

Open access • Journal Article • DOI:10.1016/S0012-8252(01)00062-9

Atmospheric methane and global change — Source link []

Donald J. Wuebbles, Katharine Hayhoe

Institutions: University of Illinois at Urbana–Champaign

Published on: 01 May 2002 - Earth-Science Reviews (Elsevier)

Topics: Atmospheric methane, Methane, Greenhouse gas, Atmospheric carbon cycle and Enhanced coal bed methane recovery

Related papers:

- Climate change 2007: the physical science basis
- Changing concentration, lifetime and climate forcing of atmospheric methane
- Methane emissions from lakes: Dependence of lake characteristics, two regional assessments, and a global estimate
- Biogeochemical aspects of atmospheric methane
- · Contribution of anthropogenic and natural sources to atmospheric methane variability





Earth-Science Reviews 57 (2002) 177-210



www.elsevier.com/locate/earscirev

Atmospheric methane and global change

Donald J. Wuebbles*, Katharine Hayhoe

Department of Atmospheric Sciences, University of Illinois, Urbana, IL 61801, USA

Received 21 March 2000; accepted 4 April 2001

Abstract

Methane (CH₄) is the most abundant organic trace gas in the atmosphere. In the distant past, variations in natural sources of methane were responsible for trends in atmospheric methane levels recorded in ice cores. Since the 1700s, rapidly growing human activities, particularly in the areas of agriculture, fossil fuel use, and waste disposal, have more than doubled methane emissions. Atmospheric methane concentrations have increased by a factor of 2-3 in response to this increase, and continue to rise. These increasing concentrations have raised concern due to their potential effects on atmospheric chemistry and climate. Methane is important to both tropospheric and stratospheric chemistry, significantly affecting levels of ozone, water vapor, the hydroxyl radical, and numerous other compounds. In addition, methane is currently the second most important greenhouse gas emitted from human activities. On a per molecule basis, it is much more effective a greenhouse gas than additional CO₂. In this review, we examine past trends in the concentration of methane in the atmosphere, the sources and sinks that determine its growth rate, and the factors that will affect its growth rate in the future. We also present current understanding of the effects of methane on atmospheric chemistry, and examine the direct and indirect impacts of atmospheric methane on climate. © 2002 Elsevier Science B.V. All rights reserved.

Keywords: methane; greenhouse gases; atmospheric chemistry; global change

1. Introduction

Methane (CH₄) is the most abundant organic trace gas in the atmosphere. Concentrations¹ of CH₄ have more than doubled since pre-industrial times, with a current globally-averaged mixing ratio of ~1750 ppbv² (NOAA CMDL data, Dlugokencky, personal communication). The effects of CH₄ on climate and atmospheric chemistry are the reason for concern over its high growth rate. Following water vapor and carbon dioxide (CO₂), methane is the most abundant greenhouse gas in the troposphere. Both on a molecule and a mass basis, additional methane is actually much more effective as a greenhouse gas than additional CO₂. Methane is also the most abundant reactive trace gas in the troposphere and its reactivity is important to both tropospheric and stratospheric chemistry. The oxidation of CH₄ by hydroxyl (OH) in the troposphere leads to the formation of formaldehyde (CH₂O), carbon monoxide (CO), and ozone (O₃), in the presence of sufficiently high levels of nitrogen oxides (NO_x). Along with CO, methane helps control the amount of OH in the troposphere. Methane also

^{*} Corresponding author. Tel.: +1-217-244-1568; fax: +1-217-244-4393.

E-mail address: wuebbles@atmos.uiuc.edu (D.J. Wuebbles).

¹ Concentration for methane as used here is actually volume mixing ratio (or mole fraction).

² ppbv=Parts per billion (10^9) by volume.

affects the concentrations of water vapor and ozone in the stratosphere, and plays a key role in the conversion of reactive chlorine to less reactive HCl in the stratosphere.

The purpose of this paper is to examine the past trends in the concentration of methane, the sources and sinks affecting its growth rate, and the factors that could affect its future growth rate. This study also examines the current understanding of the effects of methane on atmospheric chemistry and on climate.

2. Record and budget of atmospheric CH₄

2.1. Distribution and trend

2.1.1. Atmospheric CH_4 in the distant past

Air bubbles trapped in polar ice provide a continuous record of the atmospheric concentration of methane (e.g., Legrand et al., 1988; Chappellaz et al., 1990; Etheridge et al., 1992, 1998; Jouzel et al., 1993; Nakazawa et al., 1993; Raynaud et al., 1993; Blunier et al., 1995, 1998; Brook et al., 1996). Ice cores from Greenland and Antarctica now extend the record of atmospheric CH_4 , CO_2 and temperature back to 420,000 years BP (Petit et al., 1999), covering four glacial-interglacial cycles.

The record shown in Fig. 1 reveals two important characteristics of past CH₄ concentrations with implications for future climate. First, present-day CH₄ levels appear to be unprecedented during any time in the past. Ice core records show glacial-interglacial transitions rising from 320-350 ppbv up to a maximum of 650-780 ppbv (Petit et al., 1999), corresponding to source increases as large as 50-60 TgCH₄/year (Brook et al., 1996, based on Chappellaz et al., 1993a). This is less than half of present-day concentrations, which are currently greater than 1700 ppbv. Secondly, CH₄ is closely tied to atmospheric temperature records, decreasing and rising in phase with temperature at the inception and termination of glacial episodes (e.g., Petit et al., 1999; Chappellaz et al., 1993a,b; Raynaud et al., 1988).

The observed synchronicity between temperature and CH_4 concentrations has raised the question of the role of CH_4 in glacial–interglacial climate change. Do increasing CH_4 emissions actually help initiate climate change, or do they respond quickly to rising

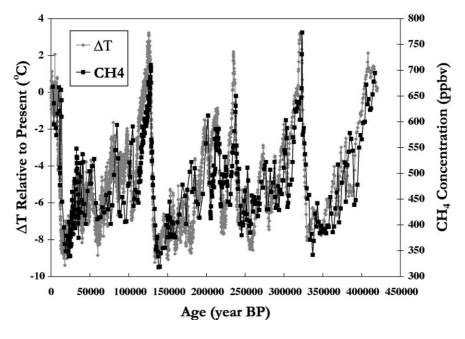


Fig. 1. Vostok ice core records of atmospheric temperature and methane concentration (volume mixing ratio) from 420,000 years BP to present (source: Petit et al., 1999).

atmospheric temperatures? Theories of CH₄ as an instigator of glacial terminations point to the potential destabilization of methane clathrates contained in sediments on continental shelves, as sea level fell due to growing ice sheets. This destabilization could have led to abrupt large-scale releases of CH4 into the atmosphere, capable of inducing a global temperature increase of up to $1-2^{\circ}C$ over a few decades (Dickens et al., 1997; Thorpe et al., 1996). Such a hypothesis has been shown to be consistent with the geological record of observed perturbations in carbon isotopes (Dickens et al., 1997; Bains et al., 1999), and is not inconsistent with the absence of an observed spike in CH₄ concentrations from ice cores with a 300-year sampling interval³ (Thorpe et al., 1996). However, results from several ice core studies support the hypothesis that increased emissions from low-latitude, followed by higher latitude, wetlands in response to climate change were responsible for the majority of CH₄ increases (Chappellaz et al., 1993a,b; Thompson et al., 1993; Blunier et al., 1995; Brook et al., 1996). The importance of wetland emissions is emphasized by an observed drop in CH₄ concentrations during the mid-Holocene, when tropical lakes may have dried (Blunier et al., 1995), and interactions between climatic influences and ecosystem dynamics would have reduced wetland emissions (Velichko et al., 1998). A recent modeling study (Dällenbach et al., 2000) also finds large changes in northern latitude sources during warm Dansgaard/Oeschger events during the last glacial period.

An innovative method for extracting temperature data from ice cores has recently shown that CH_4 increases at the end of the Younger Dryas cold period began somewhere between 0 and 30 years following an abrupt temperature increase (Severinghaus et al., 1998). A successive study of four rapid climate transitions over the past 50,000 years reveals that following each transition, temperature increased more rapidly than methane concentrations (Brook et al.,

2000). The fact that the majority of CH_4 increase lagged the warming thus seems to rule out the primarily causal role of CH_4 in glacial-interglacial climate change.

2.1.2. Pre-industrial to present concentrations

Measurements from polar ice cores over the past 1000 years reveal several notable features in global CH₄ concentrations. These include: (1) variations of 10–15 ppbv about a pre-industrial level of 700 ppb; (2) a pole-to-pole gradient of 30-60 ppb increasing to around 150 ppb in this century, indicating consistently larger emissions in the Northern Hemisphere, but variable tropical emissions; (3) a continued correlation with temperature during the Little Ice Age around the middle of the millenium; and (4) a significant and monotonic increase beginning between 1750 and 1800 (Etheridge et al., 1998; Chappellaz et al., 1997; Blunier et al., 1993; Nakazawa et al., 1993; Khalil and Rasmussen, 1987; Craig and Chou, 1982). This increase is confirmed by glacial ice cores from the early 1800s (e.g., Dibb et al., 1993), and solar spectra from 1950 (e.g., Rinsland et al., 1985). As seen in Fig. 2, ice core records and observations from ground-based stations show that the global level of methane has more than doubled since preindustrial times, and continues to increase (Rasmussen and Khalil, 1981; Blake and Rowland, 1988; Steele et al., 1992; Dlugokencky et al., 1994a, 1995, 1998; scientific reviews can be found in IPCC, 1995, 1996).

Numerous modelling studies (e.g., Khalil and Rasmussen, 1985; Thompson, 1992; Crutzen and Bruhl, 1993; Khalil and Rasmussen, 1994b; Osborn and Wigley, 1994; Crutzen, 1995; Brasseur et al., 1998; Lelieveld et al., 1998; Wang et al., 1998) show a decline in the hydroxyl radical sink for CH₄ from pre-industrial times to the present. Modelled OH decreases center around 10-30%, depending on the model used, with the range also caused by large uncertainties in past emissions of other trace gases such as CO, NO_x and NMHCs. This range, combined with analyses of past emissions (Khalil and Rasmussen, 1985, 1994a; Lelieveld et al., 1993; Subak, 1994; Stern and Kaufmann, 1996), shows that the majority of the observed increase in methane since pre-industrial times is due to the increase in emissions from anthropogenic sources, as shown in Fig. 3.

³ Detailed temporal resolution of changes in temperature and CH_4 concentrations has been complicated by the fact that air trapped in ice and snow continues to be influenced by atmospheric concentrations until sealed into impermeable bubbles. This produces a difference between gas age and ice age that can range from 30 years up to 7000 years, depending on the site's accumulation rate (Bender et al., 1997).

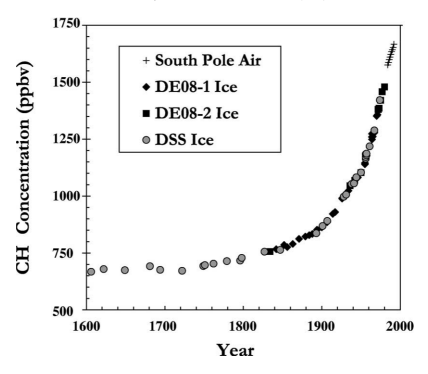


Fig. 2. Ice core and atmospheric records of the increase in atmospheric methane from pre-industrial times to the present (source: Etheridge et al., 1998).

Khalil and Rasmussen (1987, 1994a) find good agreement between increasing population, agricultural emissions, and the observed rise in emissions over the past 200 years, and it is also likely that increasing

fossil fuel use has contributed significantly (Etheridge et al., 1992, 1998; Dlugokencky et al., 1994a). However, studies of human populations and activities over the past millenium suggest that discernable anthropo-

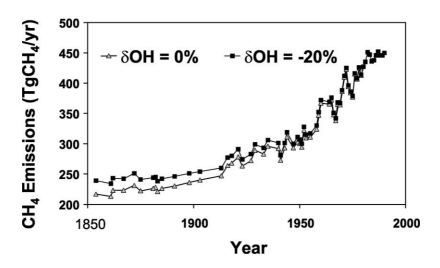


Fig. 3. Increase in methane emissions from pre-industrial times to the present. Based on a top-down analysis by Khalil and Rasmussen (1994a).

genic influence on CH_4 may extend back considerably further than the beginning of the pre-industrial age (Subak, 1994; Blunier et al., 1993).

2.1.3. Decadal changes in methane growth rates

The rate of increase for methane throughout the late 1970s averaged 20 ppbv/year (Blake and Rowland, 1988). However, during the 1980s and 1990s, the rate of methane increase slowed steadily, by an average of -1 ppb/year/year (Fig. 4). Continuous monitoring of methane trends in ambient air from 1979 to 1989 indicated a decreasing trend from 20 to just over 10 ppbv, with an average growth rate of 8.9 ppbv/year from 1984 to 1996 (Dlugokencky et al., 1994a, 1998; Khalil and Rasmussen, 1993, 1994b; Steele et al., 1992). When the data were examined for regional and seasonal effects, it was found that the slowing in the rate of increase is higher between 30 and 90°N (Dlugokencky et al., 1998; Steele et al., 1992), but that the amplitude of the seasonal cycle has exhibited no significant global trends over this time, with typical amplitudes around 30 ppb in the high

Southern latitudes, 60 ppb in the high Northern latitudes, and only a small change in the tropics (Khalil et al., 1993a; Dlugokencky et al., 1994a, 1997).

Following the 1992–1993 anomaly (discussed in the following section), the rate of methane increase continued to fall to a value of 3 ppbv/year in 1996, with a globally-averaged atmospheric concentration of 1730 ppbv and an interhemispherical gradient of 140 ppbv (Dlugokencky et al., 1994a, 1998). If the current decrease in trend had continued, zero growth was expected to occur before 2006, as concentrations slowly stabilized around 1800 ppbv (Etheridge et al., 1998; Dlugokencky et al., 1998).

The long-term slowing in the rate of CH_4 increase over the last few decades must be due to either an increase in methane removal, or declining methane emissions. Observed trends in the seasonal cycle of methane and global OH concentrations inferred from methylchloroform measurements do not support a significant change in OH over the past decades (Dlugokencky et al., 1997; Prinn et al., 1995). Similarly,

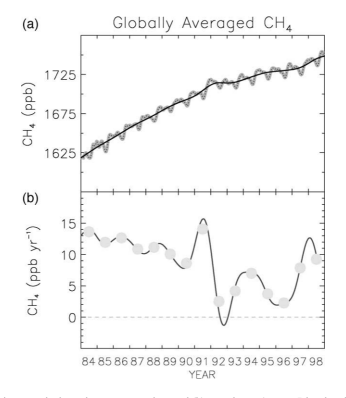


Fig. 4. (a) Observed increase in atmospheric methane concentrations and (b) growth rate (source: Dlugokencky, personal communication).

modelling studies investigating the impact of observed stratospheric ozone depletion and temperature change over the past decade found that, although the resulting increase in OH is compatible with the observed decrease in trend, by themselves these two factors are insufficient to account for the entire decrease in growth rate (Bekki and Law, 1997; Fuglestvedt et al., 1994). Isotopic analysis shows the growth rate of δ^{13} C over that time to be inconsistent with a substantial increase in the CH₄ sink (Etheridge et al., 1998). It appears, then, that decreasing changes in emissions rather than increasing sink strengths must be responsible for the major part of the observed slowing.

In the absence of any mechanism to explain a longterm decrease in natural sources such as wetlands, the answer appears to lie with anthropogenic emissions. Khalil and Rasmussen (1994a,b) examine the changing relationship between population and agricultural sources, thought to be a major contributor to CH₄ increases from pre-industrial to present times, to show that the growth in agricultural sources may be slowing. Steele et al. (1992) suggest that human sources, particularly those sources which can be reduced quickly, may be the cause of the observed rapid deceleration of the growth rate in the Northern relative to the Southern Hemisphere. Although the exact cause of this long-term decline in the global methane rate is still unresolved, Dlugokencky et al. (1998) suggested that what had been observed was in fact an approach to steady state, as CH₄ net emissions and OH levels were nearly constant over the past decade.

Optimistic projections of stabilizing atmospheric CH4 levels in the near future have been set back by the latest measurements of CH₄ growth rates that show a jump back up to growth rates of 8-12 ppbv/ year in 1997 and 1998 (Fig. 4, Dlugokencky, personal communication). Although net methane sources may have been relatively constant over the past two decades, it is likely that changes in the distribution of methane sources were occurring during that time, with increases in some sources equalling decreases in others (Dlugokencky et al., 1998). What may be occurring now is an increase in emissions from CH₄ source(s) that is larger than can be balanced by concurrent decrease in other source(s). It has been speculated, supported by ¹³C isotope observations, that the recently observed increase could in fact be the first sign of a biospheric feedback to climate change, as wetland emissions increase in response to the climate warming (Manning, personal communication). Observations of anonymously high temperatures and the lack of an obvious reason for a decrease in the OH sink of methane over this time provide indirect evidence in support of this hypothesis. However, it is equally true that the observed increase in growth rate could also be merely another temporary interannual variation as has been observed before (Dlugokencky, personal communication). Whatever the true cause of this increase, though, it serves at this time to emphasize our limited understanding of the factors that influence methane sources and sinks, and the large uncertainty this introduces in projections of future concentrations.

2.1.4. 1991–1994 drop in growth rates

In 1991, the growth rate of methane increased sharply over a period of 6 months to 1 year following the eruption of Mt. Pinatubo in June. The growth rate then dropped abruptly to very low values and even zero in some locations from 1992 to 1993. Average growth rate in 1992 was only 1.8 ppb/year in the Northern Hemisphere, and 7.7 ppb/year in the Southern Hemisphere (Dlugokencky et al., 1994a,b; 1998), producing a decrease in atmospheric concentrations that has recently been observed in lower stratospheric CH₄ concentrations, which lag 4 years behind the troposphere (Randel et al., 1999). In 1994, global methane growth rates recovered back up to about 8 ppbv/year, before continuing the previously observed long-term decrease (Dlugokencky et al., 1998). A number of theories suggesting reductions in emissions from anthropogenic or natural sources or a slowing in the rate of CH₄ removal have been proposed to explain this short-term anomaly.

Aerosols released by the Pinatubo eruption blocked UV radiation to the troposphere, which may have led to a drop in OH levels and a consequent sharp increase in the methane growth rate. Dlugokencky et al. (1996) show the initial short-lived increased growth rate to be consistent with this theory; however, the causes for the subsequent decrease in the growth rate have not been resolved.

Increases in tropospheric OH have also been proposed as causes for the short-term drop, through mechanisms such as El Niño-induced increases in tropospheric water vapor, an important source of OH (Khalil and Rasmussen, 1993), or aerosol-induced stratospheric ozone depletion, which would cause an increase in tropospheric UV radiation and a subsequent rise in OH formation (Bekki et al., 1994). Enhanced mixing of stratospheric air, with its lower CH_4 levels, has also been proposed to result from increased stratospheric temperatures following the Pinatubo eruption (Schauffler and Daniel, 1994). Reductions in emissions from tropical biomass burning, Northern Hemisphere fossil fuel sources, or wetlands have also been put forward as possible contributors to the brief but abrupt slowing (Lowe et al., 1994, 1997; Dlugokencky et al., 1994).

The ability of these theories to match the observed drop have been constrained by a number of observational and modelling studies comparing source strengths, mixing ratios, and isotopic abundances. Comparison of regional mixing ratios with δ^{13} C isotopic measurements (as discussed in the following section) suggest that changes in atmospheric chemistry or wetland emissions played a minimal role. Instead, observations support a decrease in δ^{13} C-enriched sources such as biomass burning in the Southern Hemisphere or fossil fuel use in the Northern Hemisphere (Lowe et al., 1994, 1997; Gupta et al., 1996). However, using isotopic measurements from six sites, Quay et al. (1999) were unable to distinguish between the possible impacts of changes in biogenic sources, fossil sources, or OH concentration over this time. In addition, it has been shown that a change in the atmospheric CH₄ sink may not be detectable in δ^{13} C for some time (Tans, 1997; Lassey et al., 1999).

Both modelling studies and recent isotopic analyses therefore conclude that several of these mechanisms may have combined to produce the observed drop in growth rates. Estimated changes in individual sources or sinks, with moderate assumptions of their sensitivity to Pinatubo-induced chemistry and temperature changes and to human activities, are individually insufficient to account for the entire decrease (Bekki and Law, 1997; Lelieveld et al., 1998).

2.2. Sources and sinks

2.2.1. Sources

Methane differs from carbon dioxide in that it is released into the atmosphere by a wide variety of sources, both natural and anthropogenic. Anthropogenic emissions arise from biogenic sources related to agriculture and waste disposal, including enteric fermentation, animal and human wastes, rice paddies, biomass burning, and landfills (Fig. 5). Methane is also emitted by the extraction of fossil fuels such as natural gas, coal, and petroleum. Natural methane emissions arise from wetlands, termites, other wild ruminants, oceans, and hydrates (Fig. 6). Based on recent estimates, current human-related biogenic and fossil fuel-related sources for methane are approximately 275 and 100 TgCH₄/year, while total natural sources are around 160 TgCH₄/year (IPCC, 1996).

Due to the variety of methane sources, emissions are affected by numerous factors, including energy use, human population distributions, agricultural practices, and climate. As discussed in the previous section, there is strong evidence that methane emis-

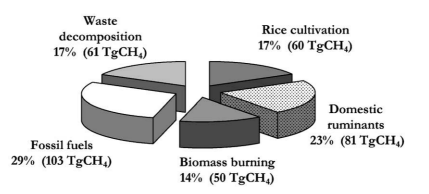


Fig. 5. Contribution of individual sources to total anthropogenic methane emissions (based on Khalil, 2000).

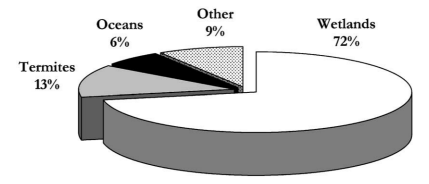


Fig. 6. Contribution of individual sources to total natural methane emissions (based on Khalil, 2000).

sions are changing over time. Here, we briefly describe the main factors influencing the magnitude of emissions from each major source, and how those factors could change under greenhouse gas controls or climate change.

2.2.1.1. Biogenic sources. Methane emissions from biogenic sources result from anaerobic decomposition and reduction of organic material by bacteria-in flooded soils, in landfills or other waste disposal sites, and in the digestive tracts of domestic ruminants such as cattle or sheep, wild ruminants such as buffalo or termites, and even humans. It has been determined that methane production is temperature-dependent, with maximum growth temperature ranging from 37°C to 45°C (Boone, 2000). This temperature dependence is an important factor with implications of future changes for emissions from flooded soils, such as wetlands or rice paddies. It represents an important positive feedback on climate change-as the atmosphere warms, emissions from these sources will increase, raising the concentration of CH₄ in the atmosphere and enhancing the warming.

Emissions from natural wetlands are influenced by a wide range of environmental parameters. These include soil characteristics such as the availability of organic carbon and nutrients (Cao et al., 1996b; Miller et al., 1999; Smith et al., 2000; Yavitt et al., 2000), vegetation type and cover (King et al., 1998; Bellisario et al., 1999; Joabsson et al., 1999; Van der Nat and Middelburg, 2000), and, most importantly, water table depth and soil temperature. A number of studies have demonstrated the high sensitivity of wetland emissions to seasonal and interannual variations in water table and

temperature (e.g., Moosavi and Crill, 1997, 1998; Oberbauer et al., 1998; Van den Pol-Van Dasselaar et al., 1999; Wickland et al., 1999; Nakano et al., 2000; Otter and Scholes, 2000; Worthy et al., 2000; see review in Matthews, 2000). This sensitivity will play a significant role in determining future wetland emissions, as local temperature and precipitation patterns are altered under global climate change. Based on projected temperature increases alone, CH₄ emissions are expected to increase, particularly at high latitudes (Cao et al., 1996b; Matthews, 2000; Worthy et al., 2000). However, the influence of soil moisture and water table level complicates predictions of future wetland emissions. The impact of global climate change on the hydrological cycle at regional levels is highly uncertain; thus, future changes in soil water may result in either increases or decreases in current wetland areas and their emissions (e.g., Wahlen and Reeburgh, 1992; Roulet et al., 1993; Moosavi et al., 1996; Moosavi and Crill, 1997; Van den Pol-Van Dasselaar et al., 1999; Hilbert et al., 2000).

In rice fields, emissions are dependent on the conditions and agricultural practices employed during cultivation, such as climate, soil characteristics, and cultivation practices including water management, fertilizers, other additives, and different varieties of rice plants (e.g., Minami and Neue, 1994; Cao et al., 1995, 1996a; Huang et al., 1997, 1998; Neue et al., 1997; Sass and Fisher, 1997; Khalil et al., 1998; Neue and Roger, 2000). Emissions are also determined by the number of crops per year and the total area under cultivation, which is a function of the demand for rice, and the amount of rice produced per hectare, or 'yield'. Practices such as occasional draining of the

fields, the addition of oxidents or other mineral fertilizers, and selection of low CH₄ cultivars have been found to reduce emissions by approximately 40-55%, 20-70% and up to 60%, respectively (EPA, 1993a,b; Neue et al., 1997; Sigren et al., 1997; Yagi et al., 1997; Butterbachbahl et al., 1997; Cole et al., 1997; Minami, 1997; Sass and Fisher, 1997; Mitra et al., 1999). Alternatively, the addition of organic fertilizers has the potential to increase emissions by over 50% relative to non-organic fertilizers (e.g., Buendia et al., 1997; Denier van der Gon and Neue, 1995; Yagi et al., 1997).

Direct emissions from ruminants, including cattle, sheep, buffalo, goats, and other domestic animals, are affected by diet (Johnson et al., 2000; EPA, 1993a,b; Cole et al., 1997; Harper et al., 1999). Methane emissions are a byproduct of incomplete digestion. In general, higher quality diets will allow animals to digest their food more completely, improving protein uptake while reducing CH₄ emissions. This is particularly true of ruminant diets in developing countries, where improving the relatively poor quality of cattle feed could lead to emission reductions per kg of milk produced of up to 75% (Ward et al., 1993; Crutzen et al., 1986). The addition of production-enhancing supplements to cattle diet has also been suggested as a means to improve digestion and reduce emissions by up to 40% (EPA, 1993a,b; Cole et al., 1997 and references therein). Animal manure is an additional source of methane. If animal waste is allowed to remain in the field, the manure dries quickly and methane emissions are minimal. However, if the waste is pooled and stored, methane emissions can rise by a factor of 10 (Bogner et al., 1995).

Methane is also emitted by the decay of biogenic waste in the anaerobic environments provided by landfills and wastewater pools. Emissions from landfills are affected by numerous environmental and technical variables, including temperature, soil moisture, and CH₄ concentration within the landfill; the amount, organic composition, and age of the waste; and the thickness of the overlying layer. (Bogner and Spokas, 1993; Peer et al., 1993; Czepiel et al., 1996). Emissions from landfills, wastewater, and manure collection lagoons can be reduced by trapping the methane before it is released to the atmosphere, and either flaring it or using it as an energy source. Depending on the capture efficiency, this method can

reduce emissions by over 90% (e.g., Bogner et al., 1995; Borjesson and Svensson, 1997).

Biomass burning releases numerous pollutants into the atmosphere. When combustion is complete, most of the emissions consist of carbon dioxide. However, if the fires smolder and combustion is incomplete, large amounts of CH_4 and other higher-order hydrocarbons can be produced. Methane emissions from biomass burning depend on the stage of combustion reached, as well as on the carbon content of the biomass, and the amount of biomass burned (Levine et al., 2000).

2.2.1.2. Non-biogenic sources. The main source of fossil-fuel related methane emissions are the leaks that occur during natural gas processing, transmission, and distribution, since natural gas is over 90% methane (e.g., Beck et al., 1993; EPA, 1993a). The magnitude of this source depends directly on the loss rate, which has been estimated to average between 1% to 2% of natural gas production for developed countries (GRI, 1997; Beck et al., 1993; EPA, 1993b; Barns and Edmonds, 1990). Uncertainty remains substantial, however, as these values are based mainly on the analysis of gas systems in developed countries where transportation is likely to be more tightly controlled than in less developed nations (Kirchgessner, 2000). Regional leakage percentages have been estimated to range from 1% to 15% of total natural gas production, depending on the quality of the pipelines, the extraction process, leakage control, the method used to estimate gas losses, and other factors (e.g., Mitchell, 1993; Matthews, 1994; Bazhin, 1994; Beck et al., 1993).

Methane is also emitted from coal mines, when gas that has been trapped between layers of coal during its formation is released as the coal is mined. Global and even regional estimates of CH_4 emissions from coal and mines depend on many assumptions regarding the type of coal, the depth of the mine, mining practices, the methane content of the coal seam, and whether methane is flared or released (Beck et al., 1993; Kirchgessner, 2000, and references therein).

Lacking viable sources of alternative energy, methane emissions from coal mining and natural gas use may be more easily reduced by altering mining practices and improving gas leakage controls than by avoiding coal or gas use. If methane emitted from these sources can be trapped, it can be used as an energy source, providing additional benefits.

2.2.2. CH_4 sinks

In contrast to the numerous sources of methane, there are only one major and two minor sinks for tropospheric methane. Reaction with the hydroxyl radical (OH) is responsible for the removal of approximately 500 TgCH₄/year (almost 90% of the total sink), making the concentration of OH the most important determinant of the rate at which methane is removed from the atmosphere. The remainder of the CH₄ is removed through dry soil oxidation, 30 TgCH₄/year (~ 5%), and transport to the stratosphere, estimated at 40 TgCH₄/year (~ 7%) in IPCC (1996), but recently found to be only 27 TgCH₄/year based on data from the Upper Atmosphere Research Satellite (Gettelman et al., 1997).

OH is formed from the photodissociation of tropospheric ozone and water vapor. It is the primary oxidant for most tropospheric pollutants, including carbon monoxide, NO_x species, and organic compounds (e.g, Crutzen, 1995). As the most abundant organic species in the atmosphere, methane plays an influential role in determining the tropospheric oxidizing capacity, initiating an important series of chemical reactions (see discussion in Wuebbles et al., 2000). The oxidation of methane by OH is so effective that almost 90% of methane destruction currently occurs in the troposphere due to these reactions (e.g., IPCC, 1996). The amount of methane removed is constrained by OH levels and the rate constant of the reaction.

With sharp spatial and temporal gradients, and average concentrations of one part per 50 trillion, the actual concentration of OH is difficult to determine, much less its rate of formation or destruction, which depends non-linearly on the other factors mentioned above (e.g., Thompson, 1992; Crutzen, 1995). A few measurements of OH have been made at certain locations using long-path absorption spectroscopy (e.g., Hofzumahaus et al., 1991; Poppe et al., 1995), and confidence in the accuracy of direct measurements is growing (Brune, 1992). However, most estimates of OH concentration have been obtained through measuring the concentration of a tracer molecule such as methylchloroform (CH₃CCl₃) that

can be used to infer the OH concentration with a chemical transport model. Using CH₃CCl₃ data from 1978 to 1994 obtained at five ALE/GAGE stations around the globe, modelling studies deduced the average globally integrated concentration of OH to be 9.7×10^5 (Prinn et al., 1995) and 1.07×10^6 molecules/cm³ (Krol et al., 1998). Over the same time period, the studies by Prinn and Krol obtained a change in OH concentration of $0.0\pm0.2\%$ /year and $0.46 \pm 0.5\%$ /year, respectively. Based on changes in emissions of CO, NO_x, and NMHCs evaluated with a three-dimensional model of the troposphere, Karlsdottir and Isaksen (2000) estimated that tropospheric OH concentrations have increased by about 7% from 1980 to 1996, or 0.43% /year. According to Karlsdottir and Isaksen (2000), the corresponding decrease in atmospheric lifetime of CH₄ when combined with the observed record would imply overall CH₄ emissions increasing at a rate of 0.67% /year over this period.

OH concentrations are affected not only by direct emissions of methane but by its oxidation products, especially carbon monoxide. Methane has an atmospheric lifetime of 8.9 ± 0.6 years based on CH₃CCl₃ analyses and good knowledge of the loss rate with OH (Prinn et al., 1995). This is lower than the previous estimate of 10 years (WMO, 1991, 1995). Although most trace gases have a turnover time which describes the length of time they are present in the atmosphere, methane is removed by OH on an 'adjustment' time scale that exceeds the atmospheric lifetime by approximately 3 years (e.g., Prather, 1994; IPCC, 1996). The feedback factor that describes this relationship, which is the ratio of the adjustment time of methane to its atmospheric lifetime, varies from 1.2 to 1.7, depending on the model used in the evaluation. Comparisons of 2D and 3D models suggest that the factor does not depend strongly on the background atmospheric OH levels (IPCC, 1996).

This discrepancy between adjustment time and chemical lifetime is due to the fact that the hydroxyl radical is removed not only by interactions with CH_4 , but also with its product, CO. CH_4 , CO, and OH are involved in a chemical feedback cycle whereby even a small rise in the background level of methane — due to growing emissions — will reduce OH and produce CO. The carbon monoxide produced by CH_4 oxidation will in turn be oxidized by OH,

causing a further drop in OH levels. This extra decrease in OH results in an increase in the lifetime of methane, thereby enhancing the original perturbation. The result is a positive feedback, whereby increasing methane emissions can lead to an overall decrease in the oxidizing capacity of the troposphere, a slowing in the removal of methane, and a consequent build-up of methane concentrations. Thus, an incremental perturbation in CH_4 will return to an equilibrium concentration at a slower rate than expected from the atmospheric lifetime, even though there may be no apparent measurable change in OH.

2.2.3. Isotopic composition

The isotopic composition of atmospheric methane is determined by specific chemical and physical processes that, when understood, can yield valuable information concerning the strength of methane sources and sinks, and their contribution to changes in atmospheric methane. The most commonly measured isotope of atmospheric methane is ¹³C, which is generally expressed as δ^{13} C, the ratio of 13 C to 12 C relative to a reference standard. The δ^{13} C ratios can be used to differentiate between methane from bacterial sources, which are ¹³C-depleted relative to atmospheric methane, and the ¹³C-enriched methane emitted from non-bacterial sources such as natural gas and biomass burning, which is generally 20-30 % greater (Tyler, 1986; Tyler et al., 1987; Cicerone and Oremland, 1988; Quay et al., 1988; Stevens and Engelkemeir, 1988; Conny and Currie, 1996, and references therein).

The global average atmospheric δ^{13} C ranges from -47.3% to -46.2% (Stevens and Rust, 1982; Stevens, 1993; Quay et al., 1991, 1999; Tyler et al., 1999). Atmospheric δ^{13} C is enriched relative to the global composite of methane sources, which is approximately -53%, based on estimates of source strengths (Quay et al., 1991; Lassey et al., 1999). This δ^{13} C enrichment is due to the kinetic isotope effect (KIE) during the reaction of CH₄ with OH, which preferentially removes 12 C with a fractionation effect at least 1.0054 (Cantrell et al., 1990). Soil oxidation also preferentially removes 12 C, as do reactions with Cl in the stratosphere. Fractionation effects for these sinks are thought to be larger than those for OH (King et al., 1989, Tyler et al., 1994, 2000; Brenninkmeijer et al., 1995; Saueressig et

al., 1995; Crowley et al., 1999; Snover and Quay, 2000).

The radioactive isotope ¹⁴C also provides valuable information on CH₄ sources. ¹⁴C from bomb tests in the 1960s and current ¹⁴C emissions from pressurized light-water reactors enables differentiation between modern (enriched) sources of methane and fossil (depleted) sources. Based on ¹⁴C measurements, fossil sources are estimated to range from 16 to 25% of total CH₄ emissions (Wahlen et al., 1989; Manning et al., 1990; Quay et al., 1991, 1999). Together with the δ^{13} C ratio, this information can be used to deduce the contribution of biomass burning—a modern but non-bacterial source—to the current budget. Estimates center around 10–11% (Wahlen et al., 1989; Fung et al., 1991; Quay et al., 1991; Stevens, 1993).

Systematic δ^{13} C measurements have shown a seasonal cycle that increases with latitude, 180° out of phase with the seasonal change in CH₄ concentrations (Quay et al., 1991, 1999; Lassey et al., 1993; Lowe et al., 1994, 1997; Francey et al., 1999). In the tropical latitudes the cycle may be due to seasonal variations in OH concentrations, whereas at higher latitudes it is thought to be source-driven (Quay et al., 1999). In the Northern Hemisphere, seasonal isotopic variability may be caused by a strong pulse of δ^{13} C-depleted CH₄ emissions together with possible transport from lower latitudes, while seasonal variations in biomass burning are thought to be responsible for the Southern Hemisphere cycle (Quay et al., 1991, 1999; Lassev et al., 1993; Lowe et al., 1994; Tyler, 1999). The role of biomass burning is emphasized in a recent study by Lowe et al. (1999), who find seasonal variations to be divided into three distinct latitude bands, two of whose cycles-tropical and Southern Hemisphere extra-tropical - correlate strongly with seasonal emissions from biomass burning and their transport to higher latitudes.

 δ^{13} C records also indicate long-term trends. Isotopic measurements from ice cores give a pre-industrial atmospheric δ^{13} C of -49.7% (Craig et al., 1988). Together with an estimate of the sink fractionation factor, this implies an increase in the global composite δ^{13} C of approximately 2% since preindustrial times (Quay et al., 1988; Etheridge et al., 1998). A lower δ^{13} C suggests that historical emissions were controlled primarily by natural sources. Comparison of present-day atmospheric isotope ratios with ice core data gives flux estimates for current bacterial emissions around 70% of the total budget (Quay et al., 1991).

Systematic δ^{13} C measurements taken for over two decades at various sites distributed around the globe indicate that the atmosphere is becoming progres-

sively more enriched in ¹³C, as shown in Fig. 7 (Stevens, 1993; Etheridge et al., 1998; Francey et al., 1999; Quay et al., 1999). Studies suggest that the global increase may be source-driven, caused by increases in δ^{13} C-enriched emissions from biomass burning in the Southern Hemisphere, and a possible

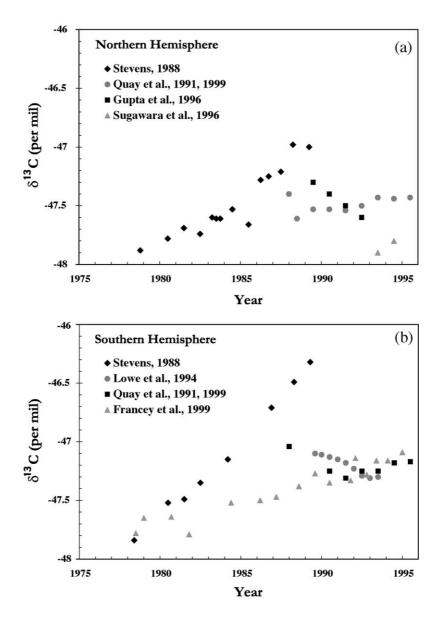


Fig. 7. Measurements of the carbon-13 isotopic composition of atmospheric methane from 1978 to 1995. Hemispherically and annually averaged data was collected at a number of sites in (a) the Northern Hemisphere, and (b) the Southern Hemisphere (Source: Stevens and Wahlen, 2000, and references therein).

decrease in ¹³C-depleted Northern Hemisphere sources such as wetlands and rice paddies (Stevens, 1993). Recent Southern Hemisphere measurements covering the period from 1978 to 1996 indicate any such redistribution in sources to have been relatively small (Francey et al., 1999). Measurements over the same time period also show steady increases in ¹⁴C due to emissions from reactors; however, its growth rate began to slow in 1993 due to a shift away from nuclear power (Quay et al., 1999).

2.2.4. Budget modelling

Uncertainties in sources and sinks of methane are constrained by direct estimates of the individual magnitudes of each source and sink. Source-specific estimates are generally made by multiplying an average emission rate, determined experimentally, by an extrapolant that applies that emission rate over the globe.⁴ Although conceptually simple, this bottomup method contains numerous uncertainties. First, point measurements taken at a limited number of locations and times must be assumed to be representative of global emissions. In the past, this has led to over-estimates of emissions from a number of sources, including rice cultivation (Sass et al., 1999), livestock (Johnson et al., 2000) and possibly wetland emissions (Worthy et al., 2000). It is also primarily responsible for the large ranges in estimates of emissions from other sources such as coal mining (Kirchgessner, 2000), biomass burning (Levine et al., 2000) and waste management (Thorneloe et al., 2000) seen in Table 1. However, as data coverage improves, attention is being re-focused on a second issue, that of resolving temporal and spatial variations in those emissions. Such variations are particularly important for biogenic sources such as wetlands and rice fields whose emissions can vary by several orders of magnitude depending on the location, time of day, season, or year (Matthews, 2000; Shearer and Khalil, 2000). Although these uncertainties continue to produce a range of values in the emission estimates for individual sources (Table 1), the range of estimates for most sources has decreased significantly not only due to the increase in the amount of experimental field data, but

⁴ See Khalil and Shearer (2000) for a full discussion of the methods and constraints of bottom–up emissions estimates.

also due to the application of atmospheric CH_4 data and other types of inventory estimates to the methane budget, as described next.

The methane budget is constrained by the spatial and temporal changes in atmospheric CH_4 mixing ratios and isotopes discussed previously in Section 2.2.3. Isotopic composition of atmospheric methane is a particularly valuable source of information, as it reflects the relative strength of bacterial vs. nonbacterial and modern vs. fossil sources of CH_4 , as well as differentiating between various formation pathways (Cicerone and Oremland, 1988; Conny and Currie, 1996; Whiticar, 2000).

Mass-balance approaches use spatial and regional variations in isotopic composition as well as hemispheric and global averages as input to box models. Atmospheric observations are then compared with the flux-weighted composition of total emissions plus the fractionation effects of sinks, taking into account the effects of atmospheric transport. In this way, the magnitudes of individual sources as well as entire budgets have been estimated (e.g., Khalil and Rasmussen, 1983; Stevens and Engelkemeir, 1988; Cicerone and Oremland, 1988; Tyler, 1989; Lassey et al., 1993, 1999; Lowe et al., 1994, 1997; Quay et al., 1999).

Two- and three-dimensional chemical transport models are also used to perform calculations that match modelled emissions, concentrations, and carbon isotopes with observed abundances. In this way, uncertainties in sources and sinks can be reduced, and the relationships between methane emissions, atmospheric chemistry, and the climate system better quantified. A number of studies (Fung et al., 1991; Hein et al., 1997; Gupta et al., 1996) have used this approach, while others have developed new inversion methods (Brown, 1995; Kandlikar and McRae, 1995; Kandlikar, 1997; Houweling et al., 1999). These analyses, combined with multi-box and source-specific studies, produce comprehensive budgets of CH₄ sources and sinks such as the composite estimate shown in Table 1 (Khalil, 2000).

2.3. Projections of future concentrations

Atmospheric methane concentrations have doubled since pre-industrial times. Increases in methane emissions from human sources such as rice cultivation, domestic ruminants, and increased fossil fuel use are

Table 1

TOTAL

(a) Estimated natural and anthro	1.0		D.C.
Source	Emissions (TgCH ₄ /year)	Range of estimate	References
Wetlands	100	92-232	Fung et al., 1991; Bartlett and Harriss, 1993; Cao et al., 1996b; Hein et al., 1997; Matthews, 2000
Termites	20	2-22	Sanderson, 1996; Sugimoto et al., 1998
Oceans	4	0.2-2.0	Bates et al., 1996; Holmes et al., 2000
Marine sediments	5	0.4-12.2	Judd, 2000
Geological sources	14	12-36	Judd, 2000, after Lacroix, 1993
Wild fires	2		Levine et al., 2000
Total Natural	145		
Ruminants	81	65-100	Crutzen et al., 1986; Johnson et al., 2000; IPCC, 1995
Rice cultivation	60	25-90	Cao et al., 1996a; Neue and Sass, 1998 and Yagi and Minami, 1998 as cited in Sass et al., 1999; Shearer and Khalil, 2000
Total Agricultural	141		
Natural gas	30	25-50	Cicerone and Oremland, 1988; Fung et al., 1991; Kirchgessner, 2000
Coal mining	46	15-64	Cicerone and Oremland, 1988; Boyer et al., 1990 as cited in Kirchgessner, 2000; Fung et al., 1991; Kirchgessner et al., 1993
Other fossil fuels	30	6-60	Khalil et al., 1993b; Piccot et al., 1996
Biomass burning	50	27-80	Levine et al., 2000; based on estimates by Seiler and Crutzen, 1980; Crutzen and Andreae, 1990; and Hao and Ward, 1993
Waste disposal	61	40-100	Johnson et al., 2000; Thorneloe et al., 2000
Total Non-Agricultural Anthropogenic	217		
TOTAL	503	410-660	
(b) Estimated sinks of methane	in TgCH ₄ /year (based on IPC	CC, 1995)	
Sink	Uptake	Range of	
	(TgCH ₄ /year)	estimate	
Tropospheric OH	445	360-530	
Removal to stratosphere	40	32-48	
Soil uptake	30	15-45	

'Best guess' estimates are those given by the methane budget in Khalil (2000). Range of estimates is based on the most recent studies as cited here.

430 - 600

thought to be the primary contributors to the rapid rise that has been observed through records preserved in ice cores, solar spectra, and direct atmospheric measurements.

515

It is likely that future methane concentrations, however, will be strongly affected by changes in both

sources and sinks of atmospheric methane. Projections of future methane concentrations therefore require modeling changes in both emission and removal rates of atmospheric methane, as well as considering feedbacks from climate change and emissions of other trace gases that affect both sources and sinks of CH₄.

190

2.3.1. Projections of CH₄ sources

In the past, increasing emissions have correlated well with population (e.g., Khalil and Rasmussen, 1993). However, recent variations in the methane growth rate have challenged our understanding of the many factors that affect the sources of atmospheric methane. This abundance of factors complicates the resolution of past emissions and makes predictions of future CH_4 emissions difficult.

Although methane emissions have climbed rapidly over the past four decades, the relative importance of agricultural sources may be declining, with non-agricultural sources related to fossil fuel use and waste disposal on the increase (Fig. 8). These sources are directly influenced by human activities and energy demand in ways that may not be easily predictable. In addition, they lack a well-resolved history of past emission trends to extrapolate into the future, a fact that greatly complicates methane emission prediction.

The magnitude of methane sources depends on numerous factors including human population, energy demand, crop yield, agricultural practices, land use area, temperature, precipitation, and other possible natural or anthropogenic influences not yet resolved, but that are expected to change significantly in the future. Methane concentrations are projected to continue to rise well into the next century, as shown by the "business as usual" IS92a emission scenario (Fig. 9). The "low emissions" scenario, IS92c, also shows a continued increase in methane concentration for a number of decades. However, several of the new SRES (IPCC Special Report on Emissions Scenarios) emissions scenarios for CH₄ that have been developed for the current ongoing IPCC assessment show reduced increases in emissions compared to the earlier scenarios (Fig. 10).

The majority of a projected increase in CH₄ emissions is expected to be from human sources related to agriculture, waste, or energy. Projections of future emissions are additionally uncertain because of the high level of ignorance concerning potential biospheric feedbacks to climate change. Observations of the response of biogenic sources to seasonal changes in temperature have shown that emissions from wetlands, rice paddies, and other ecosystems may increase in response to increases in global mean temperature, producing a positive feedback to climate warming (e.g., Hameed and Cess, 1983; Lashof, 1989; Roulet et al., 1992; Neue and Roger, 2000; Boone, 2000; Bartlett and Harriss, 1993; Slanina et al., 1994; Panikov, 1999). With high levels of warming, methane stored in methane hydrates and clathrates could

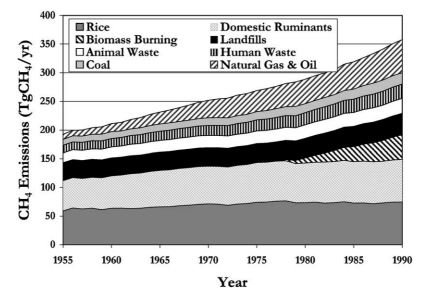


Fig. 8. Evolution of anthropogenic methane emissions, 1955-1990 (source: Hayhoe, 1997).

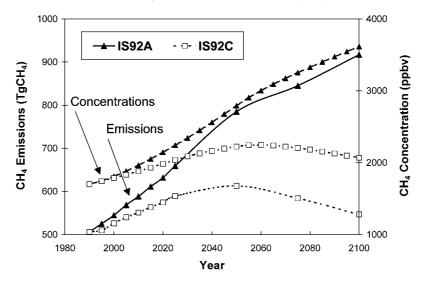


Fig. 9. Projected emissions and concentrations from IS92 scenarios A and C (source: IPCC, 1996; IPCC DDC CD-ROM).

also become destabilized, releasing large amounts of methane into the atmosphere (Nisbet, 1990a,b; Mac-Donald, 1990; Harvey and Huang, 1995), and increasing climate forcing.

Although methane concentrations are projected to continue to increase, it must be noted that recent decreases in the growth rate of methane have occurred without any controls on human emissions. If the declining trends in the growth rate over the past few decades had continued, rather than increasing as they have in the past 2 years, it is almost certain that a CH_4 reduction strategy involving reductions of more than 5% of the anthropogenic source could have stabilized or even lowered atmospheric concentrations of meth-

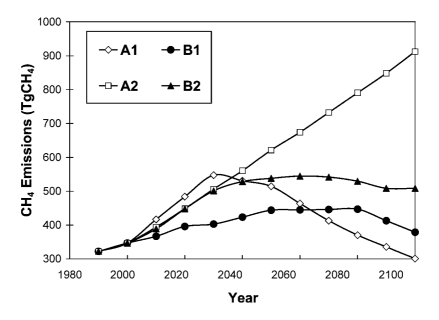


Fig. 10. New SRES emission projections for CH₄ (source: IPCC DDC CD-ROM).

ane (IPCC, 1996; Dlugokencky et al., 1998; Etheridge et al., 1998). However, as shown by Kheshgi et al. (1999), the trend in the concentration of methane is greatly dependent on any changes in the emissions of CO and NO_{x} .

2.3.2. Projections of CH₄ sinks

Concentrations of OH, the main sink of atmospheric methane, are already closely linked with CH₄ concentrations through the CH₄-CO-OH cycle. Most modelling studies conclude that OH concentrations have decreased by 10-30% since pre-industrial times, mainly due to increasing levels of CH₄ and CO in the atmosphere (e.g., Thompson, 1992; Wang and Jacob, 1998). At the present time, OH concentration appears to be relatively steady or perhaps increasing, with the effects of increasing CH₄ and CO balanced by increases in tropospheric O₃ concentrations and in emissions of NO_x and NMHCs (Prinn et al., 1995; Krol et al., 1998; Karlsdottir and Isaksen, 2000). Although OH has no significant radiative effect, it is the primary oxidizing agent of the troposphere, responsible for the removal of methane, carbon monoxide, and NMHCs as well as numerous other pollutants and long-lived greenhouse gases. Changes in its concentration will therefore impact the abundances of many trace species, affecting local air pollution in addition to global chemistry and radiative properties of the atmosphere.

Projections of OH concentrations are highly uncertain - even more so than past changes. Future changes in atmospheric composition induced by trace gas emissions and climate change are expected to have major impacts on the rate of OH production and destruction. A number of modelling studies predict up to a 25% decrease in OH concentrations by 2050 (e.g., Thompson and Cicerone, 1986; Thompson et al., 1990; Pinto and Khalil, 1991; Lu and Khalil, 1991; Osborn and Wigley, 1994; Brasseur et al., 1998; Lelieveld et al., 1998; Wang and Jacob, 1998). This small range belies the major changes in OH production and destruction that are projected to occur. These changes balance each other out to a certain extent, particularly in the studies that predict little or no change in net OH levels.

OH loss rates are projected to increase in the future due to rising anthropogenic emissions of CH_4 , CO, NMHCs, and other gases that are oxidized by OH.

However, there are a number of additional factors that may balance net OH concentrations by increasing OH production.

OH is formed when O_3 is photo-dissociated into $O(^1D)$, which in turn combines with water vapor to form two OH radicals, as shown by the following equations:

$$O_3 + hv \to O(^1D) + O_2 \tag{1}$$

$$O(^{1}D) + H_{2}O \rightarrow OH + OH$$
 (2)

Emissions of NO_x, an important component of local air pollution in urban areas, result in O₃ formation that in turn increases OH production (Eq. (1)). Increasing NO_x emissions have been modeled to result in an increase in global OH, and hence in the global methane sink (Fuglestvedt et al., 2000). A reduction in stratospheric ozone, such as that seen in recent decades, increases the flux of ultraviolet radiation reaching the troposphere, also promoting OH formation (Tang et al., 1998). As climate warms, the relationship between temperature and water vapor concentration may cause tropospheric water vapor levels to rise. This will also increase OH production (Eq. (2)). On the other hand, increases in tropospheric water vapor could also increase global mean cloud cover, which would decrease the flux of photons to the troposphere and decrease OH formation rates. Temperature increases also have an additional impact on temperature-dependent reaction coefficients, affecting many atmospheric reaction rates.

Although global average OH levels determine the average lifetime of CH_4 , several studies emphasize the importance of tropical OH levels in determining future changes in OH (Thompson et al., 1989; Crutzen and Zimmermann, 1991; Lelieveld et al., 1998). OH levels are highest in the tropics, with abundant water vapor and UV radiation ensuring high production rates. CH_4 emissions from important sources such as rice cultivation, biomass burning, and domestic ruminants are also highest in the tropics. However, the tropics are also the region where the greatest changes in land use and industrialization are expected to occur in the future. What happens in this area will to a large part determine the future of OH concentrations in the atmosphere.

The soil sink for methane, although substantially smaller than the OH sink, is also expected to change in the future. Methane oxidation in soil is sensitive to temperature and soil water content (Mancinelli, 1995; King, 1997), both of which may be altered by climate change. Perhaps more importantly, arable land has a much smaller CH₄ uptake rate than untreated soils, particularly when treated with certain fertilizers (e.g., Boeckx et al., 1997; Mosier et al., 1997b, Powlson et al., 1997). Land use change, as forests and grasslands are converted to agriculture, has already decreased the soil sink, and is likely to continue to do so in the future (King, 1997; Mosier et al., 1997a). Even when converted back to its natural state, previously cultivated land continue to have a lower oxidation rate than untouched areas (e.g., Dobbie and Smith, 1996; Hudgens and Yavitt, 1997; Prieme et al., 1997). This apparent irreversibility of human impacts on CH₄ oxidation rates has important implications for the future of land management strategies, as growing populations continue to exert pressure on natural ecosystems.

3. Impact of CH₄ on tropospheric and stratospheric chemistry

3.1. Methane and tropospheric chemistry

As discussed previously, methane oxidation is one of the major reaction pathways affecting atmospheric concentrations of hydroxyl on a global scale. Depending on nitric oxide levels, methane oxidation can be either a production or destruction process for oddhydrogen (OH+HO₂). Thus, different chemically coherent regions (Thompson et al., 1989) can be distinguished on the basis of concentrations of nitrogen oxides. Polluted (high NO_x) environments where odd hydrogen is produced include the temperate zone of the Northern Hemisphere and planetary boundary layer of the tropics during the dry season. Unpolluted (low NO_x) environments where odd hydrogen is destroyed include marine areas, the free troposphere over the tropics, and most of the Southern Hemisphere (e.g., Fishman et al., 1979; Crutzen, 1988; Cicerone and Oremland, 1988; WMO, 1991, 1995). Under warm, humid conditions, nitrogen oxide levels must be substantially higher to produce odd hydrogen, conditions that generally apply to the boundary layer. Most current modeling studies suggest that an increase in methane will decrease hydroxyl and increase ozone throughout the remainder of the troposphere.

The methane oxidation cycle is an important source of carbon monoxide, accounting for roughly a quarter of the carbon monoxide in the troposphere. Carbon monoxide concentrations are a great deal more variable than methane, due to its relatively short atmospheric lifetime (approximately 1-3 months) and because of the variety of natural and anthropogenic sources that contribute to its budget. These sources include fossil fuel combustion, biomass burning, and the oxidation of natural hydrocarbons emitted by vegetation (e.g., isoprene). As in the case of methane, the carbon monoxide oxidation cycle also depends on the levels of nitric oxide present in the atmosphere.

Based on the reaction sequences for methane, formaldehyde, and carbon monoxide in polluted as well as unpolluted environments, the effect of the complete oxidation of 1 mol of methane is an increase from 0.4 to 0.5 mol of odd hydrogen in the high NO_{r} case and a decrease from -3.5 to -3.9 mol of odd hydrogen in the low NO_x case. Correspondingly, the change in ozone for complete oxidation of methane is an increase of +3.6 to 3.8 mol for the high NO_x case and a decrease of -1.7 to -1.8 mol for the low NO_x case. According to our calculations (Wuebbles et al., 2000), the catalytic subcycle involving the reaction of methylperoxyl (CH₃O₂) with hydroperoxyl (HO₂) and the reaction of methyl peroxide (CH₃O₂H) with OH has a sizeable impact on the amount of odd hydrogen loss in the low NO_x case. Crutzen (1988) included this cycle in his results, whereas lower effects determined by Cicerone and Oremland (1988) did not include its effect. Our analyses of the resulting changes in oddhydrogen and ozone are substantially in agreement with Crutzen (1988) for both cases. The resultant destruction of odd hydrogen depends on the hydroxyl concentration, the methyl peroxide chemical reaction pathways, and the heterogeneous removal rates of important intermediate species.

Formaldehyde is an important intermediate in the removal processes of methane and other hydrocarbons as well as being important in the general chemical reactivity of the troposphere. Regardless of NO_x levels, methane oxidation is responsible for the formation of

much of the formaldehyde in the atmosphere. Based on the estimate of Lowe and Schmidt (1983), the formaldehyde production rate is on the order of 10^{14} g/ year. The other major photochemical precursors of formaldehyde consist of nonmethane hydrocarbons and higher aldehydes. The non-methane hydrocarbons are emitted from natural and anthropogenic sources while the higher aldehydes are produced by in situ photochemistry. There are appreciable anthropogenic sources of formaldehyde as well (e.g., automobile exhaust).

In order to analyze the photochemical mechanisms that produce formaldehyde, we must distinguish between different regions on the basis of NO_x levels. In the methane oxidation mechanism under low NO_{x} conditions, methyl peroxide reacts with OH to produce formaldehyde. Since methyl peroxide has a long lifetime against photolysis, heterogeneous processes (which occur on aerosols) can serve as a loss mechanism for this species in unpolluted environments. This implies that formaldehyde production is controlled by aerosol concentrations in remote regions. However, in polluted environments, formaldehyde formation does not involve methyl peroxide. In this case, it would be regulated by nitric oxide concentrations because methylperoxyl is reduced to methoxyl. Methoxyl then reacts with oxygen to produce formaldehyde.

Once it is generated, formaldehyde is quickly destroyed under direct insolation. During the daytime, photolysis is the dominant loss process for formaldehyde in the remote troposphere, while in polluted atmospheres, reactions with hydroxyl and hydroperoxyl are comparable to photolytic degradation. At night, the only appreciable formaldehyde consumption is by reaction with nitrate radical (NO₃) but this is slow in comparison to the daytime reactions with hydroxyl and hydroperoxyl.

Approximately 10% of the ozone in the atmosphere is located in the troposphere. The downward transport of ozone from the stratosphere traditionally was thought to be the major source of tropospheric ozone (Crutzen, 1988 and references therein). It is now generally accepted that the net tropospheric photochemical production of ozone is of similar magnitude to the downward transport source (e.g., Fishman et al., 1979; Fishman, 1985; WMO, 1985, 1995; Isaksen, 1988; Penkett, 1988; Hough and Derwent, 1990).

Nitrogen dioxide photolysis is by far the primary photochemical mechanism for producing ozone in the troposphere. This implies that the generation rate is roughly proportional to the concentration of nitric oxide. High concentrations of NO_x over the continental boundary layer signify that this region is likely a net source of ozone. Increases in NO_x emissions may lead to further ozone increases, especially in the tropics. However, the magnitude of odd oxygen $(O+O(^{1}D)+O_{3})$ production is ultimately limited by the supply of CO, CH₄, and NMHCs. The oxidation of 1 mol of CO can form 1 mol of ozone. In contrast, the complete oxidation of a mole of methane can produce 3-4 mol of ozone. Because there is insufficient NO_x present in the background troposphere, it is estimated that only about 10% of the potential tropospheric ozone production is being realized.

3.2. Methane and stratospheric chemistry

Although ~ 90% of total emissions of methane are consumed by reaction with tropospheric hydroxyl and about 5% by soil uptake (IPCC, 1995), the remaining 5% of methane flux, on average about 27 TgCH₄/year (Gettelman et al., 1997), enters the stratosphere.

In the stratosphere and higher layers of the atmosphere, the reaction with OH continues to be the dominant sink for CH_4 , but reactions with chlorine atoms and excited oxygen atoms are also important. Reactions with chlorine atoms account for about 9% of the methane loss in these regions (Brenninkmeijer et al., 1995). Measurements also suggest that the increased oxidation of methane resulting from increasing concentrations of chlorine in the stratosphere has led to enhanced production of stratospheric OH (e.g., Burnett and Burnett, 1995).

In order to put the role of methane in stratospheric chemistry into context, it is useful to first discuss the importance of ozone and the changes occurring in its distribution. Changes in the distribution and amount of ozone in the global troposphere and stratosphere have received much attention over the past few decades. Much of the concern about ozone has centered on the importance of ozone as an absorber of ultraviolet radiation, as its concentrations determine the amount of ultraviolet radiation reaching the Earth's surface. Ozone is also a greenhouse gas and can influence climate. Approximately 90% of the ozone in the atmosphere is contained in the stratosphere. In the stratosphere, the production of ozone begins with the photodissociation of oxygen (O_2) at ultraviolet wavelengths less than 242 nm. This reaction produces two ground-state oxygen atoms that can react with oxygen to produce ozone. Since an oxygen atom is essentially the same as having an ozone molecule, it is common to refer to the sum of the concentrations of ozone and oxygen atoms (both ground state and excited state) as odd-oxygen. The primary destruction of odd-oxygen in the stratosphere comes from catalytic mechanisms involving various free radical species. Nitrogen oxides, chlorine oxides, and hydrogen oxides participate in catalytic reactions that destroy odd-oxygen.

Chlorine and bromine catalytic mechanisms are particularly efficient in destroying ozone. These mechanisms have been the subject of much study due to their effects on concentrations of stratospheric ozone over recent decades, particularly since levels of reactive chlorine in the stratosphere have been growing rapidly over this time as a result of emissions of trichlorofluoromethane (CFCl₃), difluorodichloromethane (CF₂Cl₂), bromine-containing halons, and other halocarbons. The chlorine- and bromine-catalytic cycles can turn over thousands of times before the catalyst is converted to a less reactive form. Because of this cycling, relatively small concentrations of reactive chlorine or bromine can have a significant impact on the amount and distribution of ozone in the stratosphere. The total amount of chlorine in the current strato sphere is \sim 3 ppbv, much of which is in the form of less reactive compounds like hydrochloric acid (HCl).

Methane plays an important role in the chlorine chemistry of the stratosphere, serving both as a source and a sink in key reactions affecting reactive chlorine. The direct reaction of methane with a chlorine atom is the primary source of hydrochloric acid, the main chlorine reservoir species. However, OH produced through the oxidation of methane in the stratosphere can react with the hydrochloric acid to return the chlorine atom, thus reinitiating the chlorine catalytic mechanism. Although the most important effect of increasing methane is the increasing conversion of reactive chlorine to HCl, there is also a smaller increase in the conversion rate of reactive chlorine to HCl. Thus, increasing chlorine tends to slow down the effects of chlorine on ozone. Conversely, controls on methane could exacerbate the effects of chlorine on ozone by slowing down the conversion to HCl. It should be noted that a similar mechanism is not important for bromine because of the slow reactivity of Br with CH₄.

In addition to being involved in the reaction taking reactive chlorine to the less reactive HCl, methane has several other effects on stratospheric ozone. Hydrogen oxides produced from the dissociation of methane can react catalytically with ozone, particularly in the upper stratosphere. In the lower stratosphere, the primary effect of these hydrogen oxides is to react with nitrogen oxides and reactive chlorine, reducing the effectiveness of associated ozone destruction catalytic cycles. The hydrogen oxides at these altitudes can also react catalytically to destroy ozone.

3.3. Model-calculated effects on ozone

Changing atmospheric concentrations of a number of compounds, including methane, are currently affecting the distribution of ozone in the troposphere and stratosphere. A number of research studies have used numerical models to examine the combined effects on ozone from the increases in concentrations of methane and the other gases described in the previous section (e.g., Wuebbles et al., 1983; Stordal and Isaksen, 1987; WMO, 1985, 1989, 1995, 1999). When combined with the effects of other trace gas emissions, it is difficult to evaluate the specific contribution of methane to the observed and projected ozone trends. For this reason, it is useful to examine studies that have only considered the effects of increasing methane on ozone.

Numerical models of atmospheric chemical and physical processes generally find that increasing methane concentrations result in a net ozone production in the troposphere and lower stratosphere and net ozone destruction in the upper stratosphere (Owens et al., 1982, 1985; WMO, 1985, 1991, 1995; Isaksen and Stordal, 1986). The net effect from these calculations has been that methane by itself causes a net increase in ozone. For a doubling of the methane concentration (early papers studied CH₄ increases from 1.6 to 3.2 ppmv, while recent analyses assume a globally-averaged change from 1.7 to 3.4 ppmv), published effects on the calculated change in total ozone range from +0.3% (Prather in WMO, 1985) to +4.3% (Owens et al., 1985). With radiative feedback effects included (allowing temperature changes in the stratosphere), the published model results tend to be in the upper end of this range (Owens et al., 1985; WMO, 1985; Isaksen and Stordal, 1986).

The effects on tropospheric and stratospheric ozone for a doubling of the methane surface concentrations from the UIUC zonally-averaged model (Wuebbles et al., 1998; Rahmes et al., 1998) are given in Fig. 11 (change in total ozone as a function of latitude and season) and Fig. 12 (change in ozone with altitude and latitude for July). This two-dimensional model analysis, which includes radiative feedbacks, gives a 3.4% increase in globally-averaged total ozone from a doubling of methane surface concentrations.

Calculations of the effect of methane increases on stratospheric ozone give a small percentage increase in the lower stratosphere, a larger increase near 40 km altitude, and a decrease in ozone above 45 km. As mentioned earlier, the hydrogen oxides produced by methane oxidation affect the efficiency of the nitrogen oxide and chlorine oxide catalytic ozone destruction mechanisms. However, the effect of methane in the lower stratosphere will depend on the efficiency of the nitrogen oxide catalytic cycle; if the amount of reactive odd nitrogen is reduced, then the additional hydrogen oxides from methane could destroy ozone in this region. Additional heterogeneous reactions involving N₂O₅, BrONO₂ and ClONO₂ reduce the efficiency of the nitrogen oxide catalytic cycle, and increased HO_x from increases in atmospheric methane has a larger effect on the lower stratosphere as compared to previous calculations (Wuebbles et al., 2000). In the upper stratosphere, the additional hydrogen oxides react catalytically with ozone, causing models to find a decrease in ozone at these altitudes.

The version of the model used here includes convective mixing processes in the troposphere and also includes a number of non-methane hydrocarbons. The change in column ozone resulting from a calculation

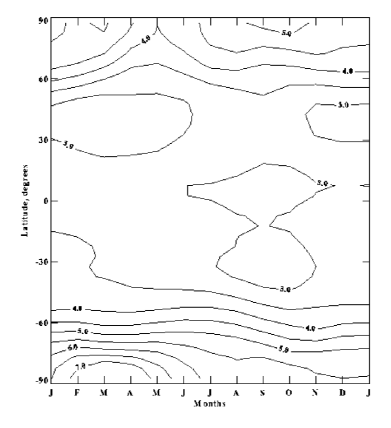


Fig. 11. Calculated percentage change of total ozone for a doubling in the concentration of atmospheric methane from 1.7 to 3.4 ppmv. Based on the UIUC two-dimensional chemical-transport model.

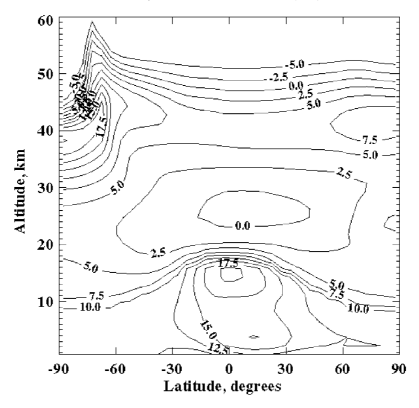


Fig. 12. Calculated percentage change of local ozone in July for a doubling in the concentration of atmospheric methane as in Fig. 10.

for doubling CH_4 without NMHCs compared to a simulation with NMHCs is an increased response ranging from 3% to 8% larger than that when the effect of NMHCs is included. The effect on the Southern Hemisphere is negligible, as there are no significant non-NMHC emissions from this region. Thus, the model response is damped for a doubling of methane surface concentration when non-methane hydrocarbons are included in the model. This damping results from the additional sources of reactive oxides available in the lower troposphere from hydrocarbons sources. The current model gives a feedback factor, defined as the percentage change in OH for a 1% change in CH_4 of 0.32%, which is well within the range of those calculated in WMO (1995).

3.4. Upper atmospheric water vapor

The spatial distribution of water vapor in the troposphere is primarily determined by evaporation, condensation and transport processes. Very little of the tropospheric water vapor penetrates into the stratosphere. The mechanisms limiting the transport of tropospheric water vapor into the stratosphere are still not well understood. Concentrations of water vapor increase with altitude in the stratosphere, from 3 ppmv in the lower stratosphere to about 6 ppmv in the upper stratosphere. This increase in concentration with altitude occurs as a result of the oxidation of methane.

Methane oxidation reactions produce roughly two mol of water vapor for each mole of methane that is destroyed. Stratospheric water vapor concentrations should increase as concentrations of methane increase. Since methane concentrations have increased from about 0.7 ppmv in the pre-industrial atmosphere to the current concentration of 1.7 ppmv, this implies that upper stratospheric water vapor concentrations have increased by roughly 2 ppmv over this time period. In actual fact, the increase in water vapor should be somewhat less than this due to methane reactivity with chlorine and oxygen atoms. Both modeling and data analysis studies (e.g., Le Texier et al., 1988; Hansen and Robinson, 1989; Randel et al., 1998; Zoger et al., 1999) are in agreement with this conclusion, indicating that the overall stratospheric water vapor yield from methane is slightly less than two. A doubling of methane surface concentration in the UIUC two-dimensional model of the global atmosphere gives an increase in the amount of water vapor at the stratopause, near 50 km, of approximately 40% or about 3 ppmv.

Increasing methane concentrations should also be leading to increasing concentrations of water vapor in the mesosphere. In conjunction with carbon dioxide, increasing methane concentrations should also lead to cooler temperatures in the stratosphere, mesosphere, and thermosphere (WMO, 1985, 1989; Brasseur and Hitchman, 1988; Roble and Dickinson, 1989). With the increase in water vapor and the cooler temperatures, there is the potential for increased occurrences of noctilucent clouds near the mesopause (Thomas et al., 1989). Roble and Dickinson (1989) also point out that other changes in mesospheric and thermospheric composition should occur due to both the cooler temperatures and direct chemical effects from increased concentrations of methane and carbon dioxide. In addition, exospheric hydrogen will increase with increasing methane (Ehhalt, 1986).

4. Impact of methane on climate

4.1. Direct effects

In the past, the concern that human activities may be affecting global climate has largely centered around carbon dioxide because of its importance as a greenhouse gas and also because of the rapid rate at which its atmospheric concentration has been increasing. However, it is clear that other greenhouse gases have also significantly affected climate over the past few centuries. Rising methane concentrations are a major contributor to the increase in radiative forcing since pre-industrial times, second only to carbon dioxide. The combined effect of methane and the other greenhouse gases has been to approximately double the overall increase in the greenhouse radiative forcing on climate relative to that from CO₂. Like other greenhouse gases, methane absorbs infrared radiation (also called longwave or terrestrial radiation)

emitted by the relatively warm planetary surface and emits radiation to space at the colder atmospheric temperatures, leading to a net trapping of infrared radiation in the atmosphere. This is called the greenhouse effect. The balance between the absorbed solar radiation and the emitted infrared radiation determines the net radiative forcing on climate.

Although its atmospheric abundance is less than 0.5% that of carbon dioxide, methane is an important greenhouse gas. Donner and Ramanathan (1980) calculated that the presence of methane at current levels causes the globally-averaged surface temperature to be about 1.3 K higher than it would be without methane. On a molar basis, an additional mole of methane in the current atmosphere is about 24 times more effective at absorbing infrared radiation and affecting climate than an additional mole of carbon dioxide (WMO, 1999). Correspondingly, on a mass basis, an additional kilogram of methane is about 66 times more effective as a greenhouse gas than a kilogram of carbon dioxide. It should be noted that the actual lifetime of carbon dioxide in the atmosphere is much longer than the atmospheric lifetime of methane (IPCC, 1990, 1996). However, it should also be recognized that the dissociation of methane eventually produces carbon dioxide, leading to additional climatic forcing from methane emissions from fossil sources.⁵

The strongest bands for absorption by methane in the infrared are in the short wavelength edge of the 'window region' ($\sim 7-13 \mu$ m, where there is little absorption by H₂O or CO₂). The most important infrared spectral feature of methane is the 7.66 µm (1306 cm⁻¹) absorption band. Due to saturation of the line cores for methane and emissions from the pressure-broadened Lorentz line wings, radiative forcing from methane increases approximately as the square root of its concentration (IPCC, 1990; Wigley, 1987). Overlap with absorption by water vapor and other species (particularly nitrous oxide) also affects the efficiency of methane absorption.

Wang et al. (1991) show that the greenhouse radiative forcing for methane has different spatial effects on climate than carbon dioxide, and that methane needs to be accounted for explicitly when attempting

⁵ The oxidation of methane from biogenic sources also results in CO_2 production, but the forcing of CO_2 from these sources has already been taken into account as part of the carbon cycle.

to predict the climate response to increasing concentrations of greenhouse gases. Calculations with threedimensional global climate models (GCMs) are beginning to consider the explicit effects of methane on climate. Hansen et al. (1988) and Wang et al. (1991) have explicitly considered the effects of methane in GCM studies, while many other studies have included methane only through accounting for its radiative forcing through use of an increase in carbon dioxide as a proxy.

Increasing concentrations of methane are thought to be a significant fraction of the increase in radiative forcing from greenhouse gases over the last two centuries. IPCC (1996) calculated that the direct radiative effect of the increase in methane since the mid-1700s has produced a radiative forcing increase of 0.47 W m^{-2.6} This is about 19% of the total change in radiative forcing due to CO₂ and other greenhouse gases over this time period.⁷ Inclusion of indirect effects of methane increases on stratospheric water vapor, ozone, and carbon dioxide (see next section) could increase this percentage by at least 10–35% (e.g., IPCC, 1995; Bruhl, 1993; Lelieveld et al., 1993, 1998; Fuglestvedt et al., 1996).

A recent analysis of the radiative forcing for methane (Jain et al., 2000), accounting for the latitudinal and seasonal variations in methane concentrations, calculates a larger radiative forcing contribution from methane than earlier studies. The globally- and annually-averaged radiative forcing based on the new calculations was 4.67×10^{-4} W m⁻² for a 1-ppb change in methane, compared to 3.7×10^{-4} W m⁻² in IPCC (1996). The estimated change in radiative forcing from pre-industrial times to 1992 was 0.55 W m⁻² compared to 0.47 in IPCC (1996). Minschwaner et al. (1998) also determined a radiative forcing for methane from 1750 to 1992 of 0.55 W m⁻² based on the observed distribution of methane from the Upper Atmosphere Research Satellite.

Over the last decade, increases in atmospheric methane concentration are calculated to have increased

radiative forcing by about 0.06 W m⁻², 11% of the total increase in radiative forcing from greenhouse gases over this time period (WMO, 1999).⁸

Various studies have evaluated the potential effects of a doubling of methane concentrations on radiative forcing and surface temperature using radiative-convective models. For an increase from 1.6 to 3.2 ppmv, resulting increases in surface temperature range 0.2-0.3 K (Owens et al., 1985; Ramanathan et al., 1987; MacKay and Khalil, 1991; IPCC, 1995), with the differences in model results primarily relating to uncertainties in the band strengths for methane infrared absorption. For a doubling from 1.7 to 3.4 ppmv, Owens et al. (1985) calculated a direct 0.34 K increase in surface temperature, along with an additional 0.26 K due to indirect effects from methane-induced effects on carbon dioxide and ozone. For a 25% increase in methane concentrations, Ramanathan et al. (1985) determine a 0.08-K increase in surface temperature when overlap with the radiative absorption with other greenhouse gases is included, and a 0.19-K increase in surface temperature without overlap.

Other modeling studies have included increasing methane concentrations in studies evaluating scenarios for potential future changes in radiative forcing and global temperatures (Wang and Molnar, 1985; Ramanathan et al., 1985, 1987; WMO, 1985; Dickinson and Cicerone, 1986; Wang et al., 1986; Wigley, 1987; Hansen et al., 1988, 1989; IPCC, 1990, 1995, 1996). In Table 2, the change in radiative forcing due to increasing methane is compared to the total change in forcing due to all greenhouse gases for three IPCC emission scenarios (high, medium and low) (IPCC, 1996). These more recent scenarios generally show a somewhat increased role for methane in determining future radiative forcing. However, these studies do not consider the significant indirect effects from chemical interactions as discussed next.

4.2. Indirect effects

In addition to its direct radiative forcing effect on climate, methane can also influence climate indirectly through chemical interactions affecting other radia-

⁶ Climate models indicate that the 4 W m⁻² forcing associated with a doubling of CO_2 from 300 to 600 ppmv would give approximately a 1.5- to 4.5-K increase in surface temperature.

⁷ Other research studies have found similar percentages for the effect of methane over this period: e.g., Rodhe (1990) derived 15%, while Hansen et al. (1989) and MacKay and Khalil (1991) calculated $\sim 22\%$.

⁸ Note that these estimates ignore the effects on radiative forcing due to changes in global ozone and aerosols over this time period.

Table 2

Change in radiative forcing (W m^{-2}) due to increasing methane, as compared to total change in forcing due to all greenhouse gases. Radiative forcing given by IPCC (1996) for their high (IS92e), medium (IS92a), and low (IS92c) emission scenarios

Time period	Scenarios									
	IS92a			IS92c			IS92e			
	CH_4	All	%	CH_4	All	%	CH_4	All	%	
1765-2025	0.66	4.01	16.4	0.59	3.63	16.2	0.69	4.31	16.0	
1765 - 2050	0.83	5.37	15.4	0.65	4.37	14.9	0.89	6.12	14.5	
1990-2025	0.19	1.61	11.8	0.12	1.23	9.7	0.22	1.91	11.5	
1990 - 2050	0.36	2.97	12.1	0.18	1.97	9.1	0.42	3.72	11.3	

tively important gases. Methane is an important influence on concentrations of hydroxyl, the primary tropospheric oxidizing agent, which in turn determines the rate at which methane is removed from the atmosphere. Methane oxidation is also a significant source of tropospheric and stratospheric ozone, stratospheric water vapor, and its eventual final product, carbon dioxide.

Resolving changes in tropospheric OH due to CH₄ emissions is difficult because of uncertainties associated with modeling the non-linear relationships in the CH₄-CO-OH cycle. Increases in the response time of methane to a perturbation can result in a much larger apparent lifetime for the perturbed methane than the overall atmospheric lifetime of methane (Prather, 1994; IPCC, 1995, 1996). Tropospheric OH concentrations are expected to change in the future due to increases in CH₄ as well as other trace gases such as CO, NO_x, O₃, NMHCs and water vapor. Depending on the emission scenario examined, trace gas emissions can either lengthen or shorten the lifetime of CH₄, resulting in a positive or negative indirect effect (Hayhoe et al., 2000; Bruhl, 1993; Lelieveld et al., 1993, 1998).

Changes in tropospheric ozone associated with increasing methane are the most important indirect effect, capable of adding $19\pm12\%$ to the direct radiative forcing from methane (IPCC, 1995; based on UIUC and other two-dimensional model studies). The large remaining uncertainty is associated with the overall uncertainties in understanding the processes affecting ozone in the troposphere.

Methane oxidation in the stratosphere is an important source of water vapor. Although the concentration of water vapor in the stratosphere is several orders of magnitude smaller than tropospheric concentrations, at the low temperatures found in the lower stratosphere, relatively small changes in water vapor levels can significantly impact radiative forcing. Therefore, an increase in stratospheric water vapor concentrations resulting from increasing methane concentrations will enhance the greenhouse effect, and provide a positive indirect effect that will enhance the direct radiative forcing from the added methane. While earlier studies assuming a mixing ratio increase in methane suggested that the increase in water vapor could enhance the radiative forcing by as much as 30% (IPCC, 1990), more recent studies (Lelieveld and Crutzen, 1991, 1992; Bruhl, 1993; Lelieveld et al., 1993; Hauglustaine et al., 1994; IPCC, 1995) using flux boundary condition changes in methane have shown the effect of the water vapor increase to be much smaller, 4-5% of the total direct radiative forcing from methane.

Increasing water vapor could also lead to an increased amount of polar stratospheric clouds. Ramanathan (1988) showed that both water and ice clouds, when formed at cold lower stratospheric temperatures, are extremely efficient in enhancing the atmospheric greenhouse effect. He also noted that there is a distinct possibility that large increases in future methane may lead to a surface warming that increases nonlinearly with the methane concentration.

Finally, methane oxidation eventually ends in the production of carbon dioxide, one of the most important greenhouse gases. However, about 500 Tg/year of methane are destroyed by reaction with hydroxyl and converted to carbon dioxide, accounting for production of only 380 TgC/year as carbon dioxide; in contrast, emissions of carbon dioxide from anthropogenic fossil fuel use and cement manufacturing are ~ 6000 TgC/year. In addition, CO₂ produced by the oxidation of methane from biogenic sources is already included in the carbon cycle budget; only CO₂ from methane emitted by fossil fuel sources contributes to the indirect effect.

4.3. Global warming potentials

Global Warming Potentials (GWPs) have been developed as an analysis tool for policy makers to assist in the evaluation of possible policy actions related to greenhouse gas controls. The Kyoto Protocol uses GWPs to weight emissions of the six greenhouse gases that can be included in national emission reduction strategies (UNFCCC, 1997). Based on emission reductions required under the Kyoto Protocol, we recently analyzed greenhouse gas abatement scenarios for the U.S. (Hayhoe et al., 1999). Using a 100-year GWP for methane, as specified in the Protocol, our results indicate that the cost of achieving a reduction of 650 GtC_{eq} by 2010 would be reduced by at least 25% if methane were included in a previously CO₂ only strategy.

The GWP of a greenhouse gas is defined as the time-integrated commitment to climatic forcing from the instantaneous release of a kilogram of the gas relative to the climatic forcing from the release of 1 kg of carbon dioxide, and is generally calculated for constant background concentrations. Published GWPs are derived for integration periods from 20 to 500 years, with the 100-year values generally thought to provide a balanced representation of the various time horizons for climatic response. Under this measure, the latest 100-year GWP for methane, including both direct and indirect effects, is 24 ± 7.5 (WMO, 1999; IPCC, 1995), compared to 1.0 for carbon dioxide. The GWPs from IPCC (1996) for direct plus indirect effects from methane for the 20- and 500-year integrations are 62 ± 20 (64 in WMO, 1999) and 7.5 ± 2.5 , respectively. As shown in IPCC (1990, 1995), indirect effects can more than double the direct methane GWP value (see also Fuglestvedt et al., 1996; Lelieveld et al., 1998). Additional uncertainties are introduced by the assumption of constant background concentrations over the length of the integration. Projected changes in atmospheric concentrations of CH₄ and other trace gases such as OH, CO and NO_x affect the removal rate of CH₄ and hence its GWP (Hayhoe et al., 2000). Uncertainties in GWPs of methane and other important greenhouse gases in turn affect their optimal contribution to emission reductions and hence to the prevention of climate change.

5. Summary and conclusions

The atmospheric concentration of methane has increased dramatically over the last century and continues to increase. While budget and isotopic analyses have confirmed the causal role of human activities in this increase, significant uncertainties remain in understanding the factors that affect emissions from various sources and how these will change over time. Uncertainties in how the sources and sinks of methane will change in the future limit our ability to develop meaningful climate change policies aimed at controlling methane emissions.

The 1998 increase in the growth rate of methane has renewed the debate about future levels of methane. This increase could just be a short-lived anomaly. However, if this increase is the first sign of one of the most important biogenic feedbacks to climate change — the response of wetlands to the increases in global temperatures — this raises urgent questions concerning the magnitude of the response of methane concentrations to the much larger increases in global temperatures projected to occur over the next century.

More research is needed to fully understand the growth rate in methane and its future projection. More complete analyses will require increased coordination between programs measuring atmospheric abundances and isotopic ratios, and modelling efforts that couple biospheric and oceanic biogeochemistry with atmospheric processes.

Acknowledgements

This study was supported in part by the U.S. Environmental Protection Agency and by the National Aeronautics and Space Administration's Atmospheric Chemistry Modeling and Analysis Program.

References

- Bains, S., Corfield, R., Norris, R., 1999. Mechanisms of climate warming at the end of the Paleocene. Science 285, 724–727.
- Barns, D.W., Edmonds, J.A., 1990. An Evaluation of the relationship between the production and use of energy and atmospheric methane emissions. U.S. Dept. of Energy, Carbon Dioxide Research Program, TR047, 223 pp.
- Bartlett, K., Harriss, R., 1993. Review and assessment of methane emissions from wetlands. Chemosphere 26, 261–320.
- Bates, T.S., Kelly, K.C., Johnson, J.E., Gammon, R.H., 1996. A reevaluation of the open ocean source of methane to the atmosphere. J. Geophys. Res. 101, 6953–6961.
- Bazhin, N., 1994. Sources and sinks of methane on the territory of the former USSR. Pure Appl. Chem. 66, 188–191.
- Beck, L.L., Piccot, S.D., Kirchgessner, D.A., 1993. Industrial sources. In: Khalil, M. (Ed.), Atmospheric Methane: Sources, Sinks and Role in Global Change. Springer-Verlag, New York, NY, pp. 341–399.

- Bekki, S., Law, K., 1997. Sensitivity of the atmospheric CH_4 growth rate to global temperature changes observed from 1980 to 1992. Tellus 49B, 409–416.
- Bekki, S., Law, K.S., Pyle, J.A., 1994. Effects of ozone depletion on atmospheric CH₄ and CO concentrations. Nature 371, 595–599.
- Bellisario, L.M., Bubier, J.L., Moore, T.R., Chanton, J.P., 1999. Controls on CH₄ emissions from a northern peatland. Global Biogeochem. Cycles 13, 81–91.
- Bender, M., Sowers, T., Brook, E., 1997. Gases in ice cores. Proc. Natl. Acad. Sci. 94, 8343–8349.
- Blake, D.R., Rowland, F.S., 1988. Continuing worldwide increase in tropospheric methane, 1978 to 1987. Science 239, 1129–1131.
- Blunier, T., Chappellaz, J., Schwander, J., Barnola, J., Desperts, T., Stauffer, B., Raynaud, D., 1993. Atmospheric methane, record from a Greenland ice core over the last 1000 years. Geophys. Res. Lett. 20, 2219–2222.
- Blunier, T., Chapellaz, J., Schwander, J., Stauffer, B., Raynaud, D., 1995. Variations in atmospheric methane concentration during the Holocene Epoch. Nature 374, 46–49.
- Blunier, T., Chappellaz, J., Schwander, J., Dällenbach, A., Stauffer, B., Stocker, T.F., Raynaud, D., Jouzel, J., Clausen, H.B., Hammer, C.U., Johnsen, S.J., 1998. Asynchrony of Antarctic and Greenland climate change during the last glacial period. Nature 394, 739–743.
- Boeckx, P., VanCleemput, O., Villaralvo, I., 1997. Methane oxidation in soils with different textures and land use. Nutr. Cycling Agroecosyst. 49, 91–95.
- Bogner, J., Spokas, K., 1993. Landfill CH₄: rates, fates and role in global carbon cycle. Chemosphere 26, 369–386.
- Bogner, J., Spokas, K., Burton, E., Sweeney, R., Corona, V., 1995. Landfills as atmospheric methane sources and sinks. Chemosphere 31, 4119–4130.
- Boone, D., 2000. Biological formation and consumption of methane. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 42–62.
- Borjesson, G., Svensson, B., 1997. Effects of a gas extraction interruption on emissions of methane and carbon dioxide from a landfill, and on methane oxidation in the cover soil. J. Environ. Qual. 26, 1182–1190.
- Brasseur, G., Hitchman, M.H., 1988. Stratospheric response to trace gas perturbations: changes in ozone and temperature distribution. Science 240, 634–637.
- Brasseur, G., Kiehl, J., Muller, J.-F., Schneider, T., Granier, C., Tie, X., Hauglustaine, D., 1998. Past and future changes in global tropospheric ozone: impact on radiative forcing. Geophys. Res. Lett. 25, 3807–3810.
- Brenninkmeijer, C.A.M., Lowe, D.C., Manning, M.R., Sparks, R.J., van Velthoven, P.F.J., 1995. The ¹³C, ¹⁴C, and ¹⁸O isotopic composition of CO, CH₄, and CO₂ in the higher southern latitudes lower stratosphere. J. Geophys. Res. 100, 26163–26172.
- Brook, E., Sowers, T., Orchardo, J., 1996. Rapid variations in atmospheric methane concentration during the past 110,000 years. Science 273, 1087–1091.
- Brook, E., Harder, S., Severinghaus, J., Steig, E., Sucher, C., 2000. On the origin and timing of rapid changes in atmospheric methane during the last glacial period. Global Biogeochem. Cycles 14, 559–572.

- Brown, M., 1995. The singular value decomposition method applied to the deduction of emissions and the isotope composition of atmospheric methane. J. Geophys. Res. 100, 11425–11446.
- Bruhl, C., 1993. The impact of the future scenarios for methane and other chemically active gases on the GWP of methane. Chemosphere 26, 731–738.
- Brune, W., 1992. Stalking the elusive atmospheric hydroxyl radical. Science 256, 1154–1155.
- Buendia, L., Neue, H., Wassmann, R., Lantin, R., Javellana, A., 1997. Understanding the nature of methane emission from rice ecosystems as basis of mitigation strategies. Appl. Energy 56, 433–444.
- Burnett, E.B., Burnett, C.R., 1995. Enhanced production of stratospheric OH from methane oxidation at elevated reactive chlorine levels in northern midlatitudes. J. Atmos. Chem. 21, 13–41.
- Butterbachbahl, K., Papen, H., Rennenberg, H., 1997. Impact of gas transport through rice cultivars on methane emission from rice paddy fields. Plant Cell Environ. 20, 1175–1183.
- Cantrell, C.A., Shetter, R.E., McDaniel, A.H., Calvert, J.G., Davidson, J.A., Lowe, D.C., Tyler, S.C., Cicerone, R.J., Greenberg, J.P., 1990. Carbon kinetic isotope effect in the oxidation of methane by hydroxyl radicals. J. Geophys. Res. 95, 22455– 22462.
- Cao, M., Dent, J., Heal, O., 1995. Modeling methane emissions from rice paddies. Global Biogeochem. Cycles 9, 183–195.
- Cao, M., Gregson, K., Marshall, S., Dent, J., Heal, O., 1996a. Global methane emissions from rice paddies. Chemosphere 33, 879– 897.
- Cao, M., Marshall, S., Gregson, K., 1996b. Global carbon exchange and methane emissions from natural wetlands: application of a process-based model. J. Geophys. Res. 101, 14399–14414.
- Chappellaz, J., Barnola, J., Raynaud, D., Korotkevich, Y., Lorius, C., 1990. Ice-core record of atmospheric methane over the past 160,000 years. Nature 345, 127–131.
- Chappellaz, J., Blunier, T., Raynaud, D., Barnola, J., Schwander, J., Stauffer, B., 1993a. Synchronous changes in atmospheric CH₄ and Greenland climate between 40-kyr and 8-kyr BP. Nature 366, 443–445.
- Chappellaz, J., Fung, I.Y., Thompson, A.M., 1993b. The atmospheric CH₄ increase since the Last Glacial Maximum. Tellus 45B, 228–241.
- Chappellaz, J., Blunier, T., Kints, S., Dallenbach, A., Barnola, J., Schwander, J., Raynaud, D., Stauffer, B., 1997. Changes in the atmospheric CH₄ gradient between Greenland and Antarctica during the Holocene. J. Geophys. Res. 102, 15987–15997.
- Cicerone, R.J., Oremland, R.S., 1988. Biogeochemical aspects of atmospheric methane. Global Biogeochem. Cycles 2, 299–327.
- Cole, C., Duxbury, J., Freney, J., Heinemeyer, O., Minami, K., Mosier, A., Paustian, K., Rosenberg, N., Sampson, N., Sauerbeck, D., Zhao, Q., 1997. Global estimates of potential mitigation of greenhouse gas emissions by agriculture. Nutr. Cycling Agroecosyst. 49, 221–228.
- Conny, J.M., Currie, L.A., 1996. The isotopic characterization of methane, non-methane hydrocarbons and formaldehyde in the troposphere. Atmos. Environ. 30, 621–638.
- Craig, H., Chou, C.C., 1982. Methane: the record in polar ice cores. Geophys. Res. Lett. 9, 1221–1224.

- Craig, H., Chou, C.C., Welhan, J.A., Stevens, C.M., Engelkemeir, A., 1988. The isotopic composition of methane in polar ice cores. Science 242, 1535–1539.
- Crowley, J.N., Saueressig, G., Bergamaschi, P., Fischer, H., Harris, G.W., 1999. Carbon kinetic isotope effect in the reaction CH₄+Cl: a relative rate study using FTIR spectroscopy. Chem. Phys. Lett. 303, 268–274.
- Crutzen, P., 1988. Tropospheric ozone: an overview. In: Isaksen, I.S.A. (Ed.), Tropospheric Ozone: Regional and Global Scale Interactions. Reidel Publishing, Boston, MA, pp. 3–11.
- Crutzen, P., 1995. Overview of tropospheric chemistry: developments during the past quarter century and a look ahead. Faraday Discuss. 100, 1–21.
- Crutzen, P.J., Andreae, M.O., 1990. Biomass burning in the tropics — Impact on atmospheric chemistry and biogeochemical cycles. Science 250, 1669–1678.
- Crutzen, P., Bruhl, C., 1993. A model study of the atmospheric temperatures and concentrations of ozone, hydroxyl, and some other photochemically active gases during the glacial, the preindustrial Holocene and the present. Geophys. Res. Lett. 20, 1047–1050.
- Crutzen, P., Zimmermann, P., 1991. The changing photochemistry of the troposphere. Tellus 43AB, 136–151.
- Crutzen, P., Aselmann, I., Seiler, W., 1986. Methane production by domestic animals, wild ruminants, and other herbivorous fauna and humans. Tellus 38B, 271.
- Czepiel, P., Mosher, B., Crill, P., Harriss, R., 1996. Quantifying the effect of oxidation on landfill methane emissions. J. Geophys. Res. 101, 16721–16729.
- Dällenbach, A., Blunier, T., Flückiger, J., Stauffer, B., Chappellaz, J., Raynaud, D., 2000. Changes in the atmospheric CH₄ gradient between Greenland and Antarctica during the Last Glacial and the transition to the Holocene. Geophys. Res. Lett. 27, 1005– 1008.
- Denier van der Gon, H., Neue, H.U., 1995. Influence of organic matter incorporation on the methane emission from a wetland rice field. Global Biogeochem. Cycles 9, 11–22.
- Dibb, J., Rasmussen, R., Mayewski, P., Holdsworth, G., 1993. Northern hemisphere concentrations of methane and nitrous oxide since 1800: results from the Mt. Logan and 20D ice cores. Chemosphere 27, 2413–2423.
- Dickens, G., Castillo, M., Walker, J., 1997. A blast of gas in the latest Paleocene: simulating first-order effects of massive dissociation of oceanic methane hydrate. Geology 25, 259–262.
- Dickinson, R.E., Cicerone, R.J., 1986. Future global warming from atmospheric trace gases. Nature 319, 109–115.
- Dlugokencky, E., Steele, L., Lang, P., Masarie, K., 1994a. The growth rate and distribution of atmospheric methane. J. Geophys. Res. 99, 17021–17043.
- Dlugokencky, E., Masaire, K., Lang, P., Tans, P., Steele, L., Nisbet, E., 1994b. A dramatic decrease in the growth rate of atmospheric methane in the northern hemisphere during 1992. Geophys. Res. Lett. 21, 45–48.
- Dlugokencky, E., Steele, L.P., Lang, P., Masarie, K., 1995. Atmospheric methane at Mauna Loa and Barrow observatories: presentation and analysis of in situ measurements. J. Geophys. Res. 100, 23103–23113.

- Dlugokencky, E., Dutton, E., Novelli, P., Tans, P., Masarie, K., Lantz, K., Madronich, S., 1996. Changes in CH₄ and CO growth rates after the eruption of Mt. Pinatubo and their link with changes in tropical tropospheric UV flux. Geophys. Res. Lett. 23, 2761–2764.
- Dlugokencky, E., Masarie, K., Tans, P., Conway, T., Xiong, X., 1997. Is the amplitude of the methane seasonal cycle changing? Atmos. Environ. 31, 21–26.
- Dlugokencky, E., Masarie, K., Lang, P., Tans, P., 1998. Continuing decline in the growth rate of the atmospheric methane burden. Nature 393, 447–450.
- Dobbie, K., Smith, K., 1996. Comparison of CH₄ oxidation rates in woodland, arable and set aside soils. Soil Biol. Biochem. 28, 1357–1365.
- Donner, L., Ramanathan, V., 1980. Methane and nitrous oxide: their effects on the terrestrial climate. J. Atmos. Sci. 37, 119– 124.
- Ehhalt, D.H., 1986. On the consequences of a tropospheric CH₄ increase to the exospheric density. J. Geophys. Res. 91, 2843.
- Environmental Protection Agency, 1993a. In: Hogan, K. (Ed.), Opportunities to Reduce Anthropogenic Methane Emissions in the United States. U.S. EPA, Office of Air and Radiation, EPA 430-R-93-012, 420 pp.
- Environmental Protection Agency, 1993b. In: Hogan, K. (Ed.), Options for Reducing Methane Emissions Internationally. U.S. EPA, Office of Air and Radiation, EPA 430-R-93-006B, 350 pp.
- Etheridge, D., Pearman, G., Fraser, P., 1992. Changes in tropospheric methane between 1841 and 1978 from a high accumulationrate Antarctic ice core. Tellus 44B, 282–294.
- Etheridge, D., Steele, L., Francey, R., Langenfelds, R., 1998. Atmospheric methane between 1000 A.D. and present: evidence of anthropogenic emissions and climatic variability. J. Geophys. Res. 103, 15979–15993.
- Fishman, J., 1985. Ozone in the troposphere. In: Whitten, R.C., Prasad, S.S. (Eds.), Ozone in the Free Atmosphere. Van Nostrand-Reinhold, New York, NY, pp. 161–194.
- Fishman, J., Solomon, S., Crutzen, P.J., 1979. Observational and theoretical evidence in support of a significant in-situ photochemical source of tropospheric ozone. Tellus 31, 432–446.
- Francey, R., Manning, M., Allison, C., Coram, S., Etheridge, D., Langenfelds, R., Lowe, D., Steele, L., 1999. A history of δ^{13} C in atmospheric CH₄ from the Cape Grim Air Archive and Antarctic firn air. J. Geophys. Res. 104, 23631–43643.
- Fuglestvedt, J., Jonson, J., Isaksen, I., 1994. Effects of reductions in stratospheric ozone on tropospheric chemistry through changes in photolysis rates. Tellus 46B, 172–192.
- Fuglestvedt, J.S., Isaksen, I.S.A., Wang, W.-C., 1996. Estimates of indirect global warming potentials for CH₄, CO and NO_x. Clim. Change 34, 405–437.
- Fuglestvedt, J.S., Bernsten, T.K., Isaksen, I.S.A., Mao, H., Liang, X.-Z., Wang, W.-C., 2000. Climatic forcing of nitrogen oxides through changes in tropospheric ozone and methane; global 3D model studies. Atmos. Environ. 33, 961–977.
- Fung, I., John, J., Lerner, J., Matthews, E., Prather, M., Steele, L., Fraser, P., 1991. Three-dimensional model synthesis of the global methane cycle. J. Geophys. Res. 96, 13033–13065.
- Gas Research Institute (GRI), 1997. Effect of Methane Emissions

on Global Warming, Appendix B. Gas Research Institute, Chicago, IL.

- Gettelman, A., Holton, J.R., Rosenlof, K.H., 1997. Mass fluxes of O₃, CH₄, N₂O and CF₂Cl₂ in the lower atmosphere calculated from observational data. J. Geophys. Res. 102, 19149–19159.
- Gupta, M., Tyler, S., Cicerone, R., 1996. Modeling atmospheric δ^{13} CH₄ and the causes of recent changes in atmospheric CH₄ amounts. J. Geophys. Res. 101, 22923–22932.
- Hameed, S., Cess, R.D., 1983. Impact of a global warming on biospheric sources of methane and its climatic consequences. Tellus 35B, 1–7.
- Hansen, A.R., Robinson, G.D., 1989. Water vapor and methane in the upper stratosphere: an examination of some of the Nimbus 7 measurements. J. Geophys. Res. 94, 8474–8484.
- Hansen, J., Fung, I., Lacis, A., Rind, D., Lebedeff, S., Ruedy, R., Russell, G., Stone, P., 1988. Global climate changes as forecast by Goddard Institute for Space studies three-dimensional model. J. Geophys. Res. 93, 9341–9364.
- Hansen, J., Lacis, A., Prather, M., 1989. Greenhouse effect of chlorofluorocarbons and other trace gases. J. Geophys. Res. 94, 16417–16421.
- Hao, W.M., Ward, D.E., 1993. Methane production from global biomass burning. J. Geophys. Res. 98, 20657–20661.
- Harper, L.A., Denmead, O.T., Freney, J.R., Byers, F.M., 1999. Direct measurements of methane emissions from grazing and feedlot cattle. J. Anim. Sci. 77, 1392–1401.
- Harvey, L.D.D., Huang, Z., 1995. Evaluation of the potential impact of methane clathrate destabilization on future global warming. J. Geophys. Res. 100, 2905–2926.
- Hauglustaine, D., Granier, C., Brasseur, G., Megie, G., 1994. The importance of atmospheric chemistry in the calculation of radiative forcing on the climate system. J. Geophys. Res. 99, 1173– 1186.
- Hayhoe, K., 1997. A modelling study of the role of methane in global climate change. MS Thesis, University of Illinois, Urbana, IL, 146 pp.
- Hayhoe, K., Jain, A., Pitcher, H., MacCracken, C., Gibbs, M., Wuebbles, D., Harvey, R., Kruger, D., 1999. Costs of multigreenhouse gas reduction targets for the U.S. Science 286, 905– 906.
- Hayhoe, K., Jain, A., Kheshgi, H., Wuebbles, D., 2000. Contribution of CH₄ to multi-gas reduction targets: the impact of atmospheric chemistry on GWPs. In: van Ham, J. (Ed.), Non-CO₂ Greenhouse Gases: Scientific Understanding, Control and Implementation. Kluwer Academic Publishing, Dordrecht, pp. 425–432.
- Hein, R., Crutzen, P.J., Heinman, R., 1997. An inverse modeling approach to investigate the global atmospheric methane cycle. Global Biogeochem. Cycles 11, 43–76.
- Hilbert, D.W., Roulet, N., Moore, T., 2000. Modelling and analysis of peatlands as dynamical systems. J. Ecol. 88, 230–242.
- Hofzumahaus, A., Dorn, H.P., Callies, J., Platt, U., Ehhalt, D., 1991. Tropospheric OH concentration measurements by laser long-path absorption spectroscopy. Atmos. Environ. 25A, 2017–2022.
- Hogan, K., Harriss, R.C., 1994. Comments on "A dramatic decrease in the growth rate of atmospheric methane in the northern hemisphere during 1992" by E.J. Dlugokencky et al. Geophys. Res. Lett. 21, 2445–2447.

- Holmes, M.E., Sansone, F.J., Rust, T.M., Popp, B.N., 2000. Methane production, consumption, and air-sea exchange in the open ocean: an evaluation based on carbon isotopic ratios. Global Biogeochem. Cycles 14, 1–10.
- Hough, A.M., Derwent, R.G., 1990. Changes in the global concentration of tropospheric ozone due to human activities. Nature 344, 645–648.
- Houweling, S., Kaminski, T., Dentener, F., Lelieveld, J., Heimann, M., 1999. Inverse modeling of methane sources and sinks using the adjoint of a global transport model. J. Geophys. Res. 104, 26137–26160.
- Huang, Y., Sass, R.L., Fisher, F.M., 1997. Methane emission from Texas rice paddy soils: 1. Quantitative multi-year dependence of CH₄ emission on soil, cultivar and grain yield. Global Change Biol. 3, 479–489.
- Huang, Y., Sass, R.L., Fisher, F.M., 1998. A semi-empirical model of methane emission from flooded rice paddy soils. Global Change Biol. 4, 247–268.
- Hudgens, D., Yavitt, J., 1997. Land-use effects on soil methane and carbon dioxide fluxes in forests near Ithaca, New York. Ecoscience 4, 214–222.
- Intergovernmental Panel on Climate Change, 1990. In: Houghton, J.T., Jenkins, G.J., Ephraums, J.J. (Eds.), Climate Change: The IPCC Scientific Assessment. Cambridge Univ. Press, Cambridge, UK.
- Intergovernmental Panel on Climate Change, 1995. In: Houghton, J.T., Meira Filho, L., Bruce, J., Lee, H., Callander, B., Haites, E., Harris, H., Maskell, K. (Eds.), Climate Change 1994. Cambridge Univ. Press, Cambridge, UK, 339 pp.
- Intergovernmental Panel on Climate Change, 1996. In: Houghton, J.T., Meira Filho, L.G., Callander, B.A., Harris, N., Kattenberg, A., Maskell, K. (Eds.), Climate Change 1995: The Science of Climate Change. Cambridge Univ. Press, Cambridge, UK, 572 pp.
- Isaksen, I.S.A., 1988. Is the oxidizing capacity of the atmosphere changing? In: Rowland, F.S., Isaksen, I.S.A. (Eds.), The Changing Atmosphere. Wiley, New York, NY, pp. 141–157.
- Isaksen, I.S.A., Stordal, F., 1986. Ozone perturbations by enhanced levels of CFCs, N₂O, and CH₄: a two-dimensional diabatic circulation study including uncertainty estimates. J. Geophys. Res. 91, 5249–5263.
- Jain, A.K., Briegleb, B.P., Minschwaner, K., Wuebbles, D.J., 2000. Radiative forcings and global warming potentials of 39 greenhouse gases. J. Geophys. Res. 105, 20773–20790.
- Joabsson, A., Christensen, T.R., Wallen, B., 1999. Vascular plant controls on methane emissions from northern peatforming wetlands. Trends Ecol. Evol. 14, 385–388.
- Johnson, D., Johnson, K., Ward, G.M., Branine, M., 2000. Ruminants and other animals. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 112–133.
- Jouzel, J., Jouzel, N.I., Barkov, J.M., Barnola, M., Bender, J., Chappellaz, C., Genthon, V.M., Kotlyakov, V., Lipenkov, C., Lorius, J.R., Petit, D., Raynaud, G., Raisbeck, C., Ritz, T., Sowers, M., Stievenard, F., Yiou, F., Yiou, P., 1993. Extending the Vostok ice-core record of palaeoclimate to the penultimate glacial period. Nature 364, 407–412.

- Judd, A.G., 2000. Geological sources of methane. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 280–303.
- Kandlikar, M., 1997. Bayesian inversion for reconciling uncertainties in global mass balances. Tellus 49B, 123–135.
- Kandlikar, M., McRae, G.J., 1995. Inversion of the global methane cycle using change constrained programming: methodology and results. Chemosphere 30, 1151–1170.
- Karlsdottir, S., Isaksen, I.S.A., 2000. Changing methane lifetime: possible cause for reduced growth. Geophys. Res. Lett. 27, 93– 96.
- Khalil, M.A.K., 2000. Atmospheric methane: an introduction. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 1–8.
- Khalil, M.A.K., Rasmussen, R.A., 1983. Sources, sinks and seasonal cycles of atmospheric methane. J. Geophys. Res. 88, 5131– 5144.
- Khalil, M.A.K., Rasmussen, R.A., 1985. Causes of increasing atmospheric methane: depletion of hydroxyl radicals and the rise of emissions. Atmos. Environ. 13, 397–407.
- Khalil, M., Rasmussen, R., 1987. Atmospheric methane: trends over the last 10,000 years. Atmos. Environ. 21, 2445–2452.
- Khalil, M., Rasmussen, R., 1993. Decreasing trend of methane: unpredictability of future concentrations. Chemosphere 26, 803– 814.
- Khalil, M., Rasmussen, R., 1994a. Global emissions of methane during the last several centuries. Chemosphere 29, 833–842.
- Khalil, M., Rasmussen, R., 1994b. Trends in atmospheric methane. Pure and Applied Chemistry. 66, Special Report: Methane in the Atmosphere, Commission on Atmospheric Chemistry.
- Khalil, M., Shearer, M.J., 2000. Sources of methane: an overview. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 98–111.
- Khalil, M., Rasmussen, R., Moraes, F., 1993a. Atmospheric methane at Cape Meares: analysis of a high-resolution data base and its environmental implications. J. Geophys. Res. 98, 14753–14770.
- Khalil, M., Rasmussen, R., Shearer, M., Ge, S., Rau, J., 1993b. Methane from coal burning. Chemosphere 26, 473–477.
- Khalil, M., Rasmussen, R., Shearer, M., Dalluge, R., Ren, L., Duan, C., 1998. Factors affecting methane emissions from rice fields. J. Geophys. Res. 103, 25219–25231.
- Kheshgi, H.S., Jain, A.K., Kotamarthi, V.R., Wuebbles, D.J., 1999. Future atmospheric methane concentrations in the context of the stabilization of greenhouse gas concentrations. J. Geophys. Res. 104, 19183–19190.
- King, G., 1997. Responses of atmospheric methane consumption by soils to global climate change. Global Change Biol. 3, 351– 362.
- King, S.L., Quay, P.D., Lansdown, J.M., 1989. The ¹³C/¹²C kinetic isotope effect for soil oxidation of methane at ambient atmospheric concentrations. J. Geophys. Res. 94, 18273–18277.
- King, J.Y., Reeburgh, W.S., Regli, S.K., 1998. Methane emission and transport by arctic sedges in Alaska: results of a vegetation removal experiment. J. Geophys. Res. 103, 29029–29083.
- Kirchgessner, D.A., 2000. Fossil fuel industries. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 263–279.

- Kirchgessner, D.A., Piccot, S.D., Winkler, J.D., 1993. Estimate of global methane emissions from coal mines. Chemosphere 26, 453–472.
- Krol, M., van Leeuwen, P., Lelieveld, J., 1998. Global OH trend inferred from methylchloroform measurements. J. Geophys. Res. 103, 10697–10711.
- Lacroix, A.V., 1993. Unaccounted-for sources of fossil and isotopically-enriched methane and their contribution to the emissions inventory—A review and synthesis. Chemosphere 26, 507– 557.
- Lashof, D., 1989. The dynamic greenhouse: feedback processes that may influence future concentrations of atmospheric trace gases and climatic change. Clim. Change 14, 213–242.
- Lassey, K.R., Lowe, D.C., Brenninkmeijer, C.A.M., Gomez, A.J., 1993. Atmospheric methane and its carbon isotopes in the Southern Hemisphere: their time series and an instructive model. Chemosphere 26, 95–109.
- Lassey, K.R., Lowe, D.C., Manning, M.R., 1999. The trend in atmospheric methane δ^{13} C and implications for isotopic constraints on the global methane budget. Global Biogeochem. Cycles 14, 41–49.
- Law, K.S., Nisbet, E.G., 1996. Sensitivity of the CH₄ growth rate to changes in CH₄ emissions from natural gas and coal. J. Geophys. Res. 101, 14387–14397.
- Legrand, M., Lorius, C., Barkov, N., Petrov, V., 1988. Vostok (Antarctic ice core): atmospheric chemistry changes over the last climatic cycle (160,000 years). Atmos. Environ. 22, 317–331.
- Lelieveld, J., Crutzen, P.J., 1991. The role of clouds in tropospheric photochemistry. J. Atmos. Chem. 12, 229–267.
- Lelieveld, J., Crutzen, P.J., 1992. Indirect chemical effects of methane on global warming. Nature 355, 339–342.
- Lelieveld, J., Crutzen, P., Bruhl, C., 1993. Climate effects of atmospheric methane. Chemosphere 26, 739–767.
- Lelieveld, J., Crutzen, P., Dentener, F., 1998. Changing concentration, lifetime and climate forcing of atmospheric methane. Tellus 50B, 128–150.
- Le Texier, L., Solomon, S., Garcia, R.R., 1988. The role of molecular hydrogen and methane oxidation in the water vapor budget of the stratosphere. Q. J. R. Meteorol. Soc. 114, 281–296.
- Levine, J.S., Cofer III, W.R., Pinto, J.P., 2000. Biomass burning. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 190–201.
- Lowe, D.C., Schmidt, U., 1983. Formaldehyde (HCHO) measurements in the nonurban atmosphere. J. Geophys. Res. 88, 10844– 10858.
- Lowe, D.C., Brenninkmeijer, C.A.M., Brailsford, G.W., Lassey, K.R., Gomez, A.J., 1994. Concentration and ¹³C records of atmospheric methane in New Zealand and Antarctica: evidence for changes in methane sources. J. Geophys. Res. 99, 16913– 16925.
- Lowe, D., Manning, M., Brailsford, G., Bromley, A., 1997. The 1991–1992 atmospheric methane anomaly: Southern Hemisphere ¹³C increase and growth rate fluctuations. Geophys. Res. Lett. 24, 857–860.
- Lowe, D.C., Allen, W., Manning, M.R., Bromley, T., Brailsford, G., Ferretti, D., Gomez, A., Knobben, R., Martin, R., Mei, Z., Moss, R., Koshy, K., Maata, M., 1999. Shipboard determinations of

the distribution of δ^{13} C in atmospheric methane in the Pacific. J. Geophys. Res. 104, 23135–26125.

- Lu, Y., Khalil, M., 1991. Tropospheric OH: model calculations of spatial, temporal, and secular variations. Chemosphere 23, 397–444.
- MacDonald, G.J., 1990. Role of methane clathrates in past and future climates. Clim. Change 16, 247–281.
- MacKay, R.M., Khalil, M.A.K., 1991. Theory and development of a one dimensional time dependent radiative convective climate model. Chemosphere 22, 383–417.
- Mancinelli, R., 1995. The regulation of methane oxidation in soil. Annu. Rev. Microbiol. 49, 581–605.
- Manning, M.R., Lowe, D.C., Melhuish, W.H., Sparks, R.J., Wallace, G., Brenninkmeijer, C.A.M., McGill, R.C., 1990. The use of radiocarbon measurements in atmospheric studies. Radiocarbon 32, 37–58.
- Matthews, E., 1994. Assessment of methane sources and their uncertainties. Pure Appl. Chem. 66, 154–161.
- Matthews, E., 2000. Wetlands. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 202–233.
- Miller, D.N., Ghiorse, W.C., Yavitt, J.B., 1999. Seasonal patterns and controls on methane and carbon dioxide fluxes in forested swamp pools. Geomicrobiol. J. 16, 325–331.
- Minami, K., 1997. Atmospheric methane and nitrous oxide: sources, sinks and strategies for reducing agricultural emissions. Nutr. Cycling Agroecosyst. 49, 203–211.
- Minami, K., Neue, H.U., 1994. Rice paddies as a methane source. Clim. Change 27, 13–26.
- Minschwaner, K., Carver, R.W., Briegleb, B.P., Roche, A.E., 1998. Infrared radiative forcing and atmospheric lifetimes of trace species based on observations from UARS. J. Geophys. Res. 103, 23243–23253.
- Mitchell, C., 1993. Methane emissions from the coal and natural gas industries in the UK. Chemosphere 26, 441–446.
- Mitra, S., Jain, M., Kumar, S., Bandyopadhyay, S., Kalra, N., 1999. Effect of rice cultivars on methane emission. Agricult., Ecosyst. Environ. 73, 177–183.
- Moosavi, S.C., Crill, P.M., 1997. Controls on CH₄ and CO₂ emissions along two moisture gradients in the Canadian boreal zone. J. Geophys. Res. 102, 29261–29277.
- Moosavi, S.C., Crill, P.M., 1998. CH₄ oxidation by tundra wetlands as measured by a selective inhibitor technique. J. Geophys. Res. 103, 29093–29106.
- Moosavi, S.C., Crill, P.M., Pullman, E.R., Funk, D.W., Peterson, K.M., 1996. Controls on CH₄ flux from an Alaskan boreal wetland. Global Biogeochem. Cycles 10, 287–296.
- Mosier, A., Delgado, J., Cochran, V., Valentine, D., Parton, W., 1997a. Impact of agriculture on soil consumption of atmospheric CH₄ and a comparison of CH₄ and N₂O flux in subarctic, temperature and tropical grasslands. Nutr. Cycling Agroecosyst. 49, 71–83.
- Mosier, A., Parton, W., Valentine, D., Ojima, D., Schimel, D., Heinemeyer, O., 1997b. CH₄ and N₂O fluxes in the Colorado shortgrass steppe: 2. Long-term impact of land use change. Global Biogeochem. Cycle 11, 29–42.

Nakano, T., Kuniyoshi, S., Fukuda, M., 2000. Temporal variation in

methane emission from tundra wetlands in a permafrost area, northeastern Siberia. Atmos. Environ. 34, 1205-1213.

- Nakazawa, T., Machida, T., Tanaka, M., Fujii, Y., Aoki, S., Watanabe, O., 1993. Differences of the atmospheric CH₄ concentration between the Arctic and Antarctic regions in preindustrial/pre-agricultural era. Geophys. Res. Lett. 20, 943– 946.
- Neue, H.-U., Roger, P.A., 2000. Rice agriculture: factors controlling emissions. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 134–169.
- Neue, H.-U., Wassmann, R., Kludze, H., Bujun, W., Lantin, R., 1997. Factors and processes controlling methane emissions from rice fields. Nutr. Cycling Agroecosyst. 49, 111–117.
- Nisbet, E., 1990a. Did the release of methane from hydrates accelerate the end of the last ice age? Can. J. of Earth Science 27, 148–157.
- Nisbet, E., 1990b. Climate change and methane. Nature 347, 23.
- Oberbauer, S., Starr, G., Pop, E., 1998. Effects of extended growing season and soil warming on carbon dioxide and methane exchange of tussock tundra in Alaska. J. Geophys. Res. 103, 29075–29082.
- Osborn, T., Wigley, T., 1994. A simple model for estimating methane concentration and lifetime variations. Clim. Dyn. 9, 181–193.
- Otter, L.B., Scholes, M.C., 2000. Methane sources and sinks in a periodically flooded South African savanna. Global Biogeochem. Cycles 14, 97–111.
- Owens, A.J., Steed, J.M., Filkin, D.L., Miller, C., Jesson, J.P., 1982. The potential effects of increased methane on atmospheric ozone. Geophys. Res. Lett. 9, 1105–1108.
- Owens, A.J., Hales, C.H., Filkin, D.L., Miller, C., Steed, J.M., Jesson, J.P., 1985. A coupled one-dimensional radiative-convective, chemistry-transport model of the atmosphere: 1. Model structure and steady state perturbation calculations. J. Geophys. Res. 90, 2283–2311.
- Panikov, N.S., 1999. Fluxes of CO₂ and CH₄ in high latitude wetlands: measuring, modelling and predicting response to climate change. Polar Res. 18, 237–244.
- Peer, R., Thorneloe, S., Epperson, D., 1993. A comparison of methods for estimating global methane emissions from landfills. Chemosphere 26, 387–400.
- Penkett, S.A., 1988. Indications and causes of ozone increase in the troposphere. In: Rowland, F.S., Isaksen, I.S.A. (Eds.), The Changing Atmosphere. Wiley, New York, NY, pp. 91–103.
- Petit, J., Jouzel, J., Raynaud, D., Barkov, N., Barnola, J.-M., Basile, I., Bender, M., Chappellaz, J., Davis, M., Delaygue, G., Delmotte, M., Kotlyakov, V., Legrand, M., Lipenkov, V., Lorius, C., Pepin, L., Ritz, C., Saltzman, E., Stievenard, M., 1999. Climate and atmospheric history of the past 420,000 years from the Vostok ice core, Antarctica. Nature 399, 429–436.
- Piccot, S.D., Beck, L., Srinivasan, S., Kersteter, S.L., 1996. Global methane emissions from minor anthropogenic sources and biofuel combustion in residential stoves. J. Geophys. Res. 101, 22757–22766.
- Pinto, J., Khalil, M., 1991. The stability of tropospheric OH during ice ages, inter-glacial epochs and modern times. Tellus 43B, 347–352.

- Poppe, D., Zimmerman, J., Dorn, H., 1995. Field data and model calculations for the hydroxyl radical. J. Atmos. Sci. 52, 3402– 3407.
- Powlson, D., Goulding, K., Willison, T., Webster, C., Hutsch, B., 1997. The effect of agriculture on methane oxidation in soil. Nutr. Cycling Agroecosyst. 49, 59–70.
- Prather, M., 1994. Lifetimes and eigenstates in atmospheric chemistry. Geophys. Res. Lett. 21, 801–804.
- Prieme, A., Christensen, S., Dobbie, K., Smith, K., 1997. Slow increase in rate of methane oxidation in soils with time following land use change from arable agriculture to woodland. Soil Biol. Biochem. 29, 1269–1273.
- Prinn, R.G., Weiss, R.F., Miller, B.R., Huang, J., Alyea, F.N., Cunnold, D.M., Fraser, P.J., Hartley, D.E., Simmonds, P.G., 1995. Atmospheric trends and lifetime of CH₃CCl₃ and global OH concentrations. Science 269, 187–198.
- Quay, P.D., King, S.L., Lansdown, J.M., Wilbur, D.O., 1988. Isotopic composition of methane released from wetlands: implications for the increase in atmospheric methane. Global Biogeochem. Cycles 2, 385–397.
- Quay, P.D., King, S.L., Stutsman, J., Wilbur, D.O., Steele, L.P., Fung, I., Gammon, R.H., Brown, T.A., Farwell, G.W., Grootes, P.M., Schmidt, F.H., 1991. Carbon isotopic composition of atmospheric CH₄: fossil and biomass burning source strengths. Global Biogeochem. Cycles 5, 25–47.
- Quay, P., Stutsman, J., Wilbur, D., Snover, A., Dlugokencky, E., Brown, T., 1999. The isotopic composition of atmospheric methane. Global Biogeochem. Cycles 13, 445–461.
- Rahmes, T.F., Omar, A.H., Wuebbles, D.J., 1998. Atmospheric distributions of soot particles by current and future aircraft fleets and resulting radiative forcing on climate. J. Geophys. Res. 103, 31657–31667.
- Ramanathan, V., 1988. The radiative and climatic consequences of the changing atmospheric composition of trace gases. In: Rowland, F.S., Isaksen, I.S.A. (Eds.), The Changing Atmosphere. Wiley, New York, NY, pp. 159–186.
- Ramanathan, V., Cicerone, R.J., Singh, H.B., Kiehl, J.T., 1985. Trace gas trends and their potential role in climate change. J. Geophys. Res. 90, 5547–5566.
- Ramanathan, V., Callis, L., Cess, R., Hansen, J., Isaksen, I., Kuhn, W., Lacis, A., Luther, F., Mahlman, J., Reck, R., Schlesinger, M., 1987. Climate-chemical interactions and effects of changing atmospheric trace gases. Rev. Geophys. 25, 1441–1482.
- Randel, W.J., Wu, F., Russell III, J.M., Roche, A., Waters, J.W., 1998. Seasonal cycles and QBO variations in stratospheric CH₄ and H₂O observed in UARS HALOE data. J. Atmos. Sci. 55, 163–185.
- Randel, W., Wu, F., Russell III, J., Waters, J. 1999. Space-time patterns of trends in stratospheric constituents derived from UARS measurements. J. Geophys. Res. 104, 3711-3727.
- Rasmussen, R.A., Khalil, M.A.K., 1981. Atmospheric methane (CH₄): trends and seasonal cycles. J. Geophys. Res. 86, 9826– 9832.
- Raynaud, D., Chappellaz, J., Barnola, J., Korotkevich, Y., Lorius, C., 1988. Climatic and CH4 cycle implications of glacial-interglacial CH₄ change in the Vostok ice core. Nature 333, 655–657.
- Raynaud, D., Jouzel, J., Barnola, J., Chapellaz, J., Delmas, R.,

Lorius, C., 1993. The ice record of greenhouse gases. Science 259, 926–934.

- Rinsland, C., Levine, J., Miles, T., 1985. Concentration of methane in the troposphere deduced from 1951 infrared solar spectra. Nature 318, 245–249.
- Roble, R.G., Dickinson, R.E., 1989. How will changes in carbon dioxide and methane modify the mean structure of the mesosphere and thermosphere? Geophys. Res. Lett. 16, 1441–1444.
- Rodhe, H., 1990. A comparison of the contribution of various gases to the greenhouse effect. Science 248, 1217–1219.
- Roulet, N., Moore, T., Bubier, J., Lafleur, P., 1992. Northern fens: methane flux and climatic change. Tellus 44B, 100–105.
- Roulet, N., Jano, A., Kelly, C., Klinger, L., Moore, T., Protz, R., Ritter, J., Rouse, W., 1993. The Hudson Bay Lowland as a source of atmospheric methane. J. Geophys. Res. 99, 1439–1454.
- Sanderson, M.G., 1996. Biomass of termites and their emissions of methane and carbon dioxide: a global database. Global Biogeochem. Cycles 10, 543–557.
- Sass, R.L., Fisher, F.M., 1997. Methane emissions from rice paddies: a process study summary. Nutr. Cycling Agroecosyst. 49, 119–127.
- Sass, R.L., Fisher, F.M., Ding, A., Huang, Y., 1999. Exchange of methane from rice fields: national, regional, and global budgets. J. Geophys. Res. 104, 26943–26951.
- Saueressig, G., Bergamaschi, P., Crowley, J.N., Fischer, H., Harris, G.W., 1995. Carbon kinetic isotope effect in the reaction of CH₄ with Cl atmos. Geophys. Res. Lett. 22, 1225–1228.
- Schauffler, S., Daniel, J., 1994. On the effect of stratospheric circulation changes on trace gas trends. J. Geophys. Res. 99, 25747–25754.
- Seiler, W., Crutzen, P.J., 1980. Estimates of gross and net fluxes of carbon between the biosphere and the atmosphere from biomass burning. Climatic Change 2, 207–247.
- Severinghaus, J., Sowers, T., Brook, E., Aley, R., Bender, M., 1998. Timing of abrupt climate change at the end of the Younger Dryas interval from thermally fractionated gases in polar ice. Nature 391, 141–146.
- Shearer, M.J., Khalil, M.A.K., 2000. Rice agriculture: emissions. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 170–189.
- Sigren, L., Lewis, S., Fisher, F., Sass, R., 1997. Effects of field drainage on soil parameters related to methane production and emission from rice paddies. Global Biogeochem. Cycles 11, 151–162.
- Slanina, J., Warneck, P., Bazhin, N., Akimoto, H., Kieskamp, W., Khalil, M., Calvert, J., Matthews, E., Barrie, L., Wahlen, M., Schwartz, S., Tang, X., Singh, O., 1994. Assessment of uncertainties in the projected concentrations of methane in the atmosphere. Pure Appl. Chem. 66, 137–140.
- Smith, L.K., Lewis, W.M., Chanton, J.P., Cronin, G., Hamilton, S.K., 2000. Methane emissions from the Orinoco River floodplain, Venezuela. Biogeochemistry 51, 113–140.
- Snover, A.K., Quay, P.D., 2000. Hydrogen and carbon kinetic isotope effects during soil uptake of atmospheric methane. Global Biogeochem. Cycles 14, 25–39.
- Steele, L., Dlugokencky, E., Lang, P., Tans, P., Martin, R., Masarie, K., 1992. Slowing down of the global accumulation of atmospheric methane during the 1980s. Nature 358, 313–316.

- Stern, D., Kaufmann, R., 1996. Estimates of global anthropogenic methane emissions 1860–1993. Chemosphere 33, 159–176.
- Stevens, C., 1993. Isotopic abundance in the atmosphere and sources. In: Khalil, M.A.K. (Ed.), Atmospheric Methane, Sources, Sinks and Role in Global Change. Springer-Verlag, New York, pp. 62–88.
- Stevens, C.M., Engelkemeir, A., 1988. Stable carbon isotopic composition of methane from some natural and anthropogenic sources. J. Geophys. Res. 93, 725–733.
- Stevens, C.M., Rust, F.E., 1982. The carbon isotopic composition of atmospheric methane. J. Geophys. Res. 87, 4879–4882.
- Stevens, C.M., Wahlen, M., 2000. The isotopic composition of athmospheric methane and its sources. In: Khalil, M. (Ed.), Atmospheric Methane: Its role in the Global Environment. Springer-Verlag, New York, pp. 25–41.
- Stordal, F., Isaksen, I.S.A., 1987. Ozone perturbations due to increases in N₂O, CH₄, and chlorocarbons: two-dimensional timedependent calculations. Tellus 39B, 333–353.
- Subak, S., 1994. Methane from the House of Tudor and the Ming Dynasty: anthropogenic emissions in the sixteenth century. Chemosphere 29, 843–854.
- Sugimoto, A., Inoue, T., Kirtibutr, N., Abe, T., 1998. Methane oxidation by termite mounds estimated by the carbon isotopic composition of methane. Global Biogeochem. Cycles 12, 595–605.
- Tang, X., Madronich, S., Wallington, T., Calamari, D., 1998. Changes in tropospheric composition and air quality. J. Photochem. Photobiol., B 46, 83–95.
- Tans, P.P., 1997. A note on isotopic ratios and the global methane budget. Global Biogeochem. Cycles 11, 77–81.
- Thomas, G., Olivero, J., Jensen, E., Schroeder, W., Toon, O., 1989. Relation between increasing methane and the presence of ice clouds at the mesopause. Nature 338, 490–492.
- Thompson, A., 1992. The oxidizing capacity of the Earth's atmosphere: probable past and future changes. Science 256, 1157–1165.
- Thompson, A., Cicerone, R., 1986. Atmospheric CH₄, CO and OH from 1860 to 1985. Nature 321, 148–150.
- Thompson, A., Stewart, R., Owens, M., Herwehe, J., 1989. Sensitivity of tropospheric oxidants to global chemical and climate change. Atmos. Environ. 23, 519–532.
- Thompson, A., Huntley, M., Stewart, R., 1990. Perturbations to tropospheric oxidants, 1985–2035: calculations of ozone and OH in chemically coherent regions. J. Geophys. Res. 95, 9829– 9844.
- Thompson, A., Chappellaz, J., Fung, I., Kucsera, T., 1993. The atmospheric CH₄ increase since the Last Glacial Maximum: 2. Interactions with oxidants. Tellus 45B, 242–257.
- Thorneloe, S.A., Barlaz, M.A., Peer, R., Huff, L.C., Davis, L., Mangino, J., 2000. Waste management. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 234–262.
- Thorpe, R., Law, K., Bekki, S., Pyle, J., Nisbet, E., 1996. Is methane-driven deglaciation consistent with the ice core record? J. Geophys. Res. 101, 28627–28635.
- Tyler, S.C., 1986. Stable carbon isotope ratios in atmospheric methane and some of its sources. J. Geophys. Res. 91, 13232– 13238.

- Tyler, S.C., 1989. ¹³C/¹²C ratios in atmospheric methane and some of its sources. Stable Isotopes in Ecological Research. Springer-Verlag, New York, pp. 395–409.
- Tyler, S.C., 1999. Measurements and modeling of atmospheric methane using stable carbon isotopes. IGACtivities, pp. 3–7, March.
- Tyler, S.C., Blake, D.R., Rowland, F.S., 1987. ¹³C/¹²C ratio in methane from the flooded amazon forest. J. Geophys. Res. 92, 1044–1048.
- Tyler, S.C., Crill, P.M., Brailsford, G.W., 1994. ¹³C/¹²C fractionation of methane during oxidation in a temperate forested soil. Geochim. Cosmochim. Acta 58, 1625–1633.
- Tyler, S.C., Ajie, H.O., Gupta, M.L., Cicerone, R.J., Blake, D.R., Dlugokencky, E.J., 1999. Stable carbon isotopic composition of atmospheric methane: a comparison of surface level and free tropospheric air. J. Geophys. Res. 104, 13895–13910.
- Tyler, S.C., Ajie, H.O., Rice, A.L., Cicerone, R.J., Tuazon, E.C., 2000. Experimentally determined kinetic isotope effects in the reaction of CH₄ with Cl: implications for atmospheric CH₄. Geophys. Res. Lett. 27, 1715–1718.
- UNFCCC, 1997. Kyoto Protocol to the United Nations Framework Convention on Climate Change. UNFCCC/CP/1997/L.7/Add.1, United Nations.
- Van den Pol-Van Dasselaar, A., Van Beusichem, M.L., Oenema, O., 1999. Methane emissions from wet grasslands on peat soil in a nature preserve. Biogeochemistry 44, 205–220.
- Van der Nat, F.J., Middelburg, J.J., 2000. Methane emission from tidal freshwater marshes. Biogeochemistry 49, 103–121.
- Velichko, A., Kremenetski, C., Borisova, O., Zelikson, E., Nechaev, V., Faure, H., 1998. Estimates of methane emission during the last 125,000 years in Northern Eurasia. Global Planet. Change 16–17, 159–180.
- Wahlen, S.C., Reeburgh, W.S., 1992. Interannual variations in tundra methane emission: a 4-year time-series at fixed sites. Global Biogeochem. Cycles 6, 139–159.
- Wahlen, M., Tanaka, N., Henry, R., Deck, B., Zeglen, J., Vogel, J.S., Southon, J., Shemesh, A., Fairbanks, R., Broecker, W.S., 1989. Carbon-14 in methane sources and in atmospheric methane: the contribution from fossil carbon. Science 245, 286–290.
- Wang, Y., Jacob, D.J., 1998. Anthropogenic forcing on tropospheric ozone and OH since preindustrial times. J. Geophys. Res. 103, 31123–31135.
- Wang, W.C., Molnar, G., 1985. A model study of the greenhouse effects due to increasing atmospheric CH₄, N₂O, CF₂C₁₂ and CFCl₃. J. Geophys. Res. 90, 12971–12980.
- Wang, W.C., Wuebbles, D.J., Washington, W.M., Isaacs, R.G., Molnar, G., 1986. Trace gases and other potential perturbations to global climate. Rev. Geophys. 24, 110–140.
- Wang, W.C., Dudek, M.P., Liang, X.Z., Kiehl, J.T., 1991. Inadequacy of effective CO₂ as a proxy in simulating the greenhouse effect of other radiatively active gases. Nature 350, 573–577.
- Wang, C., Prinn, R., Sokolov, A., 1998. A global interactive chemistry and climate model: formulation and testing. J. Geophys. Res. 103, 3399–3417.
- Ward, G., Doxtader, K., Miller, W., Johnson, D., 1993. Effects of intensification of agricultural practices on emission of greenhouse gases. Chemosphere 26, 87–93.

- Whiticar, M.J., 2000. Can stable isotopes and global budgets be used to constrain atmospheric methane budgets? In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 63–85.
- Wickland, K.P., Striegl, R.G., Schmidt, S.K., Mast, M.A., 1999. Methane flux in subalpine wetland and unsaturated soils in the southern Rocky Mountains. Global Biogeochem. Cycles 13, 101–113.
- Wigley, T.M.L., 1987. Relative contributions of different trace gases to the greenhouse effect. Clim. Monit. 16, 14–28.
- World Meteorological Organization, 1985. Scientific Assessment of Ozone Depletion: 1985. Global Ozone and Research and Monitoring Project Report 16, Geneva.
- World Meteorological Organization, 1989. Scientific Assessment of Ozone Depletion: 1989. Global Ozone and Research and Monitoring Project Report 20, Geneva.
- World Meteorological Organization, 1991. Scientific Assessment of Ozone Depletion: 1991. Global Ozone and Research and Monitoring Project Report 25, Geneva.
- World Meteorological Organization, 1995. Scientific Assessment of Ozone Depletion: 1994. Global Ozone and Research And Monitoring Project Report 37, Geneva.
- World Meteorological Organization, 1999. Scientific Assessment of Ozone Depletion: 1998. Global Ozone and Research And Monitoring Project Report 44, Geneva.

- Worthy, D., Levin, I., Hopper, F., Ernst, M., Trivett, N., 2000. Evidence for a link between climate and northern wetland methane emissions. J. Geophys. Res. 105, 4031–4038.
- Wuebbles, D.J., Luther, F.M., Penner, J.E., 1983. Effect of coupled anthropogenic perturbations on stratospheric ozone. J. Geophys. Res. 88, 1444–1456.
- Wuebbles, D.J., Wei, C.-F., Patten, K.O., 1998. Effects on stratospheric ozone and temperature during the Maunder Minimum. Geophys. Res. Lett. 25, 523–526.
- Wuebbles, D., Hayhoe, K., Kotamarthi, R., 2000. Atmospheric methane in the global environment. In: Khalil, M. (Ed.), Atmospheric Methane: Its Role in the Global Environment. Springer-Verlag, New York, NY, pp. 304–341.
- Yagi, K., Tsuruta, H., Minami, K., 1997. Possible options for mitigating methane emission from rice cultivation. Nutr. Cycling Agroecosyst. 49, 213–220.
- Yavitt, J.B., Williams, C.J., Wieder, R.K., 2000. Controls on microbial production of methane and carbon dioxide in three Sphagnum-dominated peatland ecosystems as revealed by a reciprocal field peat transplant experiment. Geomicrobiol. J. 17, 61–88.
- Zoger, M., Engel, A., McKenna, D.S., Schiller, C., Schmidt, U., Woyke, T., 1999. Balloon-borne in situ measurements of stratospheric H₂O, CH₄ and H₂ at midlatitudes. J. Geophys. Res. 104, 1817–1825.