., 2007; Moore et al., 1994). Compounding this complexity in boreal wetland emissions is the seasonal thaw that propagates from the surface to depth (Dunn et al., 2009; Wania et al., 2009).
The Hudson Bay Lowlands (HBL) is an ecologically significant and well-studied boreal wetlands region (e.g., Glooschenko et al., 1994). It is (after the West Siberian wetlands) the second largest semi-continuous wetland region in the world, covering an area of 320,000 km$^2$ or about 10% of the total area covered by boreal wetlands (Wang et al., 2008; Glooschenko et al., 1994). The ABLE-3B/NOWES airborne and ground campaign conducted in the summer of 1990 estimated an annual methane emission of $0.5 \pm 0.3$ Tg a$^{-1}$ from the HBL (Roulet et al., 1994). Worthy et al. (2000), using inverse methods to interpret observations from the Alert and Fraserdale Canadian sites, obtained a similar estimate of $0.2–0.5$ Tg a$^{-1}$. These estimates seem low compared to global inversion studies that infer total boreal wetland emissions of $27–38$ Tg a$^{-1}$ (Hein et al., 1997; Wang et al., 2004; Bousquet et al., 2006), although Chen and Prinn (2006) obtained an estimate of $7$ Tg a$^{-1}$. Typically, the HBL is assumed to contribute 10% to global boreal wetland emissions.

The above inconsistencies point to the need for a better understanding of methane emissions from the HBL as a window into the global boreal wetland source. We exploit here methane concentration measurements from the ARCTAS and Pre-HIPPO aircraft campaigns in May–July 2008 (Fig. 1), together with long-term surface data from Environment Canada at Fraserdale (81.6° W, 49.9° N) and Alert (62.5° W, 82.5° N). We interpret these data using a global bottom-up scheme for wetland emissions implemented in the GEOS-Chem chemical transport model (CTM). The ARCTAS airborne campaign based in Cold Lake, Alberta (Jacob et al., 2010) conducted three flights over the HBL in early July, while the Pre-HIPPO campaign based in Boulder, Colorado (Pan et al., 2010) conducted 2 flights in the region in May–June.

2 Model description

We use the GEOS-Chem CTM originally described by Bey et al. (2001) in a methane simulation for 2004–2008 to interpret the aircraft and surface observations. GEOS-Chem is driven by GEOS-5 analyzed meteorological data from the NASA Global Modeling and Assimilation Office (GMAO). The GEOS-5 data have $1/2^\circ \times 2/3^\circ$ (lat, lon) horizontal resolution with 72 vertical levels and 6-h temporal resolution (3-h for surface variables and mixing depths). The horizontal resolution is degraded here to $2^\circ \times 2.5^\circ$ for input to GEOS-Chem.

The methane simulation in GEOS-Chem was originally described by Wang et al. (2004), and subsequently improved and updated by Drevet et al. (2011). Major sources include anthropogenic emissions from EDGAR 4.0 (European Commission, 2009), natural emissions from wetlands as described below, and GFED2 biomass burning emissions (Giglio et al., 2006). Chemical loss of methane is computed using a global 3-D archive of monthly average OH concentrations from a GEOS-Chem simulation of tropospheric chemistry. The mean tropospheric OH concentration is $10.8 \times 10^5$ molecules cm$^{-3}$, typical of current models (a global model inter-comparison by Shindell et al. (2006) gives a mean value of $11.1 \pm 1.7 \times 10^5$ molecules cm$^{-3}$). The corresponding tropospheric lifetime of methylchloroform against oxidation by OH is 5.3 yr, within the range constrained by observations (Prinn et al., 2005). Additional minor sinks for methane in the model include stratospheric oxidation prescribed as a constant decay (Wang et al., 2004) and soil absorption (from EDGAR 4.0).

The wetlands emission scheme in GEOS-Chem is based on Kaplan (2002). The scheme applies methane emission factors to heterotrophic respiration rates in tropical and boreal wetlands, following algorithms described by Kaplan et al. (2002), Sitch et al. (2003) and Bergamaschi et al. (2007). The emission flux $E$ (molecules CH$_4$ m$^{-2}$ s$^{-1}$) for each model grid square is given by:

$$E = \delta W F \beta A e^{-T_{0} / T_{1}} \sum_{i=1}^{2} C_{i}$$

where $C_{1}$ and $C_{2}$ (mol C m$^{-2}$) are soil and litter carbon pools, respectively, specified on a $2^\circ \times 2.5^\circ$ grid by the Lund-Potsdam-Jena dynamic global vegetation model (Sitch et al., 2003). These pools have fixed residence times $T_{1} = 32$ yr and $T_{2} = 2.8$ yr from Sitch et al. (2003). The Arrhenius factor with $A = 1.0e^{+3}$, $E_{0} = 309$ K, and $T_{0} = 227$ K (Lloyd and Taylor, 1994) specifies the temperature dependence of respiration. The soil temperature $T$ is taken here as the GEOS-5 skin temperature. The methane emission factor is
$\beta = 3 \times 10^{-2}$ mol CH$_4$ per mol C respired (Christensen et al., 1996). An additional scaling factor $F$ is used to match observed ecosystem fluxes of methane, separately for tropical (T) and boreal (B) wetlands:

$$F = \alpha \cdot F_T + (1 - \alpha) \cdot F_B$$

(2)

where $\alpha = \min\{\exp[(\bar{T} - T_1)/8], 1\}$, $\bar{T}$ (K) is the mean monthly soil temperature taken here as the GEOS-5 skin temperature, and $T_1 = 303.15$ K. Scaling factors $F_T = 0.14$ and $F_B = 0.005$ were derived by J. Kaplan and J. Drevet (personal communication, 2009) to match published emission estimates for the Amazon (Melack et al., 2004) and boreal wetlands (Hein et al., 1997; Wang et al., 2004). Finally, $W$ in Eq. (1) represents the maximum potential fraction of wetland coverage for the grid square, as obtained from various databases described by Kaplan et al. (2002) and Bergamaschi et al. (2007). Whether a wetland is actually present ($\delta = 1$) or not ($\delta = 0$) over that fraction $W$ at any given time depends on the level of the water table, which we diagnose using GEOS-5 soil moisture as a proxy following Bergamaschi et al. (2007). Wetlands are present if the soil moisture exceeds a specified threshold of 0.1 for the ratio between the soil water content and the porosity of the soil (J. O. K. Kaplan, personal communication, 2009)

Annual emissions for the HBL (geographically defined as 50° N–60° N, 75° W–96° W) computed in the above manner in GEOS-Chem average 2.9 Tg a$^{-1}$ for 2004–2008, with the spatial distribution shown in Fig. 1. This estimate is much larger than the previous HBL emission estimates of Roulet et al. (1994) and Worthy et al. (2000).

3 Constraints on HBL methane emissions

We examine the consistency of the GEOS-Chem methane emissions with the aircraft data from Pre-HIPPO and ARCTAS, and the surface data from Fraserdale and Alert. The DACOM tunable diode laser instrument used in ARCTAS (Sachse et al., 1987) has an estimated accuracy/precision of 1%/0.1%. The quantum cascade laser instrument used in Pre-HIPPO has an estimated accuracy and precision of 0.25%. The surface measurements at Fraserdale are obtained by gas chromatography on samples collected from a 40-m high tower and have an accuracy/precision of 1%/0.2% (Worthy et al., 2003). Similar specifications apply to the surface measurements at Alert.

Figure 2 shows the ensemble of aircraft vertical profiles over the HBL from 12 May (Pre-HIPPO) to 5 July (ARCTAS). We excluded stratospheric air as diagnosed by a molar O$_3$/CO ratio exceeding 1.25 (Hudman et al., 2006) and fire plumes as diagnosed by CO exceeding 200 ppbv. The latter filter effectively removed biomass burning influence from the data set as inferred from correlation between methane and CO. The HBL methane enhancements in Fig. 2 can thus be reliably attributed to wetland emissions.

The ARCTAS observations on 4–5 July show strong boundary layer enhancements over the HBL. The Pre-HIPPO flight on 12 May shows no boundary layer enhancement while that on 23 June shows a moderate enhancement. Observers on the Pre-HIPPO aircraft reported snow cover over the HBL on 12 May but not on 23 June. For comparison with the aircraft we sample the model at the time and location of the flights. We see in Fig. 2 that the model provides a good simulation of the boundary layer structure for the different flights, the enhancement observed in ARCTAS, and the sharp springtime transition from May to July. However, model overestimation is evident for the 23 June profile.

To further investigate the magnitude and seasonal onset of HBL emissions we used 2004–2008 surface data at Fraserdale and Alert collected by Environment Canada, with Alert serving as an Arctic background site against which the HBL influence at the Fraserdale downwind site can be referenced (Worthy et al., 1998). For Fraserdale we sample the daily data averaged over the 1700–1900 local time window, when the surface measurements are most representative of a relatively deep mixed layer (Worthy et al., 1998), and further select for surface winds from the northern quadrants, when direct influence from the HBL can be expected (Fig. 1). Selection for northerly winds retains ~50% of the original data. We sample Alert data for the same times at Fraserdale in order to facilitate analysis of the difference between the two sites as discussed later.

Figure 3 shows the observed seasonal variations at Fraserdale and Alert for 2004–2008. The observations at Alert show a July minimum due to chemical loss in the Northern Hemisphere. The model minimum lags 4–6 weeks behind, an offset that can be attributed to background error in the seasonal variation of sources, transport, or OH concentrations. The observations at Fraserdale follow the seasonal variation at Alert in winter-spring but deviate in late May toward an August maximum, ostensibly due to emissions from the HBL. The model shows the same seasonal deviation at Fraserdale relative to Alert but shifted 6 weeks early. A model sensitivity simulation with no HBL emissions (also shown in Fig. 3) confirms that the seasonal deviation between Fraserdale and Alert is due primarily to HBL emissions. The model shows multiple seasonal peaks at Fraserdale (late June, late August, early November) compared to a single observed peak, but this fine structure reflects fluctuations in the background rather than HBL emissions as discussed below.

Although Fraserdale is at the southern end of the HBL and the data are collected at only 40 m altitude, they appear to be reasonably representative of the HBL. As a test, we partitioned the HBL into northern and southern halves and examined the model sensitivity of Fraserdale to emissions from each half. We found no significant difference. We also found no significant difference in methane concentrations when sampling the model at different altitudes over Fraserdale up to 500 m (five model layers).
Fig. 2. Methane vertical profiles from Pre-HIPPO and ARCTAS over the HBL (May–July 2008). Observations (blue) are compared to GEOS-Chem (GC) model vertical profiles sampled along the flight tracks at the flight times. The standard simulation (red) and a sensitivity simulation with no HBL emissions (black) are presented.

Fig. 3. Seasonal variation (2004–2008) of methane at Fraserdale and Alert. GEOS-Chem results are compared to observations. Also plotted is the model background concentration at Fraserdale as derived from a simulation with no HBL emissions. Data are daytime values smoothed with a 28-day moving average and then averaged over 5 yr. For Fraserdale we use only data associated with winds from the northern quadrants.

Figure 4 shows the seasonal variation of the difference in concentrations between Fraserdale and Alert ($\Delta$CH$_4$), illustrating more precisely the methane flux signature from the HBL. Here we assume that Alert provides a reasonable measure of background concentrations at Fraserdale; this is supported in the model by the comparison in Fig. 3 of the model simulation at Alert (thin blue line) and at Fraserdale in the absence of HBL emissions (thick black line). We find that the multi-peak structure of model concentrations at Fraserdale in June–November (Fig. 3) is reduced when corrected for the Alert background (Fig. 4). More importantly, we find that any residual multi-peaks in modeled $\Delta$CH$_4$ are associated more with changes in the model background at Fraserdale relative to that at Alert than in HBL model emissions. Temporal fluctuations during those two months in the model may also reflect the greater variability in surface temperatures (used in the model to compute methane emission) than in actual soil temperatures. Heat transfer in the soil column would be expected to dampen temporal variability in soil temperatures.

The observed onset of HBL methane emissions in late May, as seen in Fig. 4, is consistent with the aircraft
observations discussed above and with previous field studies in nearby James Bay peatlands that suggest an onset of emissions in mid-May (Pelletier et al., 2007). By contrast, we see from Fig. 4 that HBL emissions in the model begin in early April. In addition, the observations indicate a seasonal shutdown of HBL emissions in September whereas in the model these emissions persist into October. The early onset of model emissions was not apparent in comparison with the Pre-HIPPO profile on 12 May in Fig. 2, but that is because of delayed spring warming in 2008 and because that flight profile sampled the northern edge of the HBL (Fig. 1).

The premature onset of HBL methane emissions in the model likely reflects the use of skin temperature as proxy for soil temperature. Seasonal increases in soil temperature at depth lag behind the land surface during the spring thaw. We attempted to impose in the model a time lag for soil heating by using the standard heat transport parameterization of Campbell and Norman (1998) with thermal diffusivities from Sitch et al. (2003), but the resulting delay in the onset of emissions was insufficient. Instead we identified persistent snow cover in the GEOS-5 data well past the model onset in model emissions. Snow cover would insulate the underlying soil from warming, inhibiting methanogenesis in spring, and would also trap methane in the autumn (Friborg et al., 1997). Consequently we modified the model to restrict emissions to snow-free regions. Figure 4 shows that this mostly corrects the model biases in the spring and autumn, although there is still a small lag of 1-2 weeks in the spring. This additional delay might reflect a period of time required for the underlying peatlands to thaw before methanogenesis ensues (Dunn et al., 2009). Comparison with the aircraft profiles in Fig. 2 is unaffected by the delayed onset in model emissions. The resulting annual reduction in model HBL emissions is 20% (2.3 Tg a⁻¹ vs. 2.9 Tg a⁻¹).

The model temporal variability of methane emissions in the snow-free season is driven largely by surface temperature (Eq. 1), and this appears adequate to match the observed July–August maximum of HBL emissions (Fig. 4). Previous studies of boreal wetlands have pointed out the sensitivity of emissions to changes in the level of the water table (Moore et al., 1994; Pelletier et al., 2007). However, the flat topography of the HBL results in poor drainage and maintains persistent wetland coverage throughout the summer.

Figure 5 shows the interannual variation of HBL model emissions for 2004–2008 as driven by temperature and snow cover. The seasonal onset of emission can vary by a month from year to year. There is much less year-to-year variability in the fall shutdown of emissions. The mean annual emission for the 5 yr is 2.3 ± 0.3 Tg a⁻¹.

4 Comparison to ABLE-3B/NOWES estimates

The ABLE-3B/NOWES surface and aircraft field study in July 1990 previously reported an annual emission estimate of 0.5 ± 0.2 Tg a⁻¹ for the HBL (Harris et al., 1994; Roulet et al., 1994). This is considerably less than our best estimate of 2.3 Tg a⁻¹, and would be inconsistent with the Pre-HIPPO and ARCTAS data of Fig. 2 as well as the Fraserdale ΔCH₄ data of Fig. 4. The ABLE-3B/NOWES estimate was obtained by extrapolation of direct flux measurements at surface sites, using wetland coverage derived from satellite and aerial imagery. The surface sites and supporting aircraft eddy correlation flux measurements were located in two small study areas at the southern and northern edges of the HBL (Fig. 1). Roulet et al. (1994) reported mean June-October emission estimates for the southern and northern study areas of 3.4 g m⁻² a⁻¹ and 6.3 g m⁻² a⁻¹, respectively, from their surface measurements. Aircraft measurements over these same regions in July yielded consistent mean fluxes of 5 ± 3 g m⁻² a⁻¹ and 4 ± 6 g m⁻² a⁻¹ respectively. These values agree with our flux estimates of ∼5 g m⁻² a⁻¹ for both ABLE-3B/NOWES regions in July (Fig. 1). Roulet et al. (1994) went on to infer annual mean emissions by taking their June–October measurements to be representative of snow-free conditions and assuming zero emissions for snow-covered ground. This is consistent with our findings.

The large difference between our estimate of HBL emission estimate and that of Roulet et al. (1994) thus lies in the spatial extrapolation to the scale of the HBL. We see from Fig. 1 that our emissions are much higher in the mid-section of the HBL than in the ABLE-3B/NOWES study regions. The boundary layer methane enhancements observed from the ABLE-3B aircraft (∼30 ppbv) were indeed much lower than the mean value of 60 ppbv observed on the ARCTAS flights (Fig. 2).
5 Conclusions

Aircraft observations over the Hudson Bay Lowlands (HBL) in May–July 2008 show a seasonal onset of methane emissions in June and 60 ppbv enhancements in the boundary layer in July. Surface observations at Fraserdale (just south of the HBL), when referenced against a background Arctic site (Alert) to isolate the HBL contribution, indicate a seasonal onset of methane emission in late May, a peak emission from mid-July to the end of August, and a sharp decrease in September. The GEOS-Chem model including a standard methane emission scheme for boreal wetlands can successfully reproduce these observations except for a premature springtime onset and a delayed fall shut-off. Seasonal variation of wetland emission in the model is mainly driven by surface temperature. We find that accounting in addition for suppression of emission by snow cover corrects the model biases in spring and fall. The variability in the model is still larger than observed and this could reflect dampening of soil temperature fluctuations relative to the surface. Our resulting best estimate of HBL methane emissions is 2.3 Tg a⁻¹, much higher than previous estimates for the region (Roulet et al., 1994; Worthy et al., 2000). We argue that this reflects gradients of methane emission within the HBL that were not previously accounted for.

Acknowledgements. This work was supported by the US National Science Foundation and by the Tropospheric Chemistry Program of the National Aeronautics and Space Administration. We thank the reviewers for useful comments.

Edited by: A. Stohl

References


