Atmos. Chem. Phys. Discuss., 15, 34871–34911, 2015 www.atmos-chem-phys-discuss.net/15/34871/2015/ doi:10.5194/acpd-15-34871-2015 © Author(s) 2015. CC Attribution 3.0 License.



This discussion paper is/has been under review for the journal Atmospheric Chemistry and Physics (ACP). Please refer to the corresponding final paper in ACP if available.

Investigating Alaskan methane and carbon dioxide fluxes using measurements from the CARVE tower

A. Karion^{1,2,a}, C. Sweeney^{1,2}, J. B. Miller^{1,2}, A. E. Andrews², R. Commane³, S. Dinardo⁴, J. M. Henderson⁵, J. Lindaas^{3,b}, J. C. Lin⁶, K. A. Luus^{7,c}, T. Newberger^{1,2}, P. Tans², S. C. Wofsy³, S. Wolter^{1,2}, and C. E. Miller⁴

¹University of Colorado, Boulder, CO, USA
²NOAA Earth System Research Laboratory, Global Monitoring Division, Boulder, CO, USA
³Harvard University, Cambridge, MA, USA
⁴Jet Propulsion Laboratory, Pasadena, CA, USA
⁵Atmospheric and Environmental Research, Lexington, MA, USA
⁶Atmospheric Sciences, University of Utah, Salt Lake City, UT, USA
⁷Biogeochemical Integration, Max Planck Institute for Biogeochemistry, Jena, Germany
^anow at: National Institute of Standards and Technology, Gaithersburg, MD, USA
^bnow at: Colorado State University, Fort Collins, CO, USA



Received: 5 November 2015 – Accepted: 23 November 2015 – Published: 11 December 2015

Correspondence to: A. Karion (anna.karion@nist.gov)

Published by Copernicus Publications on behalf of the European Geosciences Union.



Abstract

Northern high-latitude carbon sources and sinks, including those resulting from degrading permafrost, are thought to be sensitive to the rapidly warming climate. Because the near-surface atmosphere integrates surface fluxes over large (~ 500–1000 km) scales, atmospheric monitoring of carbon dioxide (CO₂) and methane (CH₄) mole fractions in the daytime mixed layer is a promising method for detecting change in the carbon cycle throughout boreal Alaska. Here we use CO₂ and CH₄ measurements from a NOAA tower 17 km north of Fairbanks AK, established as part of NASA's Carbon in Arctic Reservoirs Vulnerability Experiment (CARVE), to investigate regional fluxes of CO₂ and CH₄ for 2012–2014. CARVE was designed to use aircraft and surface observations to better understand and quantify the sensitivity of Alaskan carbon fluxes to climate variability. We use high-resolution meteorological fields from the Polar Weather Research and Forecasting (WRF) model coupled with the Stochastic Time-Inverted

- Lagrangian Transport model (hereafter, WRF-STILT), along with the Polar Vegetation Photosynthesis and Respiration Model (PolarVPRM), to investigate fluxes of CO₂ in boreal Alaska using the tower observations, which are sensitive to large areas of central Alaska. We show that simulated PolarVPRM/WRF-STILT CO₂ mole fractions show remarkably good agreement with tower observations, suggesting that the WRF-STILT model represents the meteorology of the region quite well, and that the PolarVPRM flux
- ²⁰ magnitudes and spatial distribution are consistent with CO₂ mole fractions observed at the CARVE tower. CO₂ signals at the tower are larger than predicted, with significant respiration occurring in the fall that is not captured by PolarVPRM. Using the WRF-STILT model, we find that average CH₄ fluxes in boreal Alaska are somewhat lower than flux estimates by Chang et al. (2014) over all of Alaska for May–September 2012;
- we also find emissions persist during some wintertime periods, augmenting those observed during the summer and fall. The presence of significant fall and winter CO₂ and CH₄ fluxes underscores the need for year-round in-situ observations to quantify changes in boreal Alaskan annual carbon balance.



1 Introduction

The carbon cycle of the high northern latitudes has been the subject of study and research for many decades (Olefeldt et al., 2013; Harriss et al., 1992; Oechel et al., 1993; McGuire et al., 2010), with scientists and policy makers more recently focused on in its impact on global climate. This focus is in part due to the fact that global warming has affected temperatures in the high northern latitudes more significantly than any other region (IPCC, 2013). Higher temperatures could lead to a positive feedback of increased emissions of CO₂ and CH₄ (McGuire et al., 2009; O'Connor et al., 2010; Hayes et al., 2014; Schuur et al., 2015), including a possibility of large emissions from thawing Arctic permafrost. However, the timing and magnitude of such a feed-10 back remain uncertain (Schuur et al., 2008, 2015, 2009), and analysis of CH_4 and CO₂ measurements from the Global Greenhouse Gas Reference Network (GGGRN; www.esrl.noaa.gov/gmd/ccgg) do not yet show signs of enhanced Arctic to mid-latitude gradients (Bruhwiler et al., 2014; CarbonTracker, 2013). Planned future studies of ecosystems and carbon cycling in Arctic and boreal regions are intended to monitor changes in climate and carbon fluxes (e.g. NASA's Arctic-Boreal Vulnerability Exper-

iment (ABoVE), above.nasa.gov; Next-Generation Ecosystem Experiments (NGEE)
 Arctic, ngee-arctic.ornl.gov). To this end, quantification of current carbon fluxes from the northern high latitudes, including Alaska, is a crucial piece of any effort to detect
 changes in the Arctic and boreal carbon cycle.

The Carbon in Arctic Reservoirs Vulnerability Experiment (CARVE) was a 5 year NASA Earth Ventures (EV-1) airborne science investigation to quantify atmospheric mole fractions and surface–atmosphere fluxes of CO_2 and CH_4 and correlate these with key surface-state variables for terrestrial ecosystems in Arctic and boreal Alaska.

²⁵ CARVE's goal is to bridge critical gaps in our knowledge and understanding of Arcticboreal ecosystems, linkages between the hydrologic and terrestrial carbon cycles, and the feedbacks from disturbances such as thawing permafrost and fires (Miller, 2015). The principal components of CARVE were the intensive aircraft campaigns conducted



monthly from March to November for four consecutive years (2012–2015). The aircraft payload included in-situ sensors measuring CO_2 , CH_4 , and carbon monoxide (CO) throughout the flights, which are based out of the Fairbanks airport and cover several regions throughout Alaska (Chang et al., 2014). A stationary tower-based GHG mea-

- ⁵ surement site, the CARVE tower (NOAA site code CRV), was established as part of the CARVE project, in order to give year-round context to the intensive aircraft observations. These continuous observations from a single location can verify the temporal pattern of carbon cycle models, while the aircraft observations provide information on spatial accuracy.
- Measurements of CO₂ and CH₄ from towers in northern high latitudes have previously been used to analyze emissions and trends in these regions (Sasakawa et al., 2010; Winderlich et al., 2010; Worthy et al., 2015). Concentration measurements from such towers generally have large regions of influence, on scales of hundreds of kilometers, in contrast to direct flux measurements from eddy covariance flux tower sites,
- ¹⁵ which may represent spatial scales closer to tens or hundreds of meters, or chamber measurements that typically represent even smaller (~ 1 m) scales. In this sense, the tall tower measurements are able to integrate fluxes that have been shown to be spatially heterogeneous (Olefeldt et al., 2013). Such concentration or mole fraction measurements require interpretation using a model framework to quantify terrestrial fluxes, because they do not measure them directly.

One way to infer and assess fluxes from mole fraction observations is to use a Lagrangian particle dispersion model (LPDM) coupled with a meteorological model to determine the influence function, or footprint, of a given observation (Lin et al., 2012). In this study, the WRF-STILT modeling framework has been used to generate footprints

for CARVE tower observations. Henderson et al. (2015) provide details of the model configuration and validation of the meteorological simulations. We assess CO_2 fluxes from the land surface of Alaska by convolving surface fluxes from the PolarVPRM (Luus and Lin, 2015) with the footprints and comparing the resulting modeled CO_2 enhancements with tower observations. To infer CH_4 fluxes, we have convolved the footprints



with a constant (in space and time) flux model and an elevation-based flux model and scaled the results to monthly mean observed enhancements to estimate monthly average fluxes over a wide region, using similar methods as Chang et al. (2014).

In the following sections, we describe the CARVE tower site, its location, and region of influence (Sect. 2). We then describe the measurement methods and the models used to infer CO_2 and CH_4 fluxes (Sect. 3). We present the results in Sect. 4, and conclusions, including future directions, in Sect. 5.

2 Site overview

The CARVE tower site was established in October 2011 17 km north of Fairbanks, AK,
 using an existing 32 m tower at the NOAA National Environmental Satellite, Data, and Information Service (NESDIS) facility in Fox, AK (64.986° N, 147.598° W, ground elevation 611 ma.s.l.; Fig. 1). The tower was chosen for its high elevation compared to the immediate surrounding mean ground level and its relatively large region of influence, to provide temporal and spatial context for CARVE aircraft measurements in interior
 Alaska. The site was also chosen to satisfy logistical requirements, specifically that the site be easily accessible for personnel to change out flask packages year-round, and that the site be in a location that the CARVE aircraft could sample over or close to during its campaigns without impacting the flight schedules or science mission of each flight. NOAA/NESDIS personnel are stationed in a NESDIS office 5 km from the

ity throughout the year.

25

The tower is located on a ridge, and measurements from the tower represent a wide region of interior Alaska and are less likely to be dominated by very local fluxes of CO_2 or CH_4 . Indeed, surface influence fields generated from the WRF-STILT modeling framework (Henderson et al., 2015) indicate that the tower's influence region encompasses a substantial part of Alaska (Fig. 1a). It should be noted, however, that urban

34877

emissions from Fairbanks (population 32 000), at times affect measurements at the site.

3 Methods

3.1 Measurements

⁵ Three separate measurement systems for trace gases are deployed at the CRV tower site. Programmable flask packages (PFPs) are used to collect air samples from the top level of the tower at 32 ma.g.l., daily during the CARVE flight season (April–October) and twice weekly during the remainder of the year (November–March). Additionally, measurements of ¹⁴CH₄ are made from large-volume (~ 1000 L) whole-air samples
 ¹⁰ collected approximately biweekly, also from the 32 ma.g.l. level. Lastly, continuous insitu measurements of CO₂, CH₄, and CO are made by drawing air from three heights through a Picarro G2401-m or G2401 Cavity Ring-Down Spectroscopic (CRDS) analyzer. In addition to the measurements described above, a two-dimensional (2-D) sonic anemometer was deployed at the top of the tower and was operational from April 2012
 ¹⁵ through June 2014. Here we describe the continuous CO₂, CH₄ and CO measurements made from October 2011 through the present, focusing on the calendar years

2012–2014.

Two different CRDS units have been deployed at the site as part of the CARVE project: SN CFKBDS-2008 (model G2401-m, October 2011–June 2013 and Novem-

- ²⁰ ber 2014–January 2015), and SN CFKADS-2067 (model G2401, June 2013– October 2014 and January 2015–present). The only differences between the two units as configured at the site are the flow rates (~ 550 standard cubic centimeters per minute (sccm) for CFKBDS-2008 and ~ 250 sccm for CFKADS-2067) and their precision, defined here as the standard deviation of 30 s averages, in measuring CO (1.3 ppb for CFKBDS 2008 and 4 2 ppb for CFKADS 2067). Apply are precision for CO and CH
- for CFKBDS-2008 and 4.3 ppb for CFKADS-2067). Analyzer precision for CO_2 and CH_4 is the same for both analyzers (0.03 ppm and 0.2 ppb, respectively).

The CRDS analyzer draws air through 0.635 cm (0.25 inch) outer diameter (OD) tubing (Synflex 1300) with three different inlets installed at different heights a.g.l.: 31.7 m (level 3), 17.1 m (level 2), and 4.9 m (level 1). The analyzer primarily draws from the highest level (level 3) for 50 min out of every hour, and then draws air for 5 min from each of the other levels, operating on an hourly cycle. Measurements are discarded for a time equivalent to three fluching volumes of the line after a level switch or a switch

- a time equivalent to three flushing volumes of the line after a level switch or a switch to or from a calibration tank, to allow each line to flush, because there is no separate flushing of the lines during calibrations. The sample air is not dried, and a water correction to the measurements is made in post-processing. The water correction is based on
- a laboratory experiment conducted prior to deployment of each analyzer, using methods described in Chen et al. (2013) and Rella et al. (2013). Data is collected via serial communications on a Campbell Scientific CR1000 data logger along with all auxiliary measurements (room temperature, line pressure, tank pressures, sonic anemometer measurements of temperature and 2-D winds), and averaged at 30 s increments prior
 to remote collection via the Internet connection provided by NOAA/NESDIS.
- Two standard reference gases, calibrated against NOAA standards on the WMO scales for all three gases, are each sampled every 8 h for 5 min. Mole fraction measurements of CO₂, CH₄, and CO are first corrected using a linear fit to either 5 or 6 NOAA reference tanks from a calibration performed in the laboratory prior to analyzer deployment, and then drift-corrected using the measurements of the two tanks at the site. All measurements are reported here on the WMO scales for each gas (CO₂ X2007, CH₄ X2004, and CO X2004, Dlugokencky et al., 2005; Zhao and Tans, 2006). The measurements will be updated to the newer X2004A CH₄ and X2014 CO scales (http://www.esrl.noaa.gov/gmd/ccl/ch4_scale.html; http://www.esrl. noaa.gov/gmd/ccl/ch4_scale.html; http://www.esrl. noaa.gov/gmd/ccl/ch4_scale.html; http://www.esrl. noaa.gov/gmd/ccl/ch4_scale.html; http://www.esrl. noaa.gov/gmd/ccl/ch4_scale.html; http://www.esrl. noaA.GMD Tall Tower network and described in detail in Andrews et al. (2014). The
- implementation of the uncertainty calculation for CRV is similar to that described by Andrews et al. (2014) for a Picarro CRDS instrument at Walnut Grove, CA (WGC), with

the exception of a different water vapor correction uncertainty, because the sample at WGC is dried prior to measurement by the analyzer. At CRV, the water correction uncertainty is estimated to be 0.1 ppm for CO₂, 0.5 ppb for CH₄, and 4 ppb for CO, based on analysis by Chen et al. (2013) for CO and Rella et al. (2013) for CO₂ and CH₄, and is independent of other variables, including water vapor. Total uncertainty (reproducibility and comparability to other NOAA network sites) of hourly mole fraction measurements at the site are generally < 0.2 ppm, 2 ppb, and 5 ppb for CO₂, CH₄, and CO, respectively (1-sigma). Comparisons of measurements from whole air samples in PFPs during low-variability periods show differences (median \pm 1-sigma) of -0.11 ± 0.44 ppm,

 $_{10}$ 0.8 \pm 1.2 ppb, and -1.3 ± 4.5 ppb for CO_2, CH_4, and CO, respectively, over the entire 3 year period.

3.2 Polar WRF – STILT model

The scientific analysis of CARVE atmospheric trace gas measurements is enabled through use of the Stochastic Time-Inverted Lagrangian Transport (STILT) particle dis-¹⁵ persion model (Lin et al., 2003) coupled to the Polar variant version 3.5.1 (Wilson et al., 2011) of the Advanced Research version of the Weather Research and Forecasting (WRF-ARW, Skamarock et al., 2008) numerical weather prediction model. The WRF-STILT modeling framework has been used in many studies to estimate GHG emissions using airborne, surface, and tower-based observations (Jeong et al., 2013;

- ²⁰ Kort et al., 2008; McKain et al., 2015; Chang et al., 2014; Miller et al., 2014). Atmospheric dispersion in the LPDM is simulated by advecting tracer particles by the three-dimensional gridded wind field from the WRF model, plus a turbulent velocity component represented as a stochastic process (Markov chain) (Lin et al., 2003). Time-averaged mass fluxes and convective mass fluxes from WRF are used in the
- ²⁵ dispersion calculations (Nehrkorn et al., 2010). For each observation location (i.e. "receptor"), STILT produces a two-dimensional surface influence field called a "footprint" (units of ppm(μ molm⁻²s⁻¹)⁻¹) that quantifies the influence of upwind surface fluxes on atmospheric concentrations measured at the receptor location. The footprint field

is proportional to the number of particles in a surface-influenced volume (defined as the lower half of the planetary boundary layer) and the time spent in that volume (Lin et al., 2003). As utilized in the current study, the footprint can be multiplied by an a priori flux field (units of µmol m⁻² s⁻¹) and integrated over space and time to give the incremental contribution to the mole fraction (units of ppm) as measured at the receptor location. The CARVE Polar WRF configuration consists of a triply nested grid, with the innermost domain covering mainland Alaska on a 3.3 km grid to take advantage of the improved representation at this scale of the underlying topography in this region of significant orography. The reader is directed to Henderson et al. (2015) for more detail
and validation of the meteorological fields.

STILT footprints used for this analysis were generated every 3 h during local nighttime and hourly during local daytime, for a total of 16 footprints per day, and gridded at $0.5^{\circ} \times 0.5^{\circ}$ resolution. For each footprint, 500 particles were emitted from the tower location and altitude a.s.l. and traced backwards in time for 10 days. The altitude a.s.l.

- rather than ground level was used for the location of the particle emission because the elevation of the model grid cell containing the tower site was significantly lower than the actual elevation of the site (343 vs. 611 ma.s.l.), despite use of the high-resolution grid (Henderson et al., 2015). To reduce biases induced by differences in actual and modeled topography, we use footprints generated during mid-afternoon hours (1 to 6 p.m.)
- ²⁰ local Alaska standard time (LST), UTC + 8) only for our analysis, except where noted specifically. During these hours, the lower atmosphere is generally well-mixed, and the difference between the mole fractions measured at the top level (32 m a.g.l.) and the middle level (17 m a.g.l.) average between -0.25 and 0.25 ppm for CO₂ and -0.2 and 0.3 ppb for CH₄ (maximum monthly averages for the whole time series), indicating good
- ²⁵ mixing and only a small influence from nearby sources that would cause a near-surface gradient.

We also compared measurements from the top level of the tower to CARVE aircraft measurements made above the tower site, generally during the months of March to October. We compared aircraft measurements of CO_2 and CH_4 that were made be-

low 2000 ma.s.l. (1389 ma.g.l.) and within 0.2° in latitude and longitude of the tower, between the hours of 1 and 6 p.m. LST. This allowed us to determine how well measurements made from the top level of the tower represent planetary boundary layer (PBL) average mole fractions during those times. Differences between the 29 aircraft observations and tower-based hourly means were $-0.6 \pm 2.0 \text{ ppb CH}_4$ and $0.3 \pm 0.9 \text{ ppm CO}_2$ (mean ± 1 -sigma) during the March–October air campaign period, indicating that the hourly average mole fractions at the tower are generally representative of average mole fractions in the PBL.

We expect, based on the measured gradients at the tower and the comparison with ¹⁰ aircraft measurements above the tower, that during local mid-afternoon periods the tower measurements closely represent measurements within a well-mixed PBL, and that during those times, the impact of the height difference between the modeled site elevation and the real site elevation is minimized. In our flux analysis, described in the following sections, we also specifically filter out hourly averages during which the ¹⁵ absolute value of the mole fraction gradient between 17 and 32 m a.g.l. levels in CH₄ is larger than 2 ppb.

3.3 Calculation of background mole fractions

To compare the mole fraction variability and enhancements at CRV tower to those from the modeling framework, it is necessary to determine the appropriate background
 mole fractions for both CO₂ and CH₄. We derive background mole fractions using the particle trajectories from the STILT runs and a data-based Pacific basin boundary "curtain" derived from NOAA/ESRL Global Monitoring Division GGGRN measurements using an approach similar to the one described in Jeong et al. (2013) and Miller et al. (2014). Specifically, the boundary curtain is constructed using GGGRN surface
 and aircraft vertical profile CO₂ and CH₄ observations (Sweeney et al. (2015) and www.esrl.noaa.gov/gmd/ccgg/aircraft/) to create a smoothed curtain representing the Pacific boundary. The curtain is a function of time, latitude, and altitude. For each STILT run, the 500 particles are traced back in time until they either exit a box defined by

 $[170^{\circ} \text{ W}, 130^{\circ} \text{ W}]$ and $[0^{\circ} \text{ N}, 75^{\circ} \text{ N}]$ or remain in the box for the full 10 day run. All particles are then tagged with an exit time, longitude, latitude, and altitude. Any particles whose final longitude is east of 160° W with a final latitude between 55 and 72° N and altitude below 3000 m a.s.l. are removed in order to eliminate particles that did not en-

- ter Alaska from either the western boundary or from high altitudes within the 10 days of the observation. This filter is necessary because air masses that contain surface influence from Canada or remain in Alaska for more than 10 days would not properly be represented by the Pacific boundary as background. If a given 500 particle run has more than 25 % of its particles eliminated due to the above constraints, no background
- ¹⁰ is computed for that hour, and therefore no enhancement is computed either. If at least 75% of the particles remain, these remaining particles are tagged with the mole fraction from the Pacific boundary curtain at their exit latitude, longitude, and time. The mole fractions for the particles are averaged to derive the background mole fraction for the corresponding tower measurement and WRF-STILT footprint.
- ¹⁵ Uncertainty in the background is determined similarly to Jeong et al. (2013): it is assigned the quadrature sum of the standard error of the mean mole fraction (i.e. the standard deviation of the particle mole fractions divided by the square root of the number of particles used) with the average value of the RMS residuals of the empirical background curtain of the particles. This uncertainty is dominated by the RMS residuals 20 of the boundary curtain.

3.4 CO₂ flux model

The CO₂ measurements at the CRV tower were interpreted with the assistance of biospheric CO₂ flux estimates generated by the Polar Vegetation Photosynthesis and Respiration Model (PolarVPRM, Luus and Lin, 2015). PolarVPRM captures the strong diurnal and seasonal variability of CO₂ fluxes parsimoniously, according to empirical associations between environmental conditions and eddy covariance measurements of CO₂, and regionally across Alaska (3 hourly, $1/6^{\circ} \times 1/4^{\circ}$ latitude × longitude), using data products from the North American Regional Reanalysis (NARR) (Mesinger

et al., 2006) and the Moderate Resolution Imaging Spectroradiometer (MODIS). Subnivean and growing season respiration are calculated as functions of NARR soil and air temperature, respectively; snow and growing seasons are differentiated using MODIS snow cover (Riggs and Hall, 2011). Photosynthesis is calculated as a function of NARR

⁵ air temperature, NARR shortwave radiation, water availability (via MODIS), and vegetation (via the MODIS enhanced vegetation index, Solano et al., 2010). The CO₂ fluxes from PolarVPRM were convolved with footprints from observations at the tower to derive model-based enhancement above background (Δ CO₂) for the 3 year period from 2012–2014. These modeled CO₂ enhancements were compared to CO₂ enhance-10 ments observed at CRV.

Hourly observations used for CO_2 analysis were restricted to periods between 1 and 6 p.m. LST, to minimize discrepancies between real and modeled boundary layer dynamics. In addition, as described above, samples that had no background determination for more than 25% of the released particles were omitted. Additional filters on

- ¹⁵ the data were designed to restrict analysis to periods when the PBL was most likely well mixed, as determined from the vertical gradient in CH_4 mole fractions between the 17 and 32 m levels; only data for which the CH_4 vertical gradient was less than 2 ppb were retained. Also, only data observations with low temporal variability were retained, determined as having a standard deviation of 30 s measurements in an hour below
- ²⁰ 7 ppb in CO and 3 ppb in CH₄; this filter was applied to reduce influence from local sources, a concern at this site because of the proximity of Fairbanks. Lastly, biomass-burning (and some large pollution) events were filtered out by removing observations for which the enhancement in CO (relative to the background determined using methods described in Sect. 3.2) exceeded 20 ppb. The filters described above were used
- to filter data only for the CO₂ flux-model comparison analysis described in this section with the results in Sect. 4.4, and for the CH₄ footprint and flux analyses described in Sect. 3.5 with results in Sect. 4.5 and 4.6.

3.5 CH₄ flux estimation

In the CH₄ analysis, as for CO₂, only hourly average mole fractions between 1 and 6 p.m. LST were used, with the same filters applied on the observations to limit instances of high variability, large vertical gradients, and biomass burning, as outlined in Sect. 3.4.

Average CH₄ fluxes were estimated by scaling two different flux maps. The first flux map is a uniform land-based flux (with oceanic flux set to zero, assuming the oceanic CH₄ flux contribution is negligible, Kirschke et al., 2013) similar to what was used in Chang et al. (2014) to estimate CH₄ fluxes using aircraft observations from the 2012 CARVE campaign. The second flux map pattern is based on elevation data from NOAA's NGDC, (http://www.ngdc.noaa.gov/mgg/topo/report/ globedocumentationmanual.pdf) (Fig. 1a). The elevation map was averaged to the same spatial resolution as the footprints (0.5°×0.5°) and adjusted so that the ocean and elevations higher than 1000 ma.s.l. were assumed to have zero CH₄ flux. Elevations

- between 0 and 1000 m a.s.l. were scaled linearly from 1 to 0, with areas of zero elevation (including lakes) assigned 1 and 1000 m a.s.l. assigned 0. Fluxes were assumed to be diurnally constant. A third map based on elevation gradients (in which highly sloped regions had less flux and flatter areas had higher flux) was also tested, but the results were very similar to the elevation-based map, so they are not shown here.
- ²⁰ Observed CH₄ enhancements relative to the footprint background were averaged over the mid-afternoon hours (1–6 p.m. LST) to obtain daily averages. These daily enhancements were then averaged to obtain monthly average Δ CH₄ values throughout each year. However, in many winter months, fewer than 6 days of observations remained after the data filtering; those months were omitted from the analysis. The con-
- stant flux map and the elevation-based flux map were convolved with the hourly footprints from the WRF-STILT model to obtain initial values of modeled ΔCH_4 . These were then averaged to create daily values (with the same filters as for the observations) and

then to monthly values. The monthly flux maps were then scaled to the observations, so that the simulated monthly ΔCH_4 matched the observations.

Uncertainties on monthly fluxes derived from both maps were determined from the background mole fraction uncertainty alone; uncertainty from transport errors (e.g., er-

- ⁵ ror in PBL depth or wind speed) was not accounted for. Monthly background errors for CH₄ ranged from 2–7 ppb (average 5 ppb), which was generally of the same order of magnitude as the CH₄ enhancements. Uncertainty on the monthly enhancements was calculated as the average of the uncertainty on the background for each day divided by the square root of the number of independent samples during that month, approximat-
- ing the standard error of the monthly mean enhancement. The correlation time scale of the background for CH₄ (after a 60 day smoother was subtracted to eliminate the long-term temporal correlations) was approximated at 9 days, consistent with synoptic-scale variability. The number of independent realizations for each month was therefore derived as the number of days in that month divided by 9. This represents a 1-sigma uncertainty estimate; the fractional uncertainty on the monthly CH₄ enhancement was
 - then propagated to the monthly flux.

4 Results

4.1 Diurnal cycles

The diurnal cycles of the CO_2 and CH_4 at the tower have been analyzed over the study ²⁰ period, 2012–2014. All analysis shown is based on hourly-averaged measurements from the top-most level at 32 ma.g.l. Measurements during times when the CO mole fraction exceeded 200 ppb were removed to filter out the effect of large biomass burning events. No other filters were applied on the data in this portion of the analysis. The diurnal cycle of CO_2 shows an amplitude (maximum–minimum CO_2) of 10 ppm in July, with

²⁵ a wintertime (November–April) magnitude of approximately 2 ppm (Fig. 2, top panel), with similar patterns each year. CH₄ diurnal cycle amplitudes also show a maximum in

summer (in either July or August, depending on the year) between 20 and 30 ppb. The wintertime diurnal cycle of CH_4 shows an average amplitude of 10 ppb (Fig. 2, lower panel). Shaded areas in Fig. 2 indicate the standard deviation of that month's average over all days in the 3 year period, indicating significant variability in the amplitudes for both gases, and especially for CH_4 , where the amplitude variability (one-sigma) ranges from zero to 45 ppb.

The average amplitudes of the CH_4 and CO_2 diurnal cycles at the CARVE tower are significantly smaller than those that have been reported at other Arctic and boreal measurement sites. Sasakawa et al. (2010) report CH_4 diurnal cycle magnitudes at a network of Siberian tower sites on the order of 100–200 ppb in summertime, when wetland emissions in that area are largest and accumulate in the nighttime and earlymorning planetary boundary layer (PBL). Sasakawa et al. (2013) show examples of CO_2 diurnal cycle amplitudes at a tower site in West Siberia of 30 ppm or greater. Worthy et al. (2015) compare diurnal cycle amplitudes of CH_4 at various Arctic tower

- ¹⁵ sites throughout Canada and North America, finding that summertime diurnal CH₄ amplitudes at all the Arctic and boreal Canadian sites are significantly larger than at CRV tower. Winderlich et al. (2010) also report similarly large diurnal cycle amplitudes in CH₄ (~ 200 ppb) and CO₂ (~ 25 ppm) from the lower levels of the ZOTTO tall tower in boreal Siberia; however, at the highest level (301 ma.g.l.) the average July 2009 diurnal
- ²⁰ cycle amplitude is significantly smaller at ~ 50 ppb CH_4 and ~ 5 ppm CO_2 , presumably because the top of the night-time PBL is often below this tallest level. This may be the case at CRV, which despite its low height a.g.l., is elevated above the surrounding area and likely is not affected by very local CH_4 sources, such as wetlands, that result in trapped emissions in the night-time PBL at the site. The CRV tower is surrounded by
- trees and other vegetation, however, so the CO₂ cycle is comparably larger. The diurnal cycle at the 17 and 5 m a.g.l. levels is slightly greater than at 32 m a.g.l. in summer, but not significantly so (1–2 ppb larger for CH₄ and 1–2 ppm for CO₂ on average in July and August).

4.2 Seasonality of winds and influence functions

The mid-afternoon daily average footprints from the WRF-STILT model were examined to determine the influence of different regions on the measurements at the tower throughout the year. The total magnitude of land-surface influence (ocean influence is

- ⁵ not included) on the tower measurements for each month of each of the three study years (2012, 2013, 2014) was determined (Fig. 3), along with the total influence of several sub-regions: Canada (light blue, Fig. 3), the North Slope of Alaska (defined as north of the Brooks Range, red, Fig. 3), the remainder of Alaska (dark blue, Fig. 3), and Eurasia (yellow, Fig. 3). The seasonality of the land surface influence is clear and consistent between all three years. Specifically, in all three years, the months of May
- through September show significantly less land surface influence on the tower than October through April. This stems from the smaller influence of Canada, and to a lesser extent, lower Alaska, on the measurements during the summer months. The influence of the Eurasian continent is very small throughout the years, but so is the influence
- of the North Slope of Alaska. This is also apparent when the mid-afternoon footprint influences are aggregated over seasons and years, as shown in the 80% influence range (Fig. 1a), which does not include the North Slope region. From this analysis we conclude that measurements at the CRV tower are not substantially affected by emissions north of the Brooks Range, and any emissions estimates made using the tower measurements will not apply to the North Slope.

Daytime wind measurements from the 2-D sonic anemometer at the tower support the finding of large seasonality in the footprints. Winds at the tower during May– September are predominantly from the west and southwest, with some frequency of winds from the east as well. However, from October to April, the winds are almost ex-

²⁵ clusively from the east/northeast. These wind directions support the conclusion from the model influence functions that wintertime measurements are more influenced by Canadian land than in summertime, as shown in Fig. 3. In addition, winds in any season do not generally come from the North, supporting the lack of influence from the

North Slope. Similar seasonality and lack of northern influence was found in a recent analysis of data from NOAA/ESRL Aircraft Network at Poker Flat, AK (Sweeney et al., 2015).

4.3 Background and relative enhancements of CH₄, CO₂, and CO

- ⁵ The definition of an appropriate background is a crucial aspect of analyzing the CRV tower CO₂ and CH₄ measurements. We calculate the background as described in Sect. 3.2, using the particle back trajectories and the empirical Pacific boundary curtain, and refer to this as the footprint background. We also compare this background to the value of the same Pacific curtain at 3500 ma.s.l. and 65° N, i.e. the free troposphere at the latitude of the tower. For CO₂ (middle panel, Fig. 5) the definition of the background does not have as large an effect as it does for CH₄. For CH₄ (top panel, Fig. 5) the choice of background is crucial to any analysis of the measurements, for two reasons. First, the CH₄ signal at CRV is relatively small compared to the variability of the background. Second, the CH₄ background varies depending on the latitudinal ori-
- gin of the air mass, because of the large global latitudinal gradient in CH₄ (Dlugokencky et al., 2009). Comparison of the measurements with the footprint background and free-tropospheric background (Fig. 5) illustrates that the footprint background varies at synoptic time scales as air-mass origins change, and tracks the variability in the measurements at the site. CH₄ enhancements over background are small and thus very
- ²⁰ sensitive to background choice (top panel in Fig. 5). We note that despite the small signal, however, the time series of CH₄ observations clearly shows both wintertime and summertime enhancements, with wintertime enhancements sometimes correlated with CO enhancements as well, indicating a possible anthropogenic source for these signals (see Sect. 4.6 and Table 1). Evidence of biomass burning events is also clear in the sector of biomas
- all three species, but most easily observed in the CO signals during the summers of 2012 and 2013.

4.4 CO₂ model-observation comparison

Observations of monthly mean CO_2 relative to the background (ΔCO_2) show consistent features from year to year (Fig. 6), with the sign of the enhancements showing the sign of the monthly net CO_2 fluxes, or Net Ecosystem Exchange (NEE). Positive en-

- ⁵ hancements from January–April indicate that respiration is occurring even during this coldest period of the year. In addition, all years show the highest respiration signal in October, possibly indicative of photosynthesis stopping while soil temperatures are still high enough to sustain significant respiration, although some of this signal could also be due to the seasonality in vertical mixing and/or winds. Although the maximum draw-
- ¹⁰ down occurs in July and is of similar magnitude in all years (~ 8 ppm), the transition from net respiration to net photosynthetic uptake occurs earlier in 2014 (April) than in 2012 and 2013 (May). The timing and magnitude of the ΔCO_2 observations relative to background represent a stringent test for the transport and surface flux models.

The modeled ΔCO_2 from the convolution of WRF-STILT footprints with PolarVPRM

- ¹⁵ fluxes are compared to hourly averaged observed ΔCO_2 mole fractions at the tower during the mid-afternoon in Fig. 6. (Note that the time series data in Fig. 6 (top) has not been filtered, but the monthly averages in the lower panel only use filtered data.) Both the hourly time series and monthly average comparisons between modeled and observed ΔCO_2 at the tower during mid-day hours indicate that the PolarVPRM fluxes
- ²⁰ and WRF-STILT meteorology are able to reproduce the magnitude and timing of the tower CO₂ signal remarkably well. The monthly average Δ CO₂ observations compared to the model (Fig. 6, lower panel) indicate that the PolarVPRM/WRF-STILT modeled NEE is slightly lower than observations in May and June in both 2012 and 2013. (However, in all of these months the model results, with no uncertainty estimates, overlap
- with the one-sigma data uncertainty.) Whether this small offset between model and observations results from insufficient modeled respiration or too much modeled photosynthesis during the spring is impossible to tell from CO₂ observations only. In addition, PolarVPRM systematically underestimates the magnitude of the observed late fall res-

piration flux (October to November) in all three years. This may be because model respiration is calculated as a function of air temperature when per-pixel snow cover area is < 50 %, whereas actual rates of late fall respiration are influenced by microbial activity sustained in the soil, which cools more gradually than the air.

- ⁵ Hourly observations of ΔCO_2 that satisfy the filtering conditions are well correlated with modeled ΔCO_2 in all three years (Fig. 7). The data close to the 1 : 1 line indicate that the magnitude of the fluxes is generally well captured by the model. The high correlations indicate that in addition to capturing the magnitude, the PolarVPRM and WRF-STILT models are likely capturing the timing and spatial structure of the fluxes in
- ¹⁰ boreal Alaska as well. In all years there are instances of higher observed ΔCO_2 values than predicted by the model; these are likely either anthropogenic influences to the signal (which are not in the PolarVPRM fluxes) or, more likely given the discrepancy occurs specifically in October and November, the underestimate of fall CO_2 respiration. In addition, in all years the earlier modeled spring drawdown (Figs. 6 and 7), leads to
- ¹⁵ some data points with more negative ΔCO_2 in the model than in the observations. The correlations are strong in all three years however ($R^2 = 0.61$ to 0.75), indicating that the PolarVPRM CO₂ fluxes and the WRF-STILT transport model are able to reproduce observed signals at the tower remarkably well with no adjustment to match the data.

4.5 CH₄ model-observation comparison

- ²⁰ The scaled monthly CH₄ fluxes from the elevation-based and uniform (constant) flux maps were convolved with the WRF-STILT footprints corresponding to the observations. The average daily Δ CH₄ from each model was compared with the observed enhancements (Fig. 8) for each year. The elevation-based model enhancements (lower row, Fig. 8) match the data slightly better than the uniform flux model (upper row, Fig. 8)
- in 2012 and 2014, but not 2013. We also investigated a third spatial flux map pattern that was based on the gradient in elevation, but did not find any improvement correlations over the simpler elevation-based and uniform flux models. Neither model was able to achieve good correlations between the model and the observations, a conclu-

sion that was also reached by Chang et al. (2014) when they investigated multiple different CH₄ flux models. This result is in sharp contrast to the high correlations achieved using the PolarVPRM CO₂ fluxes with the same WRF-STILT footprints, leading to the conclusion that the WRF-STILT meteorology is able to replicate observations when an accurate spatial flux map is used, and that the lack of correlation with CH₄ is due to the poor spatial representation of the simple models used here and the large uncertainty in the background.

4.6 Average scaled CH₄ fluxes

Monthly CH₄ fluxes from the uniform flux map were averaged over the state of Alaska
to obtain average fluxes for each month and for each year, and compared with the results from (Chang et al., 2014) (Fig. 9). Results from the elevation-based flux map were statistically the same as those from the uniform flux map and are not shown. CH₄ fluxes are small relative to those determined from some flux tower or chamber-based studies in arctic wetlands (Euskirchen et al., 2014; Johnston et al., 2014; Whalen and Reeburgh, 1988; Fan et al., 1992; Olefeldt et al., 2013), but very similar in magnitude to CH₄ fluxes recently reported from a black spruce forest during the snow-free seasons from 2011–2013 at a flux tower site in Fairbanks, AK (Iwata et al., 2015), as well as those reported from the Zotino Tall Tower Observatory (ZOTTO) in Siberia during the

- summers of 2009–2011 (7.7 mg $CH_4 m^{-2} d^{-1}$) (Winderlich et al., 2014). They are also smaller than fluxes estimated by Chang et al. (2014) using CARVE aircraft observations in the summer of 2012 (Fig. 9), but the two results overlap within their 1-sigma uncertainty bands. One possible reason for the lower average fluxes observed at the CARVE tower could be the region to which the CARVE tower observations are sensitive in the summer. The CARVE tower observations do not capture emissions over the
- North Slope of Alaska, where other studies have shown large (but highly variable) CH₄ emissions, at least over small areas (Euskirchen et al., 2014; Morrissey and Livingston, 1992; von Fischer et al., 2010; Olefeldt et al., 2013). The signal in CH₄ at CRV may also be small because of the CARVE tower's large region of influence, which integrates

signals from a wide variety of ecosystems that have different flux profiles, including extensive upland and mountain regions, not only low-lying wetlands and forests.

The tower observations also indicate the presence of non-zero fall and wintertime fluxes. Recently, Zona et al. (2015) reported significant natural CH_4 fluxes persisting

- ⁵ through the late fall in the North Slope of Alaska, and our results support the existence of late-fall (September–October) CH_4 fluxes in the boreal zone as well. Additionally, our analysis also shows the presence of CH_4 emissions in late winter (January–March) in some years, which are less likely to be biogenic. To understand the role that local point sources from nearby Fairbanks might play, an analysis of correlations between
- ¹⁰ ΔCH_4 and ΔCO mid-afternoon hourly enhancements was done. These enhancements indicate that some wintertime CH_4 emissions are likely anthropogenic, with correlation coefficients (R^2) generally larger in the winter months and close to zero in June, July and August of all years. Not all winter months show high correlations, and May 2012 also have highly correlated ΔCH_4 and ΔCO (Table 1).

15 5 Conclusions

The CARVE tower, located on a ridge outside Fairbanks, is well situated to provide regional year-round CO₂ and CH₄ observations that provide context to the CARVE aircraft campaign measurements, which were made throughout Alaska from March to November from 2012 to 2015. The WRF-STILT transport model was used to determine the influence region of the site and its inter-annual and seasonal variability. The model showed significantly more influence from the region east of the tower in wintertime, a pattern that repeated all three years and was confirmed by anemometer data from the site. The model also indicated that processes in the North Slope of Alaska have very little influence on the tower observations. This seasonality of transport to the region has

²⁵ been previously documented (Sweeney et al., 2015), and implies that additional longterm observing sites are required to constrain Alaskan fluxes; a site in Western Alaska, for example, would be more likely to have interior Alaska in its observation footprint in

the wintertime, and a site north of the Brooks Range would be required to investigate fluxes from the North Slope.

We calculated enhancements of CO_2 and CH_4 during local mid-afternoon times by subtracting a background also determined using the WRF-STILT model particle trajec-

tories, and found that the background choice is critical for CH₄, for which enhancements are very small, and of the same order of magnitude as the uncertainties. CO₂ enhancements at the CARVE tower site are replicated remarkably well by the WRF-STILT model when convolved with PolarVPRM biogenic CO₂ fluxes (Luus and Lin, 2015). The high correlation between modeled and observed CO₂ give confidence in the STILT foot prints and the WRF meteorological model that was used to generate them. The signal in CO₂ is larger than that for CH₄, such that the background uncertainty is not as large relative to the enhancements or depletions.

The WRF-STILT meteorological model enables us to constrain the magnitude of mean monthly CH_4 fluxes in the region of influence of the tower for all three years. ¹⁵ Using two different distribution maps of CH_4 emissions we determine that average CH_4 emissions over Alaska in summer range between 3 and 8 mg CH_4 m⁻² d⁻¹, albeit with large uncertainties stemming from the large uncertainty in the background. The tower observations also indicate that there are no significant differences between the three years. This simple analysis provides a flux estimate range that applies as an

- ²⁰ average over a very large area of Alaska (Fig. 1a). CH₄ fluxes in this region are likely highly heterogeneous, but our measurements show that the average flux over the entire region is relatively small. This result suggests that although there may be small areas with large fluxes, there are other areas with little to no emissions. For this reason, the observations at the tower give context to other flux estimates, from flux towers or
- ²⁵ chamber studies, for example, that are representative of much smaller areas and are difficult to scale to the larger domain because of high spatial and temporal variability. We also observe CH_4 fluxes persisting into the fall (September–October) in all three years, and some fluxes in winter and early spring, depending on the year, which may be partially or entirely anthropogenic, based on an analysis of correlations of CH_4 with CO.

These late fall and wintertime emissions demonstrate the need for year-round in-situ observations in the high northern latitudes.

The CARVE tower site provides a continuous observation platform that will contribute to future efforts to investigate the high-latitude carbon cycle and its response

- ⁵ to warming. As a long-term measurement site with a large regional coverage it will provide understanding of changing emissions in interior Alaska. Our analysis of the years 2012–2014 indicates no measurable change in emissions influencing this site over this period. These tower observations are sensitive to changes in emissions and provide the capability to detect such changes in the future. However, the location of
- the CARVE tower prohibits any quantification or observation of processes on the North Slope, indicating that additional long-term observation sites with large regional coverage are required north of the Brooks Range of Alaska to detect changes in emissions in the higher northern latitudes. Future efforts will combine the observations from the CARVE tower with other aircraft and ground-based observations in a formal inversion for example, the applied for an atighty and term problem and CO. for an Alaska

framework to solve for spatially and temporally resolved CH_4 and CO_2 fluxes in Alaska.

Data availability

25

All the tower observations and WRF-STILT footprints used in our analysis are publicly available on the CARVE data portal at https://ilma.jpl.nasa.gov/portal/. They will be archived at the U.S. Oak Ridge National Laboratory Distributed Active Archive Center for Biogeochemical Dynamics (ORNL DAAC, https://daac.ornl.gov).

Acknowledgements. The research described in this paper was performed for the Carbon in Arctic Reservoirs Vulnerability Experiment (CARVE), an Earth Ventures (EV-1) investigation, under contract with the National Aeronautics and Space Administration. Part of the research described in this paper was performed at the Jet Propulsion Laboratory, California Institute of Technology, under contract with the National Aeronautics and Space Administration.

The authors would like to acknowledge the invaluable assistance of the NOAA/NESDIS personnel in Fox, Alaska: Marc Meindl and Frank Holan for changing out flask packages and highpressure cylinders throughout the year and providing expert technical assistance when needed,

including repairs and replacing equipment; Tom Narow and Sean Meyn for providing the Ethernet link and assistance with site networking; Robert Cox, Bonnie Croskey, and Lisa Auvil for shipping out flask packages and cylinders; and Larry Ledlow for allowing the CARVE project to use the site. We thank Jack Higgs, Eric Moglia, Molly Crotwell, Pat Lang, and Duane Kitzis at NOAA/ESRL for providing logistics support, flask packages and shipments, data flagging assistance, and cylinder calibrations for the site. We thank Matt Pender for on-site support, Sean Hardman (JPL) for making the CARVE public data portal, and Kathryn McKain (NOAA/ESRL),

Jonathan Kofler (NOAA/ESRL), Rachel Chang (Dalhousie University) and the CARVE Science Team for valuable discussions contributing to the manuscript.

10 **References**

5

- Andrews, A. E., Kofler, J. D., Trudeau, M. E., Williams, J. C., Neff, D. H., Masarie, K. A., Chao, D. Y., Kitzis, D. R., Novelli, P. C., Zhao, C. L., Dlugokencky, E. J., Lang, P. M., Crotwell, M. J., Fischer, M. L., Parker, M. J., Lee, J. T., Baumann, D. D., Desai, A. R., Stanier, C. O., De Wekker, S. F. J., Wolfe, D. E., Munger, J. W., and Tans, P. P.: CO₂, CO,
- and CH₄ measurements from tall towers in the NOAA Earth System Research Laboratory's Global Greenhouse Gas Reference Network: instrumentation, uncertainty analysis, and recommendations for future high-accuracy greenhouse gas monitoring efforts, Atmos. Meas. Tech., 7, 647–687, doi:10.5194/amt-7-647-2014, 2014.

Bruhwiler, L., Dlugokencky, E., Masarie, K., Ishizawa, M., Andrews, A., Miller, J., Sweeney, C.,

Tans, P., and Worthy, D.: CarbonTracker-CH₄: an assimilation system for estimating emissions of atmospheric methane, Atmos. Chem. Phys., 14, 8269–8293, doi:10.5194/acp-14-8269-2014, 2014.

CarbonTracker CT2013B: available at: http://www.esrl.noaa.gov/gmd/ccgg/carbontracker/, last access: 17 October 2013.

²⁵ Chang, R. Y. W., Miller, C. E., Dinardo, S. J., Karion, A., Sweeney, C., Daube, B. C., Henderson, J. M., Mountain, M. E., Eluszkiewicz, J., Miller, J. B., Bruhwiler, L. M. P., and Wofsy, S. C.: Methane emissions from Alaska in 2012 from CARVE airborne observations, P. Natl. Acad. Sci. USA, 111, 16694–16699, doi:10.1073/pnas.1412953111, 2014.

Chen, H., Karion, A., Rella, C. W., Winderlich, J., Gerbig, C., Filges, A., Newberger, T., ³⁰ Sweeney, C., and Tans, P. P.: Accurate measurements of carbon monoxide in humid air using

34896

the cavity ring-down spectroscopy (CRDS) technique, Atmos. Meas. Tech., 6, 1031–1040, doi:10.5194/amt-6-1031-2013, 2013.

- Dlugokencky, E. J., Myers, R. C., Lang, P. M., Masarie, K. A., Crotwell, A. M., Thoning, K. W., Hall, B. D., Elkins, J. W., and Steele, L. P.: Conversion of NOAA atmospheric dry air CH₄
- ⁵ mole fractions to a gravimetrically prepared standard scale, J. Geophys. Res.-Atmos., 110, D18306, doi:10.1029/2005JD006035, 2005.
 - Dlugokencky, E. J., Bruhwiler, L., White, J. W. C., Emmons, L. K., Novelli, P. C., Montzka, S. A., Masarie, K. A., Lang, P. M., Crotwell, A. M., Miller, J. B., and Gatti, L. V.: Observational constraints on recent increases in the atmospheric CH₄ burden, Geophys. Res. Lett., 36, L18803, doi:10.1029/2009gl039780, 2009.
- Euskirchen, E. S., Edgar, C. W., Turetsky, M. R., Waldrop, M. P., and Harden, J. W.: Differential response of carbon fluxes to climate in three peatland ecosystems that vary in the presence and stability of permafrost, J. Geophys. Res.-Biogeo., 119, 1576–1595, doi:10.1002/2014JG002683, 2014.

10

25

Fan, S. M., Wofsy, S. C., Bakwin, P. S., Jacob, D. J., Anderson, S. M., Kebabian, P. L., Mc-Manus, J. B., Kolb, C. E., and Fitzjarrald, D. R.: Micrometeorological measurements of CH₄ and CO₂ exchange between the atmosphere and subarctic tundra, J. Geophys. Res.-Atmos., 97, 16627–16643, doi:10.1029/91JD02531, 1992.

Harriss, R. C., Sachse, G. W., Hill, G. F., Wade, L., Bartlett, K. B., Collins, J. E., Steele, L. P., and

Novelli, P. C.: Carbon monoxide and methane in the North American Arctic and sub-arctic troposphere – July–August 1988, J. Geophys. Res.-Atmos., 97, 16589–16599, 1992.

Hayes, D. J., Kicklighter, D. W., McGuire, A. D., Chen, M., Zhuang, Q., Yuan, F., Melillo, J. M., and Wullschleger, S. D.: The impacts of recent permafrost thaw on land–atmosphere greenhouse gas exchange, Environ. Res. Lett., 9, 045005, doi:10.1088/1748-9326/9/4/045005, 2014.

- Henderson, J. M., Eluszkiewicz, J., Mountain, M. E., Nehrkorn, T., Chang, R. Y.-W., Karion, A., Miller, J. B., Sweeney, C., Steiner, N., Wofsy, S. C., and Miller, C. E.: Atmospheric transport simulations in support of the Carbon in Arctic Reservoirs Vulnerability Experiment (CARVE), Atmos. Chem. Phys., 15, 4093–4116, doi:10.5194/acp-15-4093-2015, 2015.
- ³⁰ IPCC: Climate Change 2013: The Physical Science Basis, Contribution of Working Group I to the Fourth Assessment Report of the Intergovernmental Panel on Climate Change, Cambridge University Press, Cambridge, UK, New York, NY, USA, 1535, 2013.

- Iwata, H., Harazono, Y., Ueyama, M., Sakabe, A., Nagano, H., Kosugi, Y., Takahashi, K., and Kim, Y.: Methane exchange in a poorly-drained black spruce forest over permafrost observed using the eddy covariance technique, Agr. Forest Meteorol., 214–215, 157–168, doi:10.1016/j.agrformet.2015.08.252, 2015.
- Jeong, S., Hsu, Y.-K., Andrews, A. E., Bianco, L., Vaca, P., Wilczak, J. M., and Fischer, M. L.: A multitower measurement network estimate of California's methane emissions, J. Geophys. Res.-Atmos., 118, 2013JD019820, doi:10.1002/jgrd.50854, 2013.
 - Johnston, C. E., Ewing, S. A., Harden, J. W., Varner, R. K., Wickland, K. P., Koch, J. C., Fuller, C. C., Manies, K., and Jorgenson, M. T.: Effect of permafrost thaw on CO₂ and CH₄
- exchange in a western Alaska peatland chronosequence, Environ. Res. Lett., 9, 085004, doi:10.1088/1748-9326/9/8/085004, 2014.
 - Kirschke, S., Bousquet, P., Ciais, P., Saunois, M., Canadell, J. G., Dlugokencky, E. J., Bergamaschi, P., Bergmann, D., Blake, D. R., Bruhwiler, L., Cameron-Smith, P., Castaldi, S., Chevallier, F., Feng, L., Fraser, A., Heimann, M., Hodson, E. L., Houweling, S., Josse, B., Fraser, P. J., Krummel, P. B., Lamarque, J.-F., Langenfelds, R. L., Le Quere, C., Naik, V.,
- Fraser, P. J., Krummel, P. B., Lamarque, J.-F., Langenfelds, R. L., Le Quere, C., Naik, V., O'Doherty, S., Palmer, P. I., Pison, I., Plummer, D., Poulter, B., Prinn, R. G., Rigby, M., Ringeval, B., Santini, M., Schmidt, M., Shindell, D. T., Simpson, I. J., Spahni, R., Steele, L. P., Strode, S. A., Sudo, K., Szopa, S., van der Werf, G. R., Voulgarakis, A., van Weele, M., Weiss, R. F., Williams, J. E., and Zeng, G.: Three decades of global methane sources and sinks, Nat. Geosci., 6, 813–823, doi:10.1038/ngeo1955, 2013.
- Kort, E. A., Eluszkiewicz, J., Stephens, B. B., Miller, J. B., Gerbig, C., Nehrkorn, T., Daube, B. C., Kaplan, J. O., Houweling, S., and Wofsy, S. C.: Emissions of CH₄ and N₂O over the United States and Canada based on a receptor-oriented modeling framework and COBRA-NA atmospheric observations, Geophys. Res. Lett., 35, L18808, doi:10.1029/2008gl034031, 2008.
 - Lin, J. C., Gerbig, C., Wofsy, S. C., Andrews, A. E., Daube, B. C., Davis, K. J., and Grainger, C. A.: A near-field tool for simulating the upstream influence of atmospheric observations: the Stochastic Time-Inverted Lagrangian Transport (STILT) model, J. Geophys. Res.-Atmos., 108, 4493, doi:10.1029/2002JD003161, 2003.
- ³⁰ Lin, J. C., Brunner, D., Gerbig, C., Stohl, A., Luhar, A. K., and Webley, P. W. (Eds.): Lagrangian Modeling of the Atmosphere, AGU Geophysical Monograph (Vol. 200), Washington D.C., doi:10.1029/GM200, 349 pp., 2012.

Luus, K. A. and Lin, J. C.: The Polar Vegetation Photosynthesis and Respiration Model: a parsimonious, satellite-data-driven model of high-latitude CO₂ exchange, Geosci. Model Dev., 8, 2655–2674, doi:10.5194/gmd-8-2655-2015, 2015.

McGuire, A. D., Anderson, L. G., Christensen, T. R., Dallimore, S., Guo, L. D., Hayes, D. J.,

- Heimann, M., Lorenson, T. D., Macdonald, R. W., and Roulet, N.: Sensitivity of the carbon cycle in the Arctic to climate change, Ecol. Monogr., 79, 523–555, 2009.
 - McGuire, A. D., Hayes, D. J., Kicklighter, D. W., Manizza, M., Zhuang, Q., Chen, M., Follows, M. J., Gurney, K. R., McClelland, J. W., Melillo, J. M., Peterson, B. J., and Prinn, R. G.: An analysis of the carbon balance of the Arctic Basin from 1997 to 2006, Tellus B, 62, 455– 474. doi:10.1111/i.1600-0889.2010.00497.x. 2010.
- McKain, K., Down, A., Raciti, S. M., Budney, J., Hutyra, L. R., Floerchinger, C., Herndon, S. C., Nehrkorn, T., Zahniser, M. S., Jackson, R. B., Phillips, N., and Wofsy, S. C.: Methane emissions from natural gas infrastructure and use in the urban region of Boston, Massachusetts, P. Natl. Acad. Sci. USA, 112, 1941–1946, doi:10.1073/pnas.1416261112, 2015.

10

20

25

¹⁵ Mesinger, F., DiMego, G., Kalnay, E., Mitchell, K., Shafran, P. C., Ebisuzaki, W., Jović, D., Woollen, J., Rogers, E., Berbery, E. H., Ek, M. B., Fan, Y., Grumbine, R., Higgins, W., Li, H., Lin, Y., Manikin, G., Parrish, D., and Shi, W.: North American regional reanalysis, B. Am. Meteorol. Soc., 87, 343–360, doi:10.1175/BAMS-87-3-343, 2006.

Miller, C. E.: The Carbon in Arctic Reservoirs Vulnerability Experiment (CARVE), in preparation, 2015.

- Miller, S. M., Worthy, D. E. J., Michalak, A. M., Wofsy, S. C., Kort, E. A., Havice, T. C., Andrews, A. E., Dlugokencky, E. J., Kaplan, J. O., Levi, P. J., Tian, H., and Zhang, B.: Observational constraints on the distribution, seasonality, and environmental predictors of North American boreal methane emissions, Global Biogeochem. Cy., 28, 146–160, doi:10.1002/2013GB004580, 2014.
- MODIS Vegetation Indices (MOD13) C5 Users's Guide, Terrestrial Biophysics and Remote Sensing Lab, The University of Arizona, available at: http://www.ctahr.hawaii.edu/grem/modis-ug.pdf, last access: 7 September 2010.
- Morrissey, L. A. and Livingston, G. P.: Methane emissions from Alaska Arctic tundra: an assessment of local spatial variability, J. Geophys. Res.-Atmos., 97, 16661–16670, doi:10.1029/92JD00063, 1992.

- Nehrkorn, T., Eluszkiewicz, J., Wofsy, S., Lin, J., Gerbig, C., Longo, M., and Freitas, S.: Coupled weather research and forecasting–stochastic time-inverted lagrangian transport (WRF–STILT) model, Meteorol. Atmos. Phys., 107, 51–64, doi:10.1007/s00703-010-0068-x, 2010.
 O'Connor, F. M., Boucher, O., Gedney, N., Jones, C. D., Folberth, G. A., Coppell, R., Friedling-
- stein, P., Collins, W. J., Chappellaz, J., Ridley, J., and Johnson, C. E.: Possible role of wetlands, permafrost, and methane hydrates in the methane cycle under future climate change: a review, Rev. Geophys., 48, RG4005, doi:10.1029/2010rg000326, 2010.
 - Oechel, W. C., Hastings, S. J., Vourlitis, G., Jenkins, M., Riechers, G., and Grulke, N.: Recent change of Arctic tundra ecosystems from a net carbon dioxide sink to a source, Nature, 361, 520–523, 1993.
- 10

25

Olefeldt, D., Turetsky, M. R., Crill, P. M., and McGuire, A. D.: Environmental and physical controls on northern terrestrial methane emissions across permafrost zones, Glob. Change Biol., 19, 589–603, doi:10.1111/gcb.12071, 2013.

Rella, C. W., Chen, H., Andrews, A. E., Filges, A., Gerbig, C., Hatakka, J., Karion, A.,

Miles, N. L., Richardson, S. J., Steinbacher, M., Sweeney, C., Wastine, B., and Zellweger, C.: High accuracy measurements of dry mole fractions of carbon dioxide and methane in humid air, Atmos. Meas. Tech., 6, 837–860, doi:10.5194/amt-6-837-2013, 2013.

Riggs, G. and Hall, D.: MODIS snow and ice products, and their assessment and applications, in: Land Remote Sensing and Global Environmental Change, Remote Sensing and Digital

- ²⁰ Image Processing, 11, edited by: Ramachandran, B., Justice, C. O., and Abrams, M. J., 681–707, 2011.
 - Sasakawa, M., Shimoyama, K., Machida, T., Tsuda, N., Suto, H., Arshinov, M., Davydov, D., Fofonov, A., Krasnov, O., Saeki, T., Koyama, Y., and Maksyutov, S.: Continuous measurements of methane from a tower network over Siberia, Tellus B, 62, 403–416, doi:10.1111/j.1600-0889.2010.00494.x, 2010.
 - Sasakawa, M., Machida, T., Tsuda, N., Arshinov, M., Davydov, D., Fofonov, A., and Krasnov, O.: Aircraft and tower measurements of CO₂ concentration in the planetary boundary layer and the lower free troposphere over southern taiga in West Siberia: long-term records from 2002 to 2011, J. Geophys. Res.-Atmos., 118, 9489–9498, doi:10.1002/jgrd.50755, 2013.
- ³⁰ Schuur, E. A. G., Bockheim, J., Canadell, J. G., Euskirchen, E., Field, C. B., Goryachkin, S. V., Hagemann, S., Kuhry, P., Lafleur, P. M., Lee, H., Mazhitova, G., Nelson, F. E., Rinke, A., Romanovsky, V. E., Shiklomanov, N., Tarnocai, C., Venevsky, S., Vogel, J. G., and Zimov, S. A.:

Tall Tower Observatory (ZOTTO) in Central Siberia, Atmos. Meas. Tech., 3, 1113–1128, doi:10.5194/amt-3-1113-2010. 2010.

Arctic System Reanalysis domain: surface and upper air analysis, J. Geophys. Res.-Atmos., 116, D11112, doi:10.1029/2010JD015013, 2011. Winderlich, J., Chen, H., Gerbig, C., Seifert, T., Kolle, O., Lavrič, J. V., Kaiser, C., Höfer, A., and Heimann, M.: Continuous low-maintenance CO₂/CH₄/H₂O measurements at the Zotino

Biogeochem. Cy., 2, 399-409, doi:10.1029/GB002i004p00399, 1988. Wilson, A. B., Bromwich, D. H., and Hines, K. M.: Evaluation of Polar WRF forecasts on the

- doi:10.1029/2009JG001283, 2010. Whalen, S. C. and Reeburgh, W. S.: A methane flux time series for tundra environments, Global 25
- NOAA/ESRL Global Greenhouse Gas Reference Network, J. Geophys. Res.-Atmos., 120, 5155-5190, doi:10.1002/2014JD022591, 2015. von Fischer, J. C., Rhew, R. C., Ames, G. M., Fosdick, B. K., and von Fischer, P. E.: 20 Vegetation height and other controls of spatial variability in methane emissions from the Arctic coastal tundra at Barrow, Alaska, J. Geophys. Res.-Biogeo., 115, G00103,
- Sweeney, C., Karion, A., Wolter, S., Newberger, T., Guenther, D., Higgs, J. A., Andrews, A. E., Lang, P. M., Neff, D., Dlugokencky, E., Miller, J. B., Montzka, S. A., Miller, B. R., Masarie, K. A., 15 Biraud, S. C., Novelli, P. C., Crotwell, M., Crotwell, A. M., Thoning, K., and Tans, P. P.: Seasonal climatology of CO₂ across North America from aircraft measurements in the
- Skamarock, W. C., Klemp, J. B., Dudhia, J., Gill, D. O., Barker, D. M., Duda, M. G., Wang, X.-Y., Wang, W., and Powers, J. G.: A Description of the Advanced Research WRF Version 3, Technical Note 475+STR, MMM Division, NCAR, Boulder, CO, USA, 133, 2008.
- Schuur, E. A. G., McGuire, A. D., Schadel, C., Grosse, G., Harden, J. W., Haves, D. J., Hugelius, G., Koven, C. D., Kuhry, P., Lawrence, D. M., Natali, S. M., Olefeldt, D., Romanovsky, V. E., Schaefer, K., Turetsky, M. R., Treat, C. C., and Vonk, J. E.: Climate change and the permafrost carbon feedback, Nature, 520, 171-179, doi:10.1038/nature14338, 2015.

Vulnerability of permafrost carbon to climate change: implications for the global carbon cycle,

Schuur, E. A. G., Vogel, J. G., Crummer, K. G., Lee, H., Sickman, J. O., and Osterkamp, T. E.: The effect of permafrost thaw on old carbon release and net carbon exchange from tundra.

Bioscience, 58, 701–714, doi:10.1641/b580807, 2008.

Nature, 459, 556-559, doi:10.1038/nature08031, 2009.

10

30

5

ACPD

Discussion

Paper

Discussion

Paper

Discussion Paper

Discussion Paper

15, 34871–34911, 2015

Investigating Alaskan methane and carbon dioxide fluxes

A. Karion et al.

- Winderlich, J., Gerbig, C., Kolle, O., and Heimann, M.: Inferences from CO₂ and CH₄ concentration profiles at the Zotino Tall Tower Observatory (ZOTTO) on regional summertime ecosystem fluxes, Biogeosciences, 11, 2055–2068, doi:10.5194/bg-11-2055-2014, 2014.
- Worthy, D., Taylor, C., Dlugokencky, E. J., Chan, E., Nisbet, E. G., and Laurila, T.: Chapter
 6: Long-term monitoring of atmospheric methane, in: AMAP Assessment 2015: Methane as an Arctic climate forcer, Arctic Monitoring and Assessment Programme (AMAP), Oslo, Norway, available at: http://www.amap.no/documents/download/2499 (last access: 3 December 2015), 61–75, 2015.
 - Zhao, C. L. and Tans, P. P.: Estimating uncertainty of the WMO mole fraction scale for carbon dioxide in air, J. Geophys. Res., 111, D08S09, doi:10.1029/2005jd006003, 2006.

10

Zona, D., Gioli, B., Commane, R., Lindaas, J., Wofsy, S. C., Miller, C. E., Dengel, S., Sweeney, C., Karion, A., Chang, R., Henderson, J., Murphy, P., Moreaux, V., Liljedahl, A., Watts, J., Kimball, J., Lipson, D. A., and Oechel, W. C.: Cold season emissions dominate the Arctic tundra methane budget, P. Natl. Acad. Sci. USA, in press, 2015.

Discussion Pa	AC 15, 34871–3	ACPD 15, 34871–34911, 2015 Investigating Alaskan methane and carbon dioxide fluxes A. Karion et al.					
per Discussio	Investigati methane a dioxide A. Karie						
on Paper	Title Abstract	Page Introduction					
Discuss	Conclusions Tables	References Figures					
ion Paper	I◄ ◄ Back	►I ► Close					
Discussion Paper	Full Scree Printer-frier	een / Esc ndly Version					

Table 1. Correlation coefficient (R^2) between ΔCH_4 and ΔCO in each month. Months with $R^2 > 0.2$ are in bold.

	Jan	Feb	Mar	Apr	May	Jun	Jul	Aug	Sep	Oct	Nov	Dec
2012	0.30	0.14	0.12	0.16	0.81	0.00	0.00	0.05	0.17	0.22	0.49	NA
2013	0.66	0.14	0.19	0.14	0.26	0.01	0.07	0.01	0.19	0.09	0.01	0.15
2014	0.49	0.67	0.66	0.03	0.17	0.00	0.07	0.00	0.10	0.06	0.10	0.00

Figure 1. CO_2 and CH_4 measurements from the CARVE tower (red circle, both panels) have high sensitivity to the boreal forests and lowlands of interior Alaska as shown by the 50 % (blue) and 80 % (purple) surface influence contours for the average WRF-STILT influence functions calculated for mid-afternoon averages over the period 2012–2014 (a). The black box indicates the region for the inset shown in (b). Elevation scale is the same in both panels, as indicated on color legend at the right of (b). Elevation data in (a) is from NOAA's National Geophysical Data Center (NGDC, http://www.ngdc.noaa.gov/mgg/topo/report/globedocumentationmanual. pdf). High-resolution elevation data in (b) is from ASTER GDEM, a product of METI and NASA.

Figure 2. Mean monthly diurnal cycle amplitude of hourly averaged CO_2 (top) and CH_4 (bottom). The average over three full calendar years (2012–2014) is shown in black with the gray shading indicating one standard deviation of each month's average. The average diurnal cycles for each individual year are indicated by the blue (2012), green (2013) and red (2014) solid lines.

Figure 3. Magnitude of land surface influence on the tower measurements, in ppm (μ mol m⁻² s⁻¹)⁻¹, for average monthly mid-afternoon footprints from the WRF-STILT model, for 2012 (top), 2013 (middle), and 2014 (bottom). Colors, as indicated in the figure legend, show the average monthly surface influence of Lower Alaska (defined as any part of Alaska south of the Brooks Range, i.e. not part of the North Slope), Canada, Eurasia, and the North Slope of Alaska.

Figure 4. Wind roses for the tower averaged over **(a)** May–September and **(b)** October–April for all three years during mid-afternoon hours, from the 2-D sonic anemometer at the 32 m a.g.l. level of the tower.

Discussion Paper

ACPD

15, 34871-34911, 2015

Investigating Alaskan

Figure 5. Time series of hourly average observed mole fractions (red) and background mole fractions (blue), 1 to 6 p.m. local standard time (LST) only for CH_4 (top), CO_2 (center), and CO (lower) at the CARVE tower. Observations are indicated by solid red lines, while the background mole fractions used for this analysis are shown in blue, and are derived using the particle trajectories from the STILT model and an empirical Pacific boundary curtain, described in the text. The green line represents the value of the same Pacific boundary curtain at the site latitude (65° N) at 3500 ma.s.l., i.e. the free troposphere. The vertical scale for CO has been truncated.

Figure 6. Top: ΔCO_2 observed (i.e. observations minus background, red), along with the modeled ΔCO_2 convolution (blue). Bottom: Monthly average comparisons between the model (blue) and observations (red line and circles) for each year. Error bars on the observations represent the average background uncertainty.

Figure 7. Correlation between observed and modeled ΔCO_2 , for 2012 (left), 2013 (center), and 2014 (right), colored by month. The correlation coefficient R^2 is indicated in each plot, along with the one-to-one line. Data points represent hourly averages between 1 and 6 p.m. LST and filtered according to criteria described in the text.

Figure 8. Optimized model ΔCH_4 (ppb) for the scaled uniform flux (top row) and scaled elevation-based flux map (bottom row), for 2012 (left), 2013 (center), 2014 (right), all plotted against observed ΔCH_4 (mid-afternoon daily averages). Fluxes were scaled to match monthly average observed ΔCH_4 with monthly scaling factors. The correlation coefficient is indicated on each plot, and the 1 : 1 line is shown in red.

Figure 9. Average Alaska monthly CH_4 fluxes for 2012 (blue), 2013 (red), and 2014 (green), estimated based on a uniform Alaska-wide flux scaled to monthly mean observations at the CRV tower. Light blue bars in 2012 indicate monthly fluxes derived from CARVE aircraft observations, with error bars representing the 68 % confidence interval (CI), (calculated by dividing the 95 % CI by 1.96), from Chang et al. (2014). Error bars on the tower-derived fluxes are based on propagating background uncertainty (1-sigma) only. Months for which fluxes were based on six or fewer days were eliminated from the analysis.

