

Evaluating atmospheric methane inversion model results for Pallas, northern Finland

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A state-of-the-art inverse model, CarbonTracker Data Assimilation Shell (CTDAS), was used to optimize estimates of methane (CH₄) surface fluxes using atmospheric observations of CH₄ as a constraint. The model consists of the latest version of the TM5 atmospheric chemistry-transport model and an ensemble Kalman filter based data assimilation system. The model was constrained by atmospheric methane surface concentrations, obtained from the World Data Centre for Greenhouse Gases (WDCGG). Prior methane emissions were specified for five sources: biosphere, anthropogenic, fire, termites and ocean, of which biosphere and anthropogenic emissions were optimized. Atmospheric CH₄ mole fractions for 2007 from northern Finland calculated from prior and optimized emissions were compared with observations. It was found that the root mean squared errors of the posterior estimates were more than halved. Furthermore, inclusion of NOAA observations of CH₄ from weekly discrete air samples collected at Pallas improved agreement between posterior CH₄ mole fraction estimates and continuous observations, and resulted in reducing optimized biosphere emissions and their uncertainties in northern Finland.

Introduction

Methane (CH₄) is one of the most effective greenhouse gases. Its global warming potential over a 100 year time horizon is 28 times that of

carbon dioxide (IPCC 2013: chapter 8.7), and its atmospheric burden more than doubled since 1750 (MacFarling Meure *et al.* 2006, Rhodes *et al.* 2013). Anthropogenic emissions are the main reason for the increase up to the 1980s

(Rasmussen and Khalil 1981, Khalil and Rasmussen 1985). In the 1990s, the annual growth rate decreased, and became nearly zero in the early 2000s. Several reasons have been proposed to explain the variations: a decline in the emissions from biogenic sources, such as wetlands and rice, in the northern hemisphere (Kai *et al.* 2011), a decline in the fossil fuel emissions (Aydin *et al.* 2011), or an increase in the main sink of CH₄ reaction with atmospheric OH (Krol *et al.* 1998, Bousquet *et al.* 2006, Monteil *et al.* 2011). However, if lifetime and emission were constant, decrease in growth rate is an indicator that atmosphere is reaching a steady state (Dlugokencky *et al.*, 2011). Furthermore, the concentration started to increase again in 2006 (Rigby *et al.* 2008), and the exact reasons are still unknown (Dlugokencky *et al.* 2009, Heimann 2011).

Several inverse modeling studies assessed the reasons for the changes in the methane growth rate (Bergamaschi *et al.* 2005, 2007, 2010, 2013, Bousquet *et al.* 2005, 2011, Bruhwiler *et al.* 2014, Houweling *et al.* 2014). Inverse modeling, which optimizes emissions from different source categories, is a powerful tool for understanding changes in the emissions from both human activities and nature. The main sources of anthropogenic emissions are agriculture, waste and fossil fuels, whereas biosphere emissions are dominated by wetlands (Kirschke *et al.* 2013). Anthropogenic emissions are responsible for long-term and inter-annual variability of atmospheric methane, whereas biosphere emissions have large effects on seasonal cycles. Global methane emission is estimated to be around 500–600 Tg y⁻¹, and for Europe 35–60 Tg y⁻¹, where anthropogenic emissions also dominate (Kirschke *et al.* 2013 and references therein). However, the estimates vary substantially among models and model settings. Although the variation among inverse modeling estimates are smaller than for process models, the choice of transport model (Locatelli *et al.* 2013), observation data sets (Alexe *et al.* 2014, Villani *et al.* 2010) and prior emission distributions affect the results.

In this study, we evaluated the performance of a state-of-the-art inverse model system based on the data assimilation system CTDAS

(Carbon Tracker Data Assimilation Shell, *see* <http://www.carbontracker.eu/ctdas/>) for boreal sites of northern Finland in 2007. The system contains the latest version of the atmospheric chemistry-transport model TM5 (Krol *et al.* 2005) driven by the European Centre for Medium-Range Weather Forecasts (ECMWF) ERA-Interim meteorological fields. TM5 is used with a two-way nested European domain with a 1° × 1° grid over Europe and high northern latitudes. Here, observations of atmospheric CH₄ for 2007 from NOAA's global cooperative air sampling network and other discrete air sampling networks were used to constrain the emissions in the model. The discrete air sample data have been used extensively as constraints in surface flux inversion studies based on data assimilation (Bergamaschi *et al.* 2007, Bruhwiler *et al.* 2014, Houweling *et al.* 1999).

To examine the performance of the inverse model for boreal areas, especially northern Finland, the modeled atmospheric concentrations estimated using the emissions before and after optimization of the emissions were compared with model-independent continuous observations for 2007. Furthermore, two inversions, one with and one without assimilating the Pallas discrete air sample observations were compared to test the effects of the Pallas observations in the inverse model results.

Methods and data sets

CTDAS atmospheric inverse model

CTDAS is a pyshell version of CarbonTracker, developed by NOAA-ESRL (National Oceanic and Atmospheric Administration's Earth System Research Laboratory) and the Wageningen University. The model had originally been designed to optimize carbon dioxide fluxes (Peters *et al.* 2005), and it has been further developed for CH₄ by Bruhwiler *et al.* (2014) and in this study (European CarbonTracker-CH₄). Models developed by Bruhwiler *et al.* (2014) and in this study differ to certain extent: the emission sources that are optimized, the geographical area of focus, i.e. the zoom grid in the TM5 transport model, prior emissions such as anthropogenic and biosphere,

observations assimilated, and source region boundaries are different from their studies.

The optimal weekly mean CH₄ fluxes $F(r,t)$ in region r and time t (week) were calculated as follows:

$$F(r,t) = \lambda_{\text{bio}}(r,t)F_{\text{bio}}(r,t) + \lambda_{\text{anth}}(r,t)F_{\text{anth}}(r,t) + F_{\text{fire}}(r,t) + F_{\text{term}}(r,t) + F_{\text{oce}}(r,t), \quad (1)$$

where F_{bio} , F_{anth} , F_{fire} , F_{term} , F_{oce} are the emissions

from biosphere, anthropogenic activities, fire, termites and ocean, respectively. The scaling factors for the biosphere and anthropogenic emissions are optimized in the model.

In order to assess the influence of the NOAA Pallas flask measurements on the optimized fluxes for European and boreal regions, we performed two runs: S1 including all the measurements listed in Table 1, and S2 excluding the Pallas flask measurements.

Table 1. List of sites used in European CarbonTracker-CH₄. The model data mismatch (mdm) was used as the observation error. Observations were rejected in the assimilation if the estimated mole fractions were not within 3 × observation error. Note that only observations from discrete air samples were used.

Site code	Station name	Country/territory	Contributor	Lat. (°N)	Long. (°E)	Elevation (m a.s.l.)	mdm (ppb)
abp	Arembepe	Brazil	NOAA/ESRL	-12.77	-38.17	0	7.5
alt	Alert	Canada	NOAA/ESRL	82.72	-62.52	210.0	15.0
ams	Amsterdam Island	France	NOAA/ESRL	-37.8	77.53	55.0	7.5
amt	Argyle	USA	NOAA/ESRL	45.03	-68.68	50.0	30.0
arh	Arrival Heights	New Zealand	NIWA	-77.8	166.67	184.0	7.5
asc	Ascension Island	UK	NOAA/ESRL	7.55	14.25	54.0	7.5
ask	Assekrem	Algeria	NOAA/ESRL	23.27	5.63	2710.0	25.0
azr	Terceira Island	Portugal	NOAA/ESRL	38.77	-27.37	40.0	15.0
bal	Baltic Sea	Poland	NOAA/ESRL	55.35	17.22	28.0	75.0
bgu	Begur	Spain	LSCE	41.97	3.23	13.0	15.0
bhd	Baring Head	New Zealand	NOAA/ESRL	-41.41	174.87	85.0	7.5
bkt	Bukit Koto Tabang	Indonesia	NOAA/ESRL	-0.2	100.32	864.5	75.0
bme	St. David's Head	UK	NOAA/ESRL	32.37	-64.65	30.0	15.0
bmw	Tudor Hill	UK	NOAA/ESRL	32.27	-64.87	30.0	15.0
brw	Barrow	USA	NOAA/ESRL	71.32	-156.6	11.0	15.0
bsc	Black Sea	Romania	NOAA/ESRL	44.17	28.67	3.0	75.0
cba	Cold Bay	USA	NOAA/ESRL	55.20	-162.72	25.0	15.0
cfa	Cape Ferguson	Australia	CSIRO	-19.28	147.05	2.0	25.0
cgo	Cape Grim	Australia	NOAA/ESRL	-40.68	144.68	94.0	7.5
chr	Christmas Island	Kiribati	NOAA/ESRL	1.70	-157.17	3.0	7.5
crz	Crozet	France	NOAA/ESRL	-46.45	51.85	120.0	7.5
cya	Casey Station	Australia	CSIRO	-66.28	110.53	60.0	7.5
eic	Easter Island	Chile	NOAA/ESRL	-27.13	-109.45	50.0	7.5
esp	Estevan Point	Canada	EC	49.38	-126.55	39.0	25.0
esp	Estevan Point	Canada	CSIRO	49.38	-126.55	39.0	25.0
fik	Finokalia	Greece	LSCE	35.34	25.67	150.0	15.0
gmi	Guam	USA	NOAA/ESRL	13.43	144.78	2.0	15.0
hba	Halley Bay	UK	NOAA/ESRL	-75.57	-26.5	33.0	7.5
hpb	Hohenpeissenberg	Germany	NOAA/ESRL	-75.57	11.02	985.0	25.0
hun	Hegyhatsal	Hungary	NOAA/ESRL	46.95	16.65	248.0	75.0
ice	Heimaey	Iceland	NOAA/ESRL	63.40	-20.28	100.0	15.0
izo	Izaña	Spain	NOAA/ESRL	28.30	-16.5	2367.0	15.0
key	Key Biscayne	USA	NOAA/ESRL	25.67	-80.2	3.0	25.0
kum	Cape Kumukahi	USA	NOAA/ESRL	19.52	-154.82	3.0	7.5
kzd	Sary Taukum	Kazakhstan	NOAA/ESRL	44.45	75.57	412.0	75.0
kzm	Plateau Assy	Kazakhstan	NOAA/ESRL	43.25	77.87	2519.0	25.0
lln	Lulin	China	NOAA/ESRL	23.47	120.87	2867.0	25.0
lmp	Lampedusa	Italy	NOAA/ESRL	35.52	12.63	45.0	25.0

continued

Table 1. Continued.

Site code	Station name	Country/territory	Contributor	Lat. (°N)	Long. (°E)	Elevation (m a.s.l.)	mdm (ppb)
lpo	Ile Grande	France	LSCE	48.80	-3.58	10.0	15.0
maa	Mawson	Australia	CSIRO	-67.62	62.87	32.0	7.5
mhd	Mace Head	Ireland	NOAA/ESRL	53.33	-9.9	8.0	25.0
mid	Sand Island	USA	NOAA/ESRL	28.20	-177.37	7.7	15.0
mlo	Mauna Loa	USA	NOAA/ESRL	19.54	-155.58	3397.0	15.0
mqa	Macquarie Island	Australia	CSIRO	-54.48	158.97	12.0	7.5
nmb	Gobabeb	Namibia	NOAA/ESRL	-23.57	15.02	461.0	25.0
nwr	Niwot Ridge	USA	NOAA/ESRL	40.05	-105.59	3523.0	15.0
oxk	Ochsenkopf	Germany	NOAA/ESRL	50.03	11.80	1185.0	75.0
pal	Pallas-Sammaltunturi	Finland	NOAA/ESRL	67.97	24.12	560.0	15.0
pdm	Pic du Midi	France	LSCE	42.94	0.14	2877.0	15.0
psa	Palmer Station	USA	NOAA/ESRL	-64.92	-64	10.0	7.5
pta	Point Arena	USA	NOAA/ESRL	38.95	-123.72	17.0	25.0
puy	Puyde Dome	France	LSCE	45.77	2.97	1465.0	15.0
rpb	Ragged Point	Barbados	AGAGE	13.17	-59.43	45.0	15.0
sey	Mahe Island	Seychelles	NOAA/ESRL	-4.67	55.17	7.0	7.5
sgp	Southern Great Plains	USA	NOAA/ESRL	36.78	-97.5	314.0	75.0
shm	Shemya Island	USA	NOAA/ESRL	52.72	174.08	40.0	25.0
smo	Tutuila	USA	AGAGE	-14.24	-170.57	42.0	7.5
spo	South Pole	USA	NOAA/ESRL	-89.98	-24.8	2810.0	7.5
stm	Ocean Station M	Norway	NOAA/ESRL	66.00	2.00	5.0	15.0
sum	Summit	Denmark	NOAA/ESRL	72.58	-38.48	3238.0	15.0
syo	Syowa Station	Japan	NOAA/ESRL	-69	39.58	16.0	7.5
tap	Tae-ahn Peninsula	Korea	NOAA/ESRL	36.72	126.12	20.0	75.0
tdf	Tierradel Fuego	Argentina	NOAA/ESRL	-54.87	-68.48	20.0	7.5
ter	Teriberka	Russia	MGO	69.20	35.10	40.0	15.0
thd	Trinidad Head	USA	NOAA/ESRL	41.05	-124.15	120.0	7.5
uta	Wendover	USA	NOAA/ESRL	39.88	-113.72	1320.0	25.0
uum	Ulaan Uul	Mongolia	NOAA/ESRL	44.45	111.08	914.0	25.0
wis	Sede Boker	Israel	NOAA/ESRL	31.12	34.87	400.0	25.0
wkt	Moody	USA	NOAA/ESRL	31.32	-97.32	708.0	30.0
wlg	Mt. Waliguan	China	NOAA/ESRL	36.28	100.90	3810.0	15.0
wsa	Sable Island	Canada	EC	43.93	-60.02	5.0	25.0
zep	Zeppelinfjellet	Norway	NOAA/ESRL	78.90	11.88	475.0	15.0

TM5 atmospheric transport model

The link between atmospheric CH₄ measurements and exchange of CH₄ at the Earth's surface is the transport of CH₄ in the atmosphere. In our assimilation system, the release 3 of the TM5 chemistry transport model was used as the linearized observation operator. TM5 was run with a 1° × 1° zoom region over Europe (24–74°N, 21°W–45°E), framed by an intermediate grid of 2° × 3° for outer Europe, and 4° × 6° globally (Fig. 1), driven by ECMWF ERA-Interim meteorological fields. Atmospheric chemical loss was calculated using off-line chemistry with monthly tropospheric OH concentrations (Hou-

weling *et al.* 2014). Furthermore, stratospheric sink due to reaction with OH, Cl and O(¹D) were included by applying reaction rates based on a 2D photochemical Max-Planck-Institute (MPI) model (Bergamaschi *et al.* 2005). The global total atmospheric chemical loss, i.e. the integrating OH, Cl and O(¹D) losses during 2007, was about 511 Tg CH₄ y⁻¹, with a methane lifetime of about 9.6 years defined by the global burden divided by the loss.

In this work, atmospheric mole fractions were estimated using the TM5 forward model. The mole fractions estimated with prior emissions are henceforth called 'prior', and the estimates with posterior emissions 'posterior'.

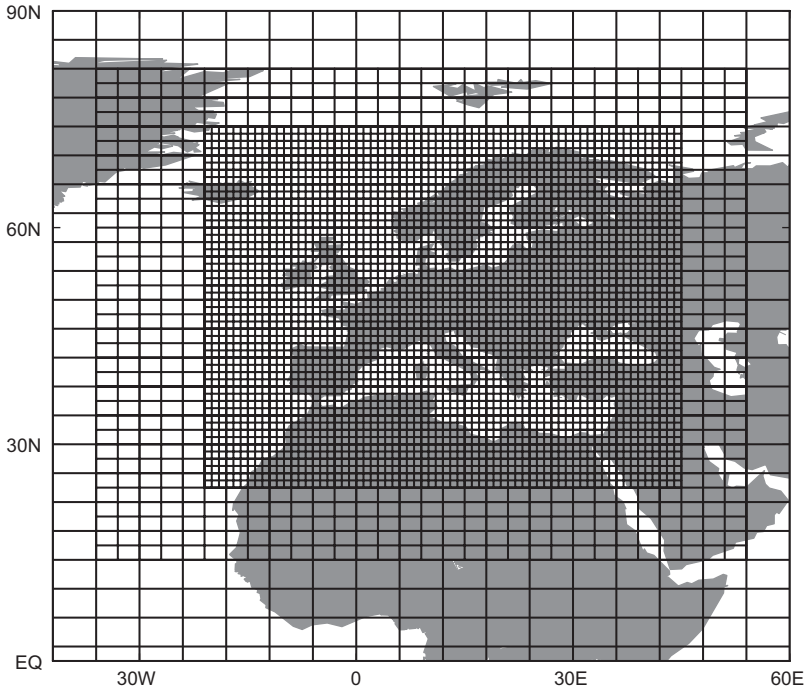


Fig. 1. TM5 zoom grid definition used in the European CarbonTracker-CH₄.

Prior CH₄ flux data sets

The prior data sets consisted of anthropogenic, biosphere, fire, termite and ocean emissions, collected from inventories and studies outside this project. Total posterior emissions were calculated using Eq. 1. All emission fields were gridded to match the finest TM5 grid, i.e., $1^\circ \times 1^\circ$, globally.

Anthropogenic methane emissions are responsible for more than half of the global methane source. For monthly mean anthropogenic emissions, the Emission Database for Global Atmospheric Research ver. 4.2 (EDGARv4.2) was used. The emissions from agricultural waste burning and large scale biomass-burning were removed, because they overlap with fire emissions described later. The annual anthropogenic emission for 2007 was 337 Tg CH₄ y⁻¹ (excluding fires), in which agricultural emissions, such as enteric fermentation (99 Tg CH₄ y⁻¹) and agricultural soils (36 Tg CH₄ y⁻¹), and fugitive emissions from oil and gas (66 Tg CH₄ y⁻¹) and solid fuels (43 Tg CH₄ y⁻¹) dominated. We did not introduce seasonal variations in the anthropogenic emissions.

Natural emissions, dominated by wetlands, are estimated to contribute about 40% of the total emissions, where inter-annual variability of emissions from wetland ecosystems is estimated to be ± 12 Tg CH₄ y⁻¹ (Spahni *et al.* 2011). For monthly-mean biosphere emissions, the estimates by the LPJ-WHyMe vegetation model (Spahni *et al.* 2011) were used. The emissions from rice fields were removed since they were already included in the anthropogenic emissions. EDGARv4.2 estimates of the emissions from rice fields were ca. 6 Tg CH₄ y⁻¹ smaller than LPJ-WHyMe estimates; no scaling was applied to the EDGARv4.2 estimates. The annual biosphere emission for 2007 was 160 Tg CH₄ y⁻¹ (excluding rice fields), with the seasonal cycle already captured in the prior.

Methane emissions from natural fires account for less than 10% of the global total (Kirschke *et al.* 2013). These emissions are an important part of the carbon cycle and their inter-annual variability can be large because of occasional intense fires and events, such as strong El Niño that lead to dry periods around the equator (Langenfelds *et al.* 2002). Also, the spatial variability of fire emissions should be taken into account

(Andreae and Merlet 2001). Monthly-mean fire emissions from the Global Fire Emissions Database ver. 3.1 (GFEDv3.1) were used. The data set contained both natural and anthropogenic fire emissions (van der Werf *et al.* 2010), including agricultural waste burning and large-scale biomass burning with seasonal cycles captured. Annual total fire emissions for 2007 were 17.44 Tg CH₄ y⁻¹ in GFEDv3.1, and 30.36 Tg CH₄ y⁻¹ in EDGARv4.2.

Methane emissions of termites accounts for about 5% of global emissions, but the spatial variability of these emission should be taken into account. Annual-mean termite emissions from Ito *et al.* (2012) were used. The emissions were estimated based on Fung *et al.*'s (1991) up-scaling method, biome-specific termite biomass density and emission factors were obtained from Fraser *et al.* (1986), and a historical land cover map based on Hurtt *et al.* (2006) was used to introduce inter-annual variability. The average termite biomass density in boreal forests was assumed to be zero. We did not introduce seasonal variations into the termites emissions. In this study, emissions for 2007 were assumed to be the same as the latest estimate, i.e. of 2006.

Methane emissions from open oceans are a relatively minor, about 0.2% of the global emissions. Monthly-mean ocean emission fields were pre-calculated based on seasonal methane saturation ratios from Bates *et al.* (1996), which were derived from measurements of seawater and atmospheric methane mixing ratios throughout the Pacific Ocean. The saturation ratios were used globally as zonal averages. The difference between air and seawater partial pressures of methane, $d_p(\text{CH}_4)$, was calculated using saturation ratios. The zonal monthly mean dry air CH₄ mixing ratios were taken from GLOBALVIEW-CH4 (Cooperative Atmospheric Data Integration Project — Methane, *see* www.esrl.noaa.gov/gmd/ccgg/globalview/ch4/ch4_intro.html). Sea level pressure, sea surface temperature, sea-ice concentration and 10 meter wind speeds were from the ECMWF ERA-interim data (Dee *et al.* 2011). Solubility of methane in seawater was calculated according to Wiesenburg and Guinasso (1979), assuming salinity of 35‰. Gas transfer velocity was parametrized using the wind speed and Schmidt number (Wanninkhof 1992).

Fluxes were then calculated as the product of gas transfer velocity, gas solubility and $d_p(\text{CH}_4)$. Sea-ice was assumed to inhibit gas transfer.

Atmospheric methane observations

Global methane atmospheric measurements were obtained from the World Data Centre for Greenhouse Gases (WDCGG). Measurements are from the NOAA, CSIRO, EC, LSCE, NIWA and MGO discrete air samples, and background measurements were selected according to each contributor's quality control flags. The location of each site is shown in Fig. 2. The model data mismatch (mdm), used for the criteria of observation rejection thresholds and observation error covariance matrix, was defined by site types: 7.5 ppb for marine boundary layer and high southern hemisphere sites, 15 ppb for mixed sites, 25 ppb for land and tower sites, 30 ppb for sites with large variability in observations, and 75 ppb for so called "problematic" sites. For the list and details of the measurement sites, *see* Table 1. The observations were rejected in the assimilation if estimated concentrations were not within three times the measurement errors. The number of measurements available varied by site, but around 70 measurements were assimilated per week.

To assess the model results in the European boreal regions, two independent observation data sets from northern Finland were compared with the TM5-estimated CH₄ mole fractions using the prior and posterior emissions. The data sets were the FMI continuous atmospheric measurements from Pallas (Aalto *et al.* 2007), and the Fourier transform infrared spectroscopy (FTIR) measurements from Sodankylä (Kivi *et al.* 2014).

Pallas is located at 67.58°N and 24.06°E (565 m a.s.l.), where its main station Sammalunturi is located on the top of a hill. About 6% of the nearest 20 km² consists of open wetland, and the area is sparsely populated. During 2007, CH₄ was measured four times per hour with an automated gas chromatographic system (Agilent 6890N) equipped with a flame ionization detector for CH₄ detection. The measurements are calibrated using standards on the WMO/CCL scale (Aalto *et al.* 2007). Hourly mean observations for day and night were used.

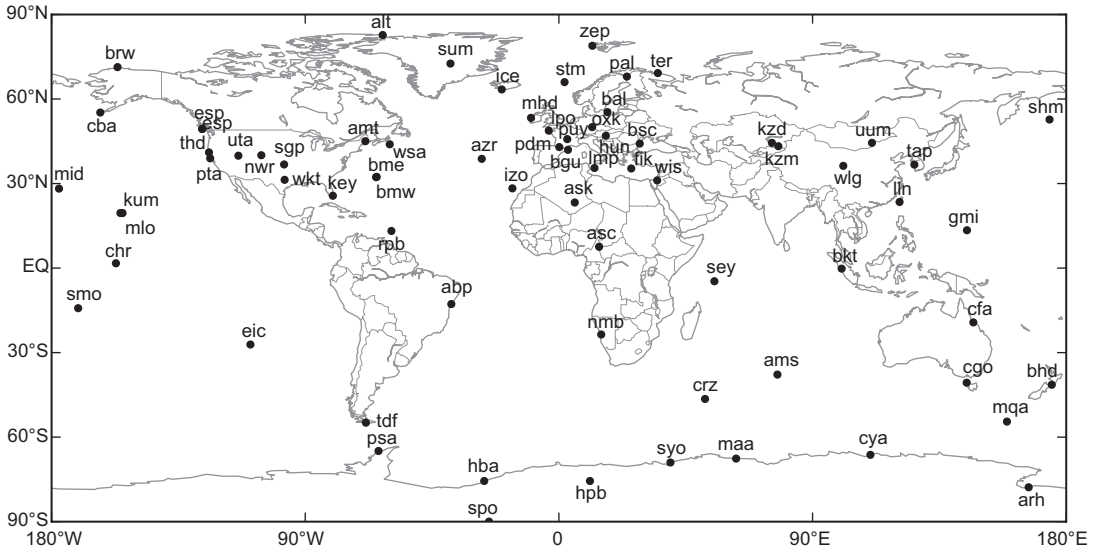


Fig. 2. Locations (black dots) of the sites from which the data were assimilated in European CarbonTracker-CH₄ for 2007. For site-name codes see Table 1.

Sodankylä is located at 67.36°N, 26.63°E (179 m above mean sea level), about 150 km southeast of Pallas, and a FTIR instrument has been operated at the site since February 2009. The FTIR instrument in Sodankylä acquires solar spectra using a Bruker 125HR Fourier transform spectrometer. The instrument is participating in the Total Carbon Column Observing Network (TCCON, <http://www.tcon.caltech.edu/>), and the total column measurements were processed using the standard approach used in the network (Wunch *et al.* 2011). Due to its high-latitude location, there is little or no sunlight during winter, FTIR observations between November and January are discontinued. Monthly means and standard deviations of the measurements were calculated using all data from 2009–2012. Note that the inverse model was run for 2007, so we compare only the shapes of the seasonal cycle with the model estimates. Note that Sodankylä is both spatially and temporally a model-independent site; i.e., no measurements from Sodankylä were used in this study.

TransCom and land-ecosystem regions

We optimized fluxes region-wise, where the

regions r in Eq. 1 were defined by TransCom and land-ecosystem maps. The global surface was divided into 16 regions, based on the TransCom regions used in Peters *et al.* (2007), except that oceans were aggregated into one region and Europe was divided into four subregions (Fig. 3). Further, terrestrial areas were divided based on soil types, because methane emissions are affected by soil properties (Matthews and Fung 1987, Yvon-Durocher *et al.* 2014). Land-ecosystem regions were defined mainly based on Prigent *et al.* (2007) and Wania *et al.* (2010), also used in LPJ-WhyMe vegetation model (Spahni *et al.* 2011). The regions consisted of six types of land ecosystems (Fig. 4): inundated wetland and peatland (IWP), wet mineral soil (WMS), rice (RIC), anthropogenic land (ANT), water (WTR) and ice (ICE). Each grid point was therefore defined by TransCom and land-ecosystem region. The land-ecosystem map was not dynamic and did not change during the year. The prior scaling factors were all equal to one: $\lambda = (\lambda_{\text{bio}}, \lambda_{\text{anth}}) = (1, 1, \dots, 1) = 1$. For land-ecosystem region IWP and WMS, biosphere emissions were optimized, i.e. $\lambda_{\text{anth}} = 1$, and for RIC, ANT and WTR, anthropogenic emissions were optimized, i.e. $\lambda_{\text{bio}} = 1$. Theoretically, this approach results in 72 (14 TransCom \times 5 land-ecosystem + ocean

Fig. 3. The land regions in European CarbonTracker-CH₄. They are defined according to the TransCom regions, except for Europe, which is divided into four sub-regions.

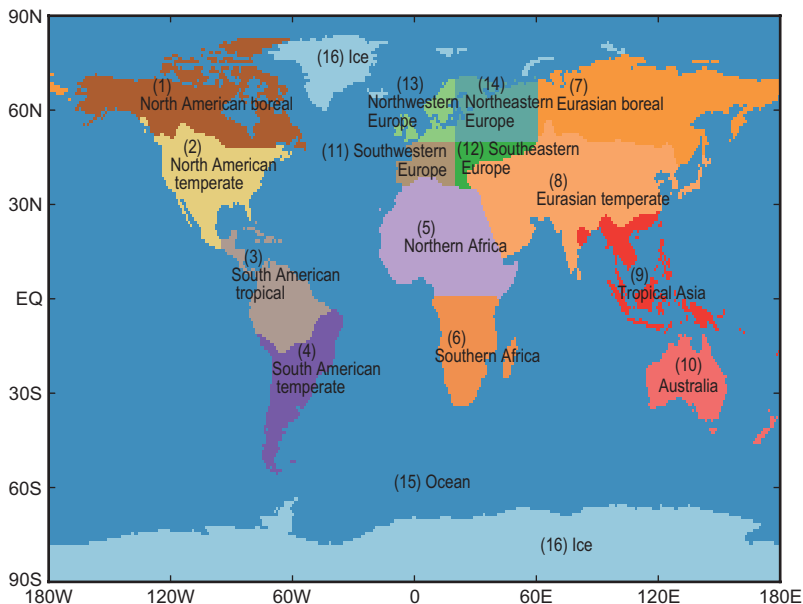
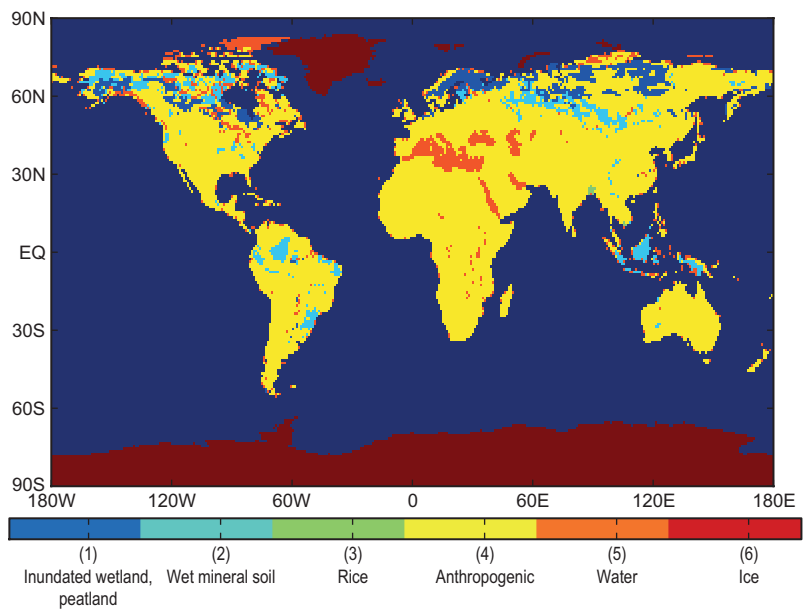


Fig. 4. Land-ecosystem map used for European CarbonTracker-CH₄.



+ ice) state-vector elements each week. However, some TransCom regions contain fewer than five ecosystems types, and for ICE, emissions were assumed to equal zero for both prior and posterior. Therefore, the actual number of scaling factors $\lambda = (\lambda_{\text{bio}}, \lambda_{\text{anth}})$ to be optimized was 49 per week globally.

Results

Comparison with Pallas continuous observations

The mismatch between simulated CH₄ and observations indicates that using the prior fluxes generally results in overestimations of CH₄

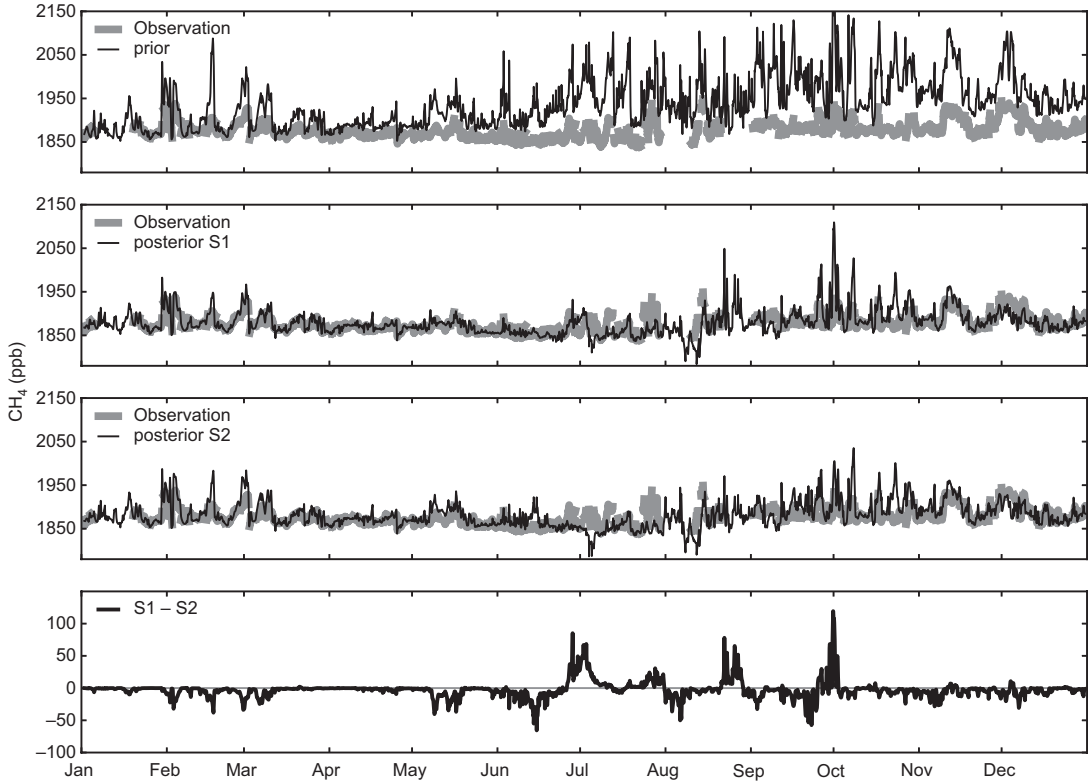


Fig. 5. Measured and modeled atmospheric methane for 2007 at Pallas. Thick gray lines are FMI continuous measurements, and thin back lines show modeled mole fractions using prior (top), weekly posterior with (S1) and without (S2) assimilating Pallas NOAA measurements from discrete air samples. The bottom panel shows differences between the S1 and S2 posterior fractions.

abundance at Pallas (Fig. 5). The prior matches the observations fairly well in the beginning of the year, but the baseline increases faster than in the observations, reaching ca. 20 ppb higher values at the end of the year. Peaks in the prior during summer and autumn were much higher than in the observations, pointing to prior CH_4 fluxes that are too large.

As expected, the hourly S1 posterior concentrations matched the FMI hourly continuous observations better than the prior concentrations (Fig. 5). The unrealistically strong seasonal cycle in the prior was improved in the posterior. As a result of the lower posterior wetland emissions, the peaks in the posterior concentrations were much lower than those in the prior, especially during summer and autumn, matching the observed concentrations better throughout the year. The large increase in the baseline of the prior concentrations in summer and autumn

was attributed to an overestimation in biosphere emissions. Wetland fractions for the region may be overestimated in the prior emission calculation (Prigent *et al.* 2007). There was a considerable mismatch between posterior concentrations and the observations around the end of September, when the posterior concentrations were much higher than the observed ones. Although better than the prior, maximum differences in the posterior concentrations were still up to 200 ppb. Note that there was often only one observation per site per week assimilated during the inversion, and during some weeks, observations from Pallas and nearby sites such as Teriberka and Zeppelinfjellet, were rejected from the assimilation (Fig. 6). The number of rejected observations was greatest in autumn, including the period when the maximum differences were estimated. For the rejected observations, the modeled concentrations were 45.0 ppb, or

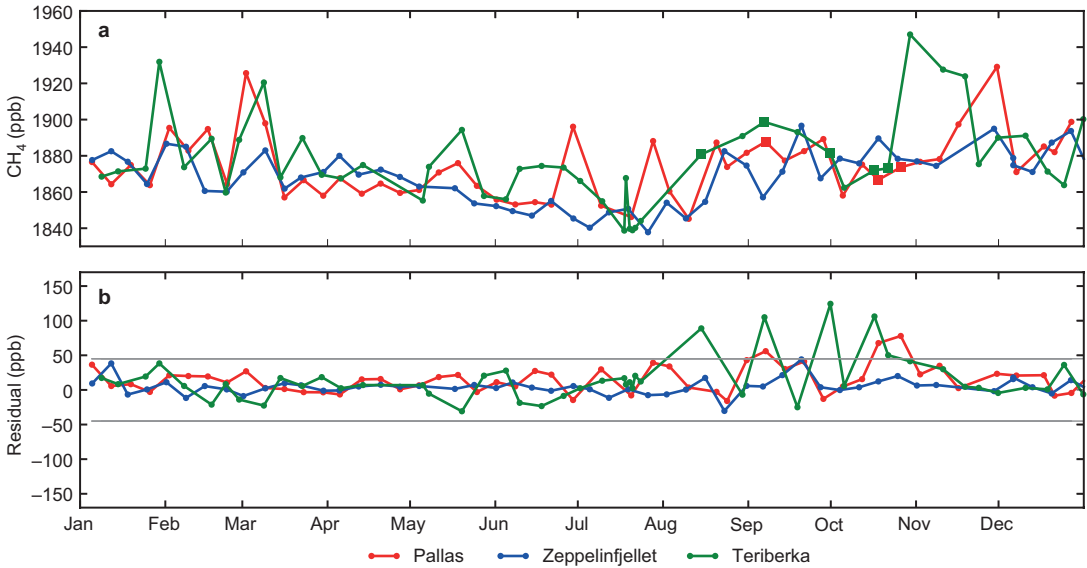


Fig. 6. (a) Time series of NOAA and MGO measurements: dots are assimilated measurements, and squares are rejected. (b) Differences between prior and measured for each measurements: gray lines are the rejection thresholds of -45 and 45 ppb for all three sites. Measurements were rejected when the difference exceeded the rejection threshold.

more, greater than the observed concentrations, probably resulted from too high prior emission estimates. The highest concentration in the observations (1947 ppb) at Teriberka in November was unintentionally added to the assimilation data set. This observation should have been removed during the preprocessing because it was rejected, according to the flag assigned by the data provider, as not being representative of background conditions.

The S2 posterior concentrations also generally better match the observations than the prior. Compared with the S1 posterior, the baseline of the two were similar during winter and spring, but S2 baseline was lower than that of S1 in summer, and slightly higher in autumn (Fig. 5).

The annual mean of the residuals between the observations and prior concentrations again showed that the prior estimates were generally higher than the observations; a histogram of the residuals was positively skewed, and the root mean square error (RMSD) was about 1.5 times greater than the standard deviation (Fig. 7). The residuals using the S1 posterior mole fractions were closer to normal distribution (the left and right tails of the histogram are almost equally long), the mean of the residuals was much closer

to zero than that of the prior, and the differences between the standard deviation and the RMSD was only about 0.2 ppb. This confirms that the S1 posterior matched the observations better than the prior. Comparing the residuals of the S1 and S2 posterior, the mean residual was about 4 ppb greater in S2, the S2 standard deviation was a little greater, and RMSD was not as close to its standard deviation as that in S1. Thus, the S1 residuals at Pallas more closely resemble a normal distribution as compared with the S2 residuals, and the S1 mole fractions matched the observations better. Some outliers were seen in the S1 residuals, which mainly resulted from the high concentrations estimated in autumn.

Next, two test runs with TM5 were performed to assess the effect of the anthropogenic and biosphere emissions on the methane concentration at Pallas. TM5 was run using the S1 posterior emissions with either the European anthropogenic emissions or the European biosphere emissions artificially set to zero (Fig. 8). Here, Europe was defined as an aggregate of four regions: northeast, northwest, southeast and southwest Europe. The posterior concentrations (Fig. 8a) indicate that the baseline was lower than the observations throughout the year

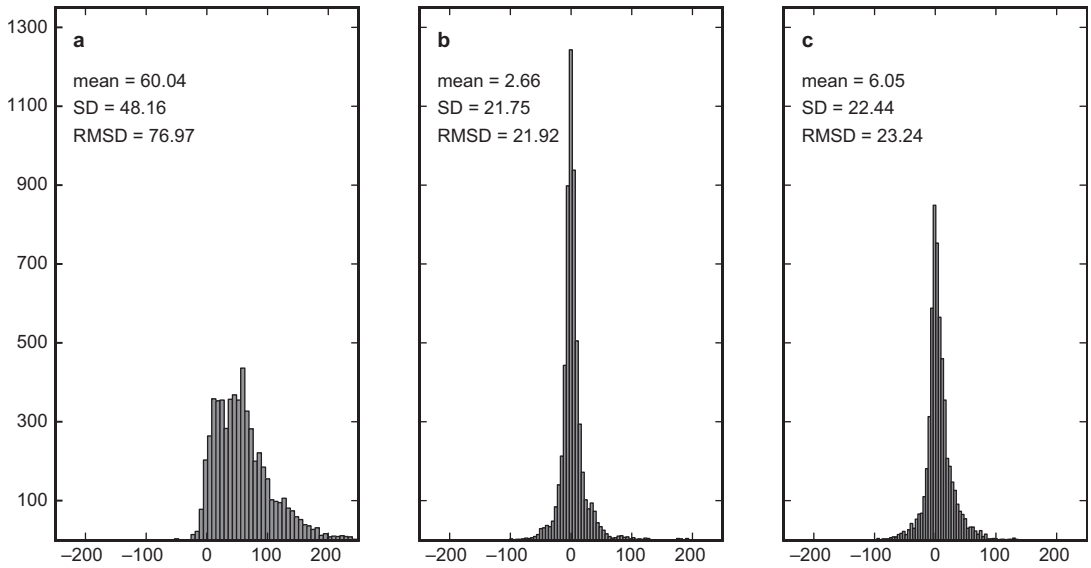


Fig. 7. Histograms and statistics of the residuals between FMI continuous observations and TM5-modeled mole fractions using (a) prior, and weekly posterior emissions optimized (b) with and (c) without NOAA Pallas observations assimilated for 2007. Note the prior emissions were monthly. The root-mean-square deviation (RMSD) = $[\sum(\text{estimate} - \text{measurement})^2/n]^{1/2}$. Units of the x-axis and the statistics are parts per billion (ppb).

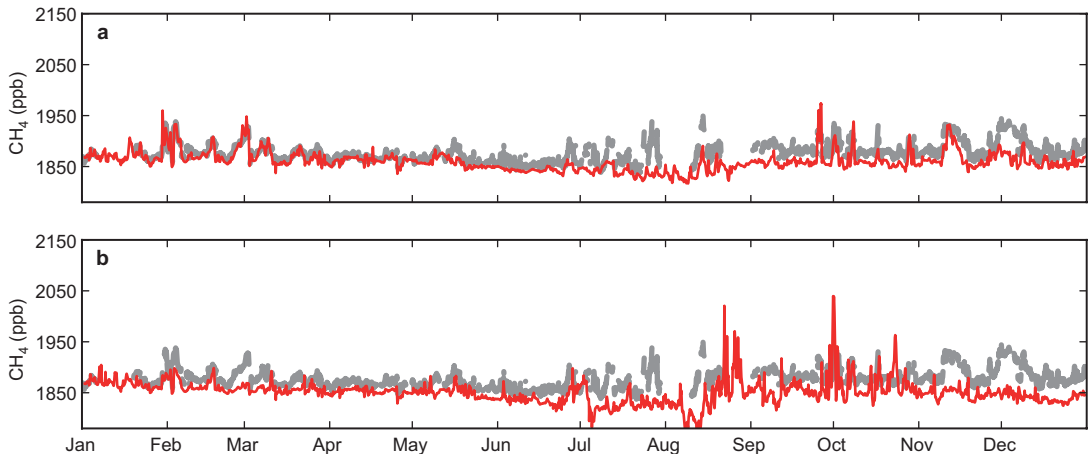


Fig. 8. Measured and modeled atmospheric methane for 2007 at Pallas. Thick grey lines are FMI continuous observations, and thin red lines show modeled mole fractions using weekly posterior emissions, in which the European (a) anthropogenic and (b) biosphere emissions were artificially set to zero.

when European anthropogenic emissions were excluded. Without European anthropogenic emissions, simulated CH_4 failed to capture most of the peaks during winter and spring, but the summer and autumn peaks were well captured. Some peaks in late autumn and small peaks during winter and spring were seen episodically at Pallas, which may be due to long-range trans-

port from Russia (Siberia). The posterior mole fractions (Fig. 8b) indicate that the baseline follows the observations well in the beginning of the year and throughout the spring, but it was too low during the rest of the year. Winter and spring peaks were well captured (see Fig. 8b), but only few of summer and autumn peaks were generated. This shows that winter and spring concen-

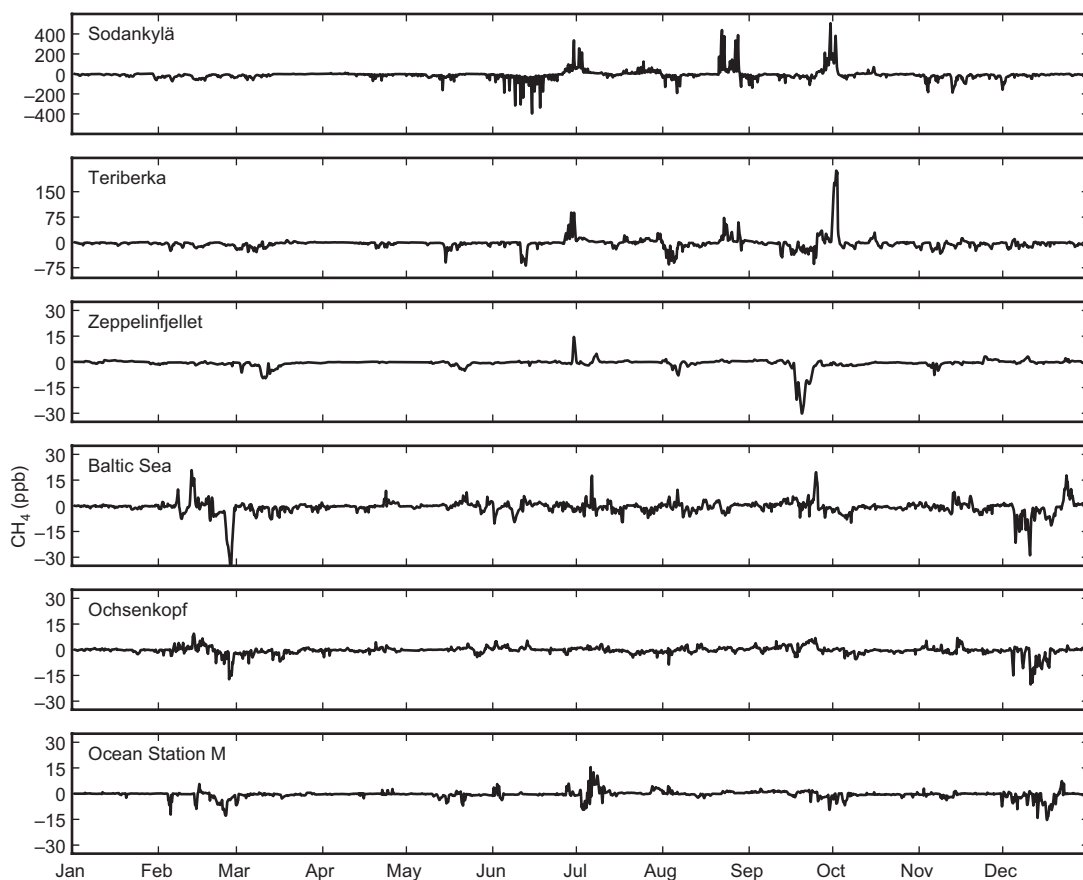


Fig. 9. Differences in estimated hourly mole fractions for 2007 using posterior emissions estimated with and without assimilating NOAA Pallas observations. The hourly mole fractions were estimated using a forward model (TM5). Note the differences in the y-axis scales.

trations at Pallas contain mainly anthropogenic signals, whereas summer and autumn concentrations are affected by the biosphere emissions.

The effect of the Pallas observations was also seen at nearby sites, such as Sodankylä and Teriberka. The differences between the S1 and S2 mole fractions at those sites showed similar features that were found for Pallas (Fig. 9). The differences were small during winter and spring, but large in summer and autumn; the differences were positive during mid-June–July and September–early October, but negative in the beginning of August and between mid-October and mid-November. Although MGO discrete air sample observations were assimilated for Teriberka, many late summer and autumn observations were rejected (Fig. 6). Therefore, the effect of Pallas observations was greater as compared

with that of other sites, such as Zeppelinfjellet, where most of NOAA discrete air sample observations were assimilated throughout the year. For other nearby sites, such as Baltic Sea, Ochsenkopf, and Ocean Station M, the differences between S1 and S2 posterior mole fractions were small. In the beginning of December, both anthropogenic and biosphere S1 emissions in northwest Europe were ca. $0.05 \text{ Tg CH}_4 \text{ week}^{-1}$ lower than those of S1, which resulted in lower S1 mole fractions at Baltic Sea, Ochsenkopf, and Ocean Station M. The differences were small as compared with those in biosphere emissions in northeast Europe, which differed as much as $0.16 \text{ Tg CH}_4 \text{ week}^{-1}$ during summer. This suggests that Pallas has only small effect on northwest European emissions also in winter.

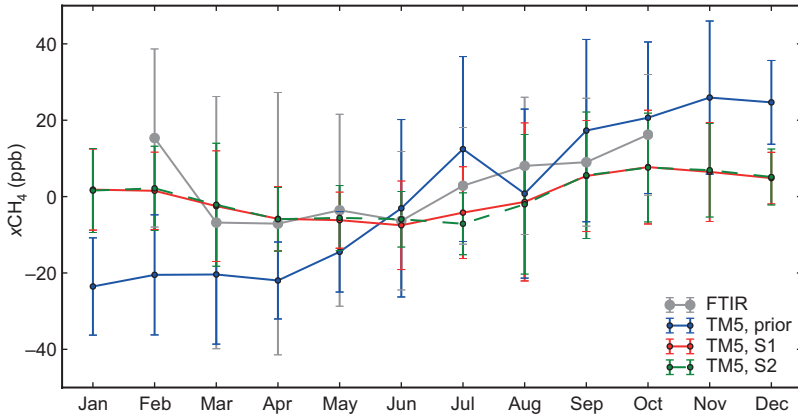


Fig. 10. Anomalies of monthly XCH₄ (column averaged mixing ratios) at Sodankylä. Filled circles are monthly means, and the vertical bars are the standard deviations. The FTIR monthly detrended seasonal cycle was calculated from all data from 2009–2012. Coloured lines are TM5 estimates for 2007, using monthly prior and weekly posterior emission fields. The posterior emissions were optimized assimilating all global flask measurements (S1), and all but the NOAA Pallas observations (S2). The nearest observations to Sodankylä are from Pallas.

The seasonal cycle in Sodankylä column averaged mixing ratio

Here, we compare Sodankylä FTIR observations with modeled column averaged mixing ratios of methane (XCH₄). The Sodankylä XCH₄ data were not assimilated in the model. Detrended monthly averages of the XCH₄ observations were calculated using all data of the period 2009–2012, and the model data from 2007. Note that the observations and the model data do not overlap.

The monthly column-averaged mixing ratios for 2007 were estimated with TM5 using the prior and posterior emissions for the $1^\circ \times 1^\circ$ grid box containing Sodankylä. Monthly statistics were calculated from daily 3D atmospheric concentration fields, and the annual mean was subtracted to compare the shapes of the seasonal cycle with that of Sodankylä FTIR measurements (Fig. 10). The mean of all the measurements was subtracted to calculate the anomalies of the measurements. The observations showed a XCH₄ decrease in spring, and an increase from the summer towards winter. The prior estimates failed to capture the shape of the seasonal cycle of the observations. The prior showed ca. 20 ppb XCH₄ increase from the beginning to the end of the year, whereas the observations showed no such increase. The shape of posterior estimates matched the observations better; the means and

standard deviations of the anomalies were within the two standard deviations of the anomalies of the measurements throughout the year.

The differences between the S1 and S2 estimates were very small. The S1 means were slightly lower in June and higher in July than the S2 means. This illustrates that the Pallas observations had no large effect on the column averages in the inversion, which is expected as the column integrates the effect of air masses that are influenced by emissions over large areas.

Emission estimates for Europe

As compared with the prior estimates, posterior estimates of annual biosphere and anthropogenic emissions for Europe showed reductions (Table 2). The magnitude of the CH₄ emission reduction was greater for the biosphere CH₄ emissions than for anthropogenic CH₄ emissions, whose estimates were reduced from 19.10 to 11.64 (± 8.76) Tg y⁻¹ in S1 and 12.64 (± 10.40) Tg y⁻¹ in S2. The greatest change was seen between June and October for the biosphere CH₄ emissions (Fig. 11), which was also illustrated in the comparison of atmospheric mole fractions at Pallas. The estimated anthropogenic CH₄ emissions for Europe were reduced from 44.67 to 35.88 (± 5.57) Tg y⁻¹ in S1 and 35.62 (± 5.62)

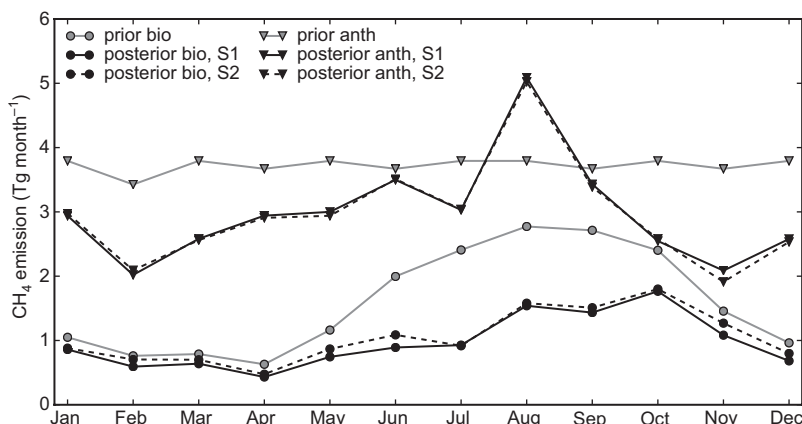


Fig. 11. Monthly biosphere and anthropogenic emissions of CH_4 in Europe estimated for 2007. Black lines show posterior emissions with (S1) and without (S2) weekly NOAA Pallas observations assimilated in the model. Prior emissions for biosphere were obtained from the LPJ-WHyMe vegetation model, and the EDGARv4.2 inventory was used for prior anthropogenic emissions. The seasonal variation in the prior anthropogenic emissions comes from the differences in the number of days per month.

Tg y^{-1} in S2 posterior emissions. Although the prior anthropogenic emissions were constant over the year, some seasonal variability was introduced in the posterior estimates (Fig. 11).

The regional emission estimates in the S1 and S2 runs were quite similar (Table 2). The S1 posterior estimate for the biosphere emissions was $1.0 \text{ Tg CH}_4 \text{ y}^{-1}$ lower than the S2 estimate. For anthropogenic emissions, the S1 estimate was $0.3 \text{ Tg CH}_4 \text{ y}^{-1}$ higher than the S2 estimate. As expected, the largest differences in the two posterior biosphere emissions as well as in the anthropogenic emissions were found

for northeastern Europe, where Pallas is located. However, these differences were not statistically significant (t -test: $n = 180, p > 0.05$). The differences in the monthly mean estimates between the S1 and S2 European biosphere emissions were also small. One of the reasons was, for example, the S1 emission estimate in July was ca. 10% smaller in northeast of Europe than the S2 estimates, but it was about the same percentage greater in northwest Europe, resulting in only small differences between the S1 and S2 estimates when estimates were aggregated over Europe (figure not shown).

Table 2. Estimated annual CH_4 emissions \pm uncertainties (Tg y^{-1}) for six TransCom regions and Europe for 2007. The prior estimate of anthropogenic emissions was from the EDGAR v4.2 inventory, and that of the biosphere was from the LPJ-WHyMe vegetation model. Two posterior emission estimates are given, with Pallas observations included (S1) and without (S2). Due to the definition of land ecosystem map, biosphere emissions for southwestern and southeastern Europe were not optimized, therefore uncertainties are not given. The last row is the sum of the four European regions.

TransCom region	Biosphere			Anthropogenic		
	Prior	Posterior S1	Posterior S2	Prior	Posterior S1	Posterior S2
North American boreal	10.16	9.18 ± 7.15	8.89 ± 7.15	0.47	0.45 ± 2.60	0.46 ± 2.60
Eurasian boreal	15.77	14.93 ± 10.65	15.25 ± 10.65	9.19	9.08 ± 4.48	8.98 ± 4.49
Southwestern Europe	1.63	1.63	1.63	10.98	8.09 ± 0.80	8.05 ± 0.8
Southeastern Europe	0.86	0.86	0.86	8.12	7.14 ± 0.73	7.15 ± 0.74
Northwestern Europe	4.83	4.08 ± 2.81	4.40 ± 2.96	9.36	5.46 ± 1.45	5.54 ± 1.47
Northeastern Europe	11.78	5.07 ± 5.94	5.75 ± 7.45	16.21	15.19 ± 2.59	14.88 ± 2.61
Europe	19.10	11.64 ± 8.76	12.64 ± 10.40	44.67	35.88 ± 5.57	35.62 ± 5.62

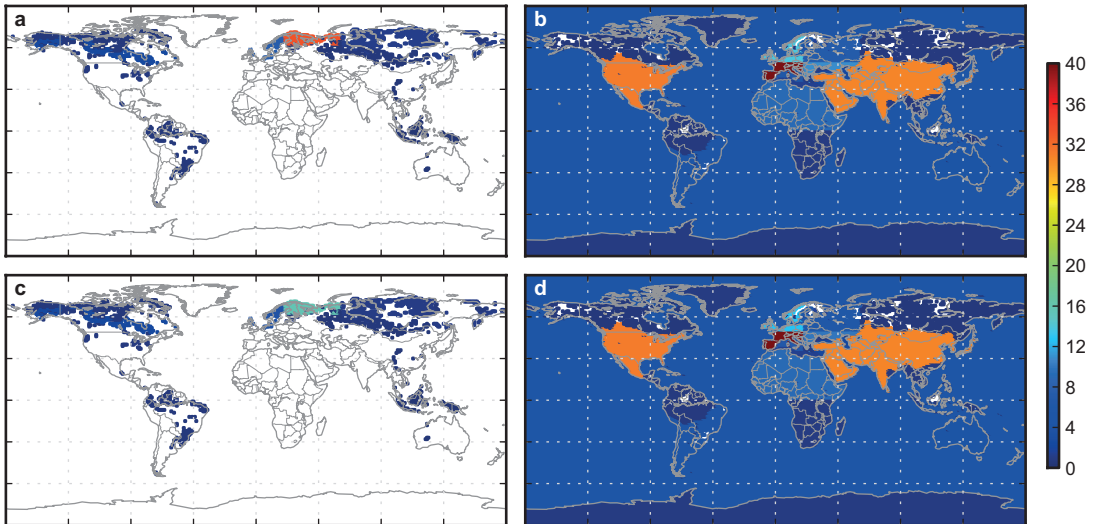


Fig. 12. Annual mean uncertainty reduction rates (%): 1 – posterior/prior. (a and b) Discrete air sample observations, and (c and d) observations except those at NOAA Pallas. (a and c) Biosphere emissions, and (b and d) anthropogenic emissions.

At high northern latitudes (North American boreal, Eurasian boreal and Europe), our S1 posterior estimates for 2007 was $81.16 (\pm 39.22)$ Tg $\text{CH}_4 \text{ y}^{-1}$, which is within the estimated range of Bruhwiler *et al.* (2014). However, our biosphere emission estimates were more than 10 Tg $\text{CH}_4 \text{ y}^{-1}$ greater than their estimates, leaving a smaller contribution from the anthropogenic emissions. The biosphere and anthropogenic emissions at high northern latitudes in our S1 estimates were $35.75 (\pm 26.56)$ and $45.42 (\pm 12.65)$ Tg $\text{CH}_4 \text{ y}^{-1}$, respectively.

Uncertainties

The annual mean uncertainty reduction ($1 - \text{posterior/prior}$) was calculated for each region (Fig. 12). Here, uncertainty refers to the standard deviation of the ensembles. As expected, the uncertainties were reduced by the inversion. The maximum reduction rates for biosphere and anthropogenic emissions were in northeast Europe (ca. 33%) and in southwest Europe (ca. 44%), respectively. Reduction rates were generally greater where the prior uncertainties were large, and enough observations were available. For example, the reduction rate was high in Asia, where the prior anthropogenic uncertainties were

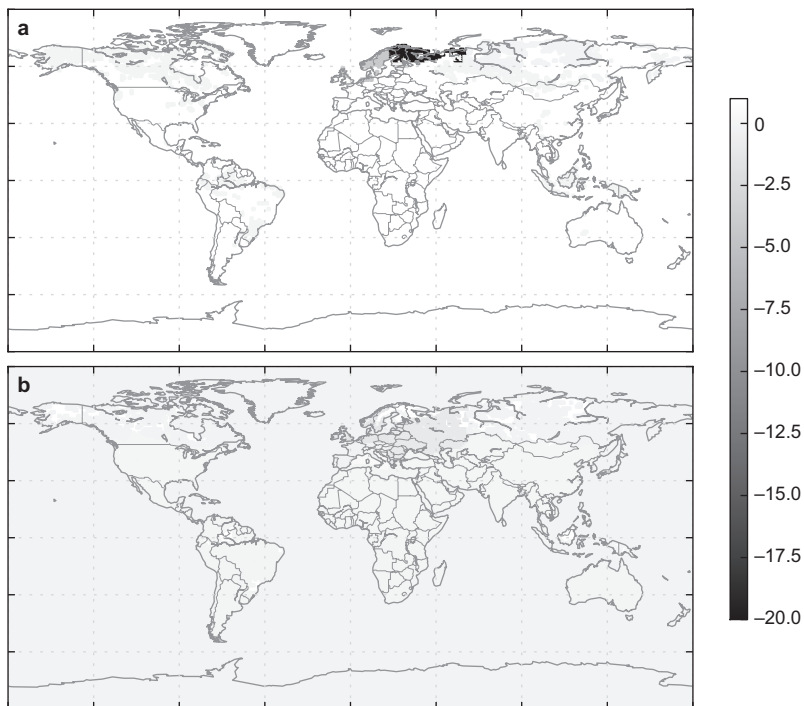
large (figure not shown), but many observations were assimilated. On the other hand, the reduction rate was small in Russian boreal, where the prior uncertainty was large, but no observations were available.

Comparison of the posterior uncertainties of the S1 and S2 emission estimates showed that they were ca. 20% smaller in S1 biosphere estimates for northeastern Europe (Fig. 13). The differences in the other regions were between -0.07% and 0.01% for the biosphere emissions and -1.2% and 0.4% for the anthropogenic emissions.

Discussion

The RMSDs between atmospheric mole fractions in the model and observations at Pallas were less than half in the posterior than in the prior estimates. The residuals between the S1 and S2 estimates showed that the effects of the Pallas observations on the inversion was substantial in summer and autumn, but low in winter and spring. Tests where European biosphere or anthropogenic emissions were set to zero showed that winter and spring concentrations were mostly affected by anthropogenic emissions, and summer and autumn concentrations

Fig. 13. Relative differences (%) in the posterior annual mean uncertainties are shown for (a) biosphere and (b) anthropogenic emissions between two runs, one with (S1) and another without (S2) NOAA Pallas observations assimilated ($1 - S1/S2$). The negative colours show that the posterior uncertainties were smaller in the S1 estimates.



were mainly affected by biosphere emissions. Since Pallas is located in a region of higher summertime biosphere emissions, this confirms the findings of Bruhwiler *et al.* (2014) that the Pallas observations constrain mostly signals from biosphere emissions, and little from the anthropogenic emissions.

The posterior uncertainties and the error reduction in the S1 and S2 estimates differed most for the biosphere emissions in northeast Europe, where Pallas is located. For other biosphere regions and anthropogenic regions, the posterior uncertainties and the reduction rates were almost equal. Thus, the effect of Pallas discrete air sample observations was large in the region where Pallas is located, but small elsewhere. This highlights the ability of the Pallas observations to reduce the uncertainties of biosphere emissions estimates in the region, but they have little influence on the anthropogenic emissions.

The effect of Pallas observations on the column averaged mixing ratios of methane was not as clearly seen as for the surface mixing ratios. The differences between shapes of seasonal cycles of S1 and S2 Sodankylä XCH₄

estimates were small, although some differences were seen in June and July. There was a decrease in the mean S1 XCH₄ from May to June, and an increase from June to July, but the mean S2 XCH₄ remained stable from May to June, and decreased from June to July. The monthly changes of S1 match the observations better during the period. The differences between the observations and S1 and S2 estimates were within the estimates by other studies, such as Saito *et al.* (2012), who estimated a spring model bias of 23.6 ppb. The spring overestimation of the modeled XCH₄ in this study may be due to the TM5 model bias in the stratosphere (Alexe *et al.* 2014, Bergamaschi *et al.* 2013).

As continuous observations bring a wealth of additional information to constrain the inversion, and may substantially improve the accuracy of optimised emission estimations, we will in our next study use the continuous observations that are available globally.

Furthermore, another useful assessment would be to carry similar tests for other sites. This study suggests that removing Teriberka observations would have little impact on the biosphere emissions, as Pallas observations con-

strain biosphere emissions in northeast Europe well. Also, the differences between S1 and S2 mole fractions at Teriberka suggest that it has little influence on winter anthropogenic emissions. Removing all observations from Baltic Sea, Ochsenkopf, and Ocean Station M would influence anthropogenic emission estimates for northern Europe, especially during summer, due to their location, and because Pallas has little influence on summertime anthropogenic emissions. However, this study is limited for assessing the effects of the Baltic Sea, Ochsenkopf, and Ocean and Zeppelinfjellet observations on surface fluxes in the inversion.

Conclusions

In this study, we explored the ability of European CarbonTracker-CH₄ build with the TM5 chemistry-transport model as an observation operator, to estimate CH₄ emissions in Europe and boreal region. We specifically looked at the role of the Pallas Station observations by analysing the simulated CH₄ time series with and without assimilating NOAA Pallas discrete air sample observations. The present analysis shows that European CarbonTracker-CH₄ is able to estimate the emissions in northern Finland well. Using optimized emissions, simulated Pallas surface CH₄ mole fractions agree well with independent observations. The simulation without Pallas discrete air sample observations underestimate posterior mole fractions during summer and early autumn. Emission estimates in Europe show that the influence of Pallas discrete air sample observations on uncertainty reduction is larger in European biosphere emissions than in anthropogenic emissions. The influences in other boreal regions were small. Those indicate that Pallas observations mostly constrain biosphere emissions in the European boreal region. This shows that a dense observation network that constrains different emission sources is important for further development of emission estimates using inverse models. Model performance at other sites and the influence of other observations such as continuous measurements, are an extension of this work to be considered.

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