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Estimating regional fluxes of CO₂ and CH₄ using space-borne observations of XCH₄ : XCO₂

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Abstract

We use the GEOS-Chem global 3-D atmospheric chemistry transport model to interpret XCH₄ : XCO₂ column ratios retrieved using a proxy method from the Japanese Greenhouse gases Observing SATellite (GOSAT). The advantage of these data over CO₂ and CH₄ columns retrieved independently using a full physics optimal estimation algorithm is that they suffer less from scattering-related regional bias. We show the model is able to reproduce observed global and regional spatial (mean bias = 0.7 %) and temporal variations (global $r^2 = 0.92$) of this ratio with model bias < 2.5 %. We also show these variations are driven by emissions of CO₂ and CH₄ that are typically six months out of phase which may reduce the sensitivity of the ratio to changes in either gas. To simultaneously estimate fluxes of CO₂ and CH₄ we use a formal Bayesian inverse model infrastructure. We use two approaches to independently resolve flux estimates of these two gases using GOSAT observations of XCH₄ : XCO₂: (1) the a priori error covariance between CO₂ and CH₄ describing common source from biomass burning; and (2) also fitting independent surface atmospheric measurements of CH₄ and CO₂ mole fraction that provide additional constraints, improving the effectiveness of the observed GOSAT ratio to constrain fluxes. We demonstrate the impact of these two approaches using Observing System Simulation Experiments. A posteriori flux estimates inferred using only the GOSAT ratios and taking advantage of the error covariance due to biomass burning are not consistent with the true fluxes in our experiments, as the inversion system cannot judge which species' fluxes to adjust. This can result in a posteriori fluxes that are further from the truth than the a priori fluxes. We find that adding the surface data to the inversion dramatically improves the ability of the GOSAT ratios to infer both CH₄ and CO₂ fluxes. We show that using real GOSAT XCH₄ : XCO₂ ratios together with the surface data during 2010 outcompetes inversions using the individual XCH₄ or the full-physics XCO₂ data products. Regional fluxes that show the greatest improvements have model minus observation differences with a large seasonal cycle such as Tropical South America for which we report a small but significant

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annual source of CO₂ compared to a small annual sink inferred from the XCO₂ data. Based on our analysis we argue that using the ratios we may be reaching the limitations on the precision of these data.

1 Introduction

5 Space-borne atmospheric column measurements of CO₂ and CH₄ have the potential to improve our quantitative understanding of their surface fluxes and to underpin the development of testable climate policies. For these data to address these potential applications the column measurements have to meet strict precision requirements, reflecting small signals from surface fluxes (a few percent of the column amount) compared
10 to the variations due to atmospheric transport. Any uncharacterized systematic error in these measurements compromises the ability of these data to infer surface fluxes. The CO₂ inverse problem is particularly sensitive to these systematic errors acting on length scales 10³–10⁴ km, in between the spatial scales of numerical models and those observed by the sparse network of well characterized upward-looking Fourier transform spectrometers, regional aircraft, and the network of ground-based measurements.
15 Here, we develop a method to infer simultaneous regional CO₂ and CH₄ flux estimates (Fig. 1) from XCH₄ : XCO₂ ratios, retrieved from the Japanese Greenhouse gases Observing SATellite (GOSAT) using the proxy approach (based on University of Leicester proxy XCH₄ v4.0), which are less prone to systematic error from aerosols
20 (Schepers et al., 2012).

Two methods have been used to retrieve CO₂ and CH₄ columns from calibrated GOSAT L1B spectra: the “full physics” and the “proxy” methods (Cogan et al., 2012; Parker et al., 2011). The full physics method uses an optimal estimation approach and incorporates a rigorous treatment of the atmospheric radiative transfer including the
25 effects of clouds and aerosols. This method uses optimized spectral windows to fit CO₂ and CH₄. The main advantage of this approach is the error characterization of the a posteriori state vector, and the main disadvantage is having to accurately character-

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ize the atmospheric aerosol for the radiative transfer calculation. The proxy method, used to infer CH₄ columns, fits both gases in nearby spectral windows with the assumption that any fitting artefacts common to both gases (e.g. aerosol and clouds) will be removed by taking the ratio of the two gases. This method is simpler than the
5 full physics approach and more robust against scattering and as a result many more retrievals are possible from the GOSAT spectra. We also believe that these measurements are less compromised by systematic errors on regional scales. Interpretation of this ratio has in the past relied on scaling it with a model CO₂ column so that any erroneous model information about CO₂ can influence the interpretation of the GOSAT
10 CH₄ columns (e.g. Parker et al., 2011; Fraser et al., 2013). We propose a method to simultaneously optimize CH₄ and CO₂ fluxes using the retrieved XCH₄ : XCO₂ ratio. This eliminates the need for a CO₂ model, removing the impact of model uncertainty on the retrieved methane columns, and increases the number of observations available to constrain CO₂ fluxes (Fig. 2).

15 In the following section we describe the space-borne and ground-based data used in our experiments. In Sect. 3 we describe the GEOS-Chem chemical transport model, and the data assimilation scheme developed for this work. In Sect. 4 we report the GOSAT and model spatial and temporal distributions of XCH₄ : XCO₂ ratios (Sect. 4.1), we test the assimilation scheme using a series of Observing System Simulation Experiments (OSSEs, Sect. 4.2) and present inversion results (Sect. 4.3). We conclude the
20 paper in Sect. 5.

2 Data

2.1 GOSAT CO₂ and CH₄ atmospheric column mole fraction measurements

GOSAT was launched in 2009 by the Japanese Space Agency in a sun-synchronous
25 orbit with an equatorial local overpass time of 13:00 LT, providing global coverage every three days (Kuze et al., 2009). GOSAT includes two instruments: TANSO-FTS (Thermal

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inversion largely reflect the larger volume of $XCH_4 : XCO_2$ ratio data resulting in better spatial and temporal coverage (Fig. 2). We may also expect the largest differences where we believe there are the greatest biases in the other single gas retrievals. We find that the associated error reductions for the CO_2 fluxes inferred from the $XCH_4 : XCO_2$ ratio data are typically larger than those for CH_4 , and are different from those inferred from the EnKF inversion.

5 Concluding remarks

We have interpreted measurements of $XCH_4 : XCO_2$ from GOSAT in which XCH_4 and XCO_2 are retrieved in nearby spectral windows under the assumption that their ratio will largely remove common sources of biases. By interpreting the ratio directly we minimize any bias introduced by model XCO_2 ; although we acknowledge other sources of model bias remain. A major advantage of the ratio is this data product does not suffer from the measurement bias that befalls the full physics XCO_2 data. Another advantage is that the volume of these data is greater than their full physics counterpart. While the ratio benefits from these three advantages the difference between model and observed quantities are much smaller (typically $< \pm 2\%$) than either XCO_2 or XCH_4 becoming comparable in magnitude to other sources of error, e.g. model transport error, that cannot easily be characterized and removed; by using the ratio we may be reaching the limitations on the precision of these data and our ability to interpret them using current-day transport models. However, over particular geographical regions we find there are seasonally varying GOSAT minus model ratio differences that are large enough to be exploited, e.g., Tropical South America and Tropical Asia.

Using a series of numerical experiments we showed that the simultaneous estimation of CO_2 and CH_4 fluxes using the GOSAT ratio is possible with the information split as a function of the a priori uncertainties, however the inversion system returns unphysical fluxes in some regions. We showed that including surface mole fraction measurements of CO_2 and CH_4 in the measurement vector provides an “anchor” for the inversion, and

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we showed that the combined GOSAT and surface data can distinguish between CO_2 and CH_4 fluxes.

Using real data for 2010 we showed that the combination of the GOSAT $XCH_4 : XCO_2$ ratio and the surface mole fraction data outcompeted inversions using the individual XCH_4 and XCO_2 GOSAT data and corresponding surface data. We found the greatest differences between the two approaches over the regions where we found that the GOSAT minus model difference had a seasonal variation that was larger than a few percent. For instance, we found that Tropical South America was a small but significant source of CO_2 while analysis of full physics XCO_2 showed a small sink term. Analysis of the ratio led to slightly larger reductions globally, and in some regions, primarily in the tropics, much larger reductions in uncertainty of CO_2 and CH_4 .

The main reason for using the $XCH_4 : XCO_2$ ratio is that it minimizes scattering and potentially other biases and significantly increases geographical coverage. Although CO_2 and CH_4 do not share many common sources that result in significant atmospheric covariance we have shown that: (1) the combined information from these two gases can be disentangled using other data, and (2) the result is an improvement over what can be achieved using observations of either full-physics XCO_2 or XCH_4 . Consequently, the use of space-borne observations of the $XCH_4 : XCO_2$ ratio will be of particular interest for estimating CO_2 surface fluxes over regions that are characterized by frequent cloud cover and high aerosol loading such as the tropics where the quality and coverage of full-physics XCO_2 retrieval approaches will be limited even for missions with spatial footprints smaller than GOSAT. This ratio approach could also be used in combination with other atmospheric tracers that help improve the source attribution of CO_2 , e.g. carbon monoxide, where the ensuing correlation is driven by incomplete combustion (Palmer et al., 2006). Space-borne mission concept development related to the carbon cycle should not only focus on the primary compound but also on any secondary compound that will help interpret the observed variability of that primary gas.

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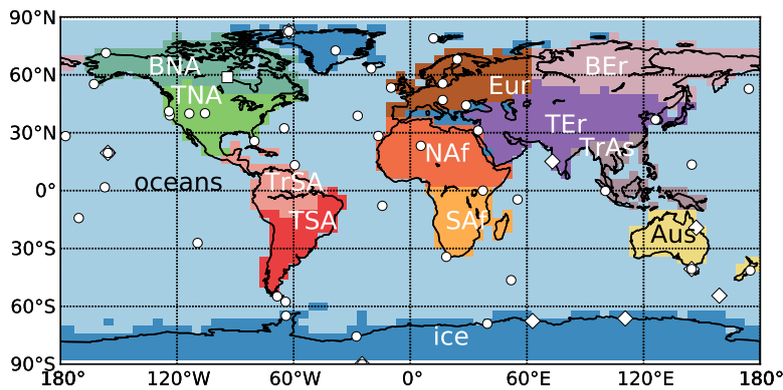


Figure 1. Distribution of the 13 geographical regions for which we estimate CO₂ and CH₄ fluxes, and the location of 57 co-operative flask sampling sites with data covering the study period, January–December 2010. The land regions, informed by previous work (Gurney et al., 2002) include: Boreal North America (BNA), Temperate North America (TNA), Tropical South America (TrSA), Temperate South America (TSA), Northern Africa (NAf), Southern Africa (SAf), Boreal Eurasia (BEr), Temperate Eurasia (TEr), Tropical Asia (TrAs), Australasia (Aus), and Europe (Eur). The ground-based measurement sites run by NOAA ESRL, CSIRO GASLAB, and EC are denoted by white circles, white diamonds, and white squares, respectively.

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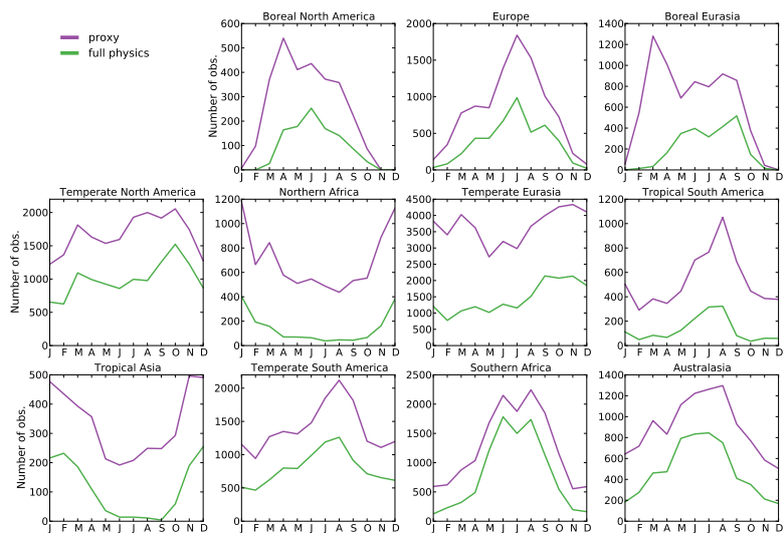


Figure 2. The number of GOSAT observations available per month during 2010 over specific geographical regions (Fig. 1) from the full-physics and proxy retrieval algorithms.

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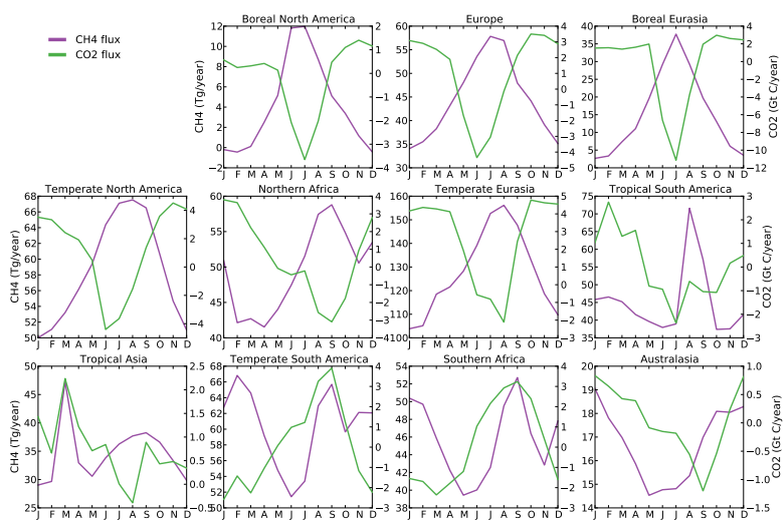


Figure 3. Monthly a priori emissions for CH₄ (Tg CH₄ year⁻¹) and CO₂ (Gt C year⁻¹) for the land regions shown in Fig. 1. Note the different y-scales.

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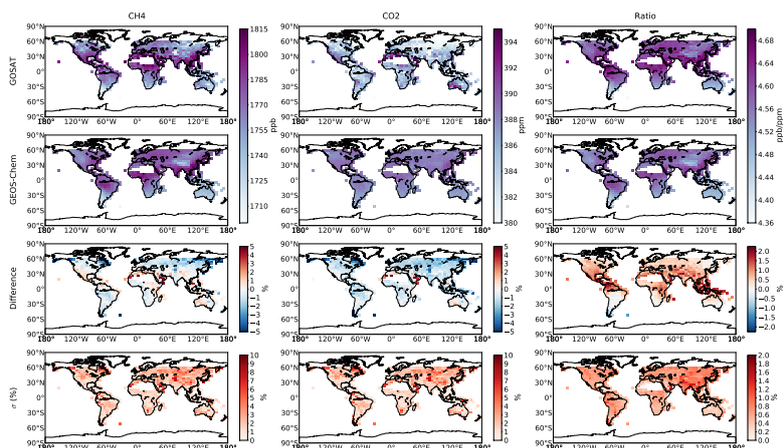


Figure 4. Annual mean GOSAT (top row) and GEOS-Chem model (second row) XCH₄, XCO₂, and XCH₄ : XCO₂ ratio measurements from GOSAT during 2010 averaged on the model 4° × 5° grid. The third row shows the percentage difference between them (GOSAT minus GEOS-Chem). For XCH₄ and XCO₂, we truncate at the mean ± 2-σ. The bottom row shows the 1-σ value in the difference as a percentage about the mean GOSAT XCH₄, XCO₂, and XCH₄ : XCO₂ data. The model has been sampled at the time and location of the GOSAT observations, and convolved with scene-dependent averaging kernels.

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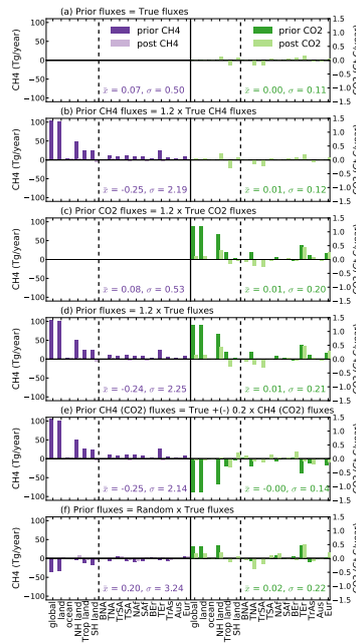


Figure 7. As Fig. 6 but all experiments use CH₄ and CO₂ surface flask data and GOSAT data. Experiment (a) for which a priori fluxes are equal to the true fluxes; experiment (b) for which CH₄ a priori fluxes are 20% larger than the true fluxes; experiment (c) for which CO₂ a priori fluxes are 20% larger than the true fluxes; experiment (d) for which CH₄ and CO₂ a priori fluxes are 20% larger than their true fluxes; experiment (e) for which CH₄ a priori fluxes are 20% larger and CO₂ a priori fluxes are 20% smaller than their true fluxes; and experiment (f) for which all a priori fluxes are perturbed stochastically, ranging from -20% to 20%, from the true fluxes.

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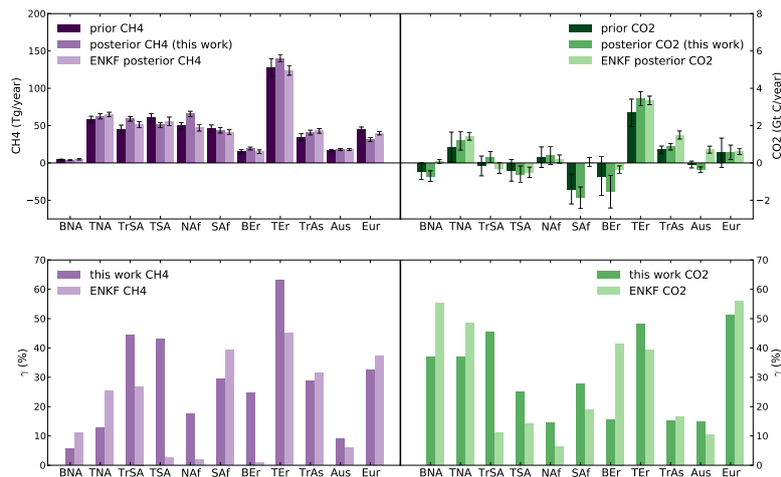


Figure 8. A priori and a posteriori CO₂ and CH₄ regional land fluxes inferred from GOSAT XCH₄ : XCO₂ and surface measurements of CO₂ and CH₄ and from XCO₂ or XCH₄ using an ensemble Kalman filter (top) (Feng et al., 2011; Fraser et al., 2013), and the corresponding reduction in uncertainty (bottom), during 2010. Error bars atop of emission estimates represents the 1- σ uncertainty.

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