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CarbonTracker-CH₄: an assimilation system for estimating emissions of atmospheric methane

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Discussion Paper

Discussion Paper

Discussion Pape

CarbonTracker-CH₄

ACPD

14, 2175–2233, 2014

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures













Full Screen / Esc

Printer-friendly Version



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We describe an assimilation system for atmospheric methane (CH₄), CarbonTracker-CH₄, and demonstrate the diagnostic value of global or zonally averaged CH₄ abundances for evaluating the results. We show that CarbonTracker-CH₄ is able to simulate the observed zonal average mole fractions and capture inter-annual variability in emissions quite well at high northern latitudes (53-90° N). CarbonTracker-CH₄ estimates of total fluxes at high northern latitudes are about 81 TgCH₄ yr⁻¹, about 12 TgCH₄ yr⁻¹ (13%) lower than prior estimates, a result that is consistent with other atmospheric inversions. Emissions from European wetlands are decreased by 30 %, a result consistent with previous; however, emissions from wetlands in Boreal Eurasia are increased relative to the prior estimate. Although CarbonTracker-CH₁ does not estimate increases in emissions from high northern latitudes for 2000 through 2010, significant inter-annual variability in high northern latitude fluxes is recovered. During the exceptionally warm Arctic summer of 2007, estimated emissions were greater than the decadal average by 4.4 Tg CH₄ yr⁻¹. In 2008, temperatures returned to more normal values over Arctic North America while they stayed above normal over Arctic Eurasia. CarbonTracker-CH₄ estimates were 2.4 Tg CH₄ yr⁻¹ higher than the decadal average, and the anomalous emissions occurred over Arctic Eurasia, suggesting that the data allow discrimination between these two source regions. Also, the emission estimates respond to climate variability without having the system constrained by climate parameters.

CarbonTracker- CH_4 estimates for temperate latitudes are only slightly increased over prior estimates, but about $10\, Tg\, CH_4\, yr^{-1}$ is redistributed from Asia to North America. We used time invariant prior flux estimates, so for the period from 2000 to 2006, when the growth rate of global atmospheric CH_4 was very small, the assimilation does not produce increases in natural or anthropogenic emissions in contrast to bottom-up emission datasets. After 2006, when atmospheric CH_4 began its recent increases, CarbonTracker- CH_4 allocates some of the increases to anthropogenic emissions at temperate latitudes, and some to tropical wetland emissions. For temperate North

Discussion Paper

Discussion Paper

Discussion Pape

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions

References

Tables

Figures

I∢

►I

< 1



Back



Full Screen / Esc

Printer-friendly Version

Interactive Discussion



2176

America the prior flux increases by about $4 \, \text{TgCH}_4 \, \text{yr}^{-1}$ during winter when biogenic emissions are small. Examination of the residuals at some North American observation sites suggests that increased gas and oil exploration may play a role since sites near fossil fuel production are particularly hard for the inversion to fit.

The tropics are not currently well resolved by CarbonTracker- CH_4 due to sparse observational coverage and a short assimilation window. However, there is a small uncertainty reduction and posterior emissions are about 18 % higher than prior estimates. Most of this increase is allocated to tropical South America rather than being distributed among the global tropics.

1 Introduction

Methane (CH₄) is second in importance to carbon dioxide (CO₂) among greenhouse gases with significant anthropogenic sources. It contributes about half the radiative forcing of CO₂, $0.48 \pm 0.05 \, \text{W m}^{-2}$ in 2010 (Hofmann et al., 2006; updated at http://www.esrl.noaa.gov/gmd/aggi/). Over a 100 yr time horizon it is 25 times more efficient per mass as a greenhouse gas than CO₂.

Global emissions of CH_4 are between 500 and 600 $TgCH_4$ yr $^{-1}$ (Dlugokencky et al., 2011), and about 40% of this is due to natural sources, mainly wetlands. The other 60% of global emissions are due to microbial emissions associated with rice agriculture, livestock and waste, and fugitive emissions from fossil fuel production and use (Denman et al., 2007). Global emissions have recently been approximately in balance with global sinks, mainly chemical destruction by reaction with OH. The lifetime of CH_4 in the atmosphere is between 8.5 and 10.1 yr (Holmes et al., 2013) with CO_2 the eventual product of its oxidation.

 CH_4 has increased from a preindustrial abundance of $722 \pm 4 \,\mathrm{ppb}$ (Etheridge et al., 1998 after conversion to the NOAA 2004 CH_4 standard scale, Dlugokencky et al., 2005) to current values of about 1800 ppb in 2010 (about 2.5 times), and it is likely that human activity is responsible for most of this increase. Current levels are unprecedented

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

<u>Co</u>nclusions

References

Tables

Figures

14



•



Back

Close

Full Screen / Esc

Printer-friendly Version



Discussion Paper

Back

over the last 800 kyr (Loulerge et al., 2008). NOAA atmospheric network observations extend back to the 1980s, and show that global CH₄ increased rapidly through the late 1990s, leveled off during the early 2000s and have recently begun to increase since 2007 (Rigby et al., 2008; Dlugokencky et al., 2009). The subject of the causes of the 5 recent increase is the topic of much recent work (e.g. Bousquet et al., 2011), including this study.

An important aspect of atmospheric CH₄ is the sensitivity of natural wetland emissions to climate change. Emissions from the Arctic, in particular, have the potential increase significantly as temperatures rise and the vast stores of soil carbon thaw (e.g. Schuur et al., 2011; Harden et al., 2012). Schaefer et al. (2010) pointed out that potential carbon emissions from the Arctic could have important implications for policies aimed at reducing or stabilizing emissions. This clearly highlights the importance of maintaining long-term measurements of atmospheric CH₄ in the Arctic, and in this study we hope to further the case for atmospheric inverse techniques as a tool to diagnose observed atmospheric records (see previous studies by Hein et al., 1997; Houweling et al., 1999; Chen and Prinn, 2006; Bergamaschi et al., 2005; Bousquet et al., 2006).

Atmospheric CH₄ is also influenced by diverse human activities, ranging from food production (ruminants and rice) to waste (sewage and landfills) to fossil fuel production (coal, oil and gas). Future increases in population could increase emissions from agriculture and waste as demand for more food production rises, while the current boom in shale oil/gas exploitation has focused attention on leakage from drilling, storage and transport of fossil fuel (e.g. Pétron et al., 2012). An obvious use of an atmospheric assimilation system is to quantify changes in anthropogenic emissions and attribute increases at policy relevant spatial scales, something that is possible only with adequate spatial coverage of observations. In this study we will discuss the degree to which this is currently possible given the coverage of the current observational network.

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page **Abstract** Introduction Conclusions References **Tables Figures** [■ \triangleright

Full Screen / Esc

Close

2 The CarbonTracker ensemble data assimilation system

5 The total emission of CH₄ in time and space may be described by:

$$F(x, y, t) = \lambda_1 F_{\text{natural}}(x, y, t) + \lambda_2 F_{\text{fossil}}(x, y, t) + \lambda_3 F_{\text{agriculture/waste}}(x, y, t) + \lambda_4 F_{\text{fire}}(x, y, t) + \lambda_5 F_{\text{ocean}}(x, y, t)$$
(1)

Where λ_i represents a set of linear scaling factors to be estimated in the assimilation that are applied to the fluxes (F) by multiplying prior estimates of CH₄ fluxes to produce the posterior flux estimates. A total of 121 parameters are estimated; 10 terrestrial emission processes for 12 continental regions (corresponding to the TRANSCOM 3 continental regions but with the addition of a tropical African region, see http://transcom.project.asu.edu for a map), and fluxes from the global ocean. The terrestrial emissions include fugitive emissions from coal, oil and gas production; agriculture and waste emissions (rice production, for example); livestock and their waste; and emissions from landfills and wastewater. Natural emissions include contributions from wetlands, termites, uptake in soils and wild animals. The final terrestrial emission category is biomass burning, which is treated as a separate category due to the existence of strong spatial constraints coming from satellite observations of hot spots. In general, the spatial distribution of the prior flux estimates is an important constraint on the assimilation. For example, the known location of fossil fuel production from bottom-up emission data sets provides information to the assimilation system on whether a signal measured at a particular observation site could have a fossil fuel component. If production areas change over time and not captured by the prior distribution, then fossil fuels will be underestimated by the inversion.

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Discussion Paper

Introduction

Conclusions

References

Tables

Figures

14



•



Back

Close

Full Screen / Esc

Printer-friendly Version



The ensemble system (EnKF) used to solve for the scalar multiplication factors is based on that described by Peters et al. (2005), and uses the square root ensemble Kalman filter of Whitaker and Hamill (2002). The length of the smoother window is restricted to five weeks for computational efficiency. Although the posterior flux estimates in relatively densely sampled regions such as North America were found to be robust with a window this short, regions with less dense observational coverage (the tropics, for example) are likely to be poorly constrained even after more than a month of transport and therefore not well resolved. As pointed out by Bruhwiler et al. (2005), a smoother window of at least 3 months is likely to make maximal use of remote network sites, however this may come at the expense of accumulated errors in transport. Even without the problem of a short smoother window, the sparseness of the observational network makes it difficult to resolve under-sampled regions such as the tropical terrestrial biosphere (Bruhwiler et al., 2011).

Statistics for the ensemble are created from 500 members, each with its own background ${\rm CH_4}$ concentration field to represent the time history (and thus covariances) of the filter. We experimented with different numbers of ensemble members and found that the use of too few ensemble members results in solutions that stay artificially close to prior flux estimates. To dampen spurious noise due to the approximation of the covariance matrix, we apply localization (Houtekamer and Mitchell, 1998) for non-background sites. By limiting correlations between distant sites, localization ensures that random correlations between parameters do not translate into unrealistic constraints on emissions by distant measurement sites (i.e. those connections physically impossible with only 5 weeks of transport). Localization is based on the linear correlation coefficient between the 500-parameter deviations and 500 observation deviations for each parameter, with a cut-off at 95 % significance in a student's t test with a two-tailed probability distribution. This is the same strategy employed by Peters et al. (2005).

ACPD

Paper

Discussion Paper

Discussion Paper

Discussion Paper

14, 2175-2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

Back Close

Full Screen / Esc

Printer-friendly Version



Prior values for the scaling factors, λ , are all set to unity to give fluxes that are unchanged from their values predicted by the prior flux estimates. In our assimilation, the chosen 1 s error of the prior estimates is 75 % for all parameters. The prior covariance structure describes the uncertainty on each parameter, plus their correlation in space. For the current version of CarbonTracker-CH₄, we assumed a diagonal prior covariance matrix so that no prior correlations between estimated parameters exist. The effect of this choice may be strong anti-correlations among estimated parameters in regions where few observational constraints exist; however, larger-scale aggregations of these regions are expected to yield more robust estimates. For example, the total tropical source can be better determined that the individual regions between which there can be trade-offs in emissions from time step to time step. Note also that the 5-week assimilation window used by CarbonTracker limits knowledge of temporal correlations. As a result, the uncertainty on annual average emissions is difficult to estimate.

2.3 TM5 atmospheric transport model

Transport Model 5 (TM5, Krol et al., 2005) is a community supported global model with two-way nested grids. For CarbonTracker-CH₄, we ran the simulation at 4° latitude × 6° longitude resolution without zoom regions. TM5 is developed and maintained jointly by the Institute for Marine and Atmospheric Research Utrecht (IMAU, the Netherlands), the Joint Research Centre (JRC, Italy), the Royal Netherlands Meteorological Institute (KNMI, the Netherlands), and NOAA ESRL (USA). TM5 has detailed treatments of advection, convection (deep and shallow), and vertical diffusion in the planetary boundary layer and free troposphere. The winds used for transport in TM5 come from the European Center for Medium range Weather Forecast (ECMWF) operational forecast model. This "parent" model currently runs with $\sim 25\,\mathrm{km}$ horizontal resolution and 60 layers in the vertical prior to 2006 and 91 layers in the vertical from 2006 onwards.

ACPD

Discussion Paper

Discussion Paper

Discussion Paper

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

4 F

[■

Back Close
Full Screen / Esc

Printer-friendly Version

Interactive Discussion



2181

The ECMWF meteorological data are preprocessed into coarser grids and are converted from wind fields to mass conserving horizontal and vertical mass fluxes. TM5 runs at an external time step of three hours, but due to the symmetrical operator splitting between advection, diffusion, emissions and loss the effective time step over which each process is applied is shorter. The vertical resolution of TM5 used with CarbonTracker-CH₄ is 34 hybrid sigma-pressure levels (from 2006 onwards; 25 levels for 2000–2005), unevenly spaced with more levels near the surface.

2.4 Prior emission estimates for natural sources

The largest natural emissions of methane are from wetlands, defined as regions that are permanently or seasonally water logged. Wetlands are a broad category that includes both high-latitude bogs and fens and tropical swamps. Saturated soils in warm tropical environments tend to produce the most methane. However, warming Arctic temperatures raise concerns of increasing output from high-latitude wetlands and future decomposition of carbon currently stored in frozen Arctic soils (e.g. Schaefer et al., 2011).

Methane is rapidly oxidized by methanotrophic bacteria in overlying aerobic water columns or unsaturated soil, so the water table must be at or near the surface and the depth of overlying water must be shallow for large emissions to occur. Wetland plants have adapted to low oxygen environments by having hollow stems to allow delivery of oxygen and other gases to root systems. These hollow stems also allow delivery of methane directly to the atmosphere, and along with ebullition account for most of transport to the atmosphere. Diffusion also occurs but is a significantly smaller contribution to the atmosphere. (See Barlett and Harris (1993) for an extensive overview of wetland emissions.) Estimates of global emissions from wetlands are about 150–200 Tg CH₄ yr⁻¹ with most of this occurring in tropical regions (Matthews and Fung, 1989; Melton et al., 2013). Because emissions are sensitive to temperature and precipitation, they exhibit significant seasonal cycles, especially at high latitudes, as well as inter-annual variability due to moisture and temperature variability.

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

<u>Re</u>ferences

Tables

Figures

· ·ga··oo

I⁴



4



Back



Full Screen / Esc

Printer-friendly Version



Discussion Pape

Methane emissions from wetlands are difficult to quantify using assimilation systems for two reasons; their global spatial distribution is difficult to know accurately, and there is large variability in emission rates over small spatial scales (meters), which makes extrapolation to large scales difficult. Here we used the prior flux estimates of Bergamaschi et al. (2007). The global total of the prior flux estimate is $175 \, \text{Tg} \, \text{CH}_4 \, \text{yr}^{-1}$.

Other natural sources of methane include enteric fermentation in insects (mainly termites, Sanderson, 1996) and wild ruminants (Houweling et al.,1999). Prior values for both of these sources ($\sim 25\, {\rm Tg\,CH_4\,yr^{-1}}$) are much smaller than the wetland source. Oxidation of CH₄ in dry soils ($\sim 40\, {\rm Tg\,CH_4\,yr^{-1}}$, Ridgwell et al., 1999) is a natural sink of CH₄ and is treated as a negative source in the assimilation.

2.5 Prior emission estimates for fugitive emissions from fossil fuels

Methane is the principal component of natural gas, and leaks to the atmosphere associated with natural gas production and distribution are a considerable source. Natural gas is associated with oil production and is often flared, or simply vented to the atmosphere. Together, anthropogenic emissions from oil and gas production are thought to contribute about $50\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr^{-1}}$ ($\sim 10\,\%$ of the global annual methane emissions). Methane is also associated with coal deposits and can be released by extracting and pulverizing coal. It is often vented directly to the atmosphere from mines, and this source contributes an additional $\sim 20\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr^{-1}}$. As Asian economies have undergone rapid growth, coal production there has increased by a factor of about two since 2000, while remaining approximately level for most of the rest of the world. In 2010, production of coal by China increased by 9 % over the previous year (BP Statistical Energy Review, 2011).

Combustion of natural gas is currently used to generate about a quarter of the electricity produced in the US Its popularity as a fuel has recently grown because it is a relatively clean and efficient source of energy. Recent technological advances in recovery of natural gas, principally hydraulic fracturing, have led to increases in reserve estimates, and a tremendous increase in exploitation of shale oil/gas deposits (e.g. En-

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

14







Back



Printer-friendly Version



Interactive Discussion



ergy Information Administration; http://www.eia.gov). It is possible that as natural gas reserves are increasingly exploited, emissions related to its production and distribution will rise in the future.

CarbonTracker-CH₄ uses the 1° × 1° gridded emissions from EDGAR 3.2FT2000 (Eu-5 ropean Commission, JRC, 2009) as prior emission estimates for fugitive emissions from coal, oil and gas production. This data set is based on emission inventories by country and sector for 1990 and 1995 extrapolated to 2000 using production and consumption statistics. We have not extrapolated this data over the period covered by CarbonTracker-CH₄, and have instead kept prior emission estimates constant at 2000 levels. This will allow us to test whether the emission estimates suggest changes in anthropogenic emissions, for example, the large increase in emissions from coal production in Asia or the significant increase in oil and gas drilling over the last decade in North America.

2.6 Prior emission estimates for agriculture and waste

The largest source of methane emitted by human activity is agriculture and waste; emissions from rice agriculture, waste/wastewater, and animals and their waste total 230-250 Tg CH₄ yr⁻¹. Ruminants, such as cattle, goats, sheep and water buffalo are able to convert hard-to-digest forage to energy through enteric fermentation, in which microbes produce easily digested material inside the animal's gut. Emissions from enteric fermentation may be expected to increase with increasing human population and higher standards of living. Animal waste, along with wastewater and landfills produce CH₄ when conditions favor anaerobic decomposition. Organic material is decomposed in low oxygen conditions by chains of microbial processes that terminate in production of methane by methanogens.

Rice agriculture is also a significant source of methane to the atmosphere. This is because warm, waterlogged organic-rich soils in rice paddies are ideal for methanogenesis. Bottom-up estimates of emissions from rice agriculture are 50 Tg CH₄ yr⁻¹,

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Introduction

Conclusions

References

Tables

Abstract

Figures

Back

Close

Full Screen / Esc

Discussion Pape

and emissions can be significantly reduced by drainage of paddies between harvests as well as other agricultural practices (Yan et al., 2009).

CarbonTracker-CH₄ uses the 1° × 1° degree gridded emissions from the EDGAR 3.2FT2000 as prior emission estimates for rice agriculture, enteric fermentation, animal waste management, wastewater and landfills. This data set is based on emission inventories by country and sector for the years 1990 and 1995 extrapolated to 2000 using production and consumption statistics. We have not extrapolated this data over the period covered by CarbonTracker-CH₄, and have instead kept prior emission estimates constant at 2000 levels as for fossil fuel emissions.

2.7 Prior emission estimates for biomass burning

Fires are a relatively small part of the atmospheric CH_4 budget: $15-20\,Tg\,CH_4\,yr^{-1}$ out of a total of $\sim 520\,Tg\,CH_4\,yr^{-1}$, however, they are an important contribution to interannual variability of methane.

The fire prior currently used in CarbonTracker-CH $_4$ is based on the Global Fire Emissions Database (GFED), which uses the CASA biogeochemical model to estimate the carbon fuel in various biomass pools along with burned area based on MODIS satellite observations of fire counts (Giglio et al., 2006; van der Werf et al., 2006). The dataset consists of 1 $^{\circ}$ × 1 $^{\circ}$ gridded monthly burned area, fuel loads, combustion completeness, and fire emissions for numerous atmospheric constituents, including CH $_4$ for the time period spanning January 1997–December 2010.

2.8 Prior estimates for ocean fluxes

The oceans play a relatively small role in the budget of atmospheric methane, contributing only $\sim 2-3\,\%$ of global emissions ($\sim 10-15\,Tg\,CH_4\,yr^{-1}$). A significant fraction of this is assumed to come from methane seeps in shallow coastal waters ($\sim 5\,Tg\,CH_4\,yr^{-1}$). The overlying water column must be shallow for emission to the atmosphere, since CH_4 is efficiently consumed by aerobic microbial processes. The water column also

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

14

►I

4

•

Close

Back

Full Screen / Esc

Printer-friendly Version



Discussion Pape

I



needs to be shallow for bubbles to deliver methane directly to the air. Coastal waters are sometimes supersaturated in CH₄, and may emit about 6 TgCH₄ yr⁻¹ to the atmosphere, while the open may add another 3 TgCH₄ yr⁻¹ (Houweling et al., 1999; Lambert and Schmidt, 1993).

Rhee et al. (2009) have suggested that global ocean emissions excluding natural seeps is much smaller than the $\sim 9\, Tg\, CH_4\, yr^{-1}$ we have used in this version of CarbonTracker-CH₄, only about 0.6–1.2 Tg CH₄ yr^{-1}. On the other hand, recent studies conducted in the coastal waters of the eastern Siberian Arctic hint at the possibility of a significant source of methane coming from methane bubbling from the continental shelf sediments (Shakova et al., 2010). For this version of CarbonTracker-CH₄ we followed the approach of Bergamaschi et al. (2009) and used the estimates of Houweling et al. (1999) and Lambert and Schmidt (1993) as prior flux estimates. We also assumed an uncertainty on these prior flux estimates of $\pm 75\,\%$.

2.9 Atmospheric chemical loss

Methane is removed from the atmosphere mainly by reaction with hydroxyl radical (OH), but also by reaction with atomic chlorine (Cl) and excited-state oxygen (O 1 D) in the stratosphere. The chemical loss of methane over a year is about equal to the total input from sources ($\sim 520\,\mathrm{Tg\,CH_4\,yr^{-1}}$), and the mean lifetime of methane is 9–10 yr. Small differences in the emissions and losses lead to trends in atmospheric CH $_4$ abundance, while year to year changes in the balance of emissions and loss lead to inter-annual variability and possibly to trends in observed methane.

Hydroxyl radical is extremely reactive and has such a short atmospheric residence time that it is difficult to directly measure its global distribution. Instead, observations of atmospheric species that have relatively well-quantified anthropogenic emissions and are destroyed only by reaction with OH, such as methyl chloroform (CH₃CCl₃), are used, often along with atmospheric models, to estimate the abundance of atmospheric OH. Using an empirical approach, Montzka et al. (2011) noted that the inter-annual

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

→

Back Close
Full Screen / Esc

Printer-friendly Version

variability in atmospheric OH is likely to be within about $\sim 2\%$. Errors in derived OH distributions arise from uncertainty in the emissions of CH_3CCl_3 used to estimate OH and uncertainties in transport models. Krol et al. (1998) estimated that the uncertainty in globally averaged OH is $\pm 10\%$.

About 10% of total chemical loss is due to transport and chemical destruction in the stratosphere. A small amount of this methane-depleted air is returned to the troposphere and could influence interpretation of high-altitude (aircraft) measurements of methane. In addition, errors in simulating stratosphere-troposphere transport could result in biases for model simulations covering many years.

Errors in the chemical loss of methane and the inability to adequately resolve interannual variability of OH are troublesome for estimation of methane fluxes. A 2% variation in the global methane sink is equivalent to $\sim 10\,\mathrm{Tg\,CH_4\,yr^{-1}}$, about the size of estimated inter-annual variability in methane emissions.

For the present version of CarbonTracker-CH₄ we use pre-calculated OH fields from a global chemical model that have been optimized against global observations of methyl chloroform. The chemical loss fields consist of a single, repeating seasonal cycle, and result in a methane lifetime of about 9.5 yr. Details of the chemical loss fields may be found in Bergamaschi et al. (2005).

2.10 Observational constraints

This study uses measurements of air samples collected at surface sites in the NOAA ESRL Cooperative Global Air Sampling Network (http://www.esrl.noaa.gov/gmd/ccgg/flask.html) except those identified as having analysis or sampling problems, or those thought to be strongly influenced by local sources. The availability of data varies over time. Data collection, quality control and analysis methods are described in detail by Dlugokencky et al. (1994). A map of sites used in CarbonTracker-CH₄ is shown in Fig. 1. In addition, we use in situ quasi-continuous CH₄ time series from the following towers operated by Environment Canada (EC): 30 m above ground level (a.g.l.) at Candle Lake (CDL, formerly Old Black Spruce), SK, Canada, 105 m a.g.l. at East Trout

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I**⊲**







Back

Full Screen / Esc

Printer-friendly Version



Lake, SK, Canada (ETL), 40 m a.g.l. at Fraserdale, ON, Canada (FRD), and 10 m a.g.l. at Lac Labiche, AB, Canada (LLB). Other in situ quasi-continuous CH₄ time series used are from the EC Canadian sites at Alert, Nunavut (ALT), Sable Island, NS (SBL) and Egbert, ON (EGB). All observations used in CarbonTracker-CH₄ are calibrated against the WMO GAW CH₄ X2004 mole fraction scale (Dlugokencky et al., 2005).

For most quasi-continuous sampling sites, we construct an afternoon daytime average mole fraction for each day from the time series, recognizing that our atmospheric transport model does not always capture the continental nighttime stability regime while daytime well-mixed conditions are better matched. Table 1 summarizes how data from the different measurement programs are preprocessed for this study.

We further exclude non-marine boundary layer (MBL) observations that are very poorly forecasted in our framework following the strategy used with CarbonTracker- CO_2 . We use the so-called model-data mismatch in this process, a quantity that represents random error ascribed to each observation to account for measurement and modeling errors at each site. If the observed-minus-forecasted mole fraction exceeds 3 times the prescribed model-data mismatch, then the observation is not used at that time-step of the inversion. This can happen when an air sample influenced by local emissions is not captured well by our 1° × 1° fluxes, or when local meteorological conditions are not captured by our offline transport fields. A complete list of sites used in CarbonTracker- CH_4 is given in Table 2, along with the model-data mismatch used, the number of points available, the number that were excluded, and statistics on the posterior fit to each site.

3 Evaluation of CarbonTracker-CH₄

In this section we discuss the evaluation of CarbonTracker-CH₄ using three methods: comparison of prior and posterior residuals (difference between simulated and observed CH₄ concentration), comparisons to profiles measured from aircraft that were

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I4 ÞI

•

Close

Full Screen / Esc

Back

Printer-friendly Version



not used in the assimilation, and comparisons to integrated global and zonal concentration and growth rate.

3.1 Residuals

The prior and posterior residuals, calculated by subtracting the observed CH_4 mole fraction at each site constraining the inversion from the simulated prior/posterior abundances, are shown in Fig. 2. The bottom panel shows that the balance between the prior emissions and the chemical sink leads to an underestimate of CH_4 relative to observations at all latitudes. By the end of the simulation, the negative bias of the model using prior fluxes reaches values up to 75 ppb (compared to a global average of \sim 1790 ppb in 2009, about 4 %). This negative bias is considerably reduced for the posterior residuals, as is shown in the top panel, and at most sites the posterior residuals are within \sim 15 ppb of the observations. This is partly by design, since the model-data mismatch determines how closely the posterior CH_4 abundances must match the observations; however, as Table 1 shows, the posterior residuals even for some sites that have large model-data mismatch error assigned to them, can be quite small. An example is BKT, with a model-data mismatch of 75 ppb and a posterior residual of only 6.8 ppb. For sites like this, future inversions could use reduced model-data mismatch errors, allowing the observations to more strongly constrain the inversion.

Figure 2 shows that even after assimilation of observations, some sites have large low biases (implying emissions higher than prior estimates are needed to match the observations) and Fig. 3 shows the relative sizes of the residuals. WGC (Walnut Grove, CA) is located near a densely populated urban area with agriculture as well and has an average posterior residual of –118 ppb. In addition to the difficulty of using relatively coarse resolution global transport to simulate observations amidst strong local sources, it is also likely that the prior emissions are underestimated. Other sites with large biases are WKT (central Texas) and SGP (Southern Great Plains, OK) with average residuals of –49 ppb and –57 ppb. These sites see transport from polluted urban areas, and they likely also see transport of emissions from oil and gas drilling as discussed in

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I◀

►I

■



Back



Full Screen / Esc

Printer-friendly Version



3.2 Comparison to aircraft profiles

The current version of CarbonTracker-CH₄ does not assimilate observations from the NOAA GMD aircraft project. This network currently consists of 17 sites distributed over North America where air samples are collected at 12 altitudes and analyzed for a suite of atmospheric gases, including CH₄ (www.esrl.noaa.gov/gmd/ccgg/aircraft). Because the aircraft observations were not used to constrain the inversion, these data can be used as an independent check on the inversion. In addition, they provide useful insight into the performance of TM5's vertical transport.

Figure 4 shows the prior and posterior residuals for THD (Trinidad Head, CA), where the white line represents the average of the residuals and the red lines show the standard deviation of the residuals as a function of altitude. Compared to the prior residuals, the posterior residuals show a reduction in bias at all altitudes, as well as a smaller spread in the residuals. At high altitudes the surface data constraints have resulted in estimated emissions that are in good agreement with the well-mixed free tropospheric abundances. THD shows good agreement at the lowest levels because profiles at this location are more likely to sample background marine air coming off of the Pacific Ocean. In contrast, DND (Dahlen, North Dakota) shows a much larger negative bias at low altitudes during the summer but good agreement at all levels during winter (Fig. 5). This implies that local or regional-scale sources that are not included in the CarbonTracker-CH₄ prior and are not "seen" by other sites influence these summertime profiles. Future versions of CarbonTracker-CH₄ may use at least the lower levels of the aircraft observations in order to better constrain emissions.

ACPD

Paper

Discussion Paper

Discussion Paper

Discussion Pape

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Back

Full Screen / Esc

Close

Printer-friendly Version

Interactive Discussion



2190

The abundance of CH₄ integrated over the global atmosphere and its growth rate are important diagnostics of inversion performance (Rayner et al., 2004; Bruhwiler et al., 2011) because, given the ~ 10 yr lifetime of CH₄, on global scales emissions and sinks must balance in a way that produces the observed global growth of CH₄. Here we follow the approach taken by Bruhwiler et al. (2011) that uses the same sampling, filtering and smoothing procedure used to produce the observed global and zonal CH₄ abundances for both data and model output (see Masarie and Tans, 1995 web updates at http://www.esrl.noaa.gov/gmd for a description of the data extension procedure). As shown in the top panel of Fig. 6, the global posterior CH₄ abundance produced by the CarbonTracker-CH₄ assimilation is in fairly good agreement with the observed global abundance, however it is biased low by about 10 ppb. This is because the global abundance that results from use of the prior fluxes without optimization is much lower than observed, and the posterior global total represents a compromise between CH₄ abundance obtained from prior flux estimates and the observations at each site. Reducing the model-data mismatch error and/or increasing the prior flux uncertainty would improve the agreement between posterior CH₄ and the observations, but likely at the expense of having flux estimates with unrealistic spatiotemporal variability, especially in regions that are relatively unconstrained by observations. On the other hand, if the prior flux estimates are weighted too heavily in the inversion, the posterior global total more closely follows the global abundance simulated by the prior fluxes than the observations, and these may depart significantly from the actual emissions. The middle panel of Fig. 6 shows the difference between the simulated and prior CH₄ abundance and the observations, where it can be seen that the residual difference varies slightly over time as the bias resulting from prior emissions changes. In particular, between 2004 and 2006, the prior residuals are fairly constant and the residual between the posterior and the observations is smaller than over other periods. The conclusions that ACPD

Discussion Paper

Discussion Paper

Discussion Pape

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

ld ►I

4 ▶

Back Close
Full Screen / Esc

Printer-friendly Version

Interactive Discussion



2191

can be drawn from this are that better prior flux estimates are needed for future ver-

sions of CarbonTracker-CH₄, and that the global abundance is a useful way to judge whether the solution is most influenced by the prior information or by the observational constraints.

The bottom panel of Fig. 6 shows the growth rate of global atmospheric CH_4 , a quantity that is directly related to imbalances between emissions and sinks. CarbonTracker- CH_4 follows the observed growth rate fairly well, but not perfectly since there are periods for which it under- or overestimates the observed growth rate. During 2007, for example, the observed growth was underestimated by ~ 30 %, while during 2009 it was overestimated by about the same amount. These differences are an indication of global total biases in estimated emissions.

It is also informative to consider zonally averaged CH_4 mole fraction and its growth rate at sub-hemispheric scales as shown in Fig. 7 for the high northern latitudes (53.1–90° N) and the tropics (17.5° S–17.5° N). For the high northern latitudes, the posterior simulated integrated CH_4 is quite close to the observations and the growth rate agrees well with the observed growth rate. On the other hand, the simulated integrated CH_4 in the tropics is further from the observations and closer to the prior than for the high northern latitudes. The simulated growth rate in the tropics also can differ significantly from the observed growth rate, with under or over estimates reaching 5 ppb yr⁻¹ or more. As a comparison, the growth rate agreement between the observed and simulated at northern polar latitudes is usually well within a few ppb yr⁻¹. The middle panels of Fig. 7 show that when the residuals between the prior and observations decrease, the posterior residuals are also smaller.

For the high northern latitudes, a small seasonal cycle in the residuals potentially provides some information about which emission processes may be under- or overestimated by the priors. Differences between simulated and observed CH₄ are largest during the winter with the observations being higher than the simulations. This implies that mid- and high latitude emissions from anthropogenic sources may be underestimated by the priors and not completely corrected for by the inversion. Note that biogenic emissions at mid- and high latitudes are at a minimum during winter.

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I**4**

. . .

4



Back



Full Screen / Esc

Printer-friendly Version



ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page Introduction **Abstract** References

Conclusions

Back

Tables Figures

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



Anomalously high growth rates were observed in 2007 both in the Arctic and in the tropics (Dlugokencky et al., 2009), a year when the Arctic was anomalously warm and the tropics were unusually wet. The results shown in Fig. 7 suggest that the inversion is likely able to provide good estimates of flux anomalies in high latitudes, at least in the zonal average. For the tropics zonal average flux anomaly estimates for this year are likely to be underestimated. These differences in the ability of the inversion to recover and attribute variability are due mostly to differences in the distribution of network sites with the Arctic having better observational coverage than the tropics. Another factor is that the deep vertical mixing of the tropical atmosphere makes it difficult for the network sites that are mostly located on remote islands to detect signals from terrestrial CH₄ sources. A further limitation is the 5-week lag used in CarbonTracker's EnKF scheme that cuts off transport of signals that are transported to remote observing sites.

Note that an additional diagnostic of posterior emissions is the posterior error covariance and its difference from the prior covariance. If there are no observations to constrain the posterior estimates, then the posterior error covariance will be unchanged from the prior error covariance. While the posterior error covariance is a very useful diagnostic of the error reduction coming from observations, it is less useful as an indicator of the absolute accuracy of the estimated emissions because the accuracy of the prior estimates is ultimately not very well known, and there are transport errors that cannot be adequately accounted for.

Results

The high northern latitudes

Here the high northern latitudes are an aggregation of the Transcom 3 regions Boreal North America, Boreal Eurasia and Europe. The ten-year average posterior aggregated flux for the high northern latitudes is $81 \pm 41 \, \text{TgCH}_4 \, \text{yr}^{-1}$, a decrease of a little over $12 \, \text{Tg} \, \text{CH}_{\text{A}} \, \text{yr}^{-1}$ from the prior aggregated flux. Note that due to the use of a 5-week assimilation window, the uncertainty estimate does not include temporal error covariance over timescales longer than this period and it should therefore be regarded as the best estimate possible for the long term error covariance given the limitations of the current assimilation scheme. The inversion suggests that most of this decrease is a reduction in natural wetland emissions (8 Tg CH₄ yr⁻¹) with the remaining amount coming from fugitive fossil fuel emissions, although the portioning between these sources is strongly influenced by the prior distributions and relative locations of observation sites. Although the observing network could still be considered sparse at high northern latitudes, the number of existing sites is sufficient to reduce uncertainty by over 75 % from the prior uncertainty. The total posterior flux ranges from 78 Tg CH₄ yr⁻¹ in 2004 to just under 86 Tg CH₄ yr⁻¹ in 2007 (Fig. 8), a year that saw record warm temperatures throughout much of Boreal North America and Boreal Eurasia, as well as extremely low sea ice coverage (Stroeve et al., 2008).

Annual average methane emissions at high northern latitudes are approximately evenly divided between fugitive emissions from fossil fuels, agriculture and waste (coming mostly from Europe) and natural wetlands. As a whole, emissions from fossil fuel leakage are slightly decreased relative to prior estimates by about 4 Tg CH₄ yr⁻¹ from 33 Tg CH₄ yr⁻¹, a change that is slightly larger than the posterior estimated uncertainty, 3 Tg CH₄ yr⁻¹. Emissions from agriculture and waste are unchanged. Annual average wetland emissions over the high northern latitudes are reduced by 26 % from a prior of 31 TgCH₄yr⁻¹ to about 23 TgCH₄yr⁻¹. This result is in agreement with previous studies (e.g. Chen and Prinn, 2006; Bergamaschi et al., 2007; Spahni et al., 2011). Our results do not agree with the emission estimates of Bloom et al. (2010). They find that only 2% of global wetland emissions come from the high northern latitudes, while we find closer to 10%. On the other hand our results agree within uncertainties with the estimates of McGuire et al. (2012) based on flux measurements. They find a source of 25 Tg CH₄ yr⁻¹ from Arctic tundra wetlands with uncertainty ranging from 10.7 to 38.7 Tg CH₄ yr⁻¹. Applying the same spatial filter for their Arctic tundra region, CarbonTracker-CH₄ estimates a somewhat smaller $16 \pm 5 \text{Tg CH}_4 \text{ yr}^{-1}$. The fact that **ACPD**

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I₫

►I

4



Back

Full Screen / Esc

Printer-friendly Version



field studies may be biased towards larger emissions could at least partially account for the lower estimate based on atmospheric observations. On the other hand, we cannot rule out the possibility that the TM5 representation of the polar atmosphere is too stable, leading to an accumulation of methane emissions in the lower atmosphere the inversion will therefore reduce emissions in order to match observations.

The estimated flux anomaly during 2007 is 4.4 TgCH₄ with a maximum summer anomaly of 2.3 Tg CH₄ in July. If the anomaly is calculated by subtracting the 2000-2006 average annual flux the estimated 2007 anomaly is 5.3 TqCH₄, similar to the result found by Bousquet et al. (2011). Our estimates are slightly higher than the 103 Tg CH₄ yr⁻¹ found by Bergamaschi et al. (2013) for 2007. Based on zonal average analysis of network observations, Dlugokencky et al. (2009) pointed out that in 2007 the global increase of methane was equal to about a 23 Tq CH₁ imbalance between emissions and sinks, and that the largest increases in CH₄ growth occurred in the Arctic (> 15 ppb yr⁻¹). This does not necessarily imply that the largest surface flux anomalies occurred at high northern latitudes. Bousquet et al. (2011) noted that the relatively weak vertical mixing characteristic of polar latitudes results in a larger response in atmospheric CH₄ mole fractions to anomalous surface emissions than at tropical latitudes where strong vertical mixing rapidly lofts surface emissions through a deep atmospheric column. Transport models therefore can play an important role in helping to untangle surface flux signals from variability in atmospheric transport processes, although care must to be taken to also consider possible biases in modeled transport.

In 2008, the flux anomalies dropped to $2.4\,\mathrm{Tg\,CH_4}$, or $3\,\mathrm{Tg\,CH_4}$ if the anomaly is calculated by subtracting the average annual flux over 2000–2006, as was done by Bousquet et al. (2011) who obtained $2\,\mathrm{Tg\,CH_4}$ for their INV1 that is similar to CarbonTracker-CH₄, but $-3\,\mathrm{Tg\,CH_4}\,\mathrm{yr^{-1}}$ using their higher spatial resolution variational inversion (INV2). It is noteworthy that temperatures remained high in far Eastern Siberia during 2008, while returning to normal over Boreal North America (GISS surface air temperature analysis), while estimated fluxes remained high for Boreal Eurasia and returned to average levels for Boreal North America (Fig. 8). Microbial production of

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I**⊲**



Back



Full Screen / Esc

Printer-friendly Version



[■



CH₄ is very sensitive to temperature, and so it is likely that wetland emissions remained high in Siberia during 2008, while returning to normal levels over Alaska and Canada. CarbonTracker-CH₄ is able to discriminate between these two source regions located at roughly the same latitude, and the surface temperature analysis provides independent support for this since the prior wetland emission estimates do not vary inter-annually with temperature. Posterior covariance estimates support the independence of estimates for Boreal North America and Boreal Eurasia since the covariance between these two regions is small.

For the high northern latitudes CarbonTracker-CH₄ is able to distinguish between different CH₄ source processes and regions. Wetlands may be distinguished from anthropogenic sources because of the spatial separation of prior flux constraints; many high northern latitude wetland complexes are located in relatively sparsely populated areas, while fossil fuel and agricultural and waste emissions are distributed mainly in populated areas of Europe (although the Western Siberian Lowlands is also a region of intensive fossil fuel production). Ocean methane fluxes are thought to be small compared to terrestrial fluxes, and northern Eurasia and boreal North America are separated by the North Pacific Ocean. Furthermore, the stronger zonal and weaker vertical transport characteristic of the high latitudes helps to transport flux information to network sites. Both Europe and boreal North America are constrained by surface network sites, and although boreal Eurasia is not adequately covered by network sites, a number of sites exist downwind of it (Shemya, Barrow and Cold Bay). For Europe, average trajectory calculations suggest that a large region of wetlands in eastern Scandinavia and northwestern Russia is constrained by Pallas, Finland. Other sites help to constrain the anthropogenic sources from the rest of Europe.

For boreal North America, prior flux emissions from fossil fuels and agriculture and waste form an insignificant part of total methane emissions. This is not the case for Europe for which emissions are more evenly divided between anthropogenic and wetland emissions. Prior emission estimates from natural sources for Europe are about $45\,\mathrm{Tg\,CH_4\,yr^{-1}}$ during the summer months while the 11 yr average posterior summer

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

 ■
 Back

 Close

 \triangleright

Full Screen / Esc

Printer-friendly Version

Discussion Pape

Back



estimate is about 13 Tg CH₄ yr⁻¹, a large reduction. Boreal Eurasian summertime wetland emissions are increased relative to the prior flux estimates from 26 Tg CH₄ yr⁻¹ to $37 \, \text{Tg} \, \text{CH}_4 \, \text{yr}^{-1}$. For boreal North America, the average posterior summer wetland flux is only slightly below the prior flux estimate (about 19 Tg CH₄ yr⁻¹ compared to about 16 Tg CH₄ yr⁻¹). The redistribution of emissions from Europe to northern Eurasia was found by Chen and Prinn (2006) to be sensitive to the choice of sites used in the inversion, however, our results indicate that the observations imply that while prior emissions are too high for Europe, larger emissions are still needed elsewhere to match the meridional distribution of observed methane. Bergamaschi et al. (2005) also found decreased emissions for Europe relative to prior estimates, but interestingly, their inversion also reduced high-latitude emissions from prior estimates for other high latitude source regions as well. During the winter months, when biogenic emissions are low, prior estimates are decreased by the inversion for both Boreal Eurasia (-20%) and Europe (-8%) and relatively unchanged for Boreal North America. The small change in winter European emissions supports the conclusion that prior wetland emissions for Europe are indeed overestimated.

High latitude emissions of CH_4 from agriculture and waste are significant only for Europe, and estimated fluxes are unchanged from prior estimates. Fugitive emissions of CH_4 from fossil fuel production are reduced from prior estimates for Europe and boreal Eurasia by $2\,Tg\,CH_4\,yr^{-1}$ for each region; from $21\,Tg\,CH_4\,yr^{-1}$ and $12\,Tg\,CH_4\,yr^{-1}$. For boreal North America, prior estimates of fossil fuel emissions of CH_4 are very small (< $1\,Tg\,CH_4\,yr^{-1}$), and it should be noted that the tar sand production areas are in the temperate North American TransCom 3 source region rather than Boreal North America.

ica. Significant natural CH_4 emissions have recently been proposed for the high northern latitudes. Walter et al. (2007) estimated that in addition to emissions from High Northern Latitude wetlands (31 Tg CH_4 yr⁻¹ for the CarbonTracker- CH_4 prior), ebullition from lakes could add an additional 24 ± 10 Tg CH_4 yr⁻¹. In addition to organic rich

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I4 ►I

4 -

Close

Full Screen / Esc

Printer-friendly Version

sediments and subsea permafrost, CH_{Δ} is stored in ice hydrates forming at the low temperatures and high pressures in sediments at the bottom of the Arctic Ocean and subsea permafrost, and below terrestrial permafrost as well. Relatively shallow waters make it possible for bubbles to transport methane directly and rapidly to the atmosphere. The estimates of Shakhova et al. (2010) estimate the size of this source to be ~ 10 Tg CH₄ yr⁻¹ for the Arctic Ocean, although observational records are currently insufficient to establish whether these emissions are changing over time. Walter et al. (2012) have proposed that a similar process may also occur on land as permafrost thaws and glaciers melt. Total natural emissions including all of these processes would approach 65 Tg CH₄ yr⁻¹, an amount that significantly exceeds both the average prior and posterior annual natural emissions for CarbonTracker-CH₄ (31 and 23 Tg CH₄ yr⁻¹). Since the average total posterior CarbonTracker-CH₄ High Northern Latitude emissions is $\sim 81 \, \text{TgCH}_4 \, \text{yr}^{-1}$, accommodation of a $65 \, \text{TgCH}_4 \, \text{yr}^{-1}$ natural source would have to come at the expense of fossil fuel and agriculture/waste sources (average total CarbonTracker-CH₄ posterior of $\sim 57 \, \text{Tg} \, \text{CH}_4 \, \text{yr}^{-1}$), which would need to be reduced by about 75 %.

The estimated mass of carbon thought to be frozen in Arctic permafrost down to 20 m is estimated to be $\sim 1700\,\mathrm{PgC}$ (PgC = $10^{15}\,\mathrm{g}$ carbon) (Tarnocai et al., 2009), significantly more carbon than is currently in the atmosphere ($\sim 830\,\mathrm{PgC}$) and over 3 times what has already been emitted to the atmosphere from fossil fuel use since pre-industrial times. As the Arctic warms and permafrost thaws, this ancient carbon may be mobilized to the atmosphere and a small fraction ($\sim 3\,\%$) may be emitted as CH₄ (Schuur et al., 2011). Recent studies suggest that permafrost carbon will begin to enter the atmosphere during this century (e.g. Schaefer et al., 2010; Harden et al., 2012; Melton et al., 2013; Frolking et al., 2011). Harden et al. (2012) predict that 215–380 PgC will thaw by 2100. In their assessment of the carbon balance of Arctic tundra based on flux observations McGuire et al. (2012) found that between the 1990s and 2000s emissions of CH₄ doubled (from 13 to 26 Tg CH₄ yr⁻¹), results that are consistent with warmer temperature and longer growing seasons.

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I₫



- 4



Back



Full Screen / Esc

Printer-friendly Version



Detection of trends in Arctic greenhouse gas emissions is difficult using atmospheric concentration measurements alone because changes are expected to be small in comparison to transport of much larger mid-latitude emissions. Forward and inverse modeling techniques can be helpful because they provide the ability to untangle variability coming from transport from signals associated with local sources. As shown in Fig. 9, posterior CarbonTracker-CH₄ emissions do not indicate that there has been a trend in natural high northern latitude emissions over the last decade, although we see strong evidence for substantial inter-annual variability. This conclusion was also reached by Bergamaschi et al. (2013).

4.2 The Northern Hemisphere mid-latitudes

The average estimated total emissions for the northern mid-latitudes over the period 2000–2010 are greater than prior estimates by about $5\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr}^{-1}$, increasing from about $156\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr}^{-1}$ to $162\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr}^{-1}$. After the tropics, the Northern Hemisphere mid-latitudes, consisting of the temperate North America and Eurasian Transcom regions, emit the most atmospheric $\mathrm{CH_4}$. The largest mid-latitude source of $\mathrm{CH_4}$ is agriculture and waste, and this source rises from $117\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr}^{-1}$ to $119\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr}^{-1}$, a difference that is well within the posterior uncertainty estimate. Natural wetlands are a fairly small contribution to northern mid-latitude emissions, and they are increased from about 9 to $12\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr}^{-1}$. For the northern mid-latitudes as a whole, estimated fossil fuel emissions remain very close to prior estimates at about $31\,\mathrm{Tg}\,\mathrm{CH_4}$.

In general, CarbonTracker-CH $_4$ re-distributes prior estimated emissions from temperate Eurasia to North America (Fig. 10). The total prior flux estimates for temperate Eurasia and temperate North America are 124 and $32\,\mathrm{Tg}\,\mathrm{CH}_4\,\mathrm{yr}^{-1}$, respectively. The average posterior estimates are 114 and $47\,\mathrm{Tg}\,\mathrm{CH}_4\,\mathrm{yr}^{-1}$. Agriculture and waste emissions from temperate Eurasia are reduced by almost $10\,\mathrm{Tg}\,\mathrm{CH}_4\,\mathrm{yr}^{-1}$ ($\sim 9\,\%$). Fugitive fossil fuel emissions for temperate North America increase by $\sim 9\,\%$ (6.7 to $8\,\mathrm{Tg}\,\mathrm{CH}_4\,\mathrm{yr}^{-1}$), agriculture and waste emission increase by $53\,\%$ (21 to $32\,\mathrm{Tg}\,\mathrm{CH}_4\,\mathrm{yr}^{-1}$),

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I∢

►I

- 4



Back

Close

Full Screen / Esc

Printer-friendly Version



Interactive Discussion

and natural emissions increase by 66 % (4.5 to 7.5 Tg CH_4 yr⁻¹). Posterior uncertainties for both regions decrease from prior uncertainty by about 30 %.

In this version of CarbonTracker-CH₄, we used constant anthropogenic emissions representative of the year 2000 from the EDGAR v3.2 FT database as priors. A more recent version of EDGAR (version 4.3) reports that global anthropogenic emissions of methane significantly increased over the last decade, from 309 Tg CH₄ yr⁻¹ in 2000 to $364 \, \text{TgCH}_4 \, \text{yr}^{-1}$ by 2008 (an increase of about 18%). Most of this increase (37 Tg CH₄ yr⁻¹) is estimated to have occurred between 2000 and 2006 according to EDGAR. As shown in Fig. 5, the observed global total CH₄ does not change much between 2000 and 2005. Bousquet et al. (2006) proposed that increased anthropogenic emissions were balanced by decreases in wetland emissions for the early 2000s CarbonTracker-CH₄ is constrained using a time-invariant prior, and because it must follow the observed global growth rate that flat at least until 2006, it does not find trends in either total anthropogenic or natural emissions.

After 2006 the observed global annual mean CH₄ abundance increased ~ 25 ppb by 2010, equivalent to additional emissions of $\sim 70 \, \mathrm{Tg} \, \mathrm{CH_4}$ over 4 yr (18 $\, \mathrm{Tg} \, \mathrm{CH_4} \, \mathrm{yr}^{-1}$) if mixed uniformly throughout the atmosphere. As discussed in Sect. 3.3, CarbonTracker-CH₄ follows the global growth rate fairly closely. The average total emissions from 2000 to 2006 are $\sim 514 \,\mathrm{Tg\,CH_4}$, but after 2006 it is $530 \,\mathrm{Tg\,CH_4} \,\mathrm{yr}^{-1}$, an amount that is in approximate agreement with the change in total emissions implied by observations. CarbonTracker-CH₄ allocates much of this increase to anthropogenic sources. Average global total anthropogenic emissions are 316 Tg CH₄ yr⁻¹ for 2000 to 2005, increasing to 325 Tg CH₄ yr⁻¹ for 2006 to 2010. Using surface observations and satellite retrievals from SCIAMACHY and temporally varying anthropogenic emissions from EDGARv4.2, Bergamaschi et al. (2013) find that anthropogenic emissions increased by 12-26 Tg CH₄ yr⁻¹ between 2003 and 2008 making our estimates roughly consistent with theirs The lower increases in anthropogenic emissions found by this study are due to our use of constant prior emissions for the year 2000. For the period

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page Introduction **Abstract** Conclusions References

> **Tables Figures**

Back Close

Full Screen / Esc

2000 to 2005, the estimated total natural emissions are 198 Tg CH₄ yr⁻¹, increasing to 204 Tg CH₄ yr⁻¹ from 2006 to 2010. CarbonTracker-CH₄ assigns the post-2005 estimated increases in anthropogenic emissions to the populated northern temperate latitudes, while the bulk of the global increase in natural emissions is assigned to the tropics.

The EDGAR emission data imply that anthropogenic emissions of CH₄ grew rapidly over the last decade, with significant growth occurring between 2000 and 2005, a time when the observed growth rate does not support an upward trend in emissions. Could decreased emissions from wetlands have cancelled out this increase as Bousquet et al. (2006) proposed? Although the sparseness of the observational network makes it impossible to rule this scenario out, the observations and the spatial constraint they supply to the inversion do not suggest that there was a trend in wetland emissions over the first half of the decade, although there certainly was inter-annual variability. On the other hand, this result cannot be reconciled with bottom-up estimates of increasing anthropogenic emissions over this period. From 2006 through 2010, estimated emissions increased by $\sim 15 \, \text{Tg CH}_4 \, \text{yr}^{-1}$, slightly less than the $18 \, \text{Tg CH}_4 \, \text{yr}^{-1}$ estimated by EDGAR for this period, and with considerable inter-annual variability. Bergamaschi et al. (2013) also find that the EDGAR emission estimates are too high to be consistent with atmospheric observations. CarbonTracker-CH₄ divides this growth between anthropogenic and natural emissions in proportion to their contribution to the prior global atmospheric CH₄ budget (~ 60 % anthropogenic and ~ 40 % natural). Kirschke et al. (2013) also find that the increase after 2006 is due to a combination of wetland and fossil fuel emissions. Their study is a synthesis of bottom-up and top-down approaches that includes CarbonTracker-CH₄, so they are less specific about attributing changes in regions. Although it is likely that both anthropogenic and natural emissions have been increasing since 2006, this latter fact may also be interpreted as evidence of the inability of the observational network to discriminate between these categories of sources due to insufficient spatial coverage at lower latitudes. Uncertainty reductions are substantial for the global totals of both natural and anthropogenic emissions,

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I◀

►I

- 4



Back

Close

Full Screen / Esc

Printer-friendly Version



Discussion

ACPD 14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page Introduction **Abstract** References Conclusions

> **Tables Figures**

Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



~ 66 % each, and this suggests that observational constraints are consistent with the prior allocation of emissions between natural and anthropogenic processes, but does not rule out the possibility that the network cannot discriminate between the two. Anthropogenic emissions from Asia are thought to have been increasing steeply in

recent years. EDGAR v4.2 emissions dataset estimates that total anthropogenic emissions from China increased from about 50 Tg CH₄ yr⁻¹ in 2000 to over 73 Tg CH₄ yr⁻¹ in 2008, an increase of almost 50 %. CarbonTracker-CH₄ does not show a steady upward trend in emissions from temperate Asia, but there is a $5\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr}^{-1}$ increase between the average of the first and last five years, a change that is well within the estimated uncertainty for this region of 15 Tg CH₄ yr⁻¹. Bergamaschi et al. (2013) also note lower increases in emissions from Asia than implied by the EDGAR emissions. For temperate North America, there does not appear to be much change over the decade in total estimated emissions; however, fugitive emissions from fossil fuel production in temperate North America show significant increases from prior emissions during winter months, when biogenic emissions are smallest (Fig. 11). By the end of the decade, winter fossil fuel emissions from temperate North America end up higher than prior flux estimates by about $4 \, \text{Tg} \, \text{CH}_4 \, \text{yr}^{-1}$, exceeding the estimated uncertainty of $\sim 3 \, \text{Tg} \, \text{CH}_4 \, \text{yr}^{-1}$. Due to large variability of biogenic emissions, it is difficult to see evidence of this trend during the warmer seasons, and the variability also may mask evidence of increasing fossil fuel emissions in the total estimated emissions. It is interesting to note that, as shown in Table 1, many of the sites with the largest residuals are located near potential sources of fugitive emissions from fossil fuel use. An example is SGP (Southern Great Plains) located in Northern Oklahoma. Figure 12 shows that it is increasingly difficult over time for the inversion to fit this site, possibly due to increasing emissions from fossil fuel production nearby and in Northern Texas. This feature is qualitatively consistent with the results of Miller et al. (2013 - PNAS, now accepted), who calculated larger than predicted emissions of CH₄ related to fossil fuel extraction in this part of the USA. The recent expansion of oil and gas production is not included in the prior used for CarbonTracker-CH₄. Petron et al. (2012) have recently suggested that leakage from

fossil fuel production in Colorado's Denver-Julesburg Basin may be several times larger that estimated by state and local inventories. Karion et al. (2013) have recently shown that emissions from a gas field in Utah may be much higher that previous estimates as well.

4.3 The tropics

Tropical emissions are difficult to constrain because of the sparse distribution of atmospheric observations, but also due to the tendency of the tropical atmosphere to rapidly mix surface signals throughout a deep atmospheric column. Many of the observation sites in tropical latitudes are located on islands where they sample marine air from higher latitudes. Ascension Island, for example often sees air from the South Atlantic, rather than air transported westward from tropical Africa. Although the site located in the Seychelles site sometimes sees air from southern Asia, often it sees air transported from the southern Indian Ocean. In particular, it is difficult to see how the current observational network can independently constrain tropical Asia and tropical Africa. On the other hand, Pacific Ocean sites may make it possible to discriminate between tropical America and Asia.

In agreement with the results found by other studies (e.g. Houweling et al., 1999; Bergamaschi et al., 2007; Mikaloff-Fletcher et al., 2004a, b), CarbonTracker-CH₄ increases tropical emissions relative to prior estimates. The average total prior emission is $132\,\mathrm{Tg\,CH_4\,yr^{-1}}$ and the posterior total is about $157\,\mathrm{Tg\,CH_4\,yr^{-1}}$, an increase of 19%. The estimated uncertainty is $\pm 11\,\mathrm{Tg\,CH_4\,yr^{-1}}$, a decrease from the prior uncertainty of about 39%. This suggests that the observations are able to add some information about tropical emissions to the inversion. Most of the adjustment in emissions goes to wetlands (an increase of 31% from $\sim 65\,\mathrm{Tg\,CH_4\,yr^{-1}}$ to $\sim 84\,\mathrm{Tg\,CH_4\,yr^{-1}}$, with a decrease in uncertainty from 14 to $8\,\mathrm{Tg\,CH_4\,yr^{-1}}$, or 57%). Posterior anthropogenic emissions are essentially unchanged from priors with posterior emissions of $49\,\mathrm{Tg\,CH_4\,yr^{-1}}$ for agriculture and waste, and $\sim 7.5\,\mathrm{Tg\,CH_4\,yr^{-1}}$ for fossil fuel emissions,

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I4 FI

■ Back Close

Full Screen / Esc

Printer-friendly Version



Interactive Discussion



with the uncertainty in total anthropogenic emissions decreasing from 11 Tg CH₄ yr⁻¹ to about 7 TgCH₄ yr⁻¹. CASA-GFED prior flux estimates for biomass burning are increased from about $10 \text{ Tg CH}_4 \text{ yr}^{-1}$ to $11 \text{ Tg CH}_4 \text{ yr}^{-1}$.

Interestingly, the estimated increases in decadal-average emission from tropical wetlands are not evenly distributed among the tropical regions of South America, Africa and Asia. Changes are largest for South America, increasing from a prior of about 19 Tg CH₄ yr⁻¹ to almost 32 Tg CH₄ yr⁻¹ (+68 %). For Africa, posterior emissions increase from 17 to 21 TgCH₄ yr⁻¹ (+24 %) and for Asia, 29 to 35 TgCH₄ yr⁻¹ (+21 %). The estimates from CarbonTracker-CH₄ compare well with Melack et al. (2004), who estimate that the Amazon basin emits about 29 Tg CH₄ yr⁻¹ using a combination of field studies and satellite observations of wetland extent. Estimated emissions from anthropogenic sources remain very close to prior estimates for all tropical regions, and this is also the case for biomass burning. For all tropical regions, the posterior uncertainty is only slightly reduced with respect to the prior uncertainty, generally less than 15%, and the high posterior correlations between these regions make it difficult to have confidence that the inversion is able to constrain information about these regions.

Although observations indicate inter-annual variability in the CH₄ growth rate in the tropics, CarbonTracker-CH₄ is not able to capture this very well as discussed in the previous section on evaluation. Figure 13 shows both the increase in posterior CH₄ emission estimates from the prior, as well the inter-annual variability of the estimates. Total biogenic emissions (e.g. agriculture/waste and wetlands) were larger than normal during 2007 and 2008 in agreement with the analysis of Dlugokencky et al. (2009), who noted that both of these years were relatively wet in the tropics. Wet years are also years with lower fire emissions and the posterior emissions of CarbonTracker-CH₄ show a significant anti-correlation of fire and wetland emissions as shown in Fig. 14.

In addition to a greatly needed expansion of sites sensitive to the tropical biosphere (e.g. Miller et al., 2007), progress on constraining tropical emissions could be made by increasing the length of the assimilation window, allowing signals to reach existing observation sites from terrestrial tropical source regions, and by using aircraft observa**ACPD**

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Introduction **Abstract**

Conclusions References

> **Tables Figures**

I M

Back Close

4.4 The Southern Hemisphere mid-latitudes

Decadal mean CH₄ emission estimates from southern temperate latitudes (Temperate South America, South Africa and Australia) increase from a prior of 78 Tg CH₄ yr⁻¹ to 91 Tg CH₄ yr⁻¹, an increase of about 17%. The aggregated uncertainty estimate decreases from about $12 \text{ Tg CH}_4 \text{ yr}^{-1}$ to $7 \text{ Tg CH}_4 \text{ yr}^{-1}$, a decrease of about 40%. The largest estimated CH₄ emissions are from temperate South America (54 Tg CH₄ yr⁻¹), followed by temperate South Africa $(22\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr}^{-1})$ and Australia/New Zealand (15 Tq CH₄ yr⁻¹). Annual average flux estimates for the aggregated total emissions, as well as the individual processes are little changed from prior estimates for both temperate southern Africa and Australia/New Zealand. The posterior uncertainty estimates for these regions are essentially unchanged as well, indicating a lack of significant observational constraints for these regions.

On the other hand, aggregated total emissions for temperate South America are adjusted from $43\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr}^{-1}$ to $54\,\mathrm{Tg}\,\mathrm{CH_4}\,\mathrm{yr}^{-1}$, although with a relatively small uncertainty reduction of about 9 %. The adjustment to CH₄ emissions occurs as an increase from natural biogenic sources, since fossil fuel emissions and agriculture/waste prior flux estimates are small for this region. Emissions from natural wetlands increase by $7 \text{ Tg CH}_4 \text{ yr}^{-1}$ over prior estimates, and agriculture and waste by close to $4 \text{ Tg CH}_4 \text{ yr}^{-1}$. The uncertainty reduction for natural wetlands is very small, while uncertainty estimates for agriculture and waste are about 13% smaller than the priors.

The global ocean 4.5

Emissions of CH₄ from oceans are thought to make only a small contribution to the atmospheric CH₄ budget, with a prior flux estimate in CarbonTracker-CH₄ of 2205

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page **Abstract**

Introduction

Conclusions

References

Tables

Figures





Back



Full Screen / Esc

Printer-friendly Version









Discussion Pape

14, 2175–2233, 2014

L. M. Bruhwiler et al.

CarbonTracker-CH₄

ACPD

Conclusions

We have created an assimilation system for atmospheric methane, CarbonTracker-CH₄ and used it to estimates CH₄ emissions during 2000–2010 over large spatial scales. We find that simulated CH₄ mole fractions calculated using optimized emissions at each measurement site agree well with observations, generally to within 15 ppb. Also, comparison of posterior methane profiles with measurements of CH₄ vertical profiles from aircraft that were not used in the assimilation show very good agreement, giving further confidence in the estimated emissions over large scales, as well as the representation of the transport processes that maintain the free-tropospheric CH₄ abundances. Large underestimates of CH₄ abundance can sometimes occur at the lower levels of the aircraft profiles in regions where there are likely strong local/regional sources that cannot be resolved by the spatial resolution of the system, are underestimated by the prior emission estimates, and/or not "seen" by other sites. This implies that use of aircraft data could supply important constraints for future inversions.

~ 15 Tg CH₄ yr⁻¹. Posterior estimates are adjusted downwards to 12.6 Tg CH₄ yr⁻¹,

a difference that exceeds the posterior uncertainty estimate of about 1.5 Tg CH₄ yr⁻¹. Rhee et al. (2009) have proposed that earlier estimates of the oceanic methane emis-

sions were biased towards supersaturated waters, and that the emissions are much lower, about 0.6–1.2 Tg CH₄ yr⁻¹, a decrease of over a factor of 10. It is therefore pos-

sible that future versions of CarbonTracker-CH₄ will not include estimation of ocean emissions. Note that the ocean source does not include ebullition from methane seeps

and subsea permafrost (e.g. Shakhova et al., 2010). The size and variability of emis-

sions from this source are not currently well understood, but since significant flux to the atmosphere can only occur in relatively shallow waters, this source would likely be

aliased with terrestrial sources by CarbonTracker-CH₄.

We have also demonstrated the diagnostic value of globally and zonally integrated CH₄ abundances. Comparison of observed and estimated global CH₄ abundances

Title Page **Abstract** Introduction

Conclusions References

> **Tables Figures**

Back Close Full Screen / Esc

Printer-friendly Version

Interactive Discussion



2206

Discussion Paper

Back



allows determination of the relative importance of prior estimates and observational constraints to the solution. Likewise, comparison of observed and posterior CH4 mole fractions integrated over latitude zones indicates whether observational constraints are sufficient to capture observed temporal variability. Since the growth rate of globally and zonally integrated CH₄ abundance directly reflects changes in emissions and sinks, comparison of observed and simulated integrated growth provides insight into whether the inter-annual variability of fluxes is accurately recovered. Indeed we have shown that CarbonTracker-CH₄ is able to simulate the observed zonal average mole fractions and capture inter-annual variability in emissions guite well at high northern latitudes. In contrast, CarbonTracker-CH₄ is less successful in the tropics where it misses significant variability and is more influenced by prior flux estimates. This is expected given the limited number of tropical network sites and the short smoother EnKF time window.

CarbonTracker-CH₄ posterior estimates of total fluxes at high northern latitudes are about 81 Tg CH₄ yr⁻¹, about 12 Tg CH₄ yr⁻¹ (13%) lower than prior estimates, a result that is consistent with other atmospheric inversions. Emissions from European wetlands are decreased by 30%, as found by Bergmaschi et al. (2005); however, unlike their results, emissions from wetlands in Boreal Eurasia are increased. Although CarbonTracker-CH₄ does not estimate increases in emissions from high northern latitudes over the decade covered by the inversion, significant inter-annual variability is recovered. During the exceptionally warm summer of 2007, estimated emissions were higher than the decadal average by 4.4 TgCH₄. During 2008, temperatures returned to more normal values over Arctic North America while they stayed above normal over Arctic Eurasia. CarbonTracker-CH₄ estimates were 2.4 Tg CH₄ higher than the decadal average, and the anomalous emissions occurred over Arctic Eurasia, suggesting that the inversion can discriminate between these two source regions. It is also encouraging that CarbonTracker-CH₄ estimates for the Arctic agree reasonably well and within estimated uncertainties with the analysis of flux observations by McGuire et al. (2012), although they are somewhat lower (16 Tg CH₄ yr⁻¹ compared to 25 Tg CH₄ yr⁻¹).

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page Introduction **Abstract** Conclusions References **Tables Figures** \triangleright

Full Screen / Esc

Close

CarbonTracker-CH $_4$ estimates for temperate latitudes are slightly increased over prior estimates, but about $10\,\mathrm{Tg}\,\mathrm{CH}_4\,\mathrm{yr}^{-1}$ is redistributed from Asia to North America. We used time invariant prior flux estimates for 2000 through 2005 when the growth rate of global atmospheric CH $_4$ was relatively small, so the assimilation does not estimate changes in natural or anthropogenic emissions. After 2006, when atmospheric CH $_4$ began to increase again, CarbonTracker-CH $_4$ allocates some of the emission increases to anthropogenic emissions at temperate latitudes, and some to tropical wetland emissions. The impact of increases in anthropogenic emissions from Asia implied by bottom up production statistics are not seen in the posterior flux estimates, but for temperate North America, the prior flux estimates are increase by about $4\,\mathrm{Tg}\,\mathrm{CH}_4\,\mathrm{yr}^{-1}$ during winter when signals from much larger biogenic emissions are small. Examination of the residuals at North American observation sites suggests that increased CH $_4$ emissions from gas and oil exploration may play a role.

The tropics are not currently well resolved by CarbonTracker- CH_4 due to sparse observational coverage and a short smoother window. However, posterior uncertainties are slightly reduced from prior uncertainties and posterior emissions are about 18% higher than prior estimates. Most of this increase is allocated to tropical South America rather than being distributed over all tropical regions. Our estimates for tropical South America are about $32\,Tg\,CH_4\,yr^{-1}$, in reasonably good agreement with the analysis of Melack et al. (2004), who obtained $29\,Tg\,CH_4\,yr^{-1}$ for the Amazon Basin.

As we have shown using CarbonTracker- CH_4 , even with the current sparse observational network it is possible to be able to draw conclusions about continental budgets of atmospheric CH_4 and to track and attribute variability in relatively well-sampled regions. However, information about fluxes at policy relevant scales remains elusive without increased observational coverage. This is especially true in the tropics, where droughts and flooding may have significant impact on emissions.

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ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I₫

•



Back

Close

Full Screen / Esc

Printer-friendly Version



Discussion Paper

Discussion Paper

Discussion Pape

Conclusions References

> **Tables Figures**

[■

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



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ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page **Abstract** Introduction Conclusions References

Tables Figures

[■ \triangleright

Close

Full Screen / Esc

Back

Printer-friendly Version



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ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

Back Close

Full Screen / Esc

Printer-friendly Version



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14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

> **Tables Figures**

[]◀

Close

Back

Full Screen / Esc

Printer-friendly Version



Interactive Discussion

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14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page **Abstract** Introduction Conclusions References

> **Tables Figures**

[]◀ \triangleright

Back Close

Full Screen / Esc

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

- Title Page **Abstract** Introduction
- Conclusions References
 - **Tables Figures**
- []◀ \triangleright
- **Back** Close
 - Full Screen / Esc
- **Printer-friendly Version**
- Interactive Discussion

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ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I

I

I

Back Close

Printer-friendly Version

Full Screen / Esc

Table 1. CarbonTracker-CH₄ data preprocessing.

Measurement Program	Data Preprocessing
ESRL discrete surface	All valid* data. Multiple values from the same day and location are averaged. No sample time-of-day restriction (see exception below).
EC in situ sites	All valid data from highest intake. Day average using 12:00–16:00 LST.

^{*} In this context "Valid Data" means the observation is thought to be free of sampling and analytical problems and has not been locally influenced.

14, 2175-2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I∢











Full Screen / Esc

Printer-friendly Version





Table 2. Summary of the observation sites used in CarbonTracker-CH₄, and the performance of the assimilation scheme at each site. "#Obs." and "#Rej." are the number of observations available and the number of observations for which the prior simulated concentrations 3σ from the observations using a normal distribution defined with the observed value as the mean and the model-data mismatch error (**R**) as the standard deviation. The bias is the long-term mean of the posterior residuals (simulated-observed), σ is the standard deviation of the residuals for each site, and C2 is the chi-squared statistic (χ^2) calculated as the mean residual divided by the prior uncertainty (Simulated-Observed /(**HQH** + **R**); where **H** is the matrix of transport response, **Q** is the prior flux uncertainty and **R** is the model-data mismatch error).

Site Code	Lab	Lat.	Lon.	Elev. m a.s.l.	# Obs.	# Rej.	R ppb	Bias ppb	σ ppb	χ ² ppb
abp_01d0	ESRL	12.77° S	38.17° W	1.0	112	3	7.5	-8.4	7.7	2.0
alt_01d0	ESRL	82.45° N	62.51° W	200.0	532	0	15	-2.2	8.7	0.3
alt_06c0	EC	82.45° N	62.51° W	200.0	3181	10	15	-1.2	10.2	0.4
amt_01d0	ESRL	45.03° N	68.68° W	50.0	267	4	30	-6.1	22.8	0.4
amt_01p0	ESRL	45.03° N	68.68° W	50.0	174	0	30	8.0	16.5	0.3
asc_01d0	ESRL	7.97° S	14.4° W	74.5	961	79	7.5	-10.0	9.3	3.0
ask_01d0	ESRL	23.18° N	5.42° E	2728.0	491	0	25	-6.9	9.1	0.2
azr_01d0	ESRL	38.77° N	27.38° W	40.0	350	16	15	-12.0	15.9	1.7
bal_01d0	ESRL	55.35° N	17.22° E	3.0	974	0	75	1.4	29.4	0.1
bhd_01d0	ESRL	41.41° S	174.87° E	85.0	165	0	7.5	-4.1	5.4	0.7
bkt_01d0	ESRL	0.2° S	100.32° E	864.5	345	0	75	6.8	30.8	0.2
bme_01d0	ESRL	32.37° N	64.65° W	30.0	256	14	15	-13.6	17.4	2.1
bmw_01d0	ESRL	32.27° N	64.88° W	30.0	352	7	15	-13.2	12.8	1.4
brw_01d0	ESRL	71.32° N	156.61° W	11.0	514	13	15	-5.8	16.1	1.1
bsc_01d0	ESRL	44.17° N	28.68° E	3.0	501	1	75	-14.4	56.2	0.5
cba_01d0	ESRL	55.21° N	162.72° W	21.34	892	23	15	-10.6	13.4	1.1
cdl_06c0	EC	53.99° N	105.12° W	600.0	1390	77	25	-24.7	30.3	2.1
cgo_01d0	ESRL	40.68° S	144.69° E	94.0	416	0	7.5	-4.1	4.6	0.6
chr_01d0	ESRL	1.7° N	157.17° W	3.0	426	79	7.5	-14.6	9.9	5.2
crz_01d0	ESRL	46.45° S	51.85° E	120.0	453	0	7.5	-2.9	4.3	0.5
egb_06c0	EC	44.23° N	79.78° W	251.0	1810	0	75	-6.9	28.7	0.1

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page								
Abstract	Introduc							
Conclusions	Referer							
Tables	Figur							
I	►I							
18								
Dook	Class							
Back Clos Full Screen / Esc								
Full Scre	en/Esc							

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ESRL

15.0° N

Site	Lab	Lat.	Lon.	Elev.	#	#	R	Bias	σ	χ^2
Code				ma.s.l.	Obs.	Rej.	ppb	ppb	ppb	ppb
eic_01d0	ESRL	27.15° S	109.45° W	50.0	323	3	7.5	-7.3	5.3	1.4
esp_06c0	EC	49.58° N	126.37° W	7.0	403	0	25	-6.8	12.3	0.3
etl_06c0	EC	54.35° N	104.98° W	492.0	1780	135	25	-30.1	31.9	2.8
fsd_06c0	EC	49.88° N	81.57° W	210.0	3409	10	25	-9.4	18.3	0.6
gmi_01d0	ESRL	13.43° N	144.78° E	3.0	802	11	15	-10.2	13.0	1.2
hba_01d0	ESRL	75.58° S	26.5° W	30.0	506	0	7.5	0.5	4.6	0.3
hpb_01d0	ESRL	47.8° N	11.01° E	985.0	241	17	25	-13.8	35.7	1.4
hun_01d0	ESRL	46.95° N	16.65° E	248.0	530	3	75	-14.0	43.7	0.3
ice_01d0	ESRL	63.4° N	20.29° W	118.0	529	7	15	-3.3	13.1	0.6
izo_01d0	ESRL	28.31° N	16.5° W	2360.0	443	2	15	-8.5	11.4	0.9
key_01d0	ESRL	25.67° N	80.16° W	3.0	388	3	25	-7.0	20.1	0.6
kum_01d0	ESRL	19.52° N	154.82° W	3.0	720	42	7.5	-6.8	10.6	2.2
kzd_01d0	ESRL	44.06° N	76.82° E	601.0	454	4	75	5.2	44.0	0.2
kzm_01d0	ESRL	43.25° N	77.88° E	2519.0	447	2	25	-2.8	20.2	0.6
lef_01d0	ESRL	45.95° N	90.27° W	472.0	505	6	30	-9.7	28.6	8.0
lef_01p0	ESRL	45.95° N	90.27° W	472.0	341	7	30	-2.1	30.9	0.9
Ilb_06c0	EC	54.95° N	112.45° W	540.0	1152	84	75	-79.9	122.4	3.7
lln_01d0	ESRL	23.47° N	120.87° E	2862.0	222	1	25	-4.1	24.4	0.9
lmp_01d0	ESRL	35.52° N	12.62° E	45.0	206	1	25	-0.7	20.5	0.5
mhd_01d0	ESRL	53.33° N	9.9° W	5.0	416	0	25	-4.6	11.4	0.2
mid_01d0	ESRL	28.21° N	177.38° W	3.7	525	5	15	-10.7	10.9	1.0
mkn_01d0	ESRL	0.05° S	37.3° E	3897.0	146	0	25	-14.3	14.8	0.7
mlo_01d0	ESRL	19.54° N	155.58° W	3397.0	565	0	15	-2.4	10.9	0.5
nmb_01d0	ESRL	23.58° S	15.03° E	456.0	164	0	25	-7.8	11.4	0.3
nwr_01d0	ESRL	40.05° N	105.58° W	3523.0	543	18	15	-11.1	15.2	1.5
oxk_01d0	ESRL	50.03° N	11.8° E	1022.0	202	2	75	-12.5	42.9	0.3
pal_01d0	ESRL	67.97° N	24.12° E	560.0	377	54	15	16.7	35.1	0.2
poc000_01d1	ESRL	0.0° N	155.0° W	10.0	173	33	7.5	-13.9	9.5	4.7
pocn05_01D1	ESRL	5.0° N	151.0° W	10.0	174	29	7.5	-15.1	9.5	5.3
pocn10_01D1	ESRL	10.0° N	149.0° W	10.0	174	45	7.5	-16.0	14.0	7.6

ACPD

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

14, 2175-2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I◀











Back



Full Screen / Esc

Printer-friendly Version

Interactive Discussion



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168

26

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Site	Lab	Lat.	Lon.	Elev.	#	#	R	Bias	σ	χ ²
Code	Lab	Lai.	LOII.	m a.s.l.	# Obs.	# Rej.	ppb	ppb	ppb	χ ppb
	ECDI	20.0° N	141.0° W	10.0			7.5	-6.8		2.9
pocn20_01D1 pocn25_01D1	ESRL ESRL	20.0 N 25.0° N	141.0 W 139.0° W	10.0	166 155	13 14	7.5 7.5	-6.8 -7.0	11.5 11.2	2.9
pocn25_01D1	ESRL	25.0 N 30.0° N	139.0 W 135.0° W	10.0	153	18	7.5 7.5	-7.0 -4.7	13.9	2.6
. –	ESRL	30.0 N 35.0° N	135.0 W 137.0° W	10.0	5	0	7.5 7.5	-4.7 -4.0		2. <i>1</i> 1.4
pocn35_01D1		5.0°S	157.0 W 159.0° W						8.6	
pocs05_01D1	ESRL ESRL	5.0 S 10.0° S	159.0 W 161.0° W	10.0	159	31	7.5	-15.3	8.6	5.2
pocs10_01D1			161.0 W	10.0	170	41	7.5	-14.6	10.1	5.4
pocs15_01D1	ESRL	15.0° S		10.0	163	15	7.5	-10.4	9.5	3.4
pocs20_01D1	ESRL	20.0° S	174.0° W	10.0	169	8	7.5	-7.3	7.9	2.0
pocs25_01D1	ESRL	25.0° S	171.0° W	10.0	164	0	7.5	-5.3	6.3	1.1
pocs30_01D1	ESRL	30.0° S	176.0° W	10.0	166	0	7.5	-5.0	5.0	0.8
pocs35_01D1	ESRL	35.0° S	180.0° E	10.0	14	1	7.5	-0.5	8.2	0.5
psa_01d0	ESRL	64.92° S	64.0° W	10.0	542	0	7.5	-2.7	3.7	0.3
pta_01d0	ESRL	38.95° N	123.74° W	17.0	427	1	25	-4.6	16.9	0.4
rpb_01d0	ESRL	13.17° N	59.43° W	45.0	519	2	15	-10.7	10.0	0.9
sct_01p0	ESRL	33.41° N	81.83° W	115.2	351	0	75	-23.5	33.7	0.3
sey_01d0	ESRL	4.67° S	55.17° E	3.0	514	43	7.5	-6.5	12.3	3.1
sgp_01d0	ESRL	36.8° N	97.5° W	314.0	443	10	75	-56.1	57.4	0.8
shm_01d0	ESRL	52.72° N	174.1°E	40.0	482	0	25	-8.7	11.4	0.3
smo_01d0	ESRL	14.25° S	170.56° W	42.0	568	70	7.5	-10.5	9.9	3.6
spo_01d0	ESRL	89.98° S	24.8° W	2810.0	566	0	7.5	-4.1	4.7	0.7
stm_01d0	ESRL	66.0° N	2.0° E	0.0	917	9	15	-1.4	13.5	0.5
sum_01d0	ESRL	72.58° N	38.48° W	3238.0	468	0	15	-4.7	8.4	0.4
syo_01d0	ESRL	69.0° S	39.58° E	11.0	260	0	7.5	-2.6	3.6	0.3
tap_01d0	ESRL	36.73° N	126.13° E	20.0	441	3	75	10.2	61.7	0.5
tdf_01d0	ESRL	54.87° S	68.48° W	20.0	206	0	7.5	-4.2	4.1	0.6
thd_01d0	ESRL	41.05° N	124.15° W	107.0	400	0	25	-7.0	14.6	0.4
uta_01d0	ESRL	39.9° N	113.72° W	1320.0	525	12	25	-5.5	28.7	0.4
uum_01d0	ESRL	44.45° N	111.1°E	914.0	533	1	25	-1.2	22.7	0.3
wbi_01p0	ESRL	41.72° N	91.35° W	241.7	296	12	30	-8.3	38.0	1.4
wgc_01p0	ESRL	38.27° N	121.49° W	0.0	339	53	75	-118.0	158.8	6.9
wis_01d0	ESRL	31.13° N	34.88° E	400.0	552	3	25	-6.2	23.7	8.0
wkt_01d0	ESRL	31.31° N	97.33° W	251.0	409	55	30	-48.6	43.7	3.8
wkt_01p0	ESRL	31.31° N	97.33° W	251.0	298	38	30	-46.7	58.7	5.8
wlg_01d0	ESRL	36.29° N	100.9° E	3810.0	462	17	15	-1.8	20.6	8.0
wsa_06c0	EC	49.93° N	60.02° W	5.0	2314	52	25	3.8	25.5	0.9
zep_01d0	ESRL	78.9° N	11.88° E	475.0	588	11	15	2.2	14.2	0.5

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

-

Figures

14







Back



Full Screen / Esc

Printer-friendly Version





CarbonTracker-CH₄

ACPD

14, 2175-2233, 2014

L. M. Bruhwiler et al.



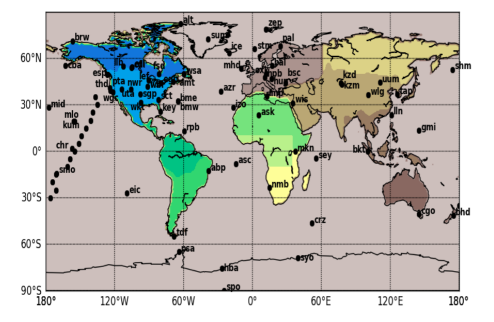


Fig. 1. Map showing locations of observations used in CarbonTracker-CH₄. The colored areas show the outlines for the Transcom 3 source regions with the addition of a tropical African region for this study.



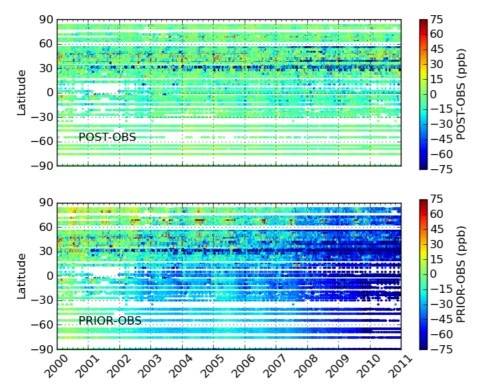


Fig. 2. The CarbonTracker posterior residuals (simulated minus observed, in nmolmol⁻¹) as a function of time and latitude (top) and prior residuals (bottom). Each dot represents the time and location of a CH₄ observation that was assimilated in CarbonTracker. Colors represent the difference between the final simulated value and the actual measurement, with warm colors indicating that CarbonTracker simulates too much methane compared to observations, and cool colors indicating that CarbonTracker estimates too little.

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I

I

I

Back Close

Printer-friendly Version

Full Screen / Esc



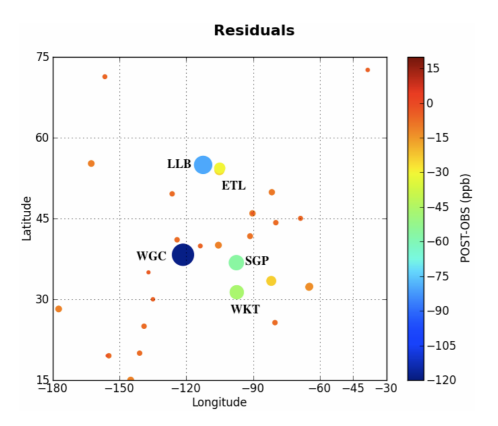


Fig. 3. The CarbonTracker posterior residuals (simulated minus observed, in nmolmol⁻¹) as a function of longitude and latitude for North America. Each bubble has a radius proportional to the size of the residual, and the values are also indicated by the color bar. The largest residuals found by CarbonTracker-CH₄ are labeled also by site code.

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I∢ ≻l

Close

■
Back

Full Screen / Esc

Printer-friendly Version



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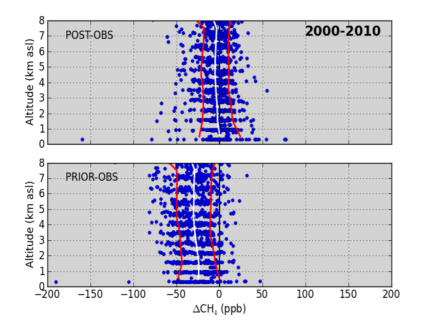


Fig. 4. Statistical summary of residuals for aircraft profiles at a site sampling marine air (Trinidad Head, CA). Units are 10⁻⁹ mol mol⁻¹ of CH₄ (ppb). The top figure shows the post-assimilation residuals (posterior-observed) and the bottom figure shows the residuals with no data assimilation (prior-observed). Aircraft data are not currently assimilated in CarbonTracker so they provide an independent evaluation of the data assimilation. Ideally, the mean of the residuals for the simulations with data assimilation should be near zero. The residuals for the simulations without data assimilation, on the other hand, tend to show large biases.

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I

I

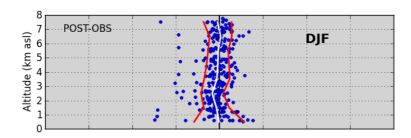
Back Close

Full Screen / Esc

Printer-friendly Version



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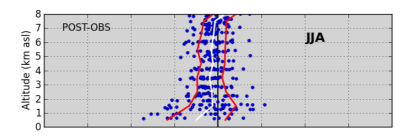


Fig. 5. Statistical summary of residuals for aircraft profiles at a site sampling marine air. Units are 10⁻⁹ mol mol⁻¹ of CH₄ (ppb). The top figure shows the post-assimilation residuals (posterior-observed) and the bottom figure shows the residuals with no data assimilation (prior-observed). Aircraft data are not currently assimilated in CarbonTracker so they provide an independent evaluation of the data assimilation. Ideally, the mean of the residuals for the simulations with data assimilation should be near zero. The residuals for the simulations without data assimilation, on the other hand, tend to show large biases.

ACPD

14, 2175-2233, 2014

iscussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ✓ ▶I

Back Close

Full Screen / Esc

Printer-friendly Version



Discussion Paper

14, 2175–2233, 2014

ACPD

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page



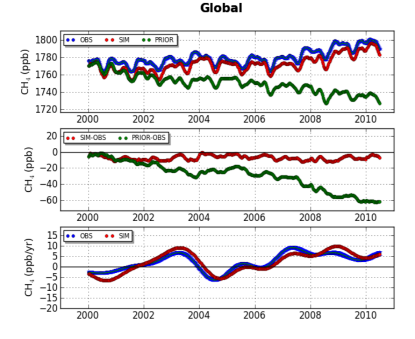
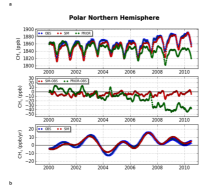


Fig. 6. (Top) time series of observed (dark blue, "OBS", with error bars), assimilated (red, "SIM") and prior (green, "PRIOR") average methane mole fraction. For the "PRIOR" simulations, prior fluxes were used to calculate CH₄ mole fractions, while for the "SIM" simulations CH₄ was calculated using fluxes that were adjusted for optimal agreement with atmospheric observations. Units are ppb (10⁻⁹ mol mol⁻¹). (Middle) the differences from observations for assimilated and prior CH₄ (ppb). (Bottom) derived growth rate of CH₄ mole fraction for observed (with error bars) and assimilated CH₄ mole fraction. The growth rate is computed by taking the first derivative of the average mole fractions shown in the top figure. Units are ppbyr⁻¹ (10⁻⁹ mol mol⁻¹ yr⁻¹).



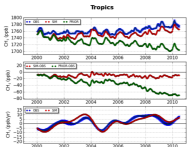


Fig. 7. (a) (Top) time series of observed (dark blue, "OBS" with error bars), assimilated (red, "SIM") and prior (green, "PRIOR") average methane mole fraction for the Polar Northern Hemisphere (53–90° N). (Middle) differences from observations for assimilated and prior CH_4 (ppb). (Bottom) derived growth rate of CH_4 mole fraction for observed and assimilated CH_4 mole fraction for the Polar Northern Hemisphere. **(b)** (Top) time series of observed (dark blue, "OBS"), assimilated (red, "SIM") and prior (green, "PRIOR") average methane mole fraction for the tropics (17.5° S–17.5° N). (Middle) differences from observations for assimilated and prior CH_4 (ppb). (Bottom) derived growth rate of CH_4 mole fraction for observed (with error bars) and assimilated CH_4 mole fraction for the tropics.

ACPD

14, 2175-2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures













Printer-friendly Version



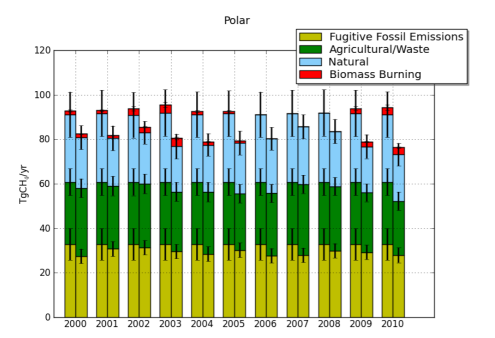


Fig. 8. The contribution to the High Northern Latitude total CH_4 flux from each category of emissions with error bars. For each pair of histogram bars, the prior flux estimates are shown on the left and the posterior estimates on the right. Note that, except for emissions from fires, the prior flux estimates are constant for each year. The units are $TgCH_4yr^{-1}$.

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures











Printer-friendly Version





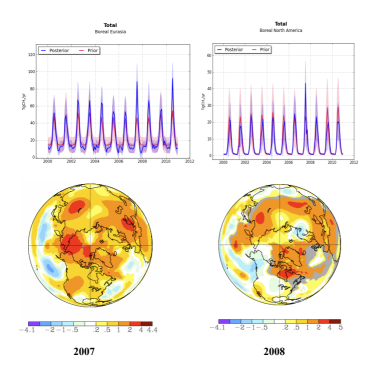


Fig. 9. (Top) Time variation of the prior and estimated CH₄ emissions. Prior estimates are shown in red, and posterior flux estimates are shown in blue. Note that only the biomass burning prior emission estimates vary from year to year; other prior estimates are constant. The posterior estimates, on the other hand, may show considerable variability due to sensitivity of biogenic emissions to year-to-year changes in temperature and precipitation. 1σ uncertainty bounds are shown as light red (prior) and light blue (post-assimilation) shaded areas. Note that microbial sources of methane, such as wetlands and agriculture, are temperaturesensitive and therefore tend to be largest during summer. Units are Tg CH₄ yr⁻¹. (Bottom) Surface temperature anomalies from the NASA GISS analysis relative to 1951-1980 in °C (http://data.giss.nasa.gov/gistemp/, Hansen et al., 2010)

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures













Printer-friendly Version



L. M. Bruhwiler et al.

Abstract

Tables

ACPD

14, 2175–2233, 2014

CarbonTracker-CH₄

Title Page

Introduction

References

Figures

 \triangleright

Close



Discussion Paper

Discussion Paper

[]◀ **Back**

Full Screen / Esc



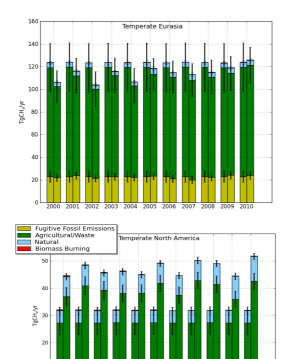


Fig. 10. The contribution to the total CH₄ flux from each category of emissions with error bars for Temperate Eurasia (top) and Temperate North America (bottom). For each pair of histogram bars, the prior flux estimates are shown on the left and the posterior estimates on the right. Note that, except for emissions from fires that are very small for these regions, the prior flux estimates are constant for each year. Units are TgCH₄ yr⁻¹.

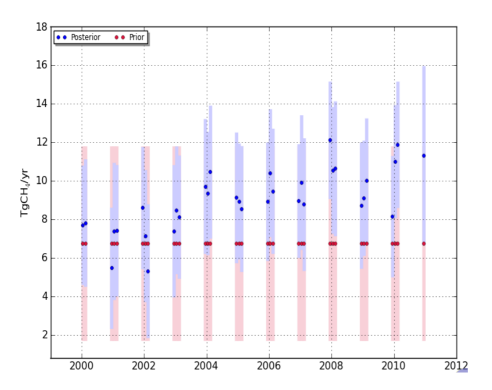


Fig. 11. Time variation of the estimated CH_4 emissions from Temperate North America for winter. Prior flux estimates are shown in red, and posterior flux estimates are shown in blue. 1σ uncertainty bounds are shown as light red (prior) and light blue (posterior) shaded areas. Units are $TgCH_4yr^{-1}$. Each point represents a monthly average estimated emission given in units of $TgCH_4yr^{-1}$.

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I

I

I

Back Close

Full Screen / Esc

Printer-friendly Version



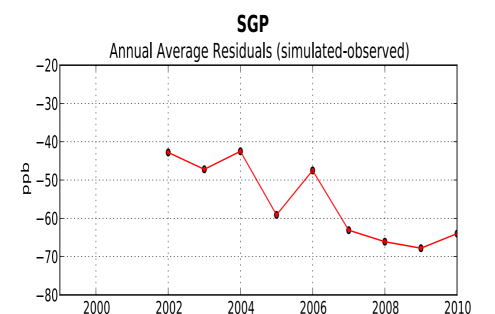


Fig. 12. Time series of residuals (the difference between the posterior and measured mole fractions). Note that the prior is constant over the length of the inversion, and the trend in the residuals can be interpreted to mean that it is increasingly difficult to fit this site over time. Units are 10^{-9} mol mol⁻¹ of CH₄ (ppb).

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Conclusions References

Introduction

Close

Abstract

Back

Tables Figures

Full Screen / Esc

Printer-friendly Version



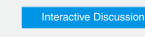
14, 2175–2233, 2014

ACPD

CarbonTracker-CH₄

L. M. Bruhwiler et al.





Tropical

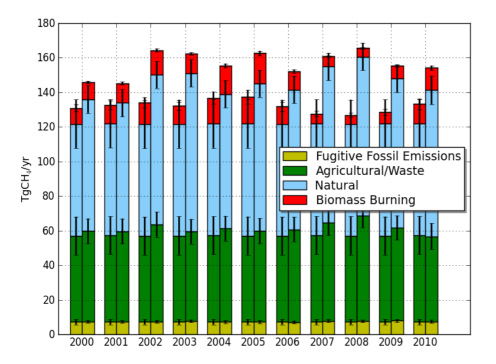


Fig. 13. The contribution to the total CH₄ flux from each category of emissions with error bars for the tropics (Tropical South America, Tropical Asia and Tropical Africa). For each pair of histogram bars, the prior flux estimates are shown on the left and the posterior estimates on the right. Note that, except for emissions from fires that are very small for these regions, the prior flux estimates are constant for each year. The units are TgCH₄ yr⁻¹.

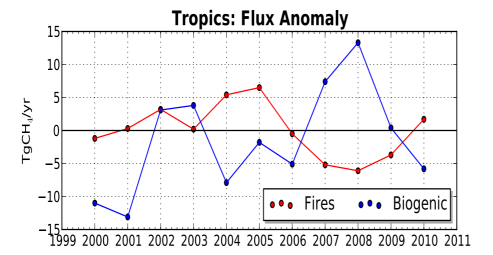


Fig. 14. Time variation of estimated total biogenic (wetlands and agriculture/waste) and fire CH_4 emission anomalies. Anomalies are calculated with respect to 10 yr average posterior emissions. The units are $TgCH_4$ yr⁻¹.

14, 2175–2233, 2014

CarbonTracker-CH₄

L. M. Bruhwiler et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures















Printer-friendly Version

