NASA-CR-196085

÷

12700 2.1

FINAL REPORT

Methane Production and Consumption in Grassland and Boreal Ecosystems

NAGW-1828

Principal Investigators:

.

David S. Schimel Ingrid C. Burke Carol Johnston John Pastor

-

(NASA-CR-196085) METHANE N94-35486 PRODUCTION AND CONSUMPTION IN GRASSLAND AND BOREAL ECOSYSTEMS Final Report (Colorado State Unclas Univ.) 25 p

G3/45 0013764

· • . .

1.01

OVERVIEW

The objectives of the this project were to develop a mechanistic understanding of methane production and oxidation suitable for incorporation into spatially explicit models for spatial extrapolation. Field studies were undertaken in Minnesota, Canada and Colorado to explore the process controls over the two microbial mediated methane transformation in a range of environments. Field measurements were done in conjunction with ongoing studies in Canada (the Canadian Northern Wetlands Projects: NOWES) and in Colorado (The Shortgrass Steppe Long Term Ecological Research Project: LTER). One of the central hypotheses of the proposal was that methane production should be substrate limited, as well as being controlled by physical variables influencing microbial activity (temperature, oxidation status, pH). Laboratory studies of peats from Canada and Minnesota (Northern and Southern Boreal) were conducted with amendments of a methanogenic substrate, at multiple temperatures and at multiple pHs (the latter by titrating samples). The studies showed control by substrate, pH and temperature, in order in anaerobic samples. Field and laboratory manipulations of natural plant litter, rather than an acetogenic substrate, showed similarly large effects. The studies concluded that substrate is an important control over methanogenesis, that substrate availability in the field is closely coupled to the chemistry of the dominant vegetation influencing its decomposition rate, that most methane is produced from recent plant litter and that landscape changes in pH are an important control, highly correlated with vegetation.

The results are published in the Journal of Geophysical Research-Atmospheres (Valentine D., E.A. Holland and D.S. Schimel 1993. Ecosystem and physiological controls over methane production in northern wetlands). The paper is attached.

Field studies on methane oxidation in Colorado utilized landscape gradients and existing field experiments at the LTER. Studies took advantage of long-term measurements and were complementary to studies initiated in the early 1980s. Measurements were made of soil inorganic N, methane oxidation in soils, and the production of nitrous oxide. Key results were that: (1) land use change dramatically lowered methane oxidation, (2) rates of methane oxidation and nitrous oxide production were negatively correlated, (3) increasing soil N tended to reduce rates of methane oxidation and, (4) preliminary evidence suggested that methane oxidation was principally limited by the diffusion rate of methane into soils, and secondarily limited by soil nitrogen. The effects of land use change (grassland to cropland) may act through these or other mechanisms. As the principal zone of methane oxidation appears to be about 10-30 cm deep in soils, in contrast to most microbial processes which occur within the top 10 cm of soils, disruption of this layer by tillage is implicated.

Results from this study were published in Nature (Mosier et al. 1991. Methane and nitrous oxide fluxes in native, fertilized and cultivated grasslands. Nature 350:330-332), and Chemosphere (Ojima et al. 1993. Effect of land use change on methane oxidation in temperate forest and grassland soils. Chemosphere 26:675-685). These papers are likewise attached.

• • .

Ecosystem and physiological controls over methane production in northern wetlands

David W. Valentine

Natural Resource Ecology Laboratory, Colorado State University, Boulder, Colorado

Elisabeth A. Holland and David S. Schimel¹

National Center for Atmospheric Research, Boulder, Colorado

Peat chemistry appears to exert primary control over methane production rates in the Canadian Northern Wetlands Study (NOWES) area. We determined laboratory methane production rate potentials in anaerobic slurries of samples collected from a transect of sites through the NOWES study area. We related methane production rates to indicators of resistance to microbial decay (peat C:N and lignin:N ratios) and experimentally manipulated substrate availability for methanogenesis using ethanol (EtOH) and plant litter. We also determined responses of methane production to pH and temperature. Methane production potentials declined along the gradient of sites from high rates in the coastal fens to low rates in the interior bogs and were generally highest in surface layers. Strong relationships between CH, production potentials and peat chemistry suggested that methanogenesis was limited by fermentation rates. Methane production at ambient pH responded strongly to substrate additions in the circumneutral fens with narrow lignin:N and C:N ratios ($\partial CH_{i}/\partial EtOH = 0.9$ -2.3mg g⁻¹) and weakly in the acidic bogs with wide C:N and lignin:N ratios ($\partial CH_{1}/\partial EtOH = -0.04-0.02$ mg g^{-1}). Observed Q_{10} values ranged from 1.7 to 4.7 and generally increased with increasing substrate availability, suggesting that fermentation rates were limiting. Titration experiments generally demonstrated inhibition of methanogenesis by low pH. Our results suggest that the low rates of methane emission observed in interior bogs during NOWES likely resulted from pH and substrate quality limitation of the fermentation step in methane production and thus reflect intrinsically low methane production potentials. Low methane emission rates observed during NOWES will likely be observed in other northern wetland regions with similar vegetation chemistry.

INTRODUCTION

Uncertainty in the global methane budget arises from the scarcity of source measurements, especially integrated over large areas [Aselmann and Crutzen, 1989; Fung et al., 1991], from uncertainty about the atmospheric destruction rate [Vaghjiani and Ravishankara, 1991] and from new information suggesting a larger soil sink [Steudler et al., 1989; Mosier et al., 1991). Three-dimensional global analyses of atmospheric methane concentrations [Fung et al., 1991] and isotopic composition [Tyler, 1986, 1989, 1991; Quay et al., 1991] are the most powerful techniques for analyzing current sources and sinks, and provide hints as to controls over emissions. These analyses and other studies have suggested an important but highly uncertain methane source from northern wetlands. The Canadian Northern Wetlands Study (NOWES) and two American Atmospheric Boundary Layer Experiments (ABLE 3A and 3B) constrain bounds on that uncertainty.

In addition, no predictive or diagnostic models of methane biogeochemistry now exist for any ecosystem type. Process models of microbial transformations and relevant transport processes have not been widely used in methane biogeochemistry

Copyright 1994 by the American Geophysical Union.

Paper number 93JD00391. 0148-0227/94/93JD-00391\$05.00 but would complement global analyses by linking edaphic, biological, and climatic controls to flux rates and provide a mechanistic basis both for spatial analysis and for projection of future changes. Process models, when coupled with data describing paleowetland extent, also provide a basis for understanding past changes in methane fluxes.

Methane production is carried out by methanogenic bacteria, which are obligate anaerobes capable of using a limited number of relatively simple substrates supplied by fermentation processes. Several reviews of the mechanisms of methanogenesis recently have been compiled [Oremland, 1988; Conrad, 1989; Knowles, Conrad [1989] enumerated the controls over 1993]. methanogenesis as (1) substrate availability, (2) temperature, (3) pH, and (4) competitive electron acceptors. He argued that population size of methanogens was not likely a major control over methane production rate. While the controls over methane production at the microbial level are reasonably well known, the relative quantitative importance of the different factors is not well known, especially under field conditions. In this paper we analyze the controls over methane production from a series of sites in northern wetland ecosystems and present results from experimental manipulations of several factors. The sites chosen contrasted in plant species composition, productivity, hydrology, and pH [Klinger et al., this issue].

Controls over methane production fall into two classes that must be considered together in predictive models of methane production: (1) controls over substrate availability (i.e., fermentation rate as affected by temperature, litter input, and litter decomposition rate); and (2) physiological controls over methanogenic activity (pH and redox potential) from a given supply rate of substrate.

¹Also at Natural Resource Ecology Laboratory, Colorado State University, Fort Collins.

The principal substrates for methanogenesis are CO_2+H_2 and acetate [Oremland, 1988; Conrad, 1989; Knowles, 1993). Substrate abundance is constrained by the slowest link in a chain of events starting with inputs of detrital organic matter and including several decomposition and fermentation steps resulting in methanogenic substrates. While many studies identify the importance of substrate limitation as a control over methane production [Aselmann and Crutzen, 1989; Conrad, 1989], few have systematically analyzed its importance [Yavitt and Lang, 1990]. If most methane is produced largely from recent organic matter, as ¹⁴C results suggest [Jeris and McCarty, 1965; Wahlen et al., 1987], then methane production should ultimately be closely coupled to terrestrial primary production and decomposition rates. NOWES provides a foundation for modeling the linkage of methane production to ecosystem change (through changing plant production and chemistry) as well as to climate change.

Temperature directly affects methanogenesis rates as well as decomposition and fermentation rates, but Westermann et al. [1989] reported that the effect of increasing temperature on the rate of methane production is largely offset by a corresponding increase in K_m (decreased substrate affinity) of methanogenic enzyme systems. Thus the observed effect of temperature will likely be manifested only through increased availability of methanogenic substrate as long as readily decomposable reduced carbon compounds are available. These interactions are important because they could lead to unexpected results, e.g., temperature increases could lead to increased instantaneous rates of methanogenesis via enhanced fermentation, but these higher rates could outstrip supply rates of readily fermentable compounds and ultimately lead to reduced methane production rates.

Appropriate redox potentials for methanogenesis require saturated or nearly saturated soil conditions and the absence of competing, more electronegative electron acceptors (NO₃⁻, SO₄²⁻, Fe³⁺). Water table fluctuations influence not only the thickness of the aerobic zone in which CH₄ oxidation can occur but also the portion of the soil or peat profile in which methanogenesis is possible. Conversely, redox potential also may limit substrate production because very low redox potentials tend to retard decomposition. Since organic matter chemistry varies with depth in most profiles, water table depth will influence both the quantity of organic matter for methanogens and the maximum rate at which that organic matter may be processed. We considered this interaction only to the extent that we surveyed methane production potentials by depth and amended samples from the upper two peat depths.

In the field, methane emission is influenced by production but is modulated by transmission to the atmosphere which occurs via diffusion, solution transport, bubbles or vascular transport in plants [Keller et al., 1986; Conrad, 1989; Chanton and Dacey, 1991; Fechner and Hemond, 1992]. Transport, oxidation and water table depth were considered by other investigators in NOWES [Klinger et al., this issue; Roulet et al., this issue].

Detailed process studies of the sort described in this paper are an important complement to field experiments not only because they aid in model development but also because they aid in interpreting field observations. Methane flux rates observed in NOWES were much lower than expected [Roulet et al., this issue] and our process studies help explain why. Our studies help establish whether fluxes were low because of intrinsically low rates of methane production (determined from maximal laboratory methane production rates). This will be of importance in extrapolating NOWES results to other regions.

METHODS

Study Site

The four study sites were located in the Hudson Bay lowlands along a 95-km transect southwest of Moosonee, Ontario, Canada (Figure 1). The region has been gradually exposed by the receding Hudson Bay during isostatic rebound since the last glacial maximum [Webber et al., 1970], giving an age sequence along the transect extending back approximately 4000 years [Klinger et al., this issue]. From youngest (closest to James Bay) to oldest the four sites studied were the coastal fen, interior fen, Carling Lake, and Kinosheo Lake. The older two sites were ombrotrophic Sphagnum bogs while the younger two were sedgedominated minerotrophic fens.

The vegetation and peat characteristics of the sites are described in detail by *Klinger et al.* [this issue] so we present only a summary here (Table 1). The relative covers of sedges, non-Sphagnum bryophytes, deciduous shrubs, and pteridophytes are highest in the interior fen, decreasing slightly in the coastal fen and are very low or absent in both of the bog sites. Conversely, the relative covers of sphagnum, ericaceous shrubs, and lichens are much higher in the bog sites (especially Kinosheo Lake) than in the fen sites. The covers of forbs and coniferous shrubs and trees were fairly constant across sites. Peat depth increased approximately monotonically from about 0.7 m in the coastal fen site to about 2.9 m at Kinosheo Lake.

Procedures

Intact cores were taken during July 1990 using a 10.2-cm^2 box corer from each of the four sites. Four cores were taken from the coastal fen site, three from the interior fen, and two each from Carling Lake and Kinosheo Lake. In addition, we sampled the top 1 m of peat from the side of a peat pit in 0.1-m increments at both bog sites. All samples were vacuum sealed in 0.13-mm-thick



Fig. 1. Location of study sites. The two end-point sites; coastal fen and Kinosheo Lake were used for detailed methane process studies.

Site	Latitude and Longitude	Distance From Coast, (km)	Water Table Depth,* (m)	Soil water pH*	Peat Depth,* (m)	Basal Peat Age,* (y)
Coastal fen	51°28.2' N	18	0.14	6.86	0.71	1090
Interior fen	80"37' W 51"30 7' N	33	0.35	6.48	1.03	1960
Interior fell	80°52.8' W	55	0.55	0.40	1.05	1700
Carling Lake	51°29.8' N	99	0.17	4.11	2.48	4110
	81°45.7' W					
Kinosheo Lake	51°33' N	113	0.26	4.15	2.56	4110
	81°49.5' W					

TABLE 1. Summary of Site Characteristics for Sites Used in Regional Survey of Methane Production Potentials

*From Klinger et al. [this issue].

polyethylene bags, then kept at 4°C until analysis. The peat horizons were classified using the von Post scale of peat humification [*Bélanger et al.*, 1988] (Table 2). Moisture content was determined gravimetrically by drying to constant weight at 70°C. Total carbon and nitrogen concentrations were determined on dried and ground samples at 10 cm increments on a Carlo Erba NA 1500 CN analyzer. Lignin and cellulose contents were determined at 10-cm intervals using the Goering-Van Soest procedure [Goering and Van Soest, 1970; Kirk and Obst, 1988].

Methane production potentials in slurried incubations were assessed for all cores at 10-cm intervals at 20°C. Incubations were performed in slurries of 20 g (fresh weight) subsamples of wellmixed peat. The peat and enough deionized water (to 100 mL) were added to a 250-mL Erlenmeyer flask under a nitrogen atmosphere. The flasks were stoppered with silicone stoppers fitted with a stopcock, septum, and in a subset of replicates, redox (platinum) and pH electrodes. These allowed monitoring of redox potentials to ensure that conditions sufficiently reduced for methanogenesis prevailed in the flasks. Slurries initially were sparged with ultrahigh pure (UHP) N₂ for 2 minutes and then continuously shaken on an orbital shaker in refrigerated incubators throughout the incubation period. Samples were subsequently allowed to preincubate for 2 weeks before measurement to allow exhaustion of initially high volatile fatty acid levels, after which incubations lasted 4-6 weeks. During the incubation period, redox potentials remained very low (-300 mV) and pH generally changed less than 0.2 units.

Flask headspaces were sampled weekly during the incubation period. Before sampling, each flask was sparged with UHP He by vigorously bubbling the slurry for 2 minutes. This procedure reduces but does not eliminate degassing problems during the assay period [*Kiene and Capone*, 1985], minimizes problems of pressure variation among flasks due to CO_2 and CH_4 accumulation, establishes similar bicarbonate equilibria among sampling periods, and lessens concentration gradients driving diffusive losses across the silicone stoppers during the assay period. The assay period following sparging varied between 0.25 and 2 hours, depending on rates of CH_4 production. At the end of the assay period we used 35-mL polypropylene syringes fitted with nylon stopcocks to sample 12 mL of the headspace. If two samples were required, as when CO_2 production rates were more rapid than CH_4 rates, then 12 mL of UHP N₂ was added as a makeup gas following the first sampling.

Gas samples were analyzed within 3 hours following sampling on a Shimadzu GC-8 gas chromatograph fitted with a flame ionization detector (FID) for CH_4 and a Carle Instruments thermal conductivity detector (TCD) for CO_2 . We used a 3.175-mm OD Ni column packed with HayeSep D in an 60°C oven with a carrier gas (He) flow rate of 35 cm³ min⁻¹. The FID was kept at 200°C and the TCD at 60°C. Concentrations were corrected for trace quantities of CH_4 (generally 0.2 parts per million by volume (ppmv)) contained in the UHP N₂ used for the headspace.

During the incubation period, production potentials generally increased for the first several weeks, after which they declined modestly in most slurries. No initial rapid methane production rates were observed, suggesting that disturbance-enhanced concentrations of volatile fatty acids were not reflected in our data.

We also assessed process controls over CH_4 production in the top two von Post horizons of Coastal fen and Kinosheo Lake peats. Temperature effects were analyzed on slurries at 10°C and 20°C. Degree of substrate limitation was assessed by adding 1.64 or 8.22 mmol ethanol (39.4 mg or 197.3 mg C equivalent, respectively) as an acetate and hydrogen source [Bryant et al. 1967; Large, 1983; Svensson, 1984]. Although ethanol cannot be used directly by methanogens, it is easily converted to hydrogen and acetate by other organisms [Bryant et al., 1967; Svensson, 1984]. For our experiments, ethanol also presented two advantages over acetate: first, protonated acetic acid is toxic to microbial populations, implying that only very small amounts of acetate could be added at low pH; second, ethanol would provide a prolonged supply of substrate during the incubation. Because

TABLE 2. Depths and Von Post Indices (VPI) for Peats Used in Amended Incubations

	Coastal	Fen	Kinos	heo Lake
Peat Horizon	Von Post Index	Depth, cm	Von Post Index	Depth, cm
Surface	2-3	0-10.5	1	12
Subsurface	3-5	10.5-19	2-3	12-15.5

Peats were sampled by Von Post Index, and the top two uniform VPI horizons were used in experimental studies. Full descriptions of peat profiles are in the work of *Klinger* et al. [this issue]. Chemical characteristics of peat profiles are shown in Table 3.

acetic acid is a weak acid, moreover, a significant fraction of added acetate salts would protonate and raise pH in the low pH slurries. Substrate limitation to fermenting populations was also assessed by adding 0.5 g of dried and ground plant litter samples collected on site. Effects of pH on production potentials were assessed by lowering the pH of coastal fen slurries to ~5.5 or ~4 with H₃PO₄ and raising the pH of Kinosheo Lake slurries to ~5.5 or ~7 with KOH. We also tested the response of Kinosheo Lake slurries to direct H₂ enhancements. Additions of 10 mL of H₂ brought headspace partial pressures to 0.05 atm., but no CH₄ production response occurred (12.1 versus 19.2 and 13.4 versus 8.3 µg/g for surface and subsurface layers, respectively). Because of the lack of response we dropped H₂ treatments thereafter in favor of ethanol amendments.

Statistical significance of experimental results were determined using analysis of variance.

RESULTS AND DISCUSSION

Landscape Patterns in CH4 and CO2 Production

Methane production potentials were highest in Coastal fen slurries and dropped rapidly with distance inland to the bogs at Carling Lake and Kinosheo Lake (p<0.01, Figure 2). This trend corresponded to patterns of soil solution pH (highest at the coast), NPP (highest at the coast), and plant community composition (sedges and bryophytes in the coastal and interior fen sites, Sphagnum moss, and ericaceous shrubs in the inland bog sites)



Fig. 2. (a) Methane and (b) CO_2 production potentials under unamended conditions for all sites along the regional transect and as a function of depth. Error bars are standard errors of the mean (n=4 for CF, 2 for IF, 3 for CL, 3 for KL).

[Klinger et al., this issue]. The trend also agreed with in situ measurements of CH_4 emissions, except for anomalously low emissions from the interior fen [Klinger et al., this issue] compared to the production potentials (Figure 2). This discrepancy probably resulted from CH_4 oxidation due to the low water table and deep aerobic zone (Table 1).

In the coastal and interior fens, methane production potentials were highest in the surface peats and declined with depth to a plateau below 0.2 m (p<0.0.05, Figure 2). The depth pattern was less clear in the bog sites; low CH₄ production potentials occurred at all depths. Surface peats from the bog sites, Carling Lake, and Kinosheo Lake had slightly higher production potentials, however, than did intermediate depths. Similar depth patterns of CH₄ production have also been observed in spruce-peatlands in west central Alberta [*Hogg et al.*, 1992].

Rates of CO_2 production exhibited the same spatial pattern as methane, supporting the argument that methane production is linked to carbon substrate available for decomposition (see below) (Figure 2b). Decreases in surface peat CO_2 production potentials from coastal to inland sites were less marked than for methane, though significant. Subsurface peats had similar and low rates along the entire transect, as with methane. More methane was produced per unit CO_2 produced in the coastal than in the inland sites (p<0.005), perhaps because of more favorable physiological conditions for methanogen activity and population growth in the neutral coastal peats.

Process Controls

Organic matter quantity and quality. Methane production potentials along the transect and with depth corresponded to changes in peat chemistry and its influence on decomposition and fermentation rates. Rates of methanogenesis depend on rates of substrate supply from fermentation, and rates of fermentation depend on the amount and decomposability of organic matter [Conrad 1989; Sass et al., 1990, 1991; Svensson, 1976, 1980; Svensson and Rosswall, 1980; Taylor et al., 1991]. Decomposition of peat is slow under unperturbed conditions, and substrate for heterotrophic processes is usually limiting [Svensson, 1980]. Several investigators have found that methane production or emission was highest at sites where NPP, plant density, or biomass was highest [Klinger et al., this issue; Morrissey and Livingston, 1992; Svensson, 1976, 1980; Whiting and Chanton, 1992]. The role of NPP is clearly through the annual supply of relatively labile fractions of plant debris, though vascular transport plays a role at some sites as well.

Striker.

The decomposability of the litter produced influences rates of decomposition, fermentation, and methanogensis. The chemical availability of carbon for decomposition is usually quantified by two indices of decomposability: C:N and lignin:N ratios of the material to be decomposed [Parnas, 1975, Melillo et al., 1982; Bosatta and Berendse 1984; Agren and Bosatta, 1987; Berendse et al., 1987]. The indices have been used frequently to describe the substrate quality of litter. We use them here to describe the decomposability of peat, which is partly decomposed plant litter. We also measured cellulose content of peats, as cellulose is a relatively available C fraction, routinely measured as part of the lignin determination. Three results from this study emphasize the importance of substrate supply in controlling spatial patterns of methanogenesis: (1) CH₄ production potentials uniformly increased with substrate addition, including both EtOH and litter amendments (Figure 3); (2) CH₄ production potentials increased with increasing substrate quality (decreasing lignin:N ratios and increasing cellulose) (Table 3); (3) CH₄ production potentials



Fig. 3. The response of methane production to added EtOH and pH at two depths (see Table 2 for horizon depths and descriptions) are shown for (a) neutral coastal fen peat (n=3), (b) acid Kinosheo Lake peat (n=3), and (c) Kinosheo Lake degradational feature ("black hole") peat (low sample availability prohibited replication). "NA" indicates that a given treatment combination was not carried out, due to insufficient remaining sample. Error bars are standard errors of the mean.

increased with increasing NPP of the peat collection site (Figure 1 and *Klinger et al.* [this issue]. Both the C:N and the lignin:N ratios of surface peat increase and, presumably, substrate decomposability declines, with distance inland (Table 3). Strongly decreasing N contents inland raised the C:N ratio at Kinosheo Lake to nearly triple that of the coastal fen. Lignin:N ratios only doubled along the same transect because of partially offsetting decreases in lignin content inland. The C:N and lignin:N ratios suggest that carbon is more available for fermentation and subsequent methanogenesis at the coastal sites than at the inland sites. Higher CH₄ and CO₂ production potentials for the coastal compared to the inland sites support this.

The highest CH_4 and CO_2 production potentials always occurred in surface peats (p<0.05) and declined with depth. The low but nonzero CH_4 production potentials at lower depths in all sites indicate that these layers can serve as important, long term contributors to CH_4 stored in pore water and episodically released during water table fluctuations [*Roulet et al.*, this issue].

The surface peat layer receives all of the aboveground litter inputs as well as most of the inputs of fresh organic matter from root turnover and most closely reflects the chemistry of the existing vegetation. Changes in peat chemistry with depth result from historic microbial transformations and the chemistry of the vegetation from which the peat formed. As indicated by the

3LE 3. Chemical Characteristics of Peats From Each of the Sites Along the Chronosequence Sampled During the NOWES Experiment B h,	Carbon Lignin Cellulose Nitrogen C:N C:N Lignin:N Lignin:N pH g c	46.2 (0.6) 33.2 (1.9) 23.5 (2.1) 1.9 (0.1) 25.1 (1.5) 17.8 (0.7) 6.56 0.0 10.7 (0.1) 2.3 10.1 (1.1) 21. (0.1) 24.0 (0.5) 10.7 (0.0) 5.00 0.0	46.6 (1.8) 46.0 (6.4) 15.6 (2.1) 2.3 (0.1) 24.0 (0.0) 19.7 (0.9) 5.80 0.1 46.9 (1.8) 46.0 (6.4) 15.6 (2.1) 2.3 (0.3) 20.6 (0.8) 20.1 (1.0) 0.1 (1.1) (1.1) (1.	45.8 (1.1) 35.2 (2.9) 23.8 (2.7) 1.8 (0.3) 26.3 (3.6) 19.4 (4.9) 5.68 0.0	45.5 (1.5) 40.4 (5.1) 18.2 (0.8) 1.9 (0.4) 20.6 (0.7) 17.5 (2.9) 5.59 0.07	45.8 (1.4) 38.5 (0.5) 19.0 (2.6) 2.3 (0.2) 21.2 (0.2) 7.1 (0.5) 2.8 (0.1)	47.6 (0.8) 28.0 (1.8) 28.7 (0.4) 0.8 (0.2) 49.8 (3.5) 26.0 (0.4) 4.29 0.04	1 47.9 (1.2) 29.4 (1.1) 32.8 (0.6) 0.9 (0.1) 59.3 (9.1) 28.4 (0.7) 4.27 0.04		0.05 (0.5) 34.6 (0.5) 32.3 (1.3) 0.9 (0.1) 29.6 (2.9) 40.5 (0.5) 34.6 (0.5) 32.3 (0.5)	0.00 (0.5) 34.6 (0.5) 32.3 (1.3) 0.9 (0.1) 29.0 (3.9) 40.2 (3.0) 0.08 46.5 (0.5) 23.5 (2.2) 35.3 (5.7) 0.8 (0.1) 63.2 (13.5) 31.4 (6.2) 4.14 0.12	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
Percent	Lignin Cellulose	33.2 (1.9) 23.5 (2.1)	42.3 (1.8) 14.0 (1.4) 46.0 (6.4) 15.6 (2.1)	35.2 (2.9) 23.8 (2.7)	40.4 (5.1) 18.2 (0.8)	38.5 (0.5) 19.0 (2.6)	28.0 (1.8) 28.7 (0.4)	29.4 (1.1) 32.8 (0.6)	34.6 (05) 323 (13)		23.5 (2.2) 35.3 (5.7)	23.5 (2.2) 35.3 (5.7) 19.5 (1.0) 38.5 (3.8)
TABLE 3. Chemical Cha Depth,	e cm Carbon	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	10-20 48.7 (0.4) 20-80 46.9 (1.8)	r fen 0-10 45.8 (1.1)	10-20 45.5 (1.5)	20-80 45.8 (1.4)	g Lake 0-10 47.6 (0.8)	10-20 47.9 (1.2)	20-80 50.5 (0.5)		heo Lake 0-10 46.5 (0.5)	neo Lake 0-10 46.5 (0.5) 10-20 48.0 (0.4)

pattern of CH_4 and CO_2 production rates, decomposability of organic matter also decreases with depth. In the bog sites, production potentials decreased with depth, paralleling strong increases in C:N and lignin:N ratios with depth that may have served to restrict decomposition rates. Neither the C:N nor the lignin:N ratio reflected the depth trend in the fen sites, however. A similar lack of pattern in different sites led *Hogg et al.* [1992] to suggest that other factors, such as higher P, K, and carbohydrate contents in recently dead organic matter, controlled decomposability within the peat column. The positive correlation between CH_4 production potentials and cellulose content in the fen sites seem to support this idea.

This overall landscape pattern suggests that two different sets of controls over decomposition and methane production may dominate at the two site types. The recalcitrant chemistry of organic matter, low pH, and likely low nutrient levels severely restrict microbial activities in the bog sites, resulting in the very low overall CH_4 and CO_2 production potentials. The more favorable peat chemistry, high pH, and likely higher nutrient levels at the fen sites remove this physiological restriction on microbial activity, allowing the rapid decomposition and methanogenesis rates to be limited only by the supply of substrate.

Fermentation controls the rate of substrate supply, which in turn limits methanogenic rates. Our experimental ethanol additions indicated that substrate availability limited methane production rates under certain conditions. In coastal fen surface peat slurries, methane production rates increased (Figure 3a) and carbon dioxide production rates decreased in the presence of ethanol, although neither response was significant (p>0.1). However, the proportion of respired carbon evolving as methane (i.e., CH_4/CO_2) increased strongly and significantly (p<0.005) with added ethanol. Little or no response occurred from subsurface samples (Figure 3a). The same qualitative pattern occurred (although at much lower methane production rates) in most Kinosheo Lake slurries: an 80% increase in methane production from the surface peats, with no response from the subsurface samples (Figure 3b).

Methane production increased eightfold from ethanol-treated Kinosheo Lake "black hole" slurries (p<0.05; Figure 3c). This responsiveness to substrate amendments suggests that in the field, methanogenic processes might outstrip fermentation rates. In the black holes, fermentation and methanogensis may occur at different depths. The dark color, low albedo, and lack of vegetation cover all contribute to unusually warm surface temperatures of these features [Klinger et al., this issue], which accelerate decomposition. The close proximity of the water table to the surface in these depressions minimizes the distance substrate must descend to reach methanogenic populations and also minimizes the oxidizing zone through which CH₄ must ascend to reach the atmosphere. Downward transport of labile organic matter can explain how these peats could serve as such strong CH₄ sources in the field, despite low unamended methane production potentials in lab incubations.

Temperature. Many studies have demonstrated that temperature can be an important control over methane production, yet correlations between methane flux and temperature are typically weak [Williams and Crawford, 1985; Crill et al., 1988; Conrad, 1989; Moore et al., 1990; Moore and Knowles, 1990; Whalen and Reeburgh, 1990; Klinger et al., this issue]. We found that temperature had a greater effect on methane production potentials in sites where substrate quality was higher (Figure 4). Values of Q_{10} (the ratio of methane production at T/methane production at T-10) for the coastal fen were 4.7 and 1.7 for surface and subsurface peats, respectively (Figure 4a). At Kinosheo lake,



1

Startin.

Fig. 4. Response of methane production at two depths (see Table 2 for horizon depths and descriptions) to (a) temperature in the coastal fen peats and (b) temperature and substrate (as EtOH) in Kinosheo Lake peats.

apparent Q_{10} values were 2 and -2.1 for surface and subsurface peats. The apparent decline in methane production potentials from Kinosheo Lake subsurface peats was not significant (p>0.1) but, if real, may have resulted from rapid depletion of an initially small pool of available carbon when microbial activity was enhanced by the warmer temperature. An additional experiment examined interactions between substrate amendment and temperature in Kinosheo Lake peats. Apparent Q_{10} values increased slightly from 2 to 3 in surface peat slurries, and from -2.1 to 1.4 in subsurface peat slurries--when 39.4 mg of carbon (as ethanol) was added (Figure 4b), although the CH₄ production response to both the temperature change and the ethanol were not significant (p>0.1).

Similar patterns have been observed in pure cultures of *Methanosarcina barkeri*, in which *Westermann et al.* [1989] found that temperature had little net effect on the conversion of acetate to methane. An offsetting increase in K_m apparently cancels out nearly all of the increased methanogenic enzyme activity with increasing temperature. Following this argument, apparent temperature responses of methane production are instead responses to increased substrate availability from temperature-enhanced rates of fermentation.

Responses to pH. The response of CH_4 production potentials to pH changes is important to understanding spatial variations in CH_4 production because wetlands tend to become more acidic as their species composition and hydrology change through vegetation succession over time. Site differences in successional status are a major source of landscape variability in wetlands [Klinger et al., 1990; Klinger et al., this issue]. Methanogenesis tends to be most rapid in the circumneutral pH range [Conrad, 1989; Conrad and Schütz, 1988], although acid tolerant [Williams and Crawford, 1985; Patel et al., 1990] and moderately acidophilic [Maestrojuán and Boone, 1991] strains of methanogenic bacteria have been isolated. Conrad [1989] suggested that pH limitation may also be important in the fermentation steps supplying substrate for methanogenesis.

Our pH manipulation experiments generally suggested that neutrophilic methanogen populations dominate in northern wetland peats. Methane production generally increased with increasing pH, even in acidic peats. The pH of the peats ranged from 6.5-7 in the fen sites down to almost 4 in the bog sites (Tables 1 and 3), correlating well with CH_4 production potentials. Lowering the pH of coastal fen surface peat slurries from 7 to circa 5.5 cut CH₄ production potentials almost sevenfold (p<0.05) from 1376 to 208 μ g g⁻¹ week⁻¹, and further lowering pH to 4 cut CH₄ production potentials to $5 \mu g g^{-1}$ week⁻¹ (Figure 3*a*). Similar patterns also occurred with subsurface peat slurries. Raising the pH of Kinosheo Lake surface peat slurries from 4 to nearly 7 increased (p<0.05) CH₄ production potentials 20-fold from 0.5 to 11 µg g⁻¹ week⁻¹, while a similar pH change in subsurface peat slurries increased CH₄ potentials from 1.3 to 3.2 μ g g⁻¹ week⁻¹ (Figure 3b). Although Kinosheo Lake slurries produced more methane in response to both increased pH and ethanol additions individually, the combined treatment typically produced less CH₄ than the ethanol treatment alone (Figure 3b). The weaker landscape gradient in CO₂ compared to CH₄ production suggests an additive effect of pH above the substrate availability constraint.

The exception to the overall neutrophilic methanogenesis pattern occurred with peats sampled from dark-colored degradational features ("black holes") at Kinosheo Lake. These black holes were strong local methane sources in situ [Klinger et al., this issue; Roulet et al., this issue], but untreated slurries of these peats produced little more CH_4 than other Kinosheo Lake peats. Methane production from black hole slurries not only failed to increase with increased pH but actually decreased slightly (p<0.05, Figure 3c). Similarly, the strong ethanol response observed in these slurries was almost completely offset when pH was increased (interaction significant at p<0.05). This pattern suggests the presence of moderately acidophilic methanogens but also may have resulted from toxic effects of ethanol or the neutralizing titrant (KOH).

CONCLUSIONS

We hypothesized that rates of methane production should be constrained by substrate availability and regulated by temperature, redox potential, and pH. Results from sites other than the Kinosheo "black holes" showed consistent patterns of response to substrate and pH, but quantitative responses varied with initial peat chemistry.

The positive response of methane production with respect to substrate amendment addition at ambient pH ($\partial CH_4/\partial EtOH$; Table 4) suggests that substrate availability plays a key role in regulating methane production. Sites with more decomposable peat due to narrower lignin: N ratios (coastal fen) could more readily respond to substrate additions, perhaps because of differences in population size or maximal growth rate.

Methane production with respect to pH at ambient substrate $(\partial CH_4/\partial pH)$ was typically strong and positive (Table 4). The $\partial CH_4/\partial pH$ at the acidic Kinosheo Lake site was much smaller than at the circumneutral coastal fen site. Similarly, $\partial CH_4/\partial T$ was larger in the coastal fen than in the inland sites (Table 4). Thus the ability of the substrate-limited site to respond to an improvement in the physiological environment (pH or temperature) for methanogenesis was limited by the low rate of substrate supply.

Litter chemistry and pH play important roles in controlling spatial and temporal variations in methane production rates. Spatially, peat chemistry reflects the chemistry of the peatland plant community, with changes in lignin:N ratio reflecting the shift in vegetation from the high N graminoids of the coastal fen community to the low N sphagnum species of the interior bogs. With increasingly acidic and nutrient deficient conditions along that successional gradient, species are favored which are acid tolerant (e.g., mosses) and which have high nutrient use efficiency, though low intrinsic growth rates [Klinger, 1990; Chapin et al., 1987; Van Cleve and Vierek, 1981; Van Cleve et al., 1991]. This results in a positive feedback, further reducing nutrient availability through immobilization into the slowly decomposing peat. As noted above, these changes in peat chemistry (lignin:N) and pH have coordinated and interactive effects on methane production, linking methane production to succession through changes in plant chemistry.

Substrate limitation of methanogenesis in northern wetland ecosystems may also contribute to seasonal variations in methane emissions. It is likely that much methane production is produced from recently produced carbon (Figure 5) [Klinger et al., this issue]. The availability of fresh litter will vary seasonally with timing of NPP and senescence and with decomposition rates. Thus seasonal variations in methane production and emission rates

Factors Constant											
Site	Depth	Lignin:N	∂CH₄/∂EtOH	Standard Error							
Coastal fen	1	17.8	47,985	27,173							
Coastal fen	2	19.7	25,399	8855							
Kinosheo Lake	1	31.4	32.6	54							
Kinosheo Lake	2	23	54.9	8.9							
	Depth	Initial pH	∂СН₄/∂ <i>р</i> Н	Standard Error							
Coastal fen	1	6.56	941	960							
Coastal fen	2	5.8	29.2	7.3							
Kinosheo Lake	1	4.14	3.85	2.31							
Kinosheo Lake	2	3.97	0.7	0.36							
	Depth	Lignin:N	∂CH₄/∂T°	Standard Error							
Coastal fen	1	17.8	2.2	1.34							
Coastal fen	2	19.7	0.89	0.44							
Kinosheo Lake	1	31.4	0.05	0.06							
Kinosheo Lake	2	23	0.11	0.24							

TABLE 4. The Response of Methane Emission to Manipulation of Substrate, pH, or Temperature as a Function of Initial Lignin or pH While Holding Other

Units are μ g CH₄ (g peat)⁻¹ per g EtOH-C, °C, or *p*H unit as appropriate.

1



Fig. 5. Processes controlling methane production for the low N, low pH Kinosheo Lake, and the high N, neutral coastal fen site. The pathway of methane production is shown with experimental estimates of control functions adjacent. Solid lines indicate the flow of carbon through the methane pathway. Dashed lines indicate the point of influence of a control function. Asterisk indicates treatments not included because of insufficient sample.

will reflect not only microclimatic controls over methanogens and fermenters (temperature and water table fluctuations) but also variations in substrate supply. These variations, in addition to changes in porewater methane content and methane oxidation rates, lead to the common failure of simple correlations of methane emission rates with temperature, water table, and other microclimatic indicators [Crill et al., 1988; Roulet et al., 1992].

Our study has established quantitative relationships between ecosystem and physiological factors and methane production rates in controlled laboratory conditions. These results form the basis for our overall model of methane production (Figure 5). Temperature, organic matter quality, and supply rate act to control fermentation and hence substrate supply rates to methanogens. Substrate supply rate and pH, in turn, control methanogenic rates.

Thus we view methane production as an ecosystem process coupled to plant species composition, primary production, and overall heterotrophic metabolism. Spatial patterns in methane production also reflect plant physiological responses to successional and resource gradients that, in addition to climate and hydrology, influence the decomposition of peat and litter. The observed interacting effects of substrate supply rate and temperature on methane production rates will require validation through field studies in order to confirm their importance in controlling methane emission rates. The importance of other factors not examined here, such as competitive electron acceptors (e.g., SO_4^{2-}), nutrient availability, water table fluctuations, methane transport, and methane oxidation rates, will also need to be included in future process level methane emission studies.

Finally, our results may provide a partial explanation for the low rates of methane emissions from interior bogs observed during the NOWES study [Klinger et al., this issue; Roulet et al., this issue]. Methane production rates in these sites are severely restricted at the fermentation step by poor substrate quality and at the methanogenic step by very low pH. This suggests that similar low emission rates will also be observed in other northern wetland regions with similar vegetation floristics and chemistry.

Acknowledgments. We acknowledge the assistance of our colleagues Arvin Mosier, Nigel Roulet, John Pastor, Indy Burke, Lee Klinger, J'm Greenberg, and Pat Zimmerman for assistance with the design. execution, and interpretation of this study. By Brown provided assistance during field sample collection, and Rod Hansen, Scott Feeley, Cory Cleveland, Jason Neff, and Becky Riggle provided expert laboratory work. The comments of Mike Keller, Patrick Crill, and two anonymous reviewers helped improve this manuscript. The senior author was supported by a distinguished postdoctoral fellowship in global change awarded by the US Department of Energy and administered through Oak Ridge Associated Universities. This study was conceived when the junior author was a National Research Council senior fellow at NASA Ames Research Center and the support of the Ecosystem Science and Technology Branch is acknowledged with gratitude. This research was supported by NASA (NAGW 1828 and NAGW 2662), the National Center for Atmospheric Research, and by the Canadian Northern Wetlands Study, which provided superb logistical support. The National Center for Atmospheric Research is sponsored by the National Science Foundation. This is Scientific Contribution number 92-19 of the Canadian Institute for Research in Atmospheric Chemistry (CIRAC). The work described herein was undertaken as part of the CIRAC Northern Wetlands Study which has been generously supported by the Natural Sciences and Engineering Council of Canada. Although the research described in this article has been carried out under the auspices of CIRAC, it reflects only the views of the authors and does not necessarily reflect the official views of CIRAC.

REFERENCES

- Ågren, G. I., and E. Bosatta, Theoretical analysis of the long-term dynamics of carbon and nitrogen in soils, *Ecology*, 68, 1181-1189, 1987.
- Aselmann, I., and P. J. Crutzen, Global distribution of natural freshwater wetlands and rice paddies, their net primary productivity, seasonality and possible methane emissions, J. Atmos. Chem., 8, 307-358, 1989.
- Bélanger, A., D. Polvin, R. Cloutier, M. Caron, and G. Thériault, Peat-A Resource of the Future, Centre de Recherche Premier, Riviere-du-Loup, Quebec, 1988.
- Berendse, F., B. Berg, and E. Bosatta, The effect of lignin and nitrogen on the decomposition of litter in nutrient-poor ecosystems: A theoretical approach, Can. J. Bot., 65, 1116-1120, 1987.
- Bosatta, E., and F. Berendse, Energy or nutrient regulations of decomposition: Implications for the mineralization-immobilization response to perturbations, Soil Biol. Biochem., 16, 63-67, 1984.
- Bryant, M. P., E. A. Wolin, M. J. Wolin, and R. S. Wolfe, Methanobacillus omelianskii, a symbiotic association of two species of bacteria, Arch. Mikrobiol., 59, 20-31, 1967.
- Chanton, J. P., and J. W. H. Dacey, Effects of vegetation on methane flux, reservoirs, and carbon isotopic composition, in *Trace Gas Emissions* by *Plants*, edited by E. A. Holland and H. A. Mooney, pp. 65-92, Academic, San Diego, Calif., 1991.
- Chapin, F. S. III, A. J. Bloom, C. B. Field, and R. H. Waring, Plant responses to multiple environmental factors, *Bioscience*, 37, 49-57, 1987.
- Conrad, R., Control of methane production in terrestrial ecosystems, in Exchange of Trace Gases Between Terrestrial Ecosystems and the Atmosphere, edited by M. O. Andreae and D. S. Schimel, pp. 39-58, John Wiley, New York, Chichester, 1989.
- Conrad, R., and H. Schütz, Methods of studying methanogenic bacteria and methanogenic activities in aquatic environments, in Methods in

Aquatic Bacteriology, edited by B. Austin, pp. 301-343, John Wiley, New York, Chichester, 1988.

- Crill, P. M., K. B. Bartlett, R. C. Harriss, E. Gorham, E. S. Verry, D. I. Sebacher, L. Madzar, and W. Sanner, Methane flux from Minnesota peatlands, *Global Biogeochem. Cycles*, 2, 371-384, 1988.
- Fechner, E. J., and H. F. Hemond, Methane transport and oxidation in the unsaturated zone of a Sphagnum peatland, Global Biogeochem. Cycles, 6, 33-44, 1992.
- Fung, I., J. John, J. Lemer, E. Matthews, M. Prather, L. P. Steele, and P. J. Fraser, Three-dimensional model synthesis of the global methane cycle, J. Geophys. Res., 96 (D7), 13,033-13,065, 1991.
- Goering, H. K. and P. J. Van Soest, Forage fiber analysis, Agri. Handb. 379, Agri. Res. Serv., U. S. Dep. of Agri., Gov. Print. Off., Washington, D. C., 1970.
- Hogg, E. H., V. J. Lieffers, and R. W. Wein, Potential carbon losses from peat profiles: Effects of temperature, drought cycles, and fire, *Ecol. Appl.*, 2, 298-306, 1992.
- Jeris, J. S., and P. L. McCarty, The biochemistry of methane fermentation using ¹⁴C tracers, J. Water Pollut. Control Fed., 37, 178, 1965.
- Keller, M., W. A. Kaplan, and S. C. Wofsy, Emissions of N₂O, CH₄ and CO₂ from tropical forest soils, J. Geophys. Res., 91(D11), 11,791-11,802, 1986.
- Kiene, R. P., and D. G. Capone, Degassing of pore water methane during sediment incubations, Appl. Environ. Microbiol., 49, 143-147, 1985.
- Kirk, T. K., and J. R. Obst, Lignin determination, in *Methods in Enzymology*, edited by W. A. Wood and S. T. Kellogg, vol. 161, pp. 87-101, 1988.
- Klinger, L. F., P. R. Zimmerman, J. P. Greenberg, L. E. Heidt, and A. B. Guenther, Carbon trace gas fluxes along a successional gradient in the Hudson Bay lowland, J. Geophys. Res., this issue.
- Klinger, L. F., S. A. Elias, V. M. Behan-Pelletier, and N. E. Williams, The bog climax hypothesis: Fossil arthropod and stratigraphic evidence in peat sections from southeast Alaska, USA, *Holarctic Ecol.*, 13, 72-80, 1990.
- Knowles, R. Methane: processes of production and consumption, in Agricultural Ecosystem Effects on Radiatively Important Trace Gases and Global Climate Change, edited by L. A. Harper, A. R. Mosier, J. M. Duxbury, and D. E. Rolston, Spec. Pub. 55, Am. Soc. of Agron., in press, 1993.
- Large, P. J., Methylotrophy and Methanogenesis, Am. Soc. for Microbiol., Washington, D. C., 1983.
- Maestrojuán, G. M., and D. R. Boone, Characterization of methanosarcina barkeri strains MST and 227, Methanosarcina mazei S-6T, and Methanosarcina vacuolata Z-761T, Int. J. Syst. Bacteriol., 41, 267-274, 1991.
- Melillo, J. M., J. D. Aber, and J. F. Muratore, Nitrogen and lignin controls of hardwood leaf litter decomposition dynamics, *Ecology*, 63, 621-626, 1982.
- Moore, T. R., and R. Knowles, Methane emissions from fen, bog, and swamp peatlands in Quebec, *Biogeochemistry*, 11, 45-61, 1990.
- Moore, T., N. Roulet, and R. Knowles, Spatial and temporal variations of methane flux from subarctic/northern boreal fens, *Global Biogeochem*. *Cycles*, 4, 29-46, 1990.
- Morrissey, L.A., and A.P. Livingston, Methane emissions from Alaska Arctic tundra: An assessment of local spatial variability, J. Geophys. Res., 97, 16,661-16,670, 1992.
- Mosier, A., D. Schimel, D. Valentine, K. Bronson, and W. Parton, Methane and nitrous oxide fluxes in native, fertilized and cultivated grasslands, *Nature*, 350, 330-332, 1991.
- Oremland, R. S., Biogeochemistry of methanogenic bacteria, in *Biology* of Anaerobic Microorganisms, edited by A. J. B. Zehnder, pp. 641-705, John Wiley, New York, 1988.
- Parnas, H., Model for decomposition of organic material by microorganisms, Soil Biol. Biochem., 7, 161-169, 1975.
- Patel, G. B., G. D. Sprott, and J. E. Fein, Isolation and characterization of methanobacterium espanolae sp. nov., a mesophilic, moderately acidophilic methanogen, Int. J. Syst. Bacteriol., 40, 12-18, 1990.
- Quay, P. D., et al., Carbon isotopic composition of atmospheric CH₄: Fossil and biomass burning source strengths, *Global Biogeochem*. Cycles, 5, 25-47, 1991.
- Roulet, N. T., R. Ash, and T. R. Moore, Low boreal wetlands as a source of atmospheric methane, J. Geophys. Res., 97, 3739-3749, 1992.
- Roulet, N. T., J. Ritter, A. Jano, C. A. Kelly, L. F. Klinger, T. R. Moore, R. Protz, and W. R. Rouse, The role of the Hudson Bay lowland as a source of atmospheric methane, J. Geophys. Res., this issue.
- Sass, R. L., F. M. Fischer, and P. A. Harcombe, Methane production and emission in a Texas rice field, *Global Biogeochem. Cycles*, 4, 47-68, 1990.

- Sass, R. L., F. M. Fischer, and P. A. Harcombe, Mitigation of methane emission from rice fields: Possible adverse effects of incorporated rice straw, Global Biogeochem. Cycles, 5, 275-287, 1991.
- Steudler, P. A., R. D. Bowden, J. M. Melillo, and J. D. Aber, Influence of nitrogen fertilization on methane uptake in temperate forest soils, *Nature*, 341, 314-316, 1989.
- Svensson, B. H., Methane production in tundra peat, in Microbial Production and Utilization of Gases (H₂, CH₄, CO), edited by H. G. Schlegel, G. Gottschalk, and N. Pfennig, pp. 135-139E, Goltze KG, Göttingen, 1976.
- Svensson, B. H., Carbon dioxide and methane fluxes from the ombrotrophic parts of a subarctic mire, in Ecology of a Subarctic Mire, edited by M. Sonesson, *Ecol. Bull. (Stockholm)*, 30, 235-250, 1980.
- Svensson, B.H., Different temperature optima for methane formation when enrichments from acid peat are supplemented with acetate or hydrogen, Appl. Env. Microbiol., 48, 389-394.
- Svensson, B. H., and T. Rosswall, Energy flow through a subarctic mire at Strodalen, in Ecology of a Subarctic Mire, edited by M. Sonesson, Ecol. Bull. (Stockholm), 30, 283-301, 1980.
- Svensson, B. H., and T. Rosswall, In situ methane production from acid peat in plant communities with different moisture regimes in a subarctic mire, Oikos, 43, 341-350, 1984.
- Taylor, J. A., G. P. Brasseur, P. R. Zimmerman, and R. J. Cicerone, A study of the sources and sinks of methane and methyl chloroform using a global three-dimensional lagrangian tropospheric tracer transport model, J. Geophys. Res., 96, 3013-3044, 1991.
- Tyler, S. C., Stable carbon isotope ratios in atmospheric methane and some of its sources, J. Geophys. Res., 91, 13,232-13,238, 1986.
 Tyler, S. C., ¹³C/¹²C ratios in atmospheric methane and some of its
- Tyler, S. C., ¹³C/¹²C ratios in atmospheric methane and some of its sources., in Stable Isotopes in Ecological Research, edited by P. W. Rundel, J. R. Ehleringer, and A. K. Nagy, Springer-Verlag, New York, *Ecol. Studies*, 68, 395-409, 1989.
- Tyler, S. C., The global methane budget, in Microbial Production and Consumption of Greenhouse Gases: Methane, Nitrogen Oxides, and Halomethanes, edited by J. E. Rogers and W. B. Whitman, pp. 7-38, American Society for Microbiology, Washington, D. C., 1991.
- Vaghjiani, G. L., and A. R. Ravishankara, New measurement of the rate coefficient for the reaction of OH with methane, *Nature*, 350, 406-409, 1991.
- Van Cleve, K., and L. A. Viereck, Forest succession in relation to nutrient cycling in the boreal forest of Alaska, in *Forest Succession: Concepts* and Application, edited by D. West, H. Shugart and D. Botkin, pp. 185-211, Springer-Verlag, New York, 1981.
- Van Cleve, K., F. S. Chapin III, C. T. Dyrness, and L. A. Viereck, Element cycling in taiga forests: State-factor control, *Bioscience*, 41, 78-88, 1991.
- Wahlen, M., N. Tanaka, R. Henry, T. Yoshinari, R. G. Fairbanks, A. Shemesh, and W. S. Broecker, ¹³C, D and ¹⁴C in methane (abstract)EOS, Trans. AGU, 68, 1220, 1987.
- Webber, P. J., J. W. Richardson, and J. T. Andrews, Post glacial uplift and substrate age at Cape Henrietta Maria, southeastern Hudson Bay, Canada, Can. J. Earth Sci., 7, 317-325, 1970.
- Westermann, P., B. K. Ahring, and R. A. Mah, Temperature compensation in Methanosarcina barkeri by modulation of hydrogen and acetate affinity, Appl. Environ. Microbiol., 55, 1262-1266, 1989.
- Whalen, S. C., and W. S. Reeburgh, A methane flux transect along the trans-Alaska pipeline haul road, *Tellus*, 42(B), 237-249, 1990.
- Whiting, G. J., and J. P. Chanton, Plant-dependent CH₄ emission in a subarctic Canadian fen, Global Biogeochem. Cycles, 6, 225-231, 1992.
- Williams, R. T., and R. L. Crawford, Methanogenic bacteria, including an acid tolerant strain, from peatlands, Appl. Environ. Microbiol., 39, 194-204, 1985.
- Yavitt, J. B., and G. E. Lang, Methane production in contrasting wetland sites: Response to organic-chemical components of peat and to sulfate reduction, *Geomicrobiol. J.*, 8, 27-46, 1990.

E.A. Holland and D.S. Schimel, National Center for Atmospheric Research, P.O. Box 3000, Boulder, CO 80307.

D. Valentine, Natural Resource Ecology Laboratory, Colorado State University, Fort Collins, CO 80523.

(Received May 29, 1992; revised January 28, 1993; accepted February 10, 1993.)

•

from the mechanism outlined above. In the Antarctic Peninsula, the northern ice fronts of George VI and Larsen ice shelves have a history of recession over the past fifty years¹⁹ and are still retreating. Another at risk is Wilkins Ice Shelf. Snow temperatures indicate that this region is at most 1-2 °C colder than the Wordie²⁰ and radio echo measurements show that large areas are infiltrated by brine. If the warming trend continues for another decade, this may be the next ice shelf to start receding.

Received 27 December 1990: accepted 26 February 1991.

- 1. Mercer.). H. Neture 271, 321-325 (1978)
- Ewen Smith, B. M. Br. Antarct, Survey Sci. Reports 72 (1972).
 Reynolds, J. M. Br. Antarct, Survey Bull. 80, 57-64 (1988).
- Walford, M. E. R. Nature 239, 95-96 (1972)
- 5. Doare, C. S. M. Nature 257, 780-782 (1975). 6. Hughes, T. J. Glaciol. 28, 98-117 (1983).
- 7. MacAyeal, O. R. & Thomas, R. H. Ann. Glaciol. 3, 189-194 (1982).
- 8. Lange, M. A. & MacAyeal, O. R. Ann. Glaciol, 12, 97-103 (1989)
- 9. Swithindami, C. W. M. US Geol. Survey Professional Paper 8-1385 (1988).
- 10. Telbot, M. H. Ann. Glacov. 11, 151-154 (1988).
- 11. Morrison, S. J. Weether 48, 231-232 (1990). 12. Pollard, D. Tellus 32, 384-388 (1980).
- 13. Oerlemans, J. Nature 297, 550-553 (1982).

Ì

Same -

Â

- 14. Liu, H. W. & Miller, K. J. J. Glacol. 22, 135-143 (1979).
- 15. Sabol, S. A. & Schulson, E. M. J. Glaciol. 36, 191-192 (1989).
- 16. Robin, G de Q. / Glaciol. 13, 543 (1974).
- 17. Jezek, K. C. J. geophys. Res. 89, 1925-1931 (1984).
- 18. Thomas, R. H. Br. Antartt. Survey Sci. Reports 79 (1973). 19. Domie, C. S. M. Ann. Glaciol. 3, 77-82 (1982).
- 20. Reynolds, J. M., Br. Antarct, Survey Bull. 54, 123-133 (1981).

ACKNOWLEDGEVENTS. We thank Jane Ferrigino and Baerbei Lucchitta of the US Geological Survey for help in supplying Landsat images, some of which were obtained under the Landsat Acquisition Programme, initiated by the SCAR Working Group on Glaciology.

Methane and nitrous oxide fluxes in native, fertilized and cultivated grasslands 71A 31523

A. Mosier*, D. Schimel†, D. Valentine†, K. Bronson* & W. Parton†

* United States Department of Agriculture, Agricultural Research Service, PO Box E, Ft Collins, Colorado 80522, USA † Natural Resource Ecology Laboratory, Colorado State University, Ft Collins, Colorado 80523, USA

METHANE and nitrous oxide are long-lived, radiatively active trace gases that account for ~20% of the total anticipated atmospheric warming¹. The atmospheric concentrations of both gases have increased dramatically over the past few decades, and continue to increase at a rate of ~1.1 and 0.25% yr⁻¹ for CH₄ (ref. 2) and N₂O (ref. 3) respectively. Increased biospheric production is generally suggested as the reason for the increases, but decreases in global sinks may also be important. It has been suggested, for example, that nitrogen fertilization may decrease the rate at which tropical^{4,5} and temperate forest soils⁶ take up methane from the atmosphere. Furthermore, the recent extensive changes in land management and cultivation could be contributing to the observed increases in both atmospheric CH4 and N2O, as has been suggested for tropical soils7. Little information exists on CH₄ uptake in temperate grasslands (which currently occupy -8% of the Earth's surface), its relation to N2O production, or the effect of land management or cultivation^{2,9}. Here we report measurements of CH4 uptake and N2O emissions in native, nitrogen-fertilized and wheat-growing prairie soils from spring to late autumn, 1990. We found that nitrogen fertilization and cultivation can both decrease CH4 uptake and increase N2O production, thereby contributing to the increasing atmospheric concentrations of these gases.

We established three sets of sites in the Colorado shortgrass steppe during late winter and spring, 1990. One set was established along a toposequence (catena) to determine whether the landscape position-sandy loam upland (midslope) versus sandy clay loam lowland (swale)-in the native prairie (catena site¹⁰) affected CH₄ uptake. Previous investigations^{10,11} had found that ammonia and N₂O fluxes differed markedly with landscape position. A second set of sites was established in unamended and annually nitrogen-fertilized native grass pastures. The third set was established in a pair of wheat fields, cropped in alternating years (wheat-fallow system), and in an adjacent native grassland for comparison. All sites were in or near the United States Department of Agriculture-Agricultural Research Service Central Plains Experimental Range located north of Nunn, Colorado (latitude 40° 48' 23" N, longitude 104° 45' 15" W). Table 1 summarizes the soil characteristics.

On the catena, the fertilized midslope site was treated with 45 g m^{-2} of urea-N in July 1981, and the fertilized swale site was similarly treated in May 1982 (ref. 11). Four replicate, permanent plots for gas flux measurement were established in the fertilized and unfertilized midslope and swale sites by driving 20.3-cm diameter PVC pipes into the soil to a depth of 3 cm. Cattle have been excluded from the catena site since 1980. Weather and soil temperature data were continuously collected \sim 200 m from the site.

The annual fertilization experiment was ~1 km north of the catena site. We established six replicate plots in both nitrogenfertilized and control grazed pastures, hereafter called AFS (annual fertilized) and CAFS (control annual fertilized) sites. The terrain in these two pastures is essentially flat, and the soil is similar to the catena midslope (Table 1). The AFS received 2.2 g N m⁻² of ammonium nitrate annually from 1976 to 1989.

Finally, we established six plots in each of three adjacent sites-wheat, fallow, and native grassland-6 km east of the Central Plains Experimental Range. The wheat and fallow sites had been in continuous cultivation without irrigation or application of nitrogen fertilizer, pesticides or herbicides since the native prairie was ploughed in 1981. Winter wheat was planted in September 1989, and the fallow site was kept weed-free by cultivation following harvest in July 1989. The crop was harvested in July 1990, and the fallow site was planted with wheat in September 1990. The native grassland is 200 m east of the wheat-fallow field.

We measured methane and nitrous oxide fluxes by placing a closed chamber¹² over the established plots and taking gas samples from inside the chambers with 50-ml polypropylene syringes fitted with nylon stopcocks, usually 0, 15 and 30 minutes after the chambers were installed. We analysed the samples within six hours by gas chromatography using a Porapak N column and flame ionization detector for CH4, and Porapak Q and electron capture detector¹³ for N₂O from the same syringe. Flux measurements were made mid-morning on each sampling day, generally weekly at each site. We tested sampling time periods ranging from 10 to 60 minutes and taken at 2 or 3 time intervals, and in all cases CH, uptake was greater during the first sampling period than in subsequent periods. We therefore calculated CH, uptake rates assuming first-order kinetics rather than linear decay as had been suggested[®]. Measurements of CH₄ and N2O fluxes were made at 6:00, 12:00 and 18:00 on several days. Over a 24-hour period, changes in CH, uptake averaged 10% or less of the mean, whereas the spatial and tempora: variation in N2O emissions was much higher¹⁴.

We collected soil samples (0-15 cm) at the time of each gas flux measurement. The soils were analysed gravimetrically for soil water content and extracted with 2M KCI, and the extracts analysed colorimetrically for ammonium and nitrate content. We analysed soils from each site for total carbon and nitrogen content with a combustion C/N analyser. From July, we measured the soil temperature at a depth of 2.5 cm at each site to supplement the continuous soil temperature readings being made near the catena.

NATURE - VOL 350 - 28 MARCH 1991

.

	TABLE 1 Physical and chemical properties of soils and flux rates of CH4 and N20 from each site										
:			Soil analysis	NOT	NHT	1	CH, uptake	Gas flux	i ratesý N	20 emissio	n .
Site	рH	Total N (%	Total C)	mg N per kg	mg N per kg	n	<i>X</i> ig C ha	s.d. 1 ⁻¹ d ⁻¹)	a	x ig∉na N	s.a. a ⁻¹ a ⁻¹)
Swale ⁺ Fertilized Unfertilized	5.6 6.0	0.223 0.189	2.215 1.841	5.1 1.6	0.7 0.9	128 136	3.6 3.6	1.6 1.3	108 116	6.2 3.0	15.5 3.7
Midslope‡ Fertilized Unfertilized	5.6 6.5	0.1 38 0.141	1.216 1.245	1.3 1.1	0. 9 0.9	124 132	4.1 6.3	1.4 2.1	112 120	3.1 1.8	4.0 1.7
Pasture‡ Fertilized Unfertilized Wheat‡ Fallow‡	5.6 6.2 5.8 6.0	0.138 0.120 0.116 0.111 0.104	1.177 1.066 0.955 0.850 0.841	6.4 1.0 1.1 10.1 0.6	20.3 1.5 1.4 1.9 1.3	180 180 162 162 162	3.8 5.8 1.3 1.8 2.6	1.8 1.9 0.6 0.8 1.1	162 162 162 162 162	6.1 2.5 2.6 4.5 3.5	10.7 2.4 2.7 6.7 5.6

AES

* Data are means for soil samples (0-15 cm) collected on each gas sampling dzy.

+ The swale is a sandy clay loam soil classified as a Pachic argiustoll.

These are sandy loarn soils classified as Ustollic haplargids.

§ n, number of samples; \vec{X} mean; s.d., standard deviation.

ショリスたちではく、

A CONTRACTOR

4

F.

Methane uptake rates in the unfertilized grassland averaged 3.6, 5.8 and 6.3 g carbon per hectare day (g C ha⁻¹ d⁻¹) in the swale, CAFS and midslope sites, respectively, between March and December 1990 (Fig. 1 and Table 1). Methane uptake at the swale was significantly less (P < 0.01) than at the CAFS and midslope sites. The midslope and CAFS soils were similar in texture and N content, whereas the swale site had finer texture and higher total N content (Table 1). The soil nitrate and ammonium contents of the three unfertilized soils were similar (P > 0.05). Annual *in situ* N mineralization at the midslope is, however, substantially less than in the swale¹⁰ (41 compared with 55 kg N ha⁻¹ yr⁻¹), suggesting that it is the N turnover (mineralization and nitrification)¹⁵, rather than the mineral N content, which directly influences the CH₄ uptake.

Nitrous oxide efflux from the soils was inversely related to methane uptake (Fig. 1), averaging 3.0 and 1.8 g N ha⁻¹ d⁻¹ in unfertilized swale and midslope sites, respectively, compared to 3.6 and 6.3 g C ha⁻¹ d⁻¹. The nitrous oxide flux from all sites was highest immediately following precipitation, when CH₄ uptake was lowest. If the fluxes of N₂O (ref. 11) and CH₄ (ref. 16) were both largely mediated by ammonium-oxidizing bacteria, then CH₄ uptake may have begun when the soil moisture declined to the point where slowed ammonium diffusion limited nitrification, although our data do not preclude other explanations. Under very dry conditions, both CH₄ uptake and N₂O emissions declined, and they remained low until soil water content increased.

Nitrogen fertilization decreased CH₄ uptake (Fig. 1), as was found for temperate forest soils⁶. In the AFS plots, annual N applications caused the CH₄ uptake to decrease by an average of 41% (P < 0.01). Methane uptake by the midslope site decreased from 6.3 to 4.1 g C ha⁻¹ d⁻¹ (Fig. 1) as a result of applying 45 g m⁻² of urea-N in 1981 (P < 0.01). A similar N application to the more fertile swale soil in 1982 did not alter CH₄ uptake, which averaged 3.6 g C ha⁻¹ d⁻¹. These data suggest that high N turnover, whether native or due to fertilization, suppresses CH₄ uptake.

Nitrogen fertilization increased N₂O emissions by a factor of 2-3 at all sites (P < 0.05), as had been found earlier¹¹. Overall, N₂O emissions from the midslope and swale sites were 2-4 times greater in 1990 than in 1981 or 1982 (ref. 11), possibly due to wetter conditions in 1990. The inverse relationship between N₂O





Mic-son

•

.

~

flux and CH₄ uptake was apparent with or without nitrogen fertilization. The influence of nitrogen fertilization on N-O flux was most evident in moist soils, whereas its influence on CH. uptake was most evident as soils began to dry after rain (Fig. 1).

To determine whether cultivation affects CH₄ uptake in grassland soils, as observed in tropical soils⁷, we began monitoring CH₄ uptake in the wheat-fallow system. During 6 months of weekly measurements, CH4 uptake averaged 2.6, 1.8 and 1.3 g C ha⁻¹ d⁻¹ in native grassland, fallow and wheat sites. respectively (Fig. 2). These data indicate that disturbing the natural grassland decreased CH4 uptake (P < 0.01). Soil moisture contents averaged 8.6, 11.1 and 12.9% in grassland, fallow and wheat, respectively. Soil ammonium content did not differ among the three sites but soil nitrate was on average -10 times higher in the fallow soil than in the other two sites. Long-term experimentation and frequent flux measurements are needed to understand the reasons for differing CH, uptake rates in different soils and crop pastures.

Our data indicate that the semi-arid grasslands represent a significant global sink for CH₄ (ref. 8). Methane uptake in the grassland ranged from 6 to 61 μ g CH₄ m⁻² h⁻¹, compared with uptake rates of 6-24, 52, 0-112, and 10-160 μ g CH₄ m⁻² h⁻¹ in tropical forest¹⁷, subtropical broad-leafed savannah¹⁸, tundra¹⁹ and temperate forest soils^{6,8}, respectively. If our measurements are representative of CH4 uptake by these grasslands globally, then 0.5 to 5.6 Tg of CH₄ are removed from the atmosphere in these grasslands each year.

Out data suggest that nitrogen fertilization in natural ecosystems increases N2O emissions and decreases CH4 uptake⁶. These



FIG. 2 Effects of wheat cropping on CH₄ uptake and N₂O emissions in shortgrass prairie soils. Arrows indicate approximate dates of harvest and planting, when the wheat and fallow designations switch,

data also indicate that recent changes in use or management of grasslands, such as cultivation, have decreased CH, uptake from and increased N2O efflux to the atmosphere. In addition to increased production from rice paddies, ruminants and other sources²⁰, decreased consumption by soils may contribute to increasing atmospheric methane.

Received 8 November 1990: accepted 28 January 1991.

- 1. Rothe H. Science 248, 1217-1219 (1990)
- Cicerone, R. J. & Oremland, R. S. Global biogeoche m. Cycles 2, 299-327 (1988)
- Prinn, R. et al. J. geophys. Res. 36, 18369-18385 (1990)
- 4. Bouwman, A. F. Soils and the Greenhouse Effect (Wiley, Chichester, 1990). Andreae, M. O. & Schimel, D. S. Exchange of Trace Gases between Terrestrial Ecosystems and
- the Atmosphere. (Wiley, Chichester, 1989).
- 5. Steudler, P. A., Bowoen, R. D., Mehlio, J. M. & Aber, J. D. Nature 341, 314-315 (1989) Keller, M., Mitre, M. M. & Stallard, R. F. Global pioteochem. Cycles 4, 21-27 (1990).
- Schutz, H., Seiler, W. & Rennenberg, H. in Soils and the Greenhouse Effect (ed. Bouw
- 269-301 (Weey, Chichester, 1990) 9
- Seller, W. & Conrad, R. in The Geophysiology of Amazonia: Vegetation and Climate Interactions led. Dickinson, R. E.) 133-162 (Wiley, New York, 1987). Schimet, D., Stillwell, M. A. & Woodmansee, R. G. Ecology 66, 276-282 (1985)
- 11.
- Parton, W. J., Mosier, A. R. & Schimel, D. S., Biogeochemistry 4, 45-58 (1988). Hutchinson, G. L. & Mosier, A. R. Soil Sci. Soc. Am. J. 45, 311-316 (1981).
- Mosier, A. R. & Mack, L. Solf Sci. Soc. Am. J. 44, 1121-1123 (1980). 13
- 14. Slemr, F., Conrad, R. & Sevier, W. J. atmos. Chem. 1, 159-169 (1984). 15. Schimel, D. S. & Parton, W. J. Plant and Soil 93, 347-357 (1986).
- Melillo, J. M., Steudler, P. A., Aber, J. D. & Bowden, R. D. in Exchange of Trace Gases Betw Terrestnal Ecosystems and the Atmosphere (eds Andreae, M. O. & Schimel, D. S.) 263-280
- (Wiley, Choffmer, 1989) 17. Keller, M., Goreau, J. T., Wofsy, S. C., Kaplan, W. A. & McElroy, M. B. Geophys. Res. Lett. 10,
- 1156-1159 (1983).
- 18. Seiler, W., Conrad, R. & Schartfe, D. J. atmos. Chem. 1, 171-186 (1984). Whaten, S. C. & Reeburgh, W. S. Nature 346, 150-152 (1990). 19.
- 20
- Aseiman, I. & Crutzen P. 1 J atmos Chem. 8, 307-358 (1989)

ACKNOWLEDGEVENTS. We thank Or M. C. Shoop for access to his grassiand nitrogen-fertilization experiments, Jim Barnes for permitting use of his wheat-failow fields, and Anita Kear and Rod Hans for sample collection and analysis. Funding for this research was provided by NASA and USDA-ARS

Phases and electrical conductivity of a hydrous silicate assemblage at lower-mantle conditions

Xiaoyuan Li* & Raymond Jeanloz

Department of Geology and Geophysics, University of California, Berkeley, California 94720, USA

THE presence of a small amount of water in the lower mantle might affect in a significant way the geophysical and geochemical properties of its host mineral assemblage¹⁻⁵. Here we present experimental observations of the phase behaviour and the electrical conductivity of a hydrous silicate assemblage synthesized from a mixture of (Mgo, =Feo, 12)SiO3 pyroxene and water under the pressure and temperature conditions of the lower mantle. Previous studies have shown that anhydrous (Mg, Fe)SiO3 pyroxene transforms to a perovskite structure under these conditions⁶⁻⁹. We find that, although the hydrous assemblage is also dominated by the (Mg, Fe)SiO₃ perovskite phase, it coexists with the so-called hydrous phase D, of estimated composition (Mg, Fe)SiH₂O₄. Our measurements show that the inclusion of small amounts of water in the silicates can enhance the electrical conductivity of the lower-mantle assemblage by more than three orders of magnitude at these temperatures and pressures.

Our starting material was prepared by homogeneously mixing a $(Mg_{0.88}Fe_{0.12})SiO_3$ pyroxene with $4(\pm 2)\%$ by weight of distilled water, giving a (Mgo ss Feo 12)SiO3/H2O molar ratio of 4/1. The pyroxene was a natural enstatite from Bamble, Norway, which was ground to a grain size $\leq 1 \mu m$ before mixing. The highpressure perovskite phase synthesized from anhydrous Bamble enstatite has been carefully documented in previous studies (for example, refs 4, 7, 10).

Laboration.

^{*} Present address: Hawaii institute of Geophysics, University of Hawaii, Honolulu, Hawaii 96822, USA

•

.

.

Chemosphere, Vol.26, Nos.1-4, pp 675-685, 1993 Printed in Great Britain 0045-6535/93 \$6.00 + 0.00 Pergamon Press Ltd.

ND

EFFECT OF LAND USE CHANGE ON METHANE OXIDATION IN TEMPERATE FOREST AND GRASSLAND SOILS

D.S. Ojima^{1*}, D.W. Valentine^{1,2}, A.R. Mosier², W.J. Parton¹, and D.S. Schimel^{1,3}

¹Natural Resource Ecology Laboratory, Colorado State University, Fort Collins, CO 80523, USA ²USDA/ARS, P.O. Box E, Fort Collins, CO 80522, USA ³Climate System Modeling Program, University Corporation for Atmospheric Research, P.O. Box 3000, Boulder, CO 80307-3000, USA

(Received in USA 13 November 1991; accepted 14 April 1992)

ABSTRACT

Evidence is accumulating that land use changes and other human activity during the past 100 to 200 years have contributed to decreased CH, oxidation in the soil. Recent studies have documented the effect of land use change on CH, oxidation in a variety of ecosystems. Increased N additions to temperate forest soils in the northeastern United States decreased CH, uptake by 30 to 60%, and increased N fertilization and conversion to cropland in temperate grasslands decreased CH, uptake by 30, to 75%.

Using these data, we made a series of calculations to estimate the impact of land use and management changes which have altered soil the CH₄ sink in temperate forest and grassland ecosystems. Our study indicates that as the atmospheric mixing ratio of CH₄ has increased during the past 150 y, the temperate CH₄ sink has risen from approximately 8 Tg y⁻¹ to 27 Tg y⁻¹, assuming no loss of land cover to cropland conversion. The net effect of intensive land cover changes and extensive chronic disturbance (i.e., increased atmospheric N deposition) to these ecosystems have resulted in about 30% reduction in the CH₄ sink relative to the soil sink assuming no disturbance to any of the temperate ecosystems. This will impact the global CH₄ budget even more as atmospheric CH₄ concentrations increase and as a result of further disturbance to other biomes. Determining the reasons for the decreased CH₄ uptake due to land disturbance is necessary to understand the role of CH₄ uptake in conjunction with the increasing atmospheric CH₄ concentrations. Without accounting for this approximately 20 Tg y⁻¹ temperate soil sink, the atmospheric CH₄ concentration would be increasing about 1.5 times the current rate.

INTRODUCTION

During the past century, global methane (CH₄) concentrations have more than doubled and during the last 30 y have increased at an annual rate of about 1% (Rasmussen and Khalil, 1981; Bolle et al., 1986; Steele et al., 1987; Khalil et al., 1989). Although atmospheric CH₄ concentration is about 200 times less than CO₂, it contributes approximately 15% to current "greenhouse" forcing in the atmosphere (Hansen et al., 1989). This is because CH₄ is about 30 times more effective as CO₂ in absorbing IR radiation (Wuebbles and Edmonds,

PRECEDING PAGE BLANK NOT FILMED

1991), and is about 45 times more effective as a greenhouse gas overall than CO_2 when the longer atmospheric residence time of CH₄ is taken into account (Rodhe, 1990). This increase in CH₄ has been attributed to increasing sources such as rice paddy cultivation, landfill operations, fossil fuel processing and distribution, and livestock management (Khalil and Rasmussen, 1983; Seiler et al., 1984; Crutzen et al., 1986; Cicerone and Oremland, 1988; Aselmann and Crutzen, 1989). Current estimates of global annual CH₄ atmospheric loading are 400 to 600 Tg.

Methane emission is offset by oxidation that largely takes place in the atmosphere through reaction with the OH radical (Crutzen and Gidel, 1983; Khalil and Rasmussen, 1983; Cicerone and Oremland, 1988). The current increases in atmospheric concentrations of CO and CH, are expected to result in a decline in the OH radical over the next 50 years (Thompson and Cicerone, 1986; Vaghjiani and Ravishankara, 1991). Thompson and Cicerone (1986) estimate an increase in atmospheric CH, from 1.6 to 2.9 ppmv would occur from 1985 to 2035, based on their photochemical model experiments. Given the magnitude of CH, increases and OH losses, it is estimated that approximately 50% of the methane increase will be due to an increased lifetime of methane resulting from a smaller OH sink, with the rest due to increased emissions of methane (Thompson and Cicerone, 1986).

in interdentity it is the birth of the birth is a first of the second second second second second second second

An additional CH, sink exists, observations that aerobic soils also serve as sinks for atmospheric CH, (Keller et al., 1983; Seiler et al., 1984) and the confirmation that the phenomenon is widespread (Born et al., 1989; Steudler et al., 1989; Striegl and Ishii, 1989; Keller et al., 1990; Whalen and Reeburgh, 1990; Mosier et al., 1991) suggest that the dynamics of the soil sink term needs to be critically evaluated in estimating the global budget. Current estimates of the terrestrial soil sink for CH, tanges between 40 to 60 Tg y⁻¹ (Cicerone and Oremland, 1988; Schutz et al., 1990). Whalen et al. (1990 and 1991) suggested that CH, oxidation in tundra and taiga soils not only accounts for consumption of CH₄ produced in these soils but also consumes atmospheric CH₄. They concluded that soil CH₄ oxidation could provide a negative feedback to atmospheric CH₄ increases, and if soils dry out with climate change, the soil oxidation sink would become more pronounced. A number of factors influence the magnitude of the soil sink for methane, including soil water status, soil temperature, nitrogen dynamics, and microbial population, but quantification of these factors on the dynamics of methane uptake is still poor. In addition, little is known about how CH₄ oxidation proceeds in the major croplands that cover approximately 1500 10⁶ ha or approximately 10% of the global land surface (Houghton et al., 1983).

The objective of this paper is to assess the role of temperate forest and grassland conversion and the interaction between land cover conversion and changes in atmospheric CH₄ on soil CH4 uptake from 1850 to the present. This assessment takes into account the change in sink strength relative to estimated changes in atmospheric concentration of CH₄ during this period and changes in atmospheric deposition of N over temperate regions of the Earth.

LAND USE IMPACT ON THE SOIL SINK FOR METHANE

Land use changes can affect key biogeochemical rates, leading to changes in N_2O fluxes from land clearing in forests (Luizao et al., 1989); increased N_2O fluxes due to fertilization (Mosier et al., 1986); decreased CH₄ uptake by soils fertilized with N (Steudler et al., 1989, Mosier et al., 1991); and increased CH₄ flux due to cattle and paddy rice cultivation (Crutzen et al., 1986; Cicerone and Oremland, 1988; Aselmann and Crutzen, 1989). These impacts are locally driven, but aggregated they have an impact on the regional and global flux of key biogenic trace gases, such as CH₄ and N₂O.

The oxidation of CH₄ has been observed in temperate forest soils (Keller et al., 1983, 1986; Steudler et al., 1989; Yavitt et al., 1990), and tropical forest soils (Keller et al., 1986, Steudler et al., 1991). Methane uptake values for temperate deciduous forests ranged between 0.5 to 5.5 kg CH₄ ha⁻¹ y⁻¹ (Keller et al., 1983; Steudler et al., 1989). In relatively undisturbed tropical ecosystems, CH₄ uptake in forests ranged from 0.5 to 2.9 kg CH₄ ha⁻¹ y⁻¹ (Keller et al., 1983, 1986, 1990; Seiler et al., 1984) and in more arid savanna and woodlands was 4.6 kg CH₄ ha⁻¹ y⁻¹ (Seiler et al., 1984). And Striegl, pers. comm., estimated uptake of 2.4 kg CH₄ ha⁻¹ y⁻¹ for Mojave Desert/Southern Great Basin soil. It should be noted that most of these estimates may not represent the annual uptake of CH₄ since measurements were not taken throughout the year.

A recent study showed that semiarid grasslands are an important methane sink and conversion of these grasslands to croplands significantly reduces oxidation of CH₄ in these soils (Mosier et al., 1991). Chamber measurements of CH₄ uptake and N₂O fluxes were made from a native shortgrass steppe, an annually fertilized shortgrass steppe, a previously cultivated site that was put back into grassland in 1939 ("go-back", visually indistinguishable from the native shortgrass steppe), and a cultivated dryland wheat-fallow site (Mosier et al. 1991).

Estimates of annual methane uptake in the dryland wheat field showed 75% less CH₄ uptake relative to the undisturbed grassland site (Figure 1). Cultivation appeared to have a long-term impact on the uptake potential of temperate grasslands. The "go-back" plot consumed less than 40% of the CH₄ in the undisturbed soils. Nitrogen additions of 22 kg ammonium-nitrate ha⁻¹ y⁻¹ for 15 years to these grasslands also depressed methane uptake. The amount of organic matter present, the fertility of the soil, and other factors such as changes in soil physical properties and soil microbiological properties appear to play an important role in CH₄ oxidation (Mosier et al. 1991).

The mechanisms by which CH₄ oxidation potential is reduced following conversion to cropland or change in N dynamics are not yet clear. Factors such as water availability, fertilizer applications, atmospheric N deposition, soil structural changes, and cropping management contribute to modifications of the soil CH₄ uptake. In particular, simultaneous measurements of N₂O and CH₄ fluxes in a semiarid grassland steppe indicated an inverse relationship between N₂O emissions and CH₄ uptake under conditions of varying soil moisture (Mosier et al. 1991). The pattern of N₂O efflux relative to rainfall is similar to the pattern of Nmineralization in grassland ecosystems (Parton et al., 1988; Schimel et al., 1988) and conforms with the observed role N-fertilizers plays on reducing CH₄ uptake by soils (Steudler et al., 1989).



Figure 1. Effect of N-fertilization ("Native+N"), past cultivation ("Go Back" returned to grass in 1939), and current cropping ("Cropped") on uptake of atmospheric methane in shortgrass steppe soils ("Native").

GLOBAL PERSPECTIVE ON THE SOIL SINK FOR CH.

Estimates indicate that the global oxidation of CH₄ is exceeded by production by about 50 Tg each year, or approximately 10% of the total source. Because 80 to 90% of this oxidation of CH₄ occurs in the atmosphere, what is the role of soils in the global CH₄ budget? Measurement and modeling of the twodimensional distribution of CH₄ concentration in an unsaturated zone near buried waste (Striegl and Ishii, 1989) demonstrated that *in situ* fates of CH₄ consumption increase in response to elevated CH₄ concentrations in the soil atmosphere. Laboratory measurements of the concentration dependence of oxidation rates indicate that potential microbial CH₄ consumption far exceed potential CH₄ diffusion rates (Whalen and Reeburgh, 1990; Bender and Conrad, this issue; Striegl, this issue), and soil temperature has a minor effect on soil uptake of atmospheric CH₄ (Born et al., 1990, Mosier et al., 1991, Whalen and Reeburgh, 1990). Assuming that soil oxidation rates are CH₄ concentration dependent, as these studies suggest, then the soil CH₄ uptake has probably been increasing with the sharp rise of the atmospheric CH₄ since 1850.

In our analysis we assumed (after Striegl and Ishii, 1989; Fung et al., 1991) that current CH₄ uptake occurs by ordinary diffusion and its rate is proportional to the difference between CH₄ mixing ratios above and below the soil surface, i.e.,

Annual CH_4 Uptake = $k \times \Delta[CH_4]$

[1]

Minimum soil CH₄ concentrations are clearly variable, but measurements in several ecosystems suggest that a reasonable minimum soil CH₄ concentration would be 0.4 ppmv (Striegl, 1988; Striegl and Ishii, 1989). Because diffusion is proportional to [air-filled porosity]^{1/3}, we also assumed that broad patterns of soil moisture in 1850 were not different from current patterns. We solved for k for each of the land cover types identified in Table 1, based on the current atmospheric CH₄ concentration and observed annual uptake rates. Using the calculated k's, estimates for land-cover-specific CH₄ uptake at atmospheric CH₄ levels in 1850, 1900, 1950, and 1980 were made.

During the past 150 y, rapid changes to land cover due to conversion of natural areas to croplands and for fuel wood extraction has occurred. Using data derived from the World Resources 1987 Report (WRI 1987), we estimated land-cover changes in the mid-latitudes during the past century from North Africa and Middle East, North America, China, South Asia, Europe, and the former USSR (Table 1). New croplands between 1850 and 1950 in these temperate regions were derived primarily from grassland conversion. In the USSR and China during this time, forest conversion was also a significant contributor to cropland expansion. In North America and Europe, reversion of cropland back into forest and grassland has occurred during the past 80 y, and over the whole temperate region the amount of land cover change was approximately 1000 10⁶ ha or about 14% of the temperate forest and grassland area being considered in these six regions.

In order to assess the effect of changing atmospheric concentrations on the soil sink in combination with land cover changes, we made a retrospective analysis of CH_4 uptake in temperate soils since 1850 under three land cover scenarios. These scenarios are a "no-disturbance" (i.e., area of land cover have maximal consumption rates of undisturbed ecosystems), "intensive" disturbance by land cover change only (i.e., consumption rates are reduced in areas that are under a different land cover class due to cropland conversion), and a combined "intensive plus extensive" disturbance which includes land use changes as well as anthropogenic changes in N deposition. This last scenario probably best represents actual changes in CH_4 uptake since 1850.

For the "no-disturbance" scenario, we assumed that pristine forests and grasslands would take up CH_4 at or near the maximum rates currently observed in these systems, and that land cover changes from the pristine condition had not occurred. Uptake rates were calculated as

$$CH_4 Uptake_u = k_u \times \Delta [CH_4] \times Area_u$$
[2]

where k_u equals current "undisturbed" uptake relative to current Δ [CH₄]. Assumptions for the "intensive" land cover change scenario were similar except that land cover changes (derived from WRI 1987) decreased CH₄ uptake rates in affected areas, and uptake rates were calculated as

$$CH_{4} Uptake_{i} = CH_{4} Uptake_{u} - (k_{u} - k_{i}) \times \Delta [CH_{4}] \times \Delta Area_{i}$$

$$[3]$$

Table 1. Estimates of historic CH4 mixing ratios (Houghton et al. 1990), temperate forest and grassland cover patterns (WRI 1987), area impacted by chronic N deposition (based on Melillo et al. 1989), and consequent CH4 uptake rates.

Line			_	_	-				1		T	- T		T		1	
		Extensive		1.14	\$.	1.42	1.99		3.70	0.64		0.80	1 10		2.09		
	take	Intensive		0.68	100	c8.0	1.18		2.20	0.38		0.48	500	0.0/	1.24		
	Annual CH4 up	Undisturbed	(kg CH ₄ ha ⁻¹)	1.69	-	2.12	2.96		5.50	0.95		1.19		1.67	3 10	2110	
		<u> </u>	R.			2.85						1 60	3				
	lake ents		R.		1.69							0.95					
	CH, up	coeffic	k	2	4.24						2.38				—		
	Extensive	impact area			0	265	500	2	1323	-	0	195		781		1953	
	Land	cover	(106 ha)		C407	2537	1383	C0C7	2302		3906	3875	2022	3732		3675	
	E.	mixing	lauo Viinni	(Amidd)	0.8	0.0		1.1	17		0.8	00	6.0	1 1		1.7	
	Vear	}			1850	1900	1050	NCKI	1080	00/1	1850	Ş	MAT	1050		1980	
	Biome				Temperate Forest						Temperate	Grassland					

developed the best of a first and state of the basis is a second second shifting the basis of the basis

Ĵ

where k_i is 0.4 k_u and area, is the area that underwent cropland conversion. The k_i value is based on estimates of cultivation effects in a grassland region which reduced CH₄ uptake rates to 25 to 40% of the non-cultivated soils (i.e., "native" grassland soil, Figure 1, Mosier et al., 1991) and we assumed the less severe impact. Finally, we assumed in the combined scenario that half of the "native" temperate forests and grasslands were in fact not pristine and therefore were taking up CH₄ at rates near the minimum rates reported. This was based on Melillo *et al.*'s (1989) estimate that half of the temperate land area receives 80% of the N burden and has reduced CH₄ uptake of these soils by 33% (Stuedler et al., 1989). This uptake rate was calculated as

$$CH_4 Uptake_{i,*} = CH_4 Uptake_i - (k_u - k_e) \times \Delta [CH_4] \times \Delta Area_e$$

$$[4]$$

where k_{e} is 0.67 k_{u} .

This analysis indicates that the magnitude of the soil sink since 1850 has tripled (Figure 2). This increase in soil uptake has resulted from the increase in atmospheric CH₄ during this period. Also, during the past 150 years, areas have undergone intensive land cover changes due to forest or grassland conversion to cropland or pasture. This cropland conversion is estimated to have resulted in a reduction of approximately 2 Tg y⁻¹ in the soil CH₄ sink (Figure 2).

In addition to the conversion of native land cover types to croplands, extensive changes have also occurred that are not reflected in data on land cover changes alone. These include increased N deposition in eastern North America, Europe, and China, as well as changes in forestry practices involving disturbance by compaction, disking (plowing), biomass burning, and N fertilization. The impact of these extensive changes would lower the soil sink even more during the past 150 years. The net effect of intensive land cover disturbance and extensive chronic disturbance to terrestrial ecosystems resulted in a 7 Tg y⁻¹ reduction from the "no-disturbance" CH₄ sink to 20 Tg y⁻¹ (Figure 2). These reductions in soil sink strength will diminish the capacity of soils to act as a negative feedback mitigating future increases in atmospheric CH₄ concentration.

SUMMARY

The global CH₄ budget is affected by processes controlling both the production and consumption of CH₄. Much attention has been focussed on the factors related to land cover and land use changes affecting CH₄ production (Cicerone and Oremland, 1988; Aselmann and Crutzen, 1989) and on factors related to atmospheric oxidation of CH₄ (Crutzen and Gidel, 1983; Vaghjiani and Ravishankara, 1991). However, recent studies have documented the widespread nature of soil CH₄ consumption and have begun to question the validity of the current global CH₄ budgets.

A severe constraint on regional or global assessments of the terrestrial CH₄ sink is the paucity of measurement sites and the infrequency of measurements from which to base CH₄ budgets. The measurements used for annual estimates from particular ecosystems are rarely collected over a sufficient period of time or area to capture spatial or seasonal variability of gas efflux or uptake from a particular area. The interaction of climate variability, microbial community and ecosystem dynamics, and land surface characteristics make



Figure 2. A retrospective estimate of the global CH₄ soil sink relative to changes in atmospheric methane from 1850 to 1980. The "No disturbance" line indicates estimated soil CH₄ uptake assuming that no major land cover changes have taken place and that uptake rates were near maximal. The "Intensive" line represents estimated soil CH₄ uptake assuming that soil CH₄ uptake was altered by land cover changes resulting from forest or grassland conversion to cropland or pasture. The "plus extensive" line represents the added change in CH₄ uptake due to increased atmospheric inputs of N over large areas of the world.

quantitative estimation of the flux of trace gases very difficult. Further, the processes resulting in change over time in soil CH, oxidation are not well known. Yet, as with many trace gas species, only scanty data are available to extrapolate fluxes over large areas and to estimate how they may change in time. While this extrapolation is prone to large estimation errors, cross-checks between surface flux inventories and inverse calculations provide bounds on uncertainties and serve to guide future studies.

In this study, we have assessed the potential CH₄ sink from temperate soils. The magnitude of this sink in the temperate region considered is approximately 27 Tg y⁻¹. However, with changes in land use and other factors affecting biogeochemical rates in these systems, we estimate that this sink has been reduced by human-induced changes by 1.5 to 6 Tg y⁻¹, depending on the nature and the extent of the disturbance. The most reasonable estimate of the current temperate soil sink is 20 Tg y⁻¹, which is approximately 30 to 50% of the global soil sink for CH₄. And without this CH₄ sink, atmospheric CH₄ concentrations would be increasing about 1.5 times the current rate.

These changes in soil consumption of CH, are not only occurring in the temperate region, but have been noted in other regions as well. For instance, in a tropical forested area, Keller et al., (1990) observed a 75% decrease in methane uptake rates in pastures and tropical agricultural sites compared to forest sites. In a ropical forest recovering from a hurricane, Steudler (pers. commun., 1991) observed a 50% decrease in methane uptake during the initial stage of recovery and has related this to the accelerated N cycling. Some fundra (Whalen and Reeburgh, 1990) and taiga (Whalen et al., 1991) ecosystems may also be significant CH, sinks, but the magnitude of this sink is uncertain due to the role of human and climate change impacts on the biogeochemical processes controlling CH, production and consumption. The potential influence of these extensive changes in land cover in the tropics during the past century on the global CH, budget needs further study.

L. .

翥

LITERATURE CITED

Aselmann, I. and P.J. Crutzen (1989) Global distribution of natural freshwater wetlands and rice paddies, their net primary productivity, seasonality and possible methane emissions, <u>J. Atmos. Chem. 8</u>, 307-358.

Bender, M. and R. Conrad (in press) Kinetics of methane oxidation in soils. Chemosphere (in press).

Bolle, H.J., W. Seiler and B. Bolin (1986) Other greenhouse gases and aerosols. Assessing their role in atmospheric radiative transfer. <u>The Greenhouse Effect, Climatic Change and Ecosystems</u>, B. Bolin and B.R. Doos, Editors (Scope Vol. 29. Wiley and Sons, New York).

Born, M., H. Dorr and I. Levin (1990) Methane concentration in aerated soils in West Germany. <u>Tellus B 42</u>, 2-8.

Cicerone, R.J. and R.S. Oremland (1988) Biogeochemical aspects of atmospheric methane. <u>Global</u> <u>Biogeochem. Cycles 2(4)</u>, 299-327.

Crutzen, P.J. and L.T. Gidel (1983) A two dimensional model of the atmosphere 2: The tropospheric budgets of anthropogenic chlorocarbons, CO, CH, CH₃Cl and the effect of various NOx sources on tropospheric ozone. J. Geophys. Res. 88, 6641-666.

Crutzen, P.J., I. Aselmann and W. Seiler (1986) Methane production by domestic animals, wild ruminants, other herbivorous fauna, and humans. <u>Tellus 38B</u>, 271-284.

Fung, I., J. John, J. Lerner, E. Mathews, M. Prather, L.P. Steele and P.J. Fraser (1991) Three-dimensional model synthesis of the global methane cycle. <u>J. Geophys. Res. 96(D7)</u>, 13033-13065.

Hansen, J., A. Lacis and M. Prather (1989) Greenhouse effect of chlorofluorocarbons and other trace gases. <u>J.</u> <u>Geophys. Res. 94</u>, 16417-16421.

Houghton, J.T., G.J. Jenkins, and J.J. Ephraums, editors (1990) Climate change. <u>The International Panel on</u> <u>Climate Change (IPCC) Scientific Assessment</u>. World Meteorological Organization (WMO), Cambridge University Press, Cambridge, England.

- Houghton, R.A., J.E. Hobbie, J.M. Melillo, B. Moore, B.J. Peterson, G.R. Schaver and g.M. Woodwell (1983)
 Changes in the carbon content of terrestrial biota and soils between 1860 and 1980: A net release of
 CO, to the atmosphere. Ecol. Monogr. 53, 235-262.
- Keller, M., T.J. Goreau, S.C. Wofsy, W.A. Kaplan and M.B. McElroy (1983) Production of nitrous oxide and consumption of methane by forest soils. <u>Geophys. Res. Let. 10(12)</u>, 1156-1159.
- Keller, M., W.A. Kaplan and S.C. Wofsy (1986) Emissions of N₂O, CH₄ and CO₂ from tropical forest soils. <u>J.</u> Geophys. Res. 91(D11), 11791-11802.
- Keller, M., M.E. Mitre and R.F. Stallard (1990) Consumption of atmospheric methane in soils of central Panama: Effects of agricultural development. <u>Global Biogeochem. Cycles 4</u>, 21-27.
- Khalil, M.A.K. and R.A. Rasmussen (1983) Sources, sinks and seasonal cycles of atmospheric methane. <u>J.</u> Geophys. Res. 88, 5131-5144.
- Khalil, M.A.K., R.A. Rasmussen, M.J. Shearer and M.J. Shearer (1989) Trends of atmospheric methane during the 1960s and 1970s. J. Geophys. Res. 94(D15), 18279-18288.
- Luizao, F., P.A. Matson, G. Livingston, R. Luizao, and P.M. Vitousek (1989) Nitrous oxide flux following tropical land clearing. <u>Global Biogeochem. Cycles 3</u>, 281-285.
- Melillo, J.M., P.A. Steudler, J.D. Aber and R.D. Bowden (1989) Atmospheric deposition and nutrient cycling. <u>Exchange of Trace Gases Between Terrestrial Ecosystems and the Atmosphere</u>, M.O. Andreae and D.S. Schimel, Editors (Berlin, February 1989, pp. 263-280).
- Mosier, A.R., W.D. Guenzi and E.E. Schweizer (1986) Field denitrification estimation by nitrogen-15 and acetylene inhibition techniques. Soil Sci. Soc. Am. J. 50, 831-833.
- Mosier, A.R., D. Schimel, D. Valentine, K. Bronson and W.J. Parton (1991) Methane and nitrous oxide fluxes in native, fertilized, and cultivated grasslands. <u>Nature 350</u>, 330-332.
- Parton, W.J., A.R. Mosier and D.S. Schimel (1988) Rates and pathways of nitrous oxide production in a shortgrass steppe. <u>Biogeochemistry 6</u>, 45-58.
- Rasmussen, R.A. and M.A.K. Khalil (1981) Atmospheric methane (CH₄): Trends and seasonal cycles. <u>J.</u> <u>Geophys. Res. 86</u>, 9826-9832.
- Rodhe, H. (1990) A comparison of the contribution of various gases to the greenhouse effect. <u>Science 248</u>, 1217-1219.
- Schimel, D.S., S. Simkins, T. Rosswall, A.R. Mosier and W.J. Parton (1988) Scale and the measurement of nitrogen-gas fluxes from terrestrial ecosystems. <u>Scales and Global Change</u>. T. Rosswall, R.G. Woodmansee and P.G. Risser, Editors, Chapter 10 (John Wiley & Sons, Ltd.).
- Schutz, H., W. Seiler and H. Rennenberg (1990) Soil and land use related sources and sinks of methane (CH.) in the context of the global methane budget. <u>Soils and the Greenhouse Effect</u>, A.F. Bouwman, Editor (John Wiley & Sons, pp. 270-285).
- Seiler, W., R. Conrad and D. Scharffe (1984) Field studies of methane emission from termite nests into the atmosphere and measurement of methane uptake by tropical soils. J. Atmos. Chem. 1, 171-186.

- eele, L.P., P.J. Fraser, R.A. Rasmussen, M.A.K. Khalil, T.J. Conway, A.J. Crawford, R.H. Gammon, K.A. Masarie and K.W. Thonig (1987) The global distribution of methane in the troposphere. <u>J. Atmos.</u> Chem. 5, 125-171.
- Steudler, P.A., R.D. Bowden, J.M. Melillo and J.D. Aber (1989) Influence of nitrogen fertilization on methane uptake in temperate forest soils. <u>Nature 341</u>, 314-316.
- beudler, P.A., J.M. Melillo, R.D. Bowden, M.S. Castro and A.E. Lugo (1991) The effects of natural and human disturbances on soil nitrogen dynamics and trace gas fluxes in a Puerto Rican wet forest. <u>Biotropica 23(5a)</u>, 356-363.
- riegl, R.G. (1988) Distribution of gases in the unsaturated zone at a low-level radioactive-waste disposal site near Sheffield, Illinois. <u>U.S. Geol. Surv. Water Resour. Inv. Rep. 88-4025, 69 p</u>.
- triegl, R.G. and A.L. Ishii (1989) Diffusion and consumption of methane in an unsaturated zone in northcentral Illinois, USA. <u>J. Hydrology 111</u>, 133-143.
- iegl, R.G. (1992) Diffusional control of methane oxidation in soils. Chemosphere (this issue).
- nompson, A.M. and R.J. Cicerone (1986) Possible perturbations to atmospheric CO, CH₄, and OH. <u>J.</u> Geophys. Res. 91, 10853-10864.
- Vaghjiani, G.L. and A.R. Ravishankara (1991) New measurements of the rate coefficient for the reaction of OH with methane. <u>Nature 350</u>, 406-408.
- Juebbles, D.J. and J. Edmonds (1991) Primer on Greenhouse Gases. (Lewis Publishers, pp. 230).
- Whalen, S.C. and W.S. Reeburgh (1990) Consumption of atmospheric methane by tundra soils. <u>Nature 346</u>, 160-162.
- Whalen, S.C., W.S. Reeburgh and K.S. Kizer (1991) Methane consumption and emission by taiga. <u>Global</u> <u>Biogeochem. Cycles 5</u>, 261-273.
- World Resources Institute (1987) World Resources. An assessment of the resource base that supports the global economy. <u>World Resources Institute/International Institute for Environment and Development</u>, (Basic Books, New York).
- Yavitt, J.B., D.M. Downey, G.E. Lang and A.J. Sexstone (1990) Methane consumption in two temperate forest soils. <u>Biogeochemistry 9</u>, 39-52.

OF POOR QUALITY