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Key Points:

- A new divergence method is developed to estimate methane emissions based on satellite observations, requiring no a priori emissions
- The applicability of this method in identifying and quantifying sources is proven by a GEOS-Chem simulation with known emission inventory
- The estimated emissions over Texas (United States) based on TROPOspheric Monitoring Instrument observations are evaluated and are found to be robust

Supporting Information:

Supporting Information may be found in the online version of this article.

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A New Divergence Method to Quantify Methane Emissions Using Observations of Sentinel-5P TROPOMI

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Abstract We present a new divergence method to estimate methane (CH₄) emissions from satellite observed mean mixing ratio of methane (XCH₄) by deriving the regional enhancement of XCH₄ in the Planetary Boundary Layer (PBL). The applicability is proven by comparing the estimated emissions with its known emission inventory from a 3-month GEOS-Chem simulation. When applied to TROPOspheric Monitoring Instrument observations, sources from well-known oil/gas production areas, livestock farms and wetlands in Texas become clearly visible in the emission maps. The calculated yearly averaged total CH₄ emission over the Permian Basin is 3.06 (2.82, 3.78) Tg a⁻¹ for 2019, which is consistent with previous studies and double that of EDGAR v4.3.2 for 2012. Sensitivity tests on PBL heights, on the derived regional background and on wind speeds suggest our divergence method is quite robust. It is also a fast and simple method to estimate the CH₄ emissions globally.

Plain Language Summary Methane (CH₄) is an important greenhouse gas in the atmosphere and plays a crucial role in the global climate change. It kept increasing over the last decades. About 70% of CH₄ comes from human activities like oil/gas productions or livestock farms. The recently launched TROPOspheric Monitoring Instrument provides an opportunity to estimate the emissions of CH₄ on a regional scale. This work presents a new method to fastly derive CH₄ emissions at a fairly high spatial resolution without a priori knowledge of sources.

1. Introduction

Methane (CH₄) is the second most important anthropogenic greenhouse gas after carbon dioxide (CO₂) and is also a principal precursor of tropospheric ozone (Shindell et al., 2012). In-situ measurements show a continuous increase of methane over the last decades (Dlugokencky et al., 2009; IPCC, 2013; Saunio et al., 2016; Turner et al., 2019), with stable concentrations from 2000 to 2006 (Dlugokencky et al., 2009; Rigby et al., 2008). CH₄ has both natural (e.g., wetlands, wildfires, termites) and anthropogenic (e.g., fossil fuels, livestock, landfills, and wastewater treatments) sources. About 360 million tons (60% of the total CH₄) are released through human activities (Saunio et al., 2020). The relatively short lifetime of CH₄ (about a decade) makes it a short-term target for mitigating climate change by reducing the emissions.

Satellite observations of CH₄ provide an efficient way to analyze its variations and emissions at a regional to global scale (Buchwitz et al., 2017; Lunt et al., 2019; Maasakkers et al., 2019; Miller et al., 2019; Zhang et al., 2020). Compared to previous widely used instruments like Greenhouse gases Observing SATellite (GOSAT) and Scanning Imaging Absorption spectrometer for Atmospheric Cartography (SCIAMACHY, onboard Envisat), the TROPOspheric Monitoring Instrument (TROPOMI) on board the Sentinel 5 Precursor (S5-P) satellite measures CH₄ at an unprecedented resolution of 7 × 7 km² since its launch in October 2017 (upgraded to 5.5 × 7 km² in August 2019) (Veeffkind et al., 2012). Several studies have shown the capability of TROPOMI on identifying and quantifying the sources at a local to regional scale (e.g. (de Gouw et al., 2020; Pandey et al., 2019; Schneider et al., 2020; Zhang et al., 2020)). These studies mainly focused on oil/gas leakage events, which show strong signals that can be easily identified, or they are using an inverse modeling relying on an a priori emission inventory.

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Freshly emitted air pollutants are usually concentrated around the emission source, in the case of not too high wind speeds (Liu et al., 2020). Beirle et al. (2019) found that the strong gradients near sources of nitrogen oxides (NO_x) are preserved by averaging horizontal fluxes. Therefore, the divergence of horizontal fluxes of nitrogen dioxide (NO_2) plus a sink term can be used to estimate the emissions of NO_2 . In our study, we apply a divergence method for deriving CH_4 emissions, which has never been attempted before for long-living gases because of the complications with the strong background concentrations. For the short-living gases like NO_2 , the background concentrations are very low that are less affected by transport and orography. The sink term can be ignored for CH_4 because of its relatively long lifetime, which makes it more straightforward to link the divergence to the emission. The divergence works on the product of horizontal fluxes and wind fields, which is independent of a priori emission inventories and models and can be applied at various resolutions regionally or globally.

The retrieved CH_4 from satellite observations are the ratios of methane total vertical columns to air density columns (XCH_4), which are strongly affected by the stratospheric abundance. Thus the influence of transport in the upper atmosphere and of orography should be removed to better distinguish gradients due to emissions. XCH_4 measured by satellites reflects the abundance of the background plus the newly emitted methane because of its around 10-year lifetime. Hence the contribution from the background should be deducted when estimating the emissions.

In this study, we present a new divergence method to quantify the emission of CH_4 from satellite retrieved XCH_4 . The XCH_4 of TROPOMI is first destriped and corrected with albedos at short-wave infrared (SWIR) wavelengths (2305–2385 nm) to improve the data quality. Before applying the method to TROPOMI observations, a 3-month (from July 2012 to September 2012) hourly GEOS-Chem nested model simulation over North America is used to test the applicability of our method. The robustness and uncertainty of the resulting emissions is further analyzed with sensitivity studies and comparisons to the literature.

2. Method and Data

Figure 1 shows the flowchart of the procedure to estimate the CH_4 emissions from TROPOMI retrieved XCH_4 . It consists of three main steps. First, applying posteriori corrections on XCH_4 to reduce the systematic biases caused by across-track biases and surface albedos. Second, the mean mixing ratios of CH_4 in the PBL ($\text{XCH}_4^{\text{PBL}}$) and the corresponding regional “backgrounds” are derived by subtracting the columns above the PBL, which are estimated by XCH_4 profiles from the Atmospheric Composition Reanalysis 4 (EAC4) of the Copernicus Atmosphere Monitoring Service (CAM5) (Inness et al., 2019). The enhancements of $\text{XCH}_4^{\text{PBL}}$ are further used to calculate the spatial divergence and estimate CH_4 emissions.

2.1. Estimate Methane Emission From TROPOMI

There are two additive corrections, the stripe correction and the albedo correction, on XCH_4 to remove biases caused by the satellite retrieval. The detailed method can be found in Part A and B of Supporting Information S1.

The continuity equation connecting the divergence (D), emission (E) and sink (S) for steady state is: $D = E + S$ (Beirle et al., 2019). As the lifetime of CH_4 is around 10 years, D in the PBL actually contains the variations of its background and sources. As D is a linear operator, the daily D_d of the fluxes in the PBL can be written as:

$$D_d = D_d^B + D_d^S \quad (1)$$

where D_d^B is the daily divergence of the background flux and D_d^S is the daily divergence caused by sources, respectively. The sink term can be ignored, and assuming the background concentrations are completely homogeneous, that is: $D_d^S = E_d$. However, in most cases, the real background is inhomogeneous because (a) the surface height in two adjacent grid cells can be very different or (b) a different bias in observations caused by an albedo difference in two adjacent pixels. Thus, $D_d - D_d^B = E_d$.

The divergence D works on horizontal fluxes (F): $D = \nabla F$, where F stands for zonal (F_u) and meridional fluxes (F_v), which is the product of gridded vertical columns (V) and horizontal wind fields (\vec{w}). For each day d :

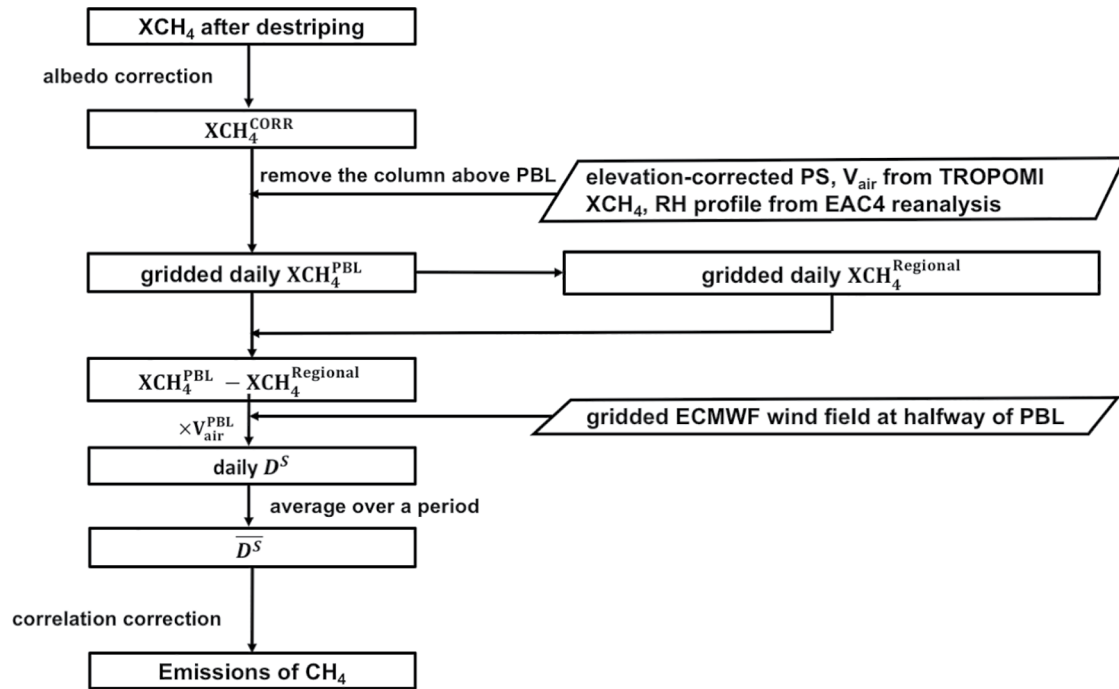


Figure 1. The flow chart of using TROPospheric Monitoring Instrument XCH_4 to derive the CH_4 emissions over a certain period. PS and V_{air} stand for the surface pressure and the total column of air density used in TROPOMI XCH_4 retrieval. RH is the relative humidity.

$$E_d = \nabla F_d = \nabla (V \cdot \vec{w}) \quad (2)$$

Numerical derivatives for D are calculated as the second-order central difference in this study, the detailed procedure can be found in Part C of Supporting Information S1. We convert XCH_4 to mean mixing ratio in the PBL, XCH_4^{PBL} (denoted by X^{PBL}), to eliminate the effects of orography and transport in upper atmosphere. The column of methane in the PBL (V^{PBL}) for day d is derived by:

$$V_d^{PBL} = X_d^{PBL} \times A_d^{PBL} \quad (3)$$

where A_d^{PBL} is the corresponding air density column in the PBL. Combining with Equations 2 and 3, Equation 1 can be written as:

$$D_d^S = \nabla \left((X_d^{PBL} - X_d^B) \right) \times A_d^{PBL} \cdot \vec{w} \quad (4)$$

where X_d^B is the background of X_d^{PBL} . It is hard to know the exact X_d^B , so we use the regional background (X_d^R) to approximate the X_d^B as will be stated in Section 2.2. Equation 4 is then written as:

$$D_d^S = \nabla \left((X_d^{PBL} - X_d^R) \times A_d^{PBL} \cdot \vec{w} \right) \quad (5)$$

Equation 5 is applied to the daily variations of CH_4 , and the emission is estimated by averaging D_d^S over a time period:

$$E_d = \overline{D_d^S} = \overline{D_d^S - D_d^R} \quad (6)$$

where D_d^R stands for the averaged divergence of the regional background. However, we found a significant correlation between $\overline{D_d^S}$ and $\overline{D_d^R}$ at some locations, which suggest that the derived emissions still contain part of the background. Strong spatial positive correlations R are typically found over areas with complicated terrain where the background is less homogenous.

The remaining background divergence is caused by local changes in the wind-fields induced by orography. Hence the mechanisms of emissions and of the regional background are independent leading to our assumption that they are uncorrelated. Therefore, if the correlation is close to 1, it is clear we have a false emission signal in $\overline{D_d^S}$ and this emission will be removed. If the correlation is zero (or negative), no

correction is needed. For positive correlations we use a first-order correction using the correlation value as regression coefficient, thus multiplying $\overline{D^S}$ by $(1-R)$.

In addition, we find that areas with negative emissions E also have negative $\overline{D^R}$ and divergence of winds ($\overline{D^W}$), implying no significant sources. Thus, the grids with negative E are set to be zero in the final estimated emissions. The practice of this posteriori correction is presented in Section 3.

2.2. Calculating the Regional Enhancement of Methane in PBL

The entire atmospheric column was divided into only 12 layers in the TROPOMI XCH_4 retrieval, which is too coarse to resolve the vertical distribution. To estimate the methane column above the PBL we use model results of EAC4 of CAMS (<https://ads.atmosphere.copernicus.eu/cdsapp#!/dataset/cams-global-reanalysis-eac4?tab=overview>). It is a global hourly reanalysis of atmospheric composition at a relative high spatial resolution, 0.75° horizontally and 60 layers vertically (Inness et al., 2019), which contains no a priori CH_4 emissions. Thus, the spatial distribution of CH_4 is solely the result of transport and orography, which will be subtracted from TROPOMI observations to estimate the PBL concentration of CH_4 . Considering the height of the planetary boundary layer (PBLH) from reanalysis or forecast data set has large uncertainties and is occasionally too shallow, we fixed the PBLH at 500 meters above the ground. XCH_4 in PBL (XCH_4^{PBL}) is derived as follow:

$$XCH_4^{PBL} = \frac{V_{CH_4} - V_{CH_4}^U}{V_{air}^{PBL}} \quad (7)$$

V_{CH_4} and V_{air}^{PBL} stand for the total column of methane and dry air density used by the retrieval of TROPOMI XCH_4 , respectively. $V_{CH_4}^U$, the vertical column of methane above the PBL, is estimated by CAMS model, in which the total dry air column is constrained by that of TROPOMI. Thus, XCH_4^{PBL} is biased because of the system difference between TROPOMI and CAMS. The XCH_4^{PBL} of each pixel is then used to build the daily gridded data at a resolution of 0.25° . In this study, for each grid, daily regional background of XCH_4^{PBL} (XCH_4^R), is defined as the average of the lower 10 percentile of its surrounding ± 5 grid cells ($11 \times 11 = 121$ grid cells in total by taking the current grid cell as the center). The difference between XCH_4^{PBL} and XCH_4^R (Equation 5) is finally used to calculate the divergence with wind speeds. Therefore, the system biases between CAMS and TROPOMI is implicitly and greatly reduced by subtracting XCH_4^R from XCH_4^{PBL} because their bias origins are the same.

The surface pressure of each pixel is adjusted by a high-resolution GMTED2010 Digital Elevation map (Hasekamp et al., 2019), and the pressure at each layer of the EAC4 XCH_4 profile is recalculated accordingly. The number of dry air molecules in the entire column of the XCH_4 profile is scaled to the total number that is used for the retrieval of the pixel. We do not interpolate the averaging kernel (AK) to the layers of EAC4, because the AK is approximately equal to 1.0 at each layer (Hasekamp et al., 2019). In this way, we ensure the conservation of air mass for each pixel as well as the high-resolution vertical distributions of methane.

The wind field halfway the PBLH close to the overpass time is obtained from the ECMWF. The divergence method works only when transport takes place, i.e., there is at least some wind. In addition, extremely high wind speeds are not favorable for the method that is based on the regional mass balance. Therefore, wind speeds are constrained between 1 m/s to 10 m/s in this study.

2.3. Using a GEOS-Chem Simulation to Test the Method

In order to evaluate the feasibility of our method, the case of a model simulated XCH_4 is suitable because of known a priori emissions. In this study, we perform a 3-month simulation starting from July 1, 2012 by the GEOS-Chem 12.5.0 (<http://geos-chem.org>) nested model over North America at a resolution of 0.5° lat. \times 0.625° lon. with 47 vertical layers extending to the mesosphere. The boundary conditions are provided by GEOS-Chem global simulation at 4° lat. \times 5° lon. using posterior methane emissions and OH levels inverted from GOSAT satellite observations (Lu et al., 2021), and therefore these boundary conditions are unbiased to GOSAT observations outside the domain. Both models are driven by MERRA-2 reanalysis meteorological fields from the NASA Global Modeling and Assimilation Office (GMAO) (Gelaro et al., 2017). The a

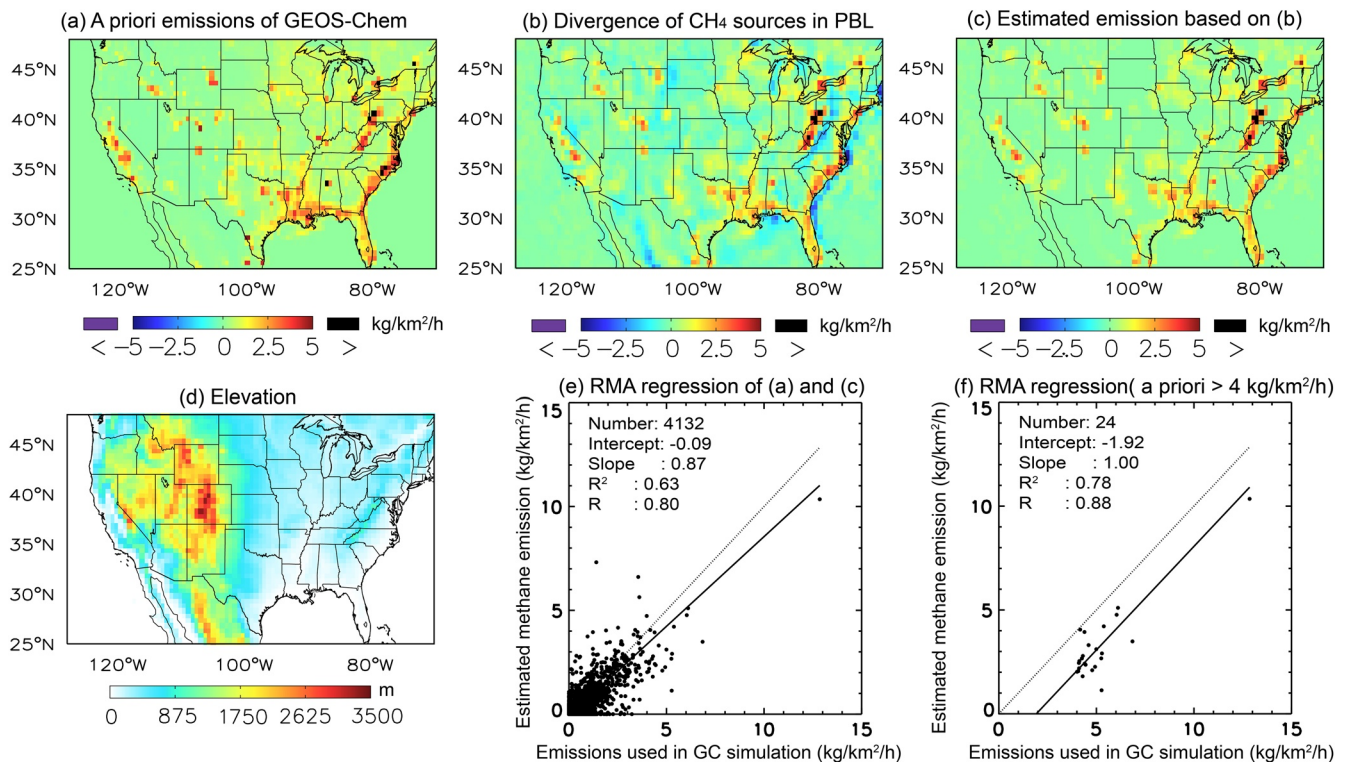


Figure 2. The spatial distributions of (a) the average of a priori CH₄ emissions used in GEOS-Chem simulation, (b) the divergence of CH₄ sources in Planetary Boundary Layer, and (c) corresponding estimated CH₄ emissions over June–August 2012 on a 0.625° lon. × 0.5° lat. grid. (d) The elevation map that is generated from GMTED2010 data set. (e) Scatter plots for emissions between a priori emissions higher than 0.0 kg/km²/h and estimated CH₄ emissions. (f) As (e) but for a priori emissions that are higher than 4.0 kg/km²/h. Each dot in (e) and (f) represents a grid cell.

priori natural emissions include wetlands, open fires, termites and seeps. The anthropogenic emissions are from EDGAR v4.3.2, with fugitive fuel emissions (oil, gas, coal) overwritten by the Scarpelli et al. (2020) inventory, and further superseded by the gridded version of Inventory of U.S. Greenhouse Gas Emissions and Sinks from the Environmental Protection Agency (EPA GHGI) over the US (Maasackers et al., 2016). More information on the model setup can be found in (Lu et al., 2021). Here we take the results at UTC 18:00, which is close to the overpass time of TROPOMI over the US. We apply our method to these simulations of XCH₄ in the PBL. The XCH₄^{PBL} is the mixing ratio of the column in PBL at the same time. The method to build regional background for each grid follows Section 2.2.

3. Results

3.1. Verification of the Method Using GEOS-Chem Simulations

Figures 2a–2c show the spatial distribution of the 3-month average of a priori emission inventory used in GEOS-Chem simulation, the divergence of XCH₄ enhancement in PBL and the estimated emission. Although the horizontal resolution of the model is much coarser than TROPOMI observations, the sources have been identified (Figures 2b and 2c), even for relatively small emissions less than 2.5 kg/km²/h. For the mountainous and coastal areas that are more complex than typical flat land terrain, the performance of the divergence works fairly well. Some fake signals caused by orography (e.g., in Mexico, convergence over oceans near the coastal) are successfully removed by the posteriori “correlation correction.” The influence from the remaining background is mostly found over the grid cells with R greater than 0.7.

We further quantitatively compare the estimated emissions with the a priori emission inventory. The grid cells with emissions >0 in the a priori inventory have been selected as the reference. The scatter plots in Figures 2d and 2e compare a priori emissions greater than zero and greater than 4 kg/km²/h with their counterparts respectively. Our estimated emissions capture the spatial variability in a priori emissions throughout

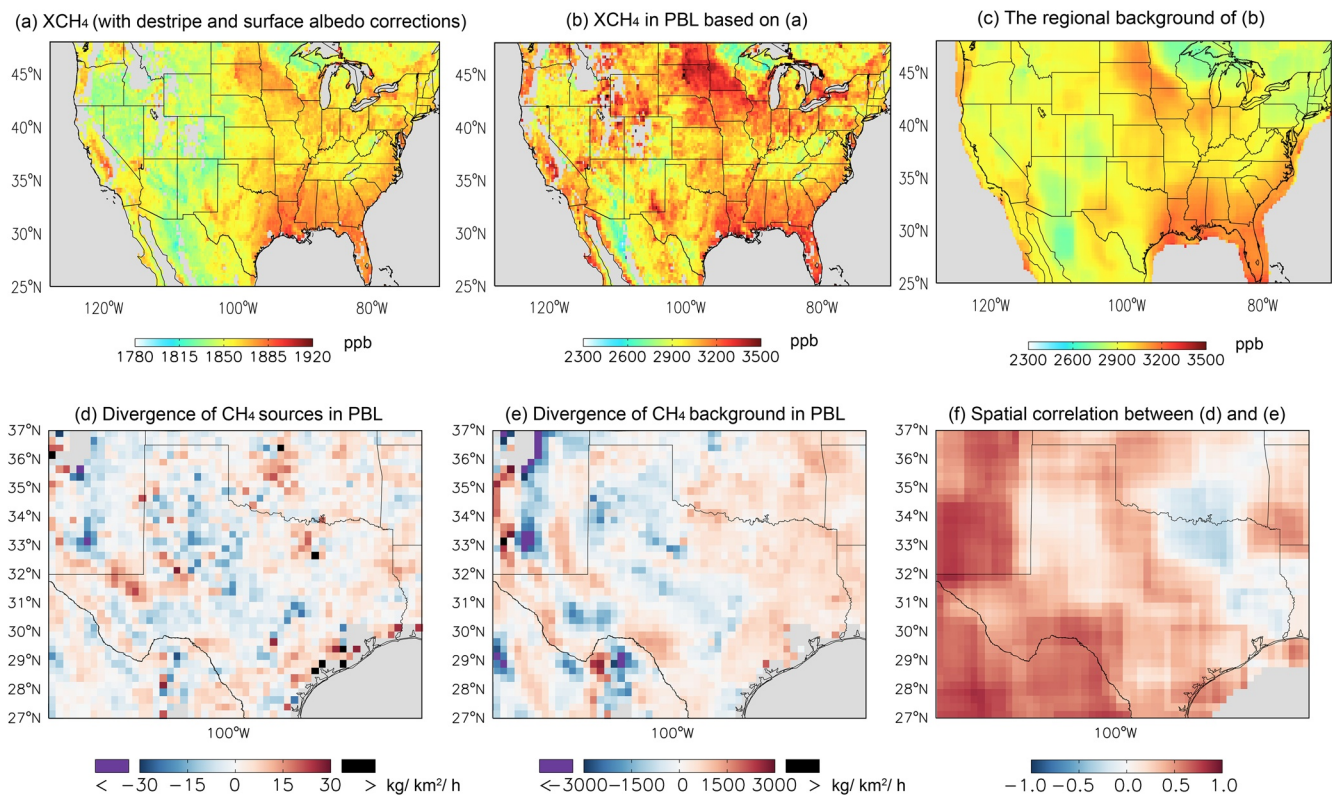


Figure 3. Spatial distributions of yearly averaged (a) XCH₄ with the stripe and surface albedo corrections, (b) the corresponding XCH₄ in Planetary Boundary Layer (PBL) and (c) its regional background. The divergences of (d) CH₄ sources in PBL and (e) of the regional background in 2019. (f) The spatial correlation between (d) and (e). For each grid cell, the correlation is calculated in a domain of 11 × 11 grid cells, taking the grid cell as center.

the full range of emissions ($R^2 = 0.63$). The Reduced Major Axis regression show a slope of 0.86 and an intercept of -0.08 , highly implying the capability of our method in retrieving model emissions using simulated columns. The biases are mainly related to the simplified regional background we used. The big sources (a priori emission greater than $4 \text{ kg/km}^2/\text{h}$) are much easier to capture by our method ($R^2 = 0.78$, $R = 0.88$). The final result shows the simple regional background removal is simplified but efficient.

We also test our method by using the enhancement in the troposphere instead of the PBL (Figure S5). The estimated emissions show a much weaker correlation with a priori emissions, especially over the areas with complicated orography. The transport in the upper troposphere are intervening with the emission estimates. Therefore, using the enhancement of XCH₄ in the PBL is more suitable to identify and quantify the emissions.

3.2. CH₄ Emissions Over the US Based on TROPOMI

Figure 3a presents the spatial distributions of TROPOMI yearly averaged XCH₄ after destripping and SWIR surface albedo corrections over North America on a 0.25° grid in 2019. After converting XCH₄ to XCH₄^{PBL}, the spatial distribution of CH₄ becomes more continuous over mountains in Figure 3b. Despite the uncertainty from surface albedo corrections (see more detailed discussion in Part B of Supporting Information S1), enhancements of CH₄ are found over Texas, California and Appalachia regions when comparing to the regional background (Figure 3c).

Figures 3d–3e show examples of the divergence of sources and of corresponding regional backgrounds in the PBL over the Texas area, one of the most prolific petroleum- and gas-producing regions in the U.S., and Figure 3f shows their spatial correlation. The areas with negative values (convergence) in Figure 3d are also negative in Figure 3e, demonstrating there are no significant sources. In addition, high positive spatial correlations mainly appear over the areas with complicated orography but few emissions. On the contrary, the

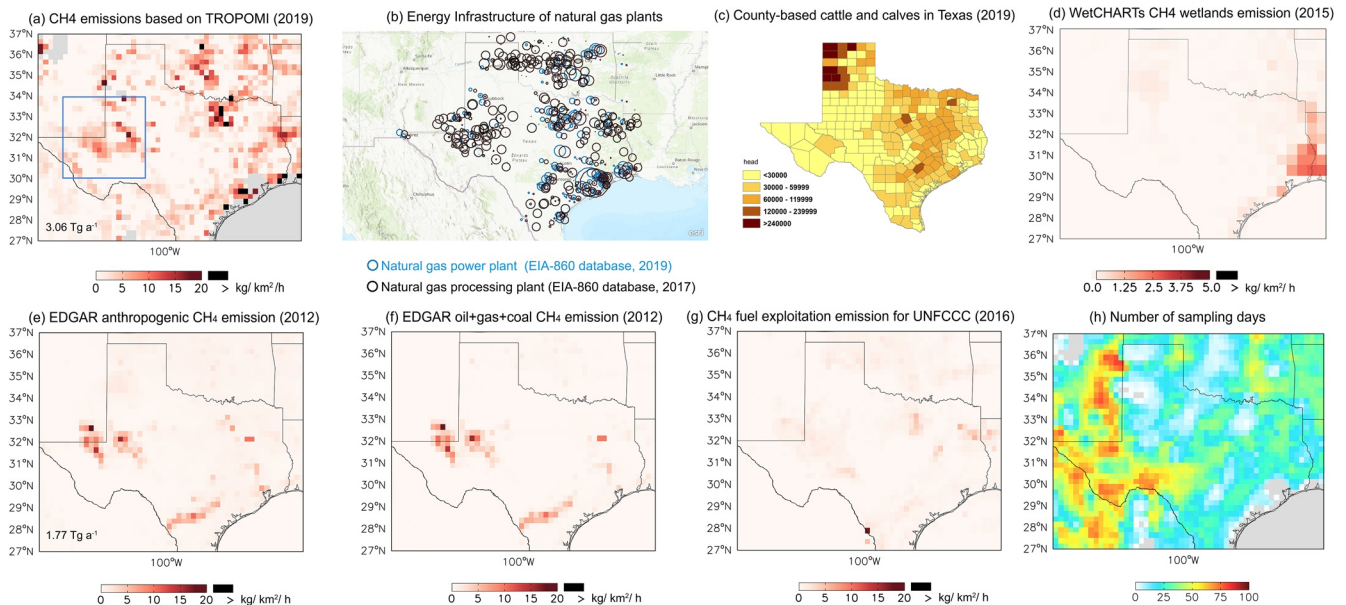


Figure 4. CH₄ emissions over the Texas area. (a) Our estimated emissions for 2019. (b) Natural gas power plants (blue circles) and processing plants (black circles) in Texas (available at: https://www.eia.gov/special/gulf_of_mexico/). The size of each circle represents the capacity of the plant. (c) County-based heads of cattle and calves in Texas in 2019 (available at: https://www.nass.usda.gov/Statistics_by_State/Texas/Publications/County_Estimates/ce_maps/ce_catt.php) (d) WeCHARTs wetland emissions for 2015 (Bloom et al., 2017), (e) EDGAR v4.3.2 anthropogenic CH₄ total emissions for 2012 (available at: https://edgar.jrc.ec.europa.eu/overview.php?vC2_GHG), (f) EDGAR v4.3.2 CH₄ oil + gas + coal emissions in 2012, and (g) a global inventory of methane emissions from oil, gas, and coal exploitation that spatially allocates the national emissions reported to the UNFCCC for 2016 (Scarpelli et al., 2020). (h) The number of observations used in the emission estimate. The area enclosed by the solid blue line is the Permian Basin (30°–34°N, 101°–105°W). The annual total emissions of CH₄ based on our estimates and EDGAR v4.3.2 over the Permian Basin are embedded in the left corner of (a) and (e).

areas with big sources have weak or negative spatial correlations between sources and regional backgrounds (Figure 3f). Here, we apply the “correlation correction” for grids with R greater than 0.0 to reduce the biases of the regional background we built.

Our method not only successfully identified the sources in abovementioned well-known oil/gas fields, but also shows the ability to capture the sources from other sectors such as livestock and wetlands. For example, the high CH₄ emissions north of the Permian Basin in Figure 4a are very likely coming from a large number of cattle farms there (Figure 4b). Dairy farms or feed yards in this region are typically open lot, and sources of CH₄ are enteric emissions from cattle and emissions of wastewater lagoons. The emission rate of cattle is estimated to be on average 0.211 kg/head/day (Todd et al., 2011). These biogenic emissions do not exist in oil/gas/coal emissions in Figures 4f and 4g but can be found as small contributions to EDGAR v4.3.2 total emissions (Figure 4e).

Figure 4h shows the number of observations used in the emission estimate. TROPOMI CH₄ retrievals are not available over water, which inevitably leads to uncertainties and limited number of observations near coasts, lakes and bays. However, the natural gas power/processing plants onshore Texas near western Gulf of Mexico (Figure 4b), which shows the energy infrastructures of U.S Energy Information Administration (EIA, 2021), are found near the locations of sources shown in Figure 4b. It implies that emissions relating to big sources like oil/gas productions in the coastal are caught by our divergence method. We should be careful about the explanation of the final emissions (Figure 4a) considering the number of sampling days (Figure 4h). Fewer samplings, especially less than about 10 days, might lead to large uncertainties. On the other hand, averaging results over a long period possibly smooth out temporary events (e.g., leakage).

We further quantify the annual average CH₄ emissions over the Permian Basin (enclosed by the solid blue boundary in Figure 4a). The number of samplings is fairly even in each season (Table S1), which is partly benefit from relatively flat orograph. Our estimated emissions in 2019 (see baseline settings in Table S2) is 3.06 Tg a⁻¹, which is 42% higher than EDGAR v4.3.2 total anthropogenic emissions in 2012 (1.77 Tg a⁻¹),

which can be due to an increase in oil production between 2012 and 2019. Zhang et al. (2020) estimated the total emission as $2.9 \pm 0.5 \text{ Tg a}^{-1}$ based on the S5P operational TROPOMI CH_4 product (Hasekamp et al., 2019; Landgraf et al., 2019) from May 2018 to March 2019 by using inverse modeling with a priori emissions. The average annual emissions for the time period 2018/2019 based on the TROPOMI/WFMD v1.2 (Schneising et al., 2019) product is reported as $3.18 \pm 1.13 \text{ Tg a}^{-1}$ by Schneising et al. (2020) using a mass balance method.

In addition to testing different surface albedo corrections (see Part B in Supporting Information S1), we designed several other sensitivity tests to discuss the uncertainties of our estimated emissions that are generated from assumptions on the PBLH, the regional background concentration and wind speeds. Table S2 shows the different results for each case and the baseline method, called REF, over the Texas area. The mean, median, maximum and minimum difference relative to REF in Texas are listed. The total emission of each case over the Permian Basin is also quantified (last column in Table S2). Figures S5–S7 are corresponding spatial distributions of estimated emissions and the difference with reference to the REF by using different assumptions of PBLH, the regional background and the wind speeds, respectively.

PBLHs varying from 300 to 1000 m were tested. The influence of the PBLH on the spatial pattern and the total amount of final emissions are small, especially for the cases below 1000 m. We also changed the size of the background region from surrounding 3 grid cells to 7 grid cells (in each direction), leading to a bias of at most -0.19 Tg a^{-1} for the total emissions of the Permian Basin. As expected, the smaller size of the regional background (e.g., Three grid cells) lead to a higher regional background over the areas with big sources. Thus, the estimated emissions are decreasing over the emissions clusters while the emissions around them often increase.

We tested various restrictions on the maximum and minimum wind speed (Figure S8). The influence of wind speed is more complicated. Unlike the tests of PBLH and regional background, different restrictions first affect the samplings of days. High wind speeds lead to large uncertainties over areas with complicated terrain. For example, large divergence values near the mountains close to the west of the Permian Basin, are not sufficiently removed with the “correlation correction” (Figure S8a). The smearing effect by high wind speeds lead to homogenous spatial distributions of XCH_4 in the PBL. The signals of sources are hard to be separated from the regional background. It also indicates that cases with high wind speeds are not handled well by our method, and are therefore excluded. In contrast, constraints on lowest wind speeds have smaller effects on final emissions (Figures S8e and S8f), because pollutants exhibit much stronger horizontal gradient in calm scenes. But the divergence method works only if transportation related to wind exists, so we set the minimum wind speed at 1 m/s.

4. Conclusions

A new divergence method has been successfully developed and applied to estimate CH_4 emissions over Texas in North America based on observations of the TROPOMI instrument. The method works fairly well to detect sources of all strengths, proven by using a GEOS-Chem model simulation as an ideal case. Applied to real TROPOMI observations it clearly identifies signals from oil/gas clusters and other sources, such as livestock and wetlands. Further quantification of annual averaged CH_4 emissions over the Permian Basin area is consistent with recent previous studies. The different spatial distributions of emissions in different inventories (ranging from 2012 to 2019) imply strong temporal variations of emissions in this area. The divergence method we built benefits from TROPOMI's high spatial resolution and provides a way to quickly estimate CH_4 emission from satellite observation. The method does not need use any a priori information on location of strength of the emissions.

Through the sensitivity tests on the PBLH, the regional background and the wind speeds, the uncertainties of estimated emissions could be reduced by constraining their values. High wind speeds cause high uncertainties over areas with complicated terrain. In future work the uncertainties caused by the winds will be reduced when longer records of background concentrations, EAC4 data set, are available. The higher spatial resolution of the estimated emissions is another aspect to be improved after the new S5P TROPOMI CH_4 data set will be released.

Conflict of Interest

The authors declare no conflicts of interest relevant to this study.

Data Availability Statement

S5P TROPOMI methane Level-2 data set is available at: <http://www.tropomi.eu/data-products/methane>. EAC4 of CAMS, which used to be estimated the column above the PBL can be accessed at: <https://ads.atmosphere.copernicus.eu/cdsapp#!/dataset/cams-global-reanalysis-eac4?tab=overview>. Natural gas power plants and processing plants in Texas are available at: https://www.eia.gov/special/gulf_of_mexico/. County-based heads of cattle and calves in Texas in 2019 is available at: https://www.nass.usda.gov/Statistics_by_State/Texas/Publications/County_Estimates/ce_maps/ce_catt.php. EDGAR v4.3.2 for the total anthropogenic emissions in 2012 is available at: https://edgar.jrc.ec.europa.eu/overview.php?v=432_GHG. WeCHARTs wetland emission in 2015 can be found at: https://daac.ornl.gov/cgi-bin/dsvviewer.pl?ds_id=1502. The GEOS-Chem simulation and its emissions can be downloaded at: https://d1qb6yzwaaq4he.cloudfront.net/data/geoschem_ch4_2012/GEOS-Chem.tar.

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