

Gradients of Column CO₂ across North America from the NOAA Global Greenhouse Gas Reference Network

Xin Lan^{1, 2}, Pieter Tans¹, Colm Sweeney^{1, 2}, Arlyn Andrews¹, Andrew Jacobson^{1, 2}, Molly Crotwell^{1, 2}, Edward Dlugokencky¹, Jonathan Kofler^{1, 2}, Patricia Lang¹, Kirk Thoning¹, Sonja Wolter^{1, 2}

¹National Oceanic and Atmospheric Administration, Earth System Research Laboratory, Boulder, 80303, Colorado, USA

²University of Colorado, Cooperative Institute for Research in Environmental Sciences, Boulder, 80309, Colorado, USA

Correspondence to: Xin.Lan (xin.lan@noaa.gov)

Abstract. This study analyzes seasonal and spatial patterns of column carbon dioxide (CO₂) over North America calculated from aircraft and tall tower measurements from the NOAA Global Greenhouse Gas Reference Network from 2004 to 2014. Consistent with expectations, gradients between the eight regions studied are larger below 2 km than above 5 km. The 11-year mean CO₂ dry mole fraction (XCO₂) in the column below ~330 hPa (~8 km above sea level) from NOAA's CO₂ data assimilation model, CarbonTracker (CT2015), demonstrates good agreement with those calculated from calibrated measurements on aircraft and towers. Total column XCO₂ was attained by combining modeled CO₂ above 330 hPa from CT2015 with the measurements. We find large spatial gradients of total column XCO₂ during June to August, and the north and northeast regions have ~3 ppm stronger summer drawdown (variation amplitude in seasonal cycle) than the south and southwest regions. The long-term averaged spatial gradients of total column XCO₂ across North America show a smooth pattern that mainly reflect large-scale circulation patterns. We have conducted a CarbonTracker experiment to investigate the impact of Eurasian long-range transport. The result suggests that the large summer time Eurasian boreal flux contributes about half of the north-south column XCO₂ gradient across North America. Our results confirm that continental-scale total column XCO₂ gradients simulated by CarbonTracker are realistic and can be used to evaluate the credibility of spatial patterns from satellite retrievals, such as the long term average spatial patterns from satellite retrievals reported for Europe which show larger spatial difference (~6 ppm) and scattered hot spots.

1 Introduction

Atmospheric measurements of carbon dioxide (CO₂) from ground and airborne platforms have greatly increased our knowledge of the global carbon cycle. Observations of CO₂, including the NOAA Global Greenhouse Gas Reference Network (GGGRN), initially emphasized ground-based measurements. These observations, started by C.D. Keeling, have monitored the CO₂ trend on both regional and global scales for over 50 years (e.g., Keeling and Rakestraw, 1960; Tans et al., 1989). In addition, the frequency and spatial distribution of airborne measurements have increased rapidly in the last two decades, providing important information about horizontal and vertical variability of atmospheric CO₂ (e.g., Gerbig et al., 2003; Choi et al., 2008; Biraud et al., 2013). Routine aircraft

36 measurements from the NOAA/ESRL GGRN monitor the large-scale distributions of a suite of trace gases,
37 including CO₂, under the influence of continental processes (Sweeney et al., 2015). A very successful approach has
38 been to employ commercial aircraft as a platform for CO₂ measurements, such as Japan's CONTRAIL
39 (Comprehensive Observation Network for TRace gases by AirLiner) project which has provided valuable
40 information for CO₂ in the high troposphere and lower stratosphere (Machida et al., 2002; Machida et al., 2008).
41 Vertical profiles of atmospheric CO₂ reflect the combined influences of surface fluxes and atmospheric mixing.
42 Vertical profiles are particularly useful for evaluating vertical mixing in atmospheric transport models that are used
43 for inverse modeling (e.g. Stephens et al., 2007) to derive estimates of regional- to continental-scale CO₂ sources
44 and sinks (e.g., Tans et al., 1990; Gurney et al., 2002; Gurney et al., 2004; Ciais et al., 2010;).

45 While CO₂ sources and sinks are better constrained at the global scale by global mass balance, it remains
46 challenging to accurately resolve CO₂ sources and sinks at regional-to continental-scale, the apportionment of which
47 depends on relatively minor variations of the observed spatial and temporal patterns of CO₂. When averaging over a
48 few months and longer the largest portion of the variations over continents results from hemispheric-scale terrestrial
49 uptake (photosynthesis)/emissions (respiration) and fossil fuel emissions, while regional net fluxes can make a
50 relatively small contribution to the signal. For example, a simple mass balance argument shows that all U.S. CO₂
51 emissions from fossil fuel burning (~1.4 Pg yr⁻¹) create a total column enhancement of only 0.6 ppm on average in
52 air parcels over the East Coast compared to the West Coast and Gulf Coast if we assume a residence time of the
53 emissions of 5 days to pass the contiguous U.S. (~8×10¹² m²).

54 With careful calibration, air handling, and analysis, the uncertainties of in-situ measurements are less than 0.1
55 ppm. However, in-situ observation networks are sparse in global and regional coverage. Remote sensing data
56 radically increase the number of observations and capture under-sampled regions. It is likely to have a valuable
57 impact on our understanding of the carbon cycle. However, both the precision and the potential of even very small
58 systematic biases in remote sensing measurements need to be carefully evaluated. Vertical profiles from in-situ CO₂
59 measurements have been used to evaluate ground-based total column XCO₂ (X stands for dry mole fraction)
60 determinations, such as those from the Total Carbon Column Observing Network (TCCON) (Washenfelder et al.,
61 2006; Wunch et al., 2010; Messerschmidt et al., 2011; Tanaka et al., 2012). The uncertainty of TCCON total column
62 CO₂ is reported to be 0.4 ppm (1σ) after comparison to aircraft measurements (Wunch et al., 2010). Vertical profiles
63 are also used to evaluate satellite retrievals of total column XCO₂, such as those from the Tropospheric Emission
64 Spectrometer (TES)(Kulawik et al., 2013) and the Greenhouse Gases Observing SATellite (GOSAT) (Inoue et al.,
65 2013, 2016; Saitoh et al., 2016). Satellite retrieval products have known and unknown biases (due to errors in
66 spectroscopy, viewing geometry, spatial differences in clouds and aerosols, surface albedo, etc.) that can result in
67 false horizontal gradients in total column XCO₂ for inverse estimates of sources (Miller et al., 2007; Crisp et al.,
68 2012; Feng et al., 2016). After correction for known biases, the mean GOSAT total column CO₂ (NIES retrievals)
69 biases range between -2.09 to 3.37 ppm (mean = 0.11 ppm, S.D.= 1.11 ppm; 20 out of 27 stations show biases lower
70 than 1 ppm) across different aircraft sites over land, compared with aircraft-based total column XCO₂ (Inoue et
71 al.,2016). By comparing with TCCON, the Orbiting Carbon Observatory-2 (OCO-2) retrieval of total column XCO₂
72 was estimated to have a mean difference less than 0.5 ppm with RMS differences typically below 1.5 ppm after bias

73 correction (Wunch et al., 2016). The overall uncertainty of satellite retrievals is relatively large compared with the
74 total column XCO₂ calculated from in-situ measurements. Total column XCO₂ calculated from vertical profiles from
75 the Japanese CONTRAIL project (Machida et al., 2008) and from the NOAA Carbon Cycle and Greenhouse Gas
76 aircraft program (Sweeney et al., 2015) complemented with simulated profiles from a chemistry–transport model
77 above the maximum altitude of the data have uncertainty less than 1 ppm (Miyamoto et al., 2013). The relatively
78 small uncertainty of the in situ-based total column XCO₂ suggests that they can be used to evaluate satellite
79 retrievals of column averaged CO₂. Since aircraft profiles co-located with satellite retrievals are rare, it is useful to
80 consider the statistics of total column XCO₂ fields derived from repeated aircraft profiles over particular locations.

81 The effect of satellite column averaging kernels and a priori profiles when comparing aircraft-based column
82 XCO₂ with GOSAT retrievals has been assessed by Inoue et al. (2013). For the case considered, application of the
83 averaging kernel and a priori profile to simulate total column XCO₂ was generally within ± 0.1 ppm of the density
84 weighted total column, suggesting that the averaging kernels can only account for small part of the overall
85 uncertainty of the GOSAT total column XCO₂ (Inoue et al., 2013).

86 Transparent and objective estimates of CO₂ sources and sinks derived from atmospheric measurements are
87 paramount for validating emissions reduction efforts and other mitigation policies, and for lowering the uncertainties
88 of carbon cycle-climate feedbacks. The latter are major ambiguities in predicting future climate, such as potential
89 uncontrolled CH₄ and CO₂ emissions from warming permafrost in Arctic regions. Satellite retrievals of total column
90 XCO₂ can significantly improve estimates of sources and sinks only if they are sufficiently precise and accurate (
91 Houweling et al., 2004; Chevallier et al., 2014), meaning that even very small systematic errors (biases) must be
92 eliminated. Here, we analyze the spatial and temporal variability of column CO₂ over North America using well-
93 calibrated CO₂ measurements from aircraft and tall tower, and we use model results from NOAA’s CarbonTracker,
94 version CT2015 (Peters et al. 2007, with updates documented at <http://carbontracker.noaa.gov>) to investigate the
95 primary drivers of variability in total column XCO₂. The aircraft data enable direct analysis of column CO₂
96 characteristics, which is the fundamental step for accurate apportionment of sources and sinks. This study focuses on
97 the long-term averaged column CO₂ gradient and the contributions of different vertical layers to the total column
98 variability. It can serve as a reference for evaluating current and future column CO₂ retrievals from both ground and
99 satellite platforms.

100 **2 Methods**

101 **2.1 Aircraft and tall tower sampling**

102 Aircraft sampling in the NOAA GGGRN intends to provide vertical profiles of long-lived trace gases to capture
103 their seasonal and interannual variability. The aircraft sampling system consists of 12 borosilicate glass flasks in
104 each programmable flask package (PFP), a stainless-steel gas manifold system, and a data logging and control.
105 These flasks (0.7 L each) are pressurized to obtain 2.2 L of sample air from each target altitude. Air samples are then
106 shipped back to NOAA/ESRL for carefully calibrated and quality-controlled measurements. Carbon dioxide is
107 measured using a nondispersive infrared analyzer. Long-term measurements at ~15 sites are carried out using light

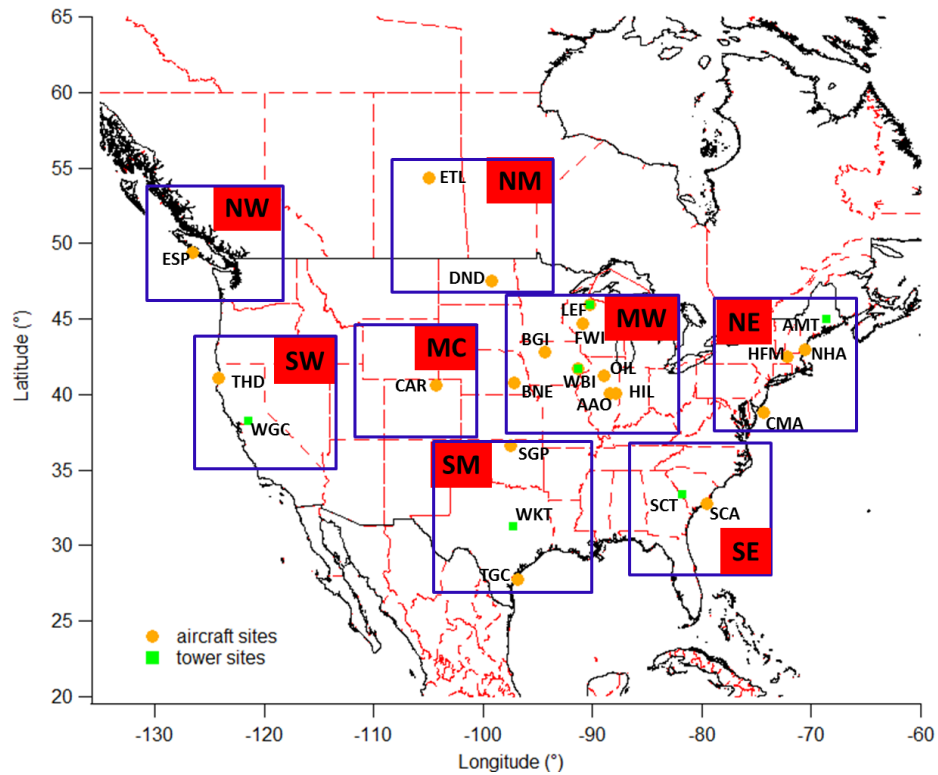
108 aircraft that can reach 8.5 km. Air samples are collected mostly during late morning to early afternoon, when the air
109 mass within the planetary boundary layer (PBL) is generally well mixed, and CO₂ enhancement near the ground
110 from plant respiration during the night has been mixed throughout the boundary layer. Normally, the aircraft follows
111 a pre-decided route such that most samples are collected within 0.1° of the site location. The sampling frequency
112 varies from site to site, currently from twice a month to once every 1.5 months. For more sampling details, quality
113 control discussions, and an evaluation of the sampling frequency, please refer to Sweeney et al. (2015). More
114 information on the aircraft sites can be found at <http://www.esrl.noaa.gov/gmd/ccgg/aircraft/>. We estimate the
115 uncertainty of individual measurements of CO₂ in flask air (68% confidence level) at 0.08 ppm. However, we have
116 seen evidence of positive biases for samples collected using older flasks that may contain contaminants. Andrews
117 et al. (2014) reported biases that increased from <0.1 ppm in 2008 to an average offset in 2013 of 0.36 ppm. The
118 aircraft sampling protocol was modified starting in August 2014 to mitigate this bias. For samples collected prior the
119 protocol change, laboratory tests showed that new/clean flasks have zero bias, but some older/dirty flasks could have
120 biases of > 1 ppm. This bias is not consistent among individual flasks and increasing over time (Andrews et al.,
121 2014), the potential bias is hard to quantify for measurements before August 2014. Thus, the high bias is not
122 corrected in our study. More recently, low bias has been found in PFP measurements when the ambient humidity is
123 high, based on comparisons of PFP measurements with data from in-situ analyzers at tall towers. We are working to
124 understand and quantify this bias, and for this study we have derived a preliminary correction factor, which shows a
125 linear trend with -1.4 ppm CO₂ offset per 1% above 1.7% of ambient water content (in mole fractions versus whole
126 air). Only ~ 4% of total aircraft measurements or ~ 12% of those below 2 km are impacted by humidity higher than
127 1.7%, for which we have applied corrections before data analysis. The mean correction applied is 0.53 ± 0.4 (1 σ)
128 ppm for the impacted data.

129 The NOAA tall tower network measures CO₂ and other trace gases within the continental boundary layer.
130 Continuous in-situ measurements are conducted using nondispersive infrared (NDIR) absorption sensors and cavity
131 ring-down analyzers. The long-term stability of these systems is typically better than 0.1 ppm for CO₂ (Andrews et
132 al., 2014). Most tall tower sites have more than one air intake height. In this study, continuous in-situ measurements
133 from the highest intake are used to minimize potential influences from local sources. More information concerning
134 the tower sites can be found at <http://www.esrl.noaa.gov/gmd/ccgg/insitu/>. For the column XCO₂ calculation, tower
135 data only from 10:00-17:00 local standard time (LST) on flight days are averaged to one data point per day, as a
136 complement to vertical profiles within the PBL.

137 **2.2 Site description**

138 We analyze data from 19 aircraft sites and 6 tall tower sites during 2004 to 2014 (see Table S1 for a summary of site
139 conditions). After considering the geographic distribution of these sites in North America, we group them into eight
140 regions for spatial comparisons (Fig. 1). The northern west (NW) and southern west (SW) regions represent the
141 inflow area in the west coast of US, directly downwind of the Pacific Ocean at both higher elevations. The northern
142 mid-continent (NM) region represents the boreal forest and agriculture region in north-central North America. The
143 mid-continent (MC) region represents a dry landscape due to its high elevation (above 1.5 km on average) and semi-

144 arid climate. The mid-west (MW) region is strongly influenced by agriculture and temperate forest. The southern
 145 mid-continent (SM) represents the south-central humid temperate region, with inflow from the Gulf of Mexico
 146 during summer. The northeast (NE) region represents the temperate forest in north-east coast of U.S., which is
 147 mostly downwind of regions to the west above the PBL, and downwind of its south-west regions within the PBL.
 148 The southeast (SE) region represents the warm temperate region in the south-east coast of U.S.
 149

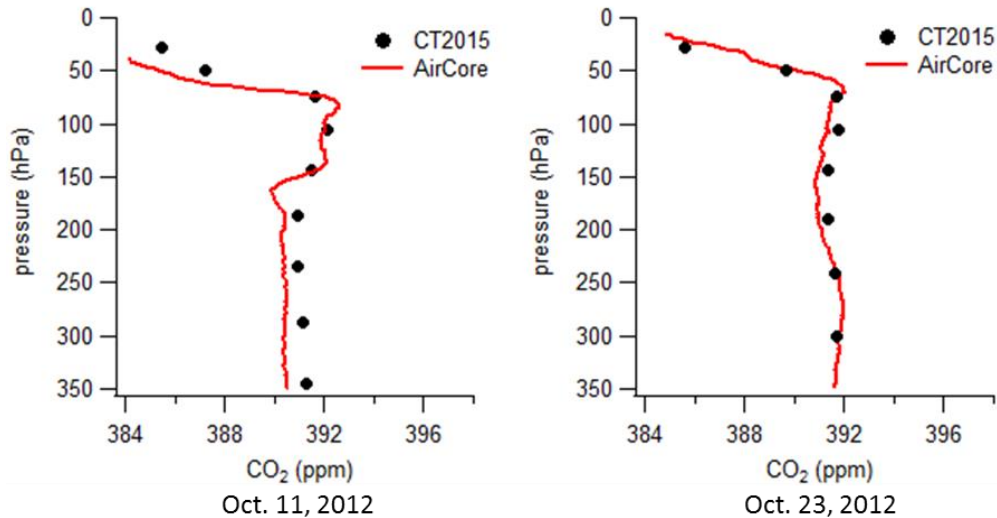


150
 151 **Fig. 1.** Aircraft, tall tower, and high elevation/tower sites in the NOAA GGRN. The eight boxes define regions
 152 that are further discussed for spatial pattern comparison.

153 2.3 Smoothing of the reference data and column XCO₂ calculation

154 We use Mauna Loa Observatory (MLO) as a reference site. The long-term trend of CO₂ measurements from this site
 155 is removed before combining multiple years of data to calculate long-term averages. MLO is located at 19.536°N,
 156 155.576°W, and 3397 m above sea level. Carbon dioxide measurements from this site are widely used to represent
 157 background CO₂ in the Northern Hemisphere. For our study, a function consisting of a quadratic polynomial and
 158 four harmonics is fitted to the MLO data, adopted from the method described by Thoning et al. (1989). Residuals of
 159 the data from this function are smoothed by a low-pass filter with full-width at half-maximum in the time domain of
 160 1.1 years. The smoothed residuals are then added back to the polynomial part of the function to produce the long-
 161 term deseasonalized trend. This trend (see Fig. 3) is subtracted from all aircraft and tall tower measurements, as well
 162 as from CarbonTracker model results (CarbonTracker - MLO deseasonalized trend, CarbonTracker results presented
 163 in this study are the differences relative to observed MLO deseasonalized trend). We use ‘Δ’ to represent detrended

164 data in the following text and figures. The choice of reference site is not important for this study, since we focus on
165 examining the relative seasonal patterns of the detrended spatial and vertical distributions of CO₂ instead of the total
166 changes in CO₂ abundance attributed to global surface fluxes.
167



168
169 **Fig. 2.** Carbon Tracker (CT2015) simulations compared with AirCore in-situ measurements in upper atmosphere.
170 AirCore profiles in the left and right panels are sampled near CAR and SGP, respectively.
171

172 We calculate partial column average CO₂ dry mole fraction using tall tower and aircraft data, and the total
173 column by adding simulations of high altitude CO₂ (above 330 hPa, ~ 8 km above sea level) from CarbonTracker.
174 Since geometric height from the onboard Global Positioning System (GPS) (after 2006) or inferred from the aircraft
175 altimeter or pressure altitude is archived with each aircraft measurement, we first convert geometric height (in
176 meter) to pressure (in hPa) for the pressure-weighted column XCO₂ calculation. This conversion uses geopotential
177 data from NOAA/NCEP North American Regional Reanalysis (NARR) (Mesinger et. al, 2004), available at
178 <https://www.esrl.noaa.gov/psd/data/gridded/data.narr.html>, in which the geopotential is a function of latitude,
179 longitude, pressure altitude and time. We interpolate the geopotential field vertically to retrieve pressure, and then
180 calculate dry pressure by incorporating specific humidity data from NARR. Eventually we use a trapezoidal method
181 to integrate over detrended vertical profiles for dry-pressure-weighted column average. For the long-term averaged
182 column Δ XCO₂ calculation, a long-term mean vertical profile is first constructed for each month by combining 11-
183 year detrended data together and then average data in each 40 hPa vertical bin. To look at the long-term averaged
184 total column Δ XCO₂ from individual aircraft sites, we combine aircraft data with upper-layer CT2015 simulations.

185 The NOAA CarbonTracker model assimilates CO₂ measurements from surface sampling networks and tall
186 towers to generate global 3D fields of atmospheric CO₂ mole fraction. The Carbon Tracker model has evolved
187 significantly since Peters et al. (2007). A detailed description of this model is provided in documents available at
188 <http://carbontracker.noaa.gov>. Our study utilizes CarbonTracker results from the 2015 release (CT2015), publicly
189 accessible at <ftp://aftp.cmdl.noaa.gov/products/carbontracker/co2/CT2015/molefractions/>. This version provides

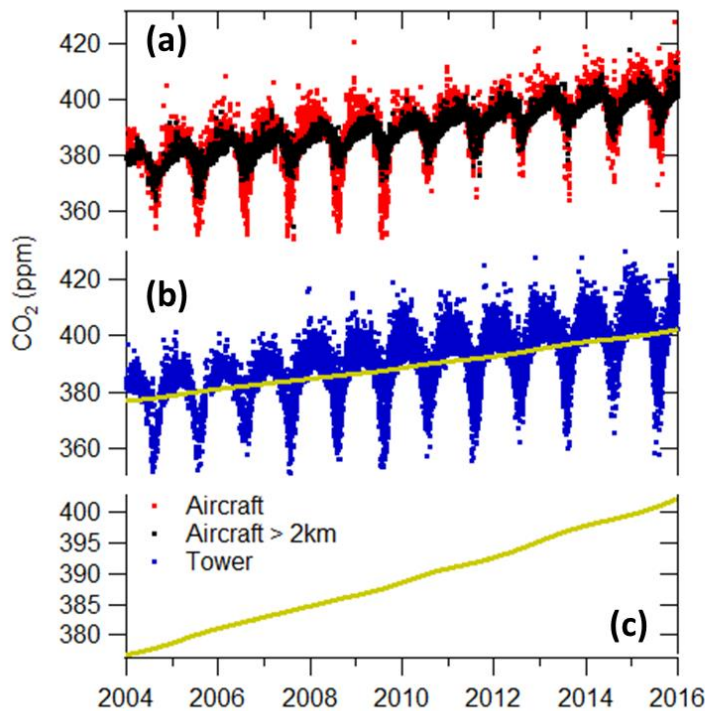
190 CO₂ mole fraction over North America with 1° × 1° spatial and 3 hour temporal resolutions, which are analyzed in
191 Sect. 3.2 and 3.3. Total column CO₂ calculated from CT2015 global data with 3° × 2° spatial resolution is also
192 presented in the supporting information (SI). We have evaluated the performance of CarbonTracker in upper
193 atmosphere (330 to 0 hPa) by comparing its simulations with in-situ measurements from 9 AirCore profiles (Karion
194 et al., 2010) sampled in 2012-2014. AirCore is a ~150 m stainless steel tube that utilizes changes in ambient
195 pressure for passive sampling of the vertical profile. It is released using balloons and it collects a continuous sample
196 as it descends. It is then measured by an analyzer after it is recovered. More information about AirCore system can
197 also be found at <https://www.esrl.noaa.gov/gmd/ccgg/aircore/>. All 9 AirCore profiles are taken near SGP and CAR
198 sites. Figure 2 shows examples of AirCore profiles compared with CT2015 in the upper atmosphere, which
199 demonstrates good agreement. We also compare partial column (330 to 0 hPa) averages from the 9 AirCore profiles
200 and CT2015. Results from CT2015 agree generally well with AirCore, with difference ranging from 0.03 to 1.22
201 ppm (mean value equals 0.66 ppm), which suggests that CT2015 may have a high bias that can contribute to
202 $0.66 \times 1/3 = 0.22$ ppm overestimate on average to the total column average. However, AirCore is in the process of
203 rigorous evaluation, the differences between AirCore and CT2015 are not well characterized yet since we only have
204 a limited amount of AirCore data. It is unclear whether the potential bias of CT2015 in this partial column is
205 dependent on time or sampling location. Adding a constant bias correction to all regions will not change the spatial
206 gradients that we focus on in this study. Thus no correction is applied when using CT2015 simulations to represent
207 the upper 1/3 of the total column.

208 For uncertainty estimates, we use a ‘bootstrap’ method that uses random resampling and repetition of individual
209 vertical profiles (low bias due to high humidity was corrected), with 100 Monte Carlo runs for each column average
210 calculation. Uncertainty is then defined as one standard deviation of the 100 Monte Carlo results.

211 **3 Results and Discussions**

212 **3.1 Seasonal patterns and spatial gradients**

213 Typically one aircraft profile contains measurements at 12 different altitudes. Column ΔXCO_2 can be computed for
214 each profile using the method described in Sect. 2.3 (Fig. S1). Figure 3 shows aircraft (at all altitudes) and tower
215 data (daily averages for 10:00-17:00 LST data) from all sites used in this study. Aircraft data above 2 km exhibit
216 much smaller seasonal variations than the full dataset, because the variations are mainly driven by CO₂ sources and
217 sinks near Earth’s surface. CO₂ concentration is enhanced in the shallow wintertime PBL primarily due to reduced
218 plant photosynthesis and ecosystem respiration combined with slightly increased fossil fuel emissions. During
219 summer the PBL is deeper, and depletions within the PBL are due to strong terrestrial uptake that dominates over
220 emissions especially during June through August. During summer of 2010 to 2012, CO₂ from aircraft measurements
221 appears higher than other years in Fig.3; however, similar characteristics are not present in tower data. This
222 difference is due to a decrease in sampling frequency at several aircraft sites that resulted in an aliased picture of the
223 full summer signals. Since we focus on climatological mean of 11 years of data in our study, this influence is
224 eliminated by combining 11 years of data together into one “average year”.



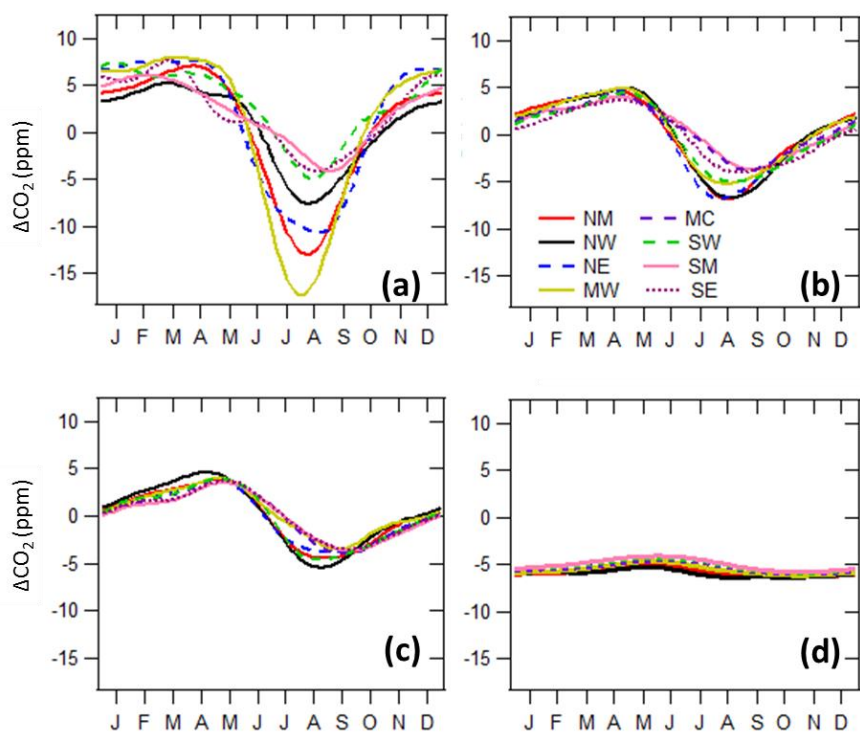
225
 226 **Fig. 3.** CO₂ observations from aircraft (a) and towers (b). The yellow line in (b) illustrates the deseasonalized trend
 227 at Mauna Loa (MLO), same as in (c), in which y-axis expanded.
 228

229 To investigate the contributions of different altitudes to spatial gradients between regions, we divided all
 230 measurement data into three layers according to their sampling altitudes: below 2 km, 2 - 5km, and 5 - 8.5 km masl
 231 (Fig. 4). Smooth seasonal curves are attained from fitting data with four harmonics using the method described by
 232 Thoning et al. (1989). The peak-to-valley amplitudes of the seasonal cycles below 2 km are the largest among the
 233 three layers for most regions, with a minimum of 10.3 ppm in SM and a maximum of 25.0 ppm in MW. The
 234 seasonal variation amplitudes decrease to 7.7-11.5 ppm in the 2 - 5 km layer, and further decrease to 7.2-10.0 ppm in
 235 the 5 - 8.5 km layer. We also observe that the seasonal cycle drawdown occurs later in the layers above 2 km (see
 236 Fig. S2, which provides similar information as Fig. 4, but seasonal curves from different vertical layers are grouped
 237 by regions to facilitate comparisons of the phases of seasonal cycles). The seasonal CO₂ drawdown below 2 km is
 238 mainly influenced by terrestrial photosynthesis and gradients are influenced by local to regional fluxes, with an
 239 earlier onset of drawdown in southern regions than in northern regions. The seasonal cycle aloft is damped and
 240 lagged compared to the PBL, with influences from throughout the Northern Hemisphere and with spatial gradients
 241 likely driven by large-scale transport. The NW, SW, SM, and SE inflow regions have significant delays of more
 242 than one month in the 2 - 5 km layer compared with the surface layer, which is likely due to the delayed phase of the
 243 seasonal cycle in well-mixed air coming from the oceans. Vertical homogeneity of air over ocean was observed
 244 during the HIPPER Pole-to-Pole Observations (HIPPO) aircraft campaign (Wofsy et al., 2011; Frankenberg et al.,

245 2016). As air masses are transported further inland, we observe reduced discrepancies of the timing of CO₂
246 drawdown between surface and upper layer air (2-5 km), which may be associated with the increased influence of
247 the land surface in the mid-troposphere due to strong convection over land. CO₂ drawdown in the 5 - 8.5 km layers
248 also occurs later than in the 2 - 5 km layers in most regions; however, differences between these two layers are
249 small. The declining amplitude and delayed phase of the seasonal cycle with altitude have been noted often (e.g.,
250 Tanaka et al., 1983; Ramonet et al., 2002; Gerbig et al., 2003, Sweeney et al. 2015). It demonstrates that there is lot
251 of important information in the vertical profile that is diminished in observations of the total column.

252 We find that the largest horizontal spatial gradients between regions occur below 2 km during summer time
253 (Fig. 4), with a maximum difference of ~15.5 ppm between MW and SM. SM and SW exhibit less pronounced
254 seasonal cycles, which is likely associated with air masses from the Gulf of Mexico and the Pacific Ocean,
255 respectively, whereas MW exhibits a deep summer drawdown (amplitude in seasonal cycles) partially as a result of
256 strong regional forest and crop uptake. Crevoisier et al. (2010) estimated the surface flux over North America using
257 vertical CO₂ measurements and average wind vectors, and reported that annually averaged land carbon flux at the
258 western (including SW region) and southern regions (including SM region) were neutral. The SE region also
259 demonstrates a less pronounced seasonal cycle with higher summertime levels compared with other northern
260 regions, which may be due to the sea-breeze influence in summer within PBL. In wintertime, CO₂ levels in NE and
261 MW are higher than in other regions, which result from regional fossil fuel and terrestrial biogenic emissions
262 combined with transport from the west and south.

263 Higher altitude data (above 2 km) exhibit only small spatial gradients. In the 2 - 5 km layer, the largest gradient
264 is 4 ppm in summer (Fig. 4b). It further decreases to less than 3 ppm in the 5 - 8.5 km layer (Fig. 4c). Figure 4d
265 shows modeled CO₂ mole fractions from CT2015 for the upper troposphere and above (330 hPa to 0 hPa), which are
266 used to fill in above the aircraft profiles for calculation of total column ΔX_{CO_2} . Spatial gradients in this layer are
267 less than 0.5 ppm, suggesting that the top third of the total column has little contribution to the spatial gradients of
268 the total column.



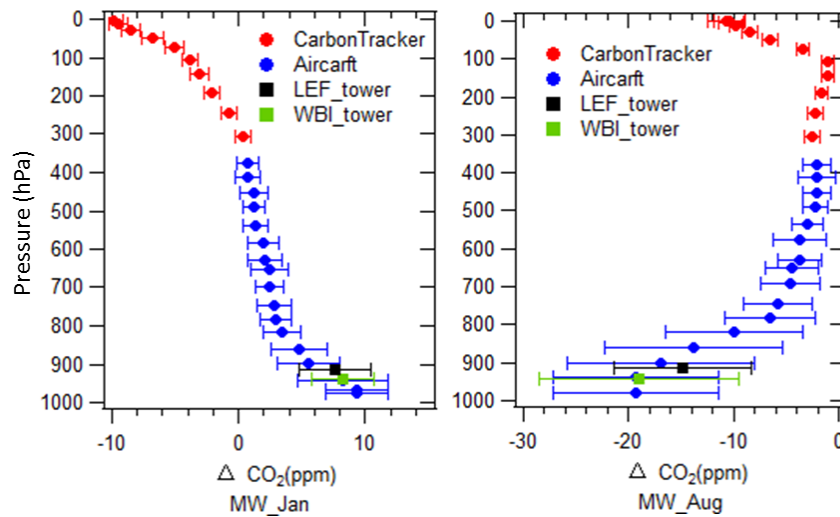
269
 270 **Fig. 4.** Multi-year (2004-2014) average smooth seasonal curves of CO₂ relative to the long-term de-seasonalized
 271 trend at Mauna Loa for different vertical layers: (a). Aircraft and tower data under 2 km, MC is not presented
 272 because only limited data were available due to high surface elevations (>1.5 km on average) in this region; (b).
 273 Aircraft data from 2 - 5 km; (c). Aircraft data from 5 - 8.5 km; (d). CT2015 model results for layers above 330 hPa
 274 (~8.5 km) to 0 hPa (~80 km).

275 3.2 Long-term mean vertical profiles

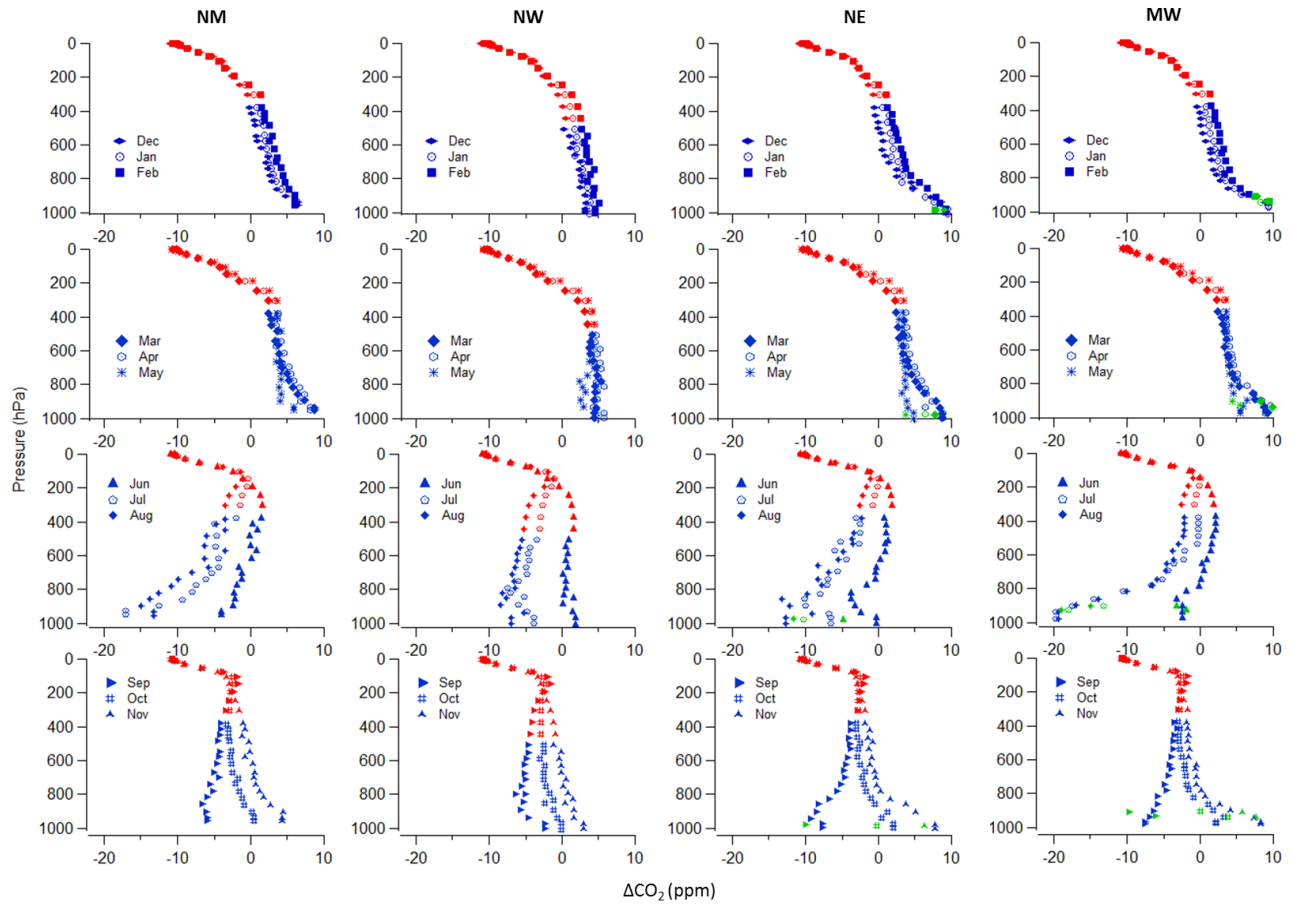
276 To investigate the mean spatial gradients, we first calculate the long-term mean monthly vertical profiles as
 277 described in Sect. 2.3. In addition, each tower serves as one additional layer in the mean profile. The long-term
 278 mean tower data generally fit well in the vertical profiles from measurements of aircraft samples (Fig. 5 and Fig. 6),
 279 suggesting that the biases described in Sect. 2.1 above do not significantly affect the long-term mean. To attain
 280 profiles of the entire atmospheric column, upper layers (330 to 0 hPa) are filled in by CT2015, and the lowest data
 281 point of the measured profile is extended to ground level, defined by the mean surface elevation in that region.

282 Figure 5 presents two examples of long term mean profiles with data variability, which is the one standard
 283 deviation for each 40 hPa bin of aircraft data or for all flight-day tower data. Variability as large as 20 ppm is seen
 284 within the PBL in the MW region in summer, which is due to strong and heterogeneous surface vegetation uptake
 285 and ecosystem respiration combined with day-to-day changes in wind direction. All long-term mean monthly
 286 vertical profiles are presented in Fig. 6, which shows the mean temporal and vertical variability of CO₂ in each
 287 season, and further demonstrates the vertical propagation of seasonal CO₂ due to changes of surface flux. In

288 wintertime, monotonic decrease of CO₂ with altitude can be observed from all regions, in which high PBL CO₂ is
 289 mainly driven by surface emissions and reduced vertical mixing (Denning et al., 1998; Stephens et al., 2007).
 290 Surface CO₂ decreases dramatically in the growing season in those regions influenced by high plant activity, such as
 291 NM and MW regions. For the summer vertical profiles in NE and SE region (east coast of the U.S.), the CO₂ mixing
 292 ratio is elevated in the layer under 900 hPa followed by significant decreases in upper layers until 750 hPa, and then
 293 increases with altitude until tropopause (Fig. 6). This is likely a feature of sea breeze influence. Lower-troposphere
 294 air from the sea, lacking terrestrial uptake of CO₂, typically has higher CO₂ in summer compared with inland air.
 295 Polluted air previously advected offshore can be brought back along with sea breeze. Without significant vertical
 296 mixing over the marine surface, high levels of pollutants remain in those air masses. The convergence of sea breeze
 297 with prevailing wind moving offshore may create a period with a stalled frontal structure that can aggregate air
 298 pollutants (Banta et al., 2005). The convective internal boundary layer structure of the sea breeze system can
 299 significantly reduce mixing height (Miller et al., 2003), and also induces higher CO₂ levels. When the sea breeze is
 300 not dominant, air advected from southwest and west (the land) can also bring in polluted air with high CO₂ since this
 301 region is downwind of continental U.S. emissions (Miller et al., 2012).

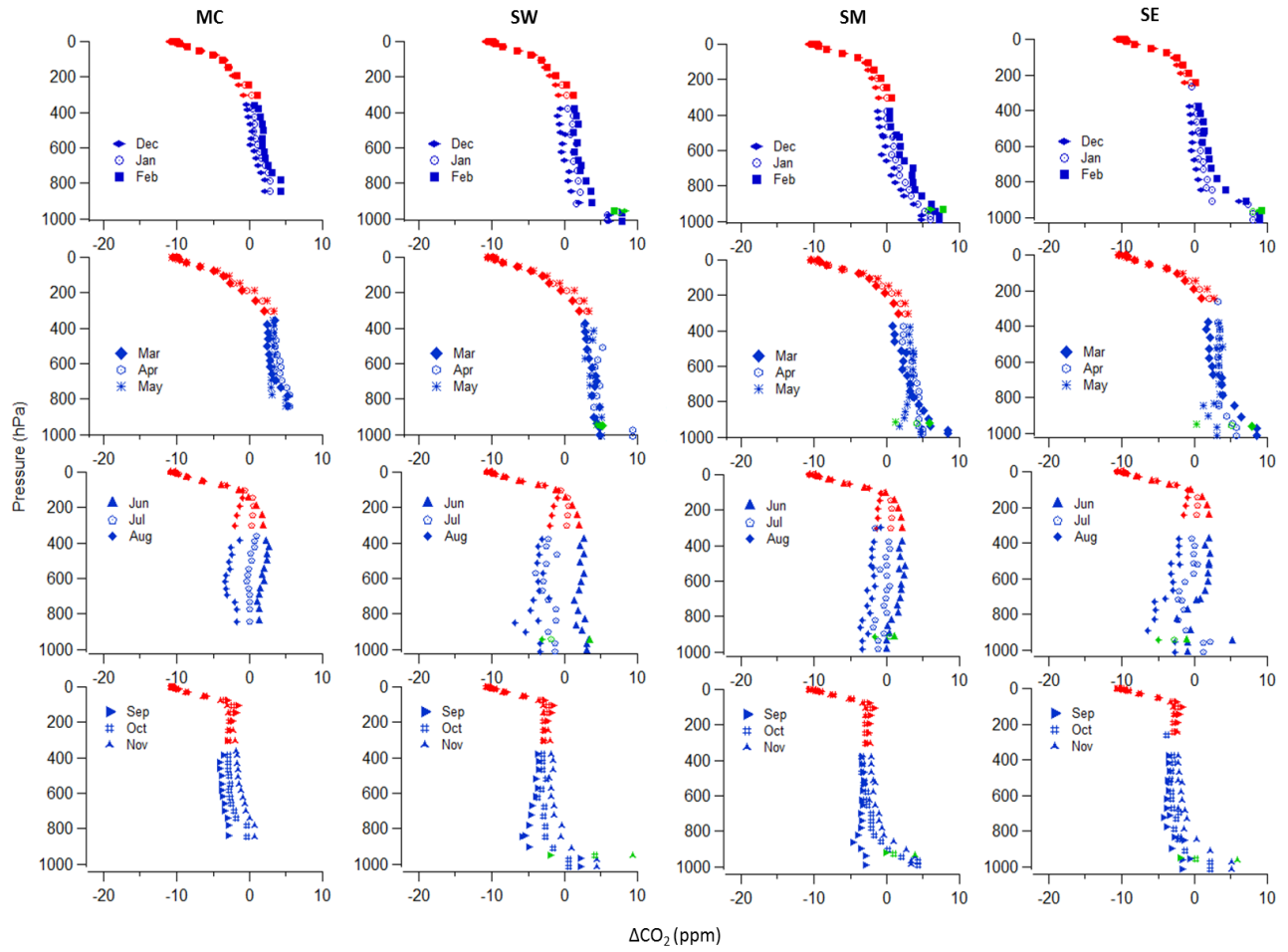


302
 303 **Fig. 5.** Long-term mean (2004-2014) average vertical profiles in January (left panel) and August (right panel) in
 304 region MW. Error bar shows one standard deviation.



305

306 **Fig. 6a.** Long-term mean (2004-2014) monthly vertical profiles in NM, NW, NE, MW (by column, from left to
 307 right). Blue points were calculated from observations, red points were calculated from CT2015, and green points
 308 were calculated from tower data.



309

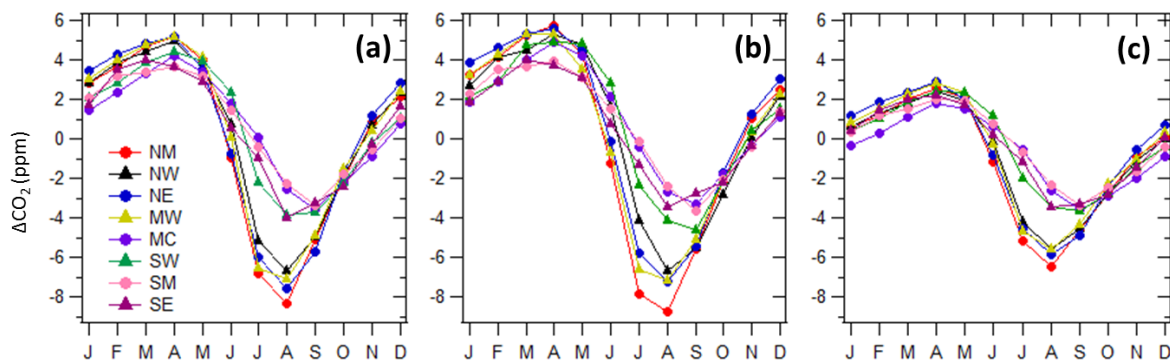
310 **Fig. 6b.** Long-term mean (2004-2014) monthly vertical profiles in MC, SW, SM, SE (by column, from left to right).

311 Blue points were calculated from observations, red points were calculated from CT2015, and green points were
 312 calculated from tower data.

313

314 **3.3 Partial column ΔXCO_2 and total column ΔXCO_2**

315 Seasonal variations of monthly averaged partial column ΔXCO_2 (below 330 hPa) demonstrate maximum values in
316 April and minimum values in August or September (Fig. 7a). The largest amplitude appears in NM, with peak-to-
317 valley difference up to 13.5 ppm. SW, SM, SE, and MC have similar amplitudes of 7-8 ppm, smaller than other
318 regions. To evaluate the performance of CT2015 on column ΔXCO_2 , CT2015 results are sampled to match the
319 latitude, longitude, altitude and time of actual measurements. Note that aircraft profiles are not assimilated in
320 CT2015, so aircraft data are independent of the CT2015 data assimilation. Figure 7b shows monthly partial columns
321 of ΔXCO_2 calculated from CT2015, which demonstrate good agreement with results from measurements. Only
322 small seasonal biases exist in CT2015, with high bias occurring mostly in spring and early summer and low bias in
323 September and October (Fig. S3). The overall differences of monthly partial column ΔXCO_2 (CT2015 -
324 measurements) mainly fall in the range of -0.64 ppm (5th percentile) to 0.84 ppm (95th percentile) with a mean
325 difference of 0.13 ppm. These differences are of similar magnitude to the uncertainties of partial column ΔXCO_2
326 calculated from the measurements (Fig. S4). It is clear that CT2015 captures the long-term mean variations of both
327 phase and amplitude of partial column XCO_2 reasonably well when compared with well-calibrated measurements
328 across North America.
329

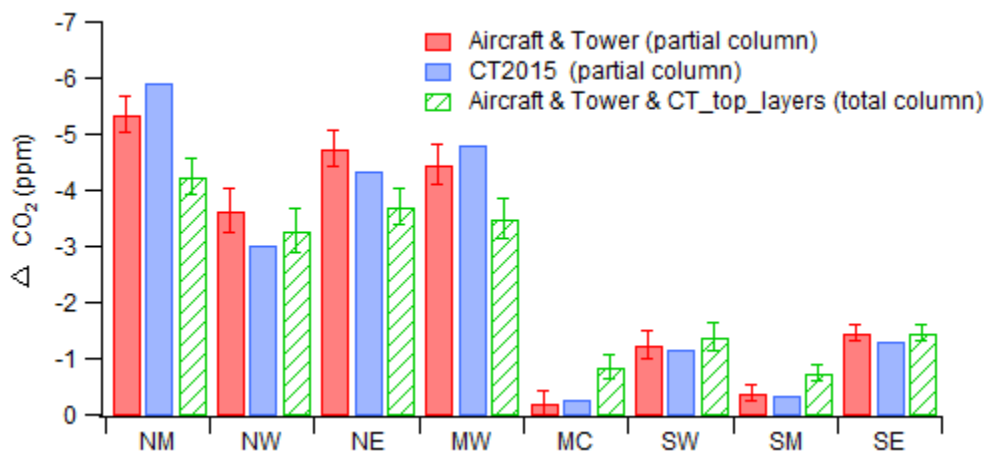


330
331 **Fig. 7.** (a). Partial column ΔXCO_2 calculated from aircraft and tower data; (b). Partial column ΔXCO_2 calculated
332 from CT2015; (c). Total column ΔXCO_2 calculated from aircraft and tower data and including the top layer data
333 from CT2015.

334

335 Total column ΔXCO_2 is presented in Fig. 7c. In regions NW, NM, NE, and MW, seasonal variations of total
336 column ΔXCO_2 are very similar in both phase and amplitude (8-9 ppm peak to valley). For SW, SM, SE, and MC,
337 amplitudes are ~5.5 ppm. The smallest spatial gradients occur during May and October, which result in maximum
338 differences among all regions of only 0.9 and 0.7 ppm, respectively. The largest spatial gradients occur during June,
339 July and August, which result in maximum differences of 2.4, 4.5, and 4.1 ppm, respectively. It is interesting that the
340 deepest drawdown (largest amplitude in seasonal cycles) is seen in region NM, not in region MW that encompasses

341 the very intensive agricultural activities in the U.S. mid-west, which suggest the possibility of strong upwind
 342 influence in the NM region. Transported signals have significant influences on total column CO₂. The summer total
 343 column ΔXCO₂, represented by the June to August average from CT2015, has a magnitude that is similar to
 344 observations with differences no more than 1 ppm (Fig. 8). Based on the seasonal patterns of total column ΔXCO₂
 345 (Fig. 7c) and the summer column ΔXCO₂ (Fig. 8), we can separate the eight regions into two groups. The group with
 346 NW, NM, NE, and MW, has ~3 ppm stronger drawdown (larger amplitude) than the group with SW, SM, SE, and
 347 MC. For winter total column ΔXCO₂ (December to February average), the maximum spatial difference is only 1.6
 348 ppm, with the highest total column ΔXCO₂ of 1.2 ppm in NE and the lowest value of -0.3 ppm in MC.
 349



350
 351 **Fig. 8.** Long-term mean (2004-2014) June to August partial and total column ΔXCO₂. Error bars represent one
 352 standard deviation from the bootstrap uncertainty calculation (see Sect. 2.3).
 353

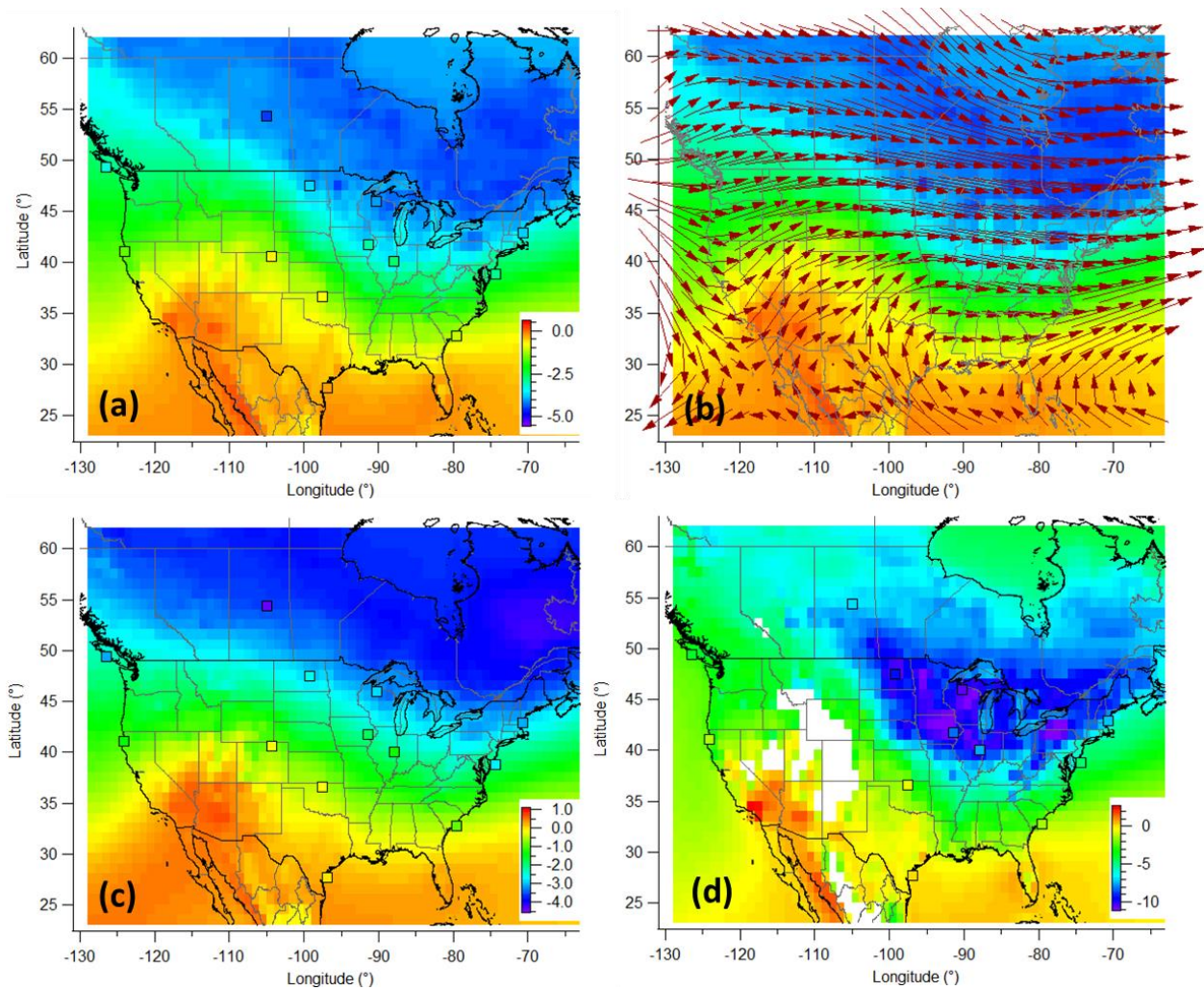
354 3.4 Influence of large scale circulation

355 Figure 9 shows long-term mean summer column ΔXCO₂ calculated from CT2015, together with full column
 356 ΔXCO₂ from individual aircraft sites (note that some aircraft sites have less than 11 years of data the CT2015 shows
 357 in Fig. 9, only aircraft sites with more than 6 years of data are presented, the actual values are provided in Table S2).
 358 The fact that total column ΔXCO₂ from CT2015 agrees well with aircraft sites also supports the performance of
 359 CT2015 on a long-term average basis. The observations show a similar summer spatial pattern, with lower column
 360 ΔXCO₂ in the north and northeast regions and higher column ΔXCO₂ in the south and southwest regions (Fig. 9a).
 361 Scattered hot spots of high column ΔXCO₂ associated with surface emissions from megacities, or cold spots
 362 associated with strong local uptake, are not or just barely visible in the long-term average column ΔXCO₂ map at
 363 1°x1° resolution. Instead, the wave-like pattern of column ΔXCO₂ over North America reflects large scale
 364 circulation. To support our hypothesis on the influence of large scale circulation, we analyze the long term mean
 365 wind pattern over North America. We can see that air masses from northwest of the continent bring in low average
 366 column ΔXCO₂, while air masses from the south (mainly the subtropical Pacific Ocean and the Gulf of Mexico)

367 bring in high column ΔXCO_2 (Fig. 9b). The zonal gradients over the continent, especially north of $40^\circ N$, also
368 reflect long-term average wind patterns; southwest wind corresponds to higher column ΔXCO_2 over the western part
369 of the continent until the wind direction shifts to west-northwest over the eastern part of the continent. This wind
370 pattern matches well with the geographic division of the over/under -3 ppm areas colored in green/blue in the
371 column ΔXCO_2 map (Fig. 9b). Figure 9c and 9d shows partial column averages for free troposphere (800-330 hPa)
372 and lower troposphere (below 800 hPa), respectively. The free troposphere spatial gradient also demonstrates a
373 wave-like pattern. A previous study on the total column CO_2 from ground based Total Carbon Column Observation
374 Network (TCCON) found strong correlation between the mid-latitude column CO_2 and synoptic-scale variation of
375 potential temperature (θ , at 700 hPa), a dynamic tracer for adiabatic air transport (Keppel-Aleks et al., 2012). Thus
376 they also propose that the variations in column CO_2 are mainly driven by large-scale flux and transport. Analysis on
377 the interannual variability of the seasonal cycle amplitudes of column CO_2 in North Hemisphere has also found
378 significant contribution of large-scale circulations to the north-south gradient (Wunch et al., 2013).

379 The strong drawdown over northeast North America in summer is a consequence of long-range transport of low
380 CO_2 from northeast Eurasia, in addition to regional terrestrial uptake. Sweeney et al. (2015) notes well-mixed
381 vertical profiles (up to 8 km) of CO_2 , CO, CH_4 , N_2O , and SF_6 from THD, ESP and PFA (Poker Flat, Alaska; 65.07° ,
382 -147.29°) sites and suggests that air coming across the Pacific was strongly influenced by Asian surface fluxes
383 before being vertically homogenized as it passed over the Pacific Ocean. This well-mixed air forms an important
384 boundary condition in the column CO_2 of air coming into the North American continent. This was best illustrated at
385 sites like PFA where the summertime minimum in CO_2 significantly preceded maximum ecosystem uptake of CO_2 ,
386 implying significant influence of transported air from lower latitude regions from Asia. We further conduct an
387 experiment using Carbon Tracker to investigate the importance of this effect. A control run and a “masked run” are
388 conducted for 2010-2012, in which the Eurasian boreal flux is turned on/off. The MLO CO_2 trend from each model
389 scenario is used as reference background and thus removed before total column ΔXCO_2 calculation. Figure 10 shows
390 the results for 2012 summer, which is an average summer when compared with the 2004-2014 mean pattern (Fig. 9
391 and Fig. 11). The maximum north-south difference reduces to ~ 2.5 ppm after we turn off the Eurasian boreal flux,
392 compared with ~ 5 ppm from the control run. In both control and masked scenarios, the free troposphere partial
393 ΔXCO_2 demonstrates similar spatial patterns as for total column ΔXCO_2 (Fig. S5). This result combined with results
394 from Sweeney et al. (2015) demonstrates that the transport of low CO_2 resulting from large summertime Eurasian
395 boreal uptake has a large contribution on the overall summer total column CO_2 drawdown in North America.

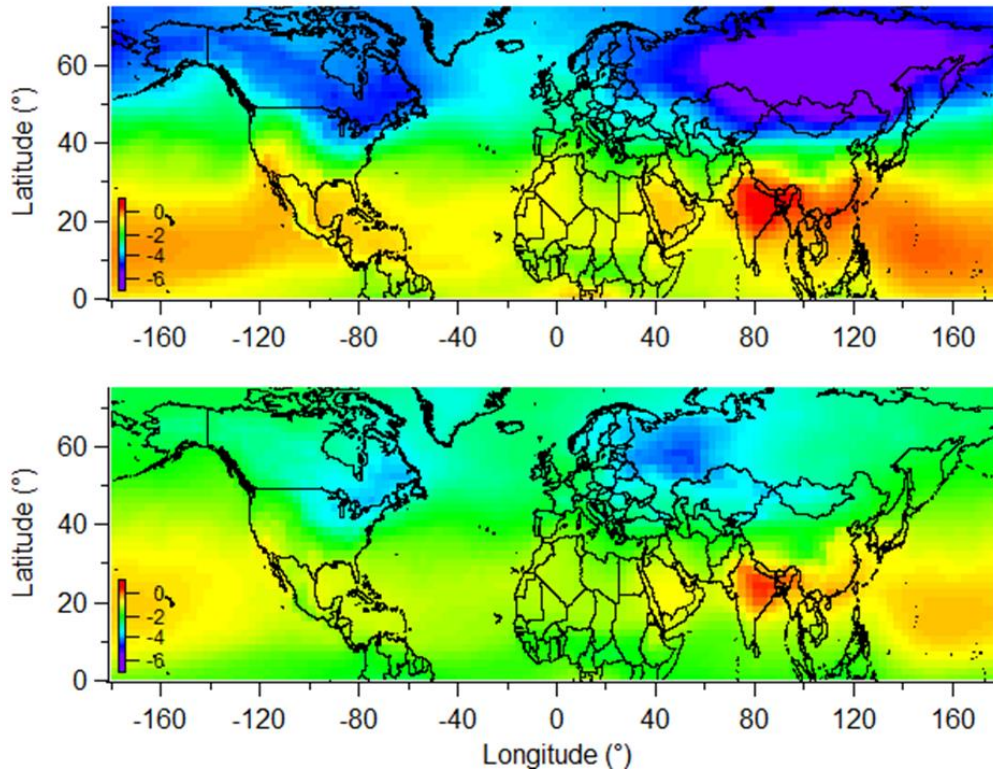
396 |
397



398

399 **Fig. 9.** Long-term mean (2004-2014) June-August total column ΔXCO_2 from CT2015 in $1^\circ \times 1^\circ$ spatial resolution
 400 with total column ΔXCO_2 for 13 individual aircraft sites in squares (a), and CT2015 column ΔXCO_2 overlaid with
 401 pressure-weighted (1000 hPa to 500 hPa) mean wind vectors for the same period (b). (c) and (d) are similar as (a),
 402 except for free troposphere (800 to 330 hPa) and lower troposphere (below 800 hPa), respectively. Note the different
 403 color scales.

404



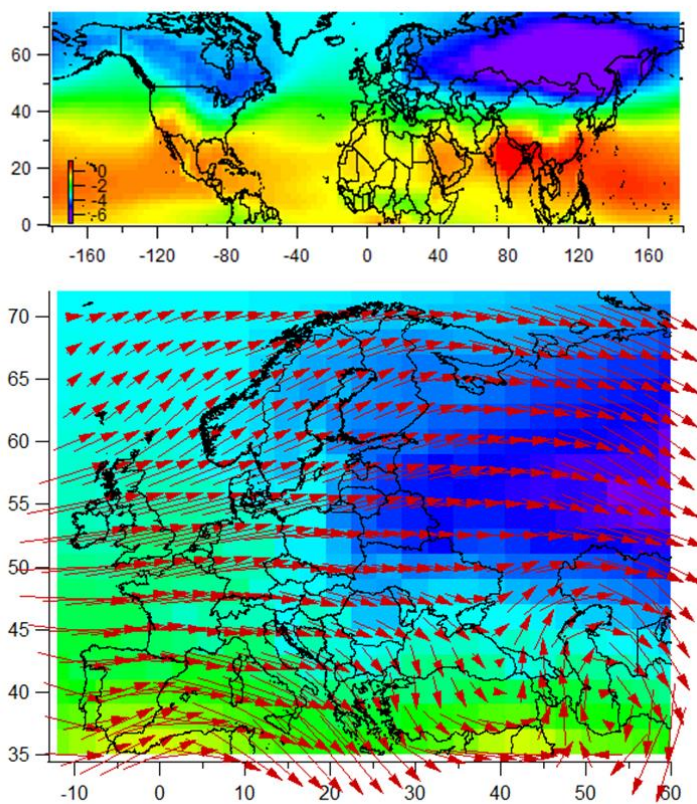
405

406 **Fig. 10.** Total column ΔXCO_2 from Carbon Tracker control (top panel) and masked (bottom panel, Eurasian boreal
 407 flux is masked) runs for 2012 June-August ($3^\circ \times 2^\circ$ spatial resolution). MLO trend from each individual scenario is
 408 removed before the ΔXCO_2 calculation. Same color scale is used as in Fig. 9a. Partial column ΔXCO_2 patterns for
 409 free troposphere (800 to 330 hPa) and lower troposphere (below 800 hPa) are provided in SI.

410 3.5 A comparison with apparent gradients over Europe

411 Figure 11 shows the climatological June - August mean modeled global column ΔXCO_2 map in $3^\circ \times 2^\circ$ spatial
 412 resolution, which presents smooth wave-like patterns. Reuter et al. (2014) use SCIAMACHY and GOSAT satellite
 413 retrievals of column CO_2 and inverse modelling to attain surface CO_2 flux over European region, and suggest a large
 414 uptake of CO_2 in this region. Column ΔXCO_2 from CT2015 (Fig. 11) exhibits a drastically different summer spatial
 415 pattern over Europe compared with the eight year mean (2003-2010) June through August satellite retrievals
 416 presented by Reuter et al. (2014, their Fig. 2a). The spatial gradient from CT2015 results in a maximum 3-4 ppm
 417 difference and a gradual pattern, instead of as much as 6 ppm from satellite retrievals. There is no sign of XCO_2 hot
 418 spots from surface emissions or removals in the CT2015 spatial pattern over Europe (Fig. 11), in contrast to several
 419 hot spots that are apparent from the 8-year averaged SCIAMACHY satellite retrievals over Ireland, U.K., northeast
 420 of France, Belgium, Netherland, north of Germany, and south of Sweden, and low spots over the Ukraine and
 421 Kazakhstan (Reuter et al., 2014). Although the NOAA/ESRL CT2015
 422 (<https://www.esrl.noaa.gov/gmd/ccgg/carbontracker/CT2015/>) assimilates fewer observations over Europe than
 423 Carbon Tracker Europe (<http://www.carbontracker.eu/>), both models produced similar fluxes over the European

424 region (see both websites for detailed fluxes). The $3^{\circ} \times 2^{\circ}$ grid from CT2015 is not likely responsible for a much
 425 smoother pattern for Carbon Tracker, compared with the $2^{\circ} \times 2^{\circ}$ grid from satellite retrievals (Reuter et al., 2014) .
 426 The North America region on the $3^{\circ} \times 2^{\circ}$ grid in Fig. 11 shows similar pattern as the $1^{\circ} \times 1^{\circ}$ grid in Fig. 9, with
 427 similar spatial difference of ~ 5 ppm. A smoother spatial distribution should be expected in Europe for the long-term
 428 mean column XCO_2 (Fig. 11) due to the influences of dominating west and southwest winds in summer. Since the
 429 satellite retrievals in Reuter et al. (2014) appear to show unrealistic column XCO_2 spatial gradients over Europe,
 430 they should not be used to derive estimates of a European carbon sink. A recent study (Feng et al., 2016) using
 431 inverse modeling suggests that satellite retrievals outside the immediate European region and a small bias of 0.5
 432 ppm were sufficient to produce the apparent large carbon sink in the study of Reuter et al. (2014). This is expected
 433 from elementary mass balance considerations as in Sec.1. Spatial gradients are the fundamental signals to infer
 434 regional fluxes. Since spatial gradients from CT2015 are realistic, boreal fluxes inferred by CT2015, which shows
 435 0.03 ± 2.33 Pg C yr⁻¹ for Europe, should be more trustworthy than fluxes estimated based on unrealistic spatial
 436 pattern. However, the European carbon sink is still inclusive; the discrepancies among different methods and results
 437 are further discussed by Reuter et al. (2017). Increasing the amount of highly precise observations such as the well-
 438 calibrated surface measurements and vertical profiles can greatly help to estimate the carbon sink.
 439
 440



441
 442 **Fig. 11.** Long-term mean (2004-2014) June - August total column ΔXCO_2 from CT2015 (top panel) in $3^{\circ} \times 2^{\circ}$
 443 spatial resolution, and zoom-in for Europe overlaid with pressure-weighted (1000 hPa to 500 hPa) mean wind

444 vectors for the same period (bottom panel). The color scale is the same as in Fig. 9a, which is scaled to reflect 6
445 ppm difference of XCO₂ to compare with satellite retrievals from Reuter et al. (their Fig. 2a, 2014).

446 **4 Conclusion**

447 Aircraft and tall tower measurements from the NOAA GGGRN provide detailed information describing the long-
448 term average temporal and spatial variations of CO₂ in the PBL and the free troposphere. These data provide
449 valuable constraints for evaluating model simulations and satellite retrievals. Seasonal cycle peak-to-peak
450 amplitudes of CO₂ are largest below 2 km, where those maximum values are about twice those in the vertical layers
451 above, indicating that most of the information on surface sources and sinks resides in the continental PBL. Large
452 spatial gradients of CO₂ over North America are observed below 2 km during summer (with maximum difference of
453 ~15.5 ppm between MW and SM), while higher altitude data (above 2 km) have much smaller contributions to
454 spatial gradients, with a maximum difference of 4 ppm. The spatial differences of CO₂ in the upper troposphere and
455 above (330 hPa to 0 hPa) are less than 0.5 ppm, according to CT2015. Comparison with Aircore measurements
456 shows that the upper troposphere and lower stratospheric simulations from CT2015 are reasonably trustworthy.

457 Our long-term mean vertical profiles show that tower data agree well with aircraft data at similar vertical levels.
458 Partial column Δ XCO₂ was calculated from the long-term mean vertical profiles. By comparing the partial column
459 Δ XCO₂ from measurements with those from CT2015, we verify that CT2015 captures the long-term mean patterns
460 of both phase and amplitude of partial Δ XCO₂.

461 Large spatial gradients of Δ XCO₂ only appeared in summer, during which time the north and northeast regions
462 had ~3 ppm stronger drawdowns than the south and southwest regions. Scattered hot spots of high column Δ XCO₂
463 associated with surface emissions from megacities, or cold spots associated with strong local uptake, are not or just
464 barely visible in the long-term average column Δ XCO₂. Instead, the wave-like pattern of column Δ XCO₂ over North
465 America matches well with large scale circulation. A CarbonTracker experiment to investigate the impact of
466 Eurasian long-range transport suggests that the large summer time Eurasian boreal flux alone contributes about half
467 of the north-south column Δ XCO₂ gradient across North America. Considering the transported signals from other
468 upwind regions, including northern Canada, we expect that the transported signals overall has major contribution to
469 the column Δ XCO₂ spatial gradient.

470 **Author contributions**

471 Xin Lan was responsible for study design, data analysis, and manuscript writing. Pieter Tans was responsible for
472 study design, data analysis, and manuscript improvement. Colm Sweeney and Arlyn Andrews provided
473 measurement data and improved manuscript. Andrew Jacobson provided modelled data and improved manuscript.
474 Edward Dlugokencky analyzed measurements and ensured data quality, and improved manuscript. Jonathan Kofler
475 conducted tower measurements and improved manuscript. Molly Crotwell, Patricia Lang, and Sonja Wolter
476 analyzed measurements and ensured data quality. Kirk Thoning provided data smoothing method.

477 **Acknowledgements**

478 We especially thank John Mund for extracting NARR meteorological variables for our measurements. This
479 research was supported by a fellowship from the National Research Council Research Associateship Programs.

480 **References**

- 481 Andrews, A. E., Kofler, J. D., Trudeau, M. E., Williams, J. C., Neff, D. H., Masarie, K. A., Chao, D. Y., Kitzis, D.
482 R., Novelli, P. C., Zhao, C. L., Dlugokencky, E. J., Lang, P. M., Crotwell, M. J., Fischer, M. L., Parker, M. J.,
483 Lee, J. T., Baumann, D. D., Desai, A. R., Stanier, C. O., De Wekker, S. F. J., Wolfe, D. E., Munger, J. W. and
484 Tans, P. P.: CO₂, CO, and CH₄ measurements from tall towers in the NOAA Earth System Research
485 Laboratory's Global Greenhouse Gas Reference Network: instrumentation, uncertainty analysis, and
486 recommendations for future high-accuracy greenhouse gas monitoring efforts, *Atmos. Meas. Tech.*, 7, 647-687,
487 2014.
- 488 Banta, R. M., Senff, C. J., Nielsen-Gammon, J., Darby, L. S., Ryerson, T. B., Alvarez, R. J., Sandberg, S. R.,
489 Williams, E. J. and Trainer, M.: A bad air day in Houston', *B. Am. Meteorol. Soc.*, 86, 657, DOI:
490 <http://dx.doi.org/10.1175/BAMS-86-5-657>, 2005.
- 491 Biraud, S. C., Torn, M. S., Smith, J. R., Sweeney, C., Riley, W. J. and Tans, P. P.: A multi-year record of airborne
492 CO₂ observations in the US Southern Great Plains, *Atmos. Meas. Tech.*, 6, 751-763, 2013.
- 493 Buchwitz, M., Reuter, M., Bovensmann, H., Pillai, D., Heymann, J., Schneising, O., Rozanov, V., Krings, T.,
494 Burrows, J. P., Boesch, H., Gerbig, C., Meijer, Y. and Loscher, A. : Carbon Monitoring Satellite (CarbonSat):
495 assessment of atmospheric CO₂ and CH₄ retrieval errors by error parameterization, *Atmos. Meas. Tech.*, 6,
496 3477-3500, 2013.
- 497 Chevallier, F., Breon, F. M. and Rayner, P. J.: Contribution of the Orbiting Carbon Observatory to the estimation of
498 CO₂ sources and sinks: Theoretical study in a variational data assimilation framework, *J. Geophys. Res.*
499 *Atmos.*, 112, D09307, doi:10.1029/2006JD007375, 2007.
- 500 Choi, Y. H., Vay, S. A., Vadrevu, K. P., Soja, A. J., Woo, J. H., Nolf, S. R., Sachse, G. W., Diskin, G. S., Blake, D.
501 R., Blake, N. J., Singh, H. B., Avery, M. A., Fried, A., Pfister, L. and Fuelberg, H. E.: Characteristics of the
502 atmospheric CO₂ signal as observed over the conterminous United States during INTEX-NA, *J. Geophys. Res.*
503 *Atmos.*, 113, D07301, doi:10.1029/2007JD008899, 2008.
- 504 Ciais, P., Rayner, P., Chevallier, F., Bousquet, P., Logan, M., Peylin, P. and Ramonet, M.: Atmospheric inversions
505 for estimating CO₂ fluxes: methods and perspectives, *Climatic Change*, 103, 69-92, 2010.
- 506 Conway, T. J., Tans, P. P., Waterman, L. S. and Thoning, K. W.: Evidence for interannual variability of the carbon-
507 cycle from the national-oceanic-and-atmospheric-administration climate-monitoring-and-diagnostics-laboratory
508 global-air-sampling-network, *J. Geophys. Res. Atmos.*, 99, 22831-22855, 1994.
- 509 Crevoisier, C., Sweeney, C., Gloor, M., Sarmiento, J. L. and Tans, P. P.: Regional US carbon sinks from three-
510 dimensional atmospheric CO₂ sampling, *Proc. Natl. Acad. Sci. U. S. A.*, 107, 18348-18353, 2010.

511 Crisp, D., Fisher, B. M., O'Dell, C., Frankenberg, C., Basilio, R., Bosch, H., Brown, L. R., Castano, R., Connor, B.,
512 Deutscher, N. M., Eldering, A., Griffith, D., Gunson, M., Kuze, A., Mandrake, L., McDuffie, J.,
513 Messerschmidt, J., Miller, C. E., Morino, I., Natraj, V., Notholt, J., O'Brien, D. M., Oyafuso, F., Polonsky, I.,
514 Robinson, J., Salawitch, R., Sherlock, V., Smyth, M., Suto, H., Taylor, T. E., Thompson, D. R., Wennberg, P.
515 O., Wunch, D., and Yung, Y. L.: The ACOS CO₂ retrieval algorithm – Part II: Global XCO₂ data
516 characterization, *Atmos. Meas. Tech.*, 5, 687–707, doi:10.5194/amt-5-687-2012, 2012.

517 Dee, D. P., Uppala, S. M., Simmons, A. J., Berrisford, P., Poli, P., Kobayashi, S., Andrae, U., Balmaseda, M. A.,
518 Balsamo, G., Bauer, P., Bechtold, P., Beljaars, A. C. M., van de Berg, L., Bidlot, J., Bormann, N., Delsol, C.,
519 Dragani, R., Fuentes, M., Geer, A. J., Haimberger, L., Healy, S. B., Hersbach, H., Holm, E. V., Isaksen, L.,
520 Kallberg, P., Kohler, M., Matricardi, M., McNally, A. P., Monge-Sanz, B. M., Morcrette, J. J., Park, B. K.,
521 Peubey, C., de Rosnay, P., Tavolato, C., Thepaut, J. N. and Vitart, F.: The ERA-Interim reanalysis:
522 configuration and performance of the data assimilation system, *Q.J.R. Meteorol. Soc.*, 137, 553-597, 2011.

523 Denning, A. S., Takahashi, T., and Friedlingstein, P.: Can a strong atmospheric CO₂ rectifier effect be reconciled
524 with a “reasonable” carbon budget?, *Tellus B*, 51, 249–253, 1999.

525 Feng, L., Palmer, P. I., Parker, R. J., Deutscher, N. M., Feist, D. G., Kivi, R., Morino, I. and Sussmann, R.:
526 Estimates of European uptake of CO₂ inferred from GOSAT X-CO₂ retrievals: sensitivity to measurement bias
527 inside and outside Europe, *Atmos. Chem. Phys.*, 16, 1289-1302, 2016.

528 Frankenberg, C., Kulawik, S. S., Wofsy, S., Chevallier, F., Daube, B., Kort, E. A., O'Dell, C., Olsen, E. T., and
529 Osterman, G.: Using airborne HIAPER Pole-to-Pole Observations (HIPPO) to evaluate model and remote
530 sensing estimates of atmospheric carbon dioxide, *Atmos. Chem. Phys.*, 16, 7867–7878, 2016. doi:10.5194/acp-
531 16-7867-2016, 2016.

532 Gerbig, C., Lin, J. C., Wofsy, S. C., Daube, B. C., Andrews, A. E., Stephens, B. B., Bakwin, P. S. and Grainger, C.
533 A.: Toward constraining regional-scale fluxes of CO₂ with atmospheric observations over a continent: 1.
534 Observed spatial variability from airborne platforms *J. Geophys. Res. Atmos.*, 108, doi:10.1029/2002JD003018,
535 2003.

536 Chevallier, F., Palmer, P. I., Feng, L., Boesch, H., O'Dell, C. W., and Bousquet, P.: Towards robust and consistent
537 regional CO₂ flux estimates from in situ and space-borne measurements of atmospheric CO₂, *Geophys. Res.*
538 *Lett.*, 41, 1065–1070, doi:10.1002/2013GL058772, 2014.

539 Gourdji, S. M., Mueller, K. L., Yadav, V., Huntzinger, D. N., Andrews, A. E., Trudeau, M., Petron, G., Nehr Korn,
540 T., Eluszkiewicz, J., Henderson, J., Wen, D., Lin, J., Fischer, M., Sweeney, C. and Michalak, A. M.: North
541 American CO₂ exchange: inter-comparison of modeled estimates with results from a fine-scale atmospheric
542 inversion, *Biogeosci.*, 9, 457-475, 2012

543 Gurney, K. R., Law, R. M., Denning, A. S., Rayner, P. J., Baker, D., Bousquet, P., Bruhwiler, L., Chen, Y. H., Ciais,
544 P., Fan, S., Fung, I. Y., Gloor, M., Heimann, M., Higuchi, K., John, J., Maki, T., Maksyutov, S., Masarie, K.,
545 Peylin, P., Prather, M., Pak, B. C., Randerson, J., Sarmiento, J., Taguchi, S., Takahashi, T. and Yuen, C. W.:
546 Towards robust regional estimates of CO₂ sources and sinks using atmospheric transport models, *Nature*,
547 415(6872), 626-630, 2002.

548 Gurney, K. R., Law, R. M., Denning, A. S., Rayner, P. J., Pak, B. C., Baker, D., Bousquet, P., Bruhwiler, L., Chen,
549 Y. H., Ciais, P., Fung, I. Y., Heimann, M., John, J., Maki, T., Maksyutov, S., Peylin, P., Prather, M. and
550 Taguchi, S.: Transcom 3 inversion intercomparison: Model mean results for the estimation of seasonal carbon
551 sources and sinks, *Global Biogeochem. Cycles*, 18, GB1010, doi:10.1029/2003GB002111, 2004.

552 Houweling, S., Breon, F. M., Aben, I., Rodenbeck, C., Gloor, M., Heimann, M. and Ciais, P.: Inverse modeling of
553 CO₂ sources and sinks using satellite data: a synthetic inter-comparison of measurement techniques and their
554 performance as a function of space and time, *Atmos. Chem. Phys.*, 4, 523-538, 2004.

555 Inoue, M., Morino, I., Uchino, O., Miyamoto, Y., Yoshida, Y., Yokota, T., Machida, T., Sawa, Y., Matsueda, H.,
556 Sweeney, C., Tans, P. P., Andrews, A. E., Biraud, S. C., Tanaka, T., Kawakami, S. and Patra, P. K.: Validation
557 of XCO₂ derived from SWIR spectra of GOSAT TANSO-FTS with aircraft measurement data, *Atmos. Chem.*
558 *Phys.*, 13, 9771-9788, 2013.

559 Inoue, M., Morino, I., Uchino, O., Nakatsuru, T., Yoshida, Y., Yokota, T., Wunch, D., Wennberg, P. O., Roehl, C.
560 M., Griffith, D. W. T., Velazco, V. A., Deutscher, N. M., Warneke, T., Notholt, J., Robinson, J., Sherlock, V.,
561 Hase, F., Blumenstock, T., Rettinger, M., Sussmann, R., Kyrö, E., Kivi, R., Shiomi, K., Kawakami, S., De
562 Mazière, M., Arnold, S. G., Feist, D. G., Barrow, E. A., Barney, J., Dubey, M., Schneider, M., Iraci, L.,
563 Podolske, J. R., Hillyard, P., Machida, T., Sawa, Y., Tsuboi, K., Matsueda, H., Sweeney, C., Tans, P. P.,
564 Andrews, A. E., Biraud, S. C., Fukuyama, Y., Pittman, J. V., Kort, E. A., and Tanaka, T.: Bias corrections of
565 GOSAT SWIR XCO₂ and XCH₄ with TCCON data and their evaluation using aircraft measurement data,
566 *Atmos. Meas. Tech.* 9, 3491–3512, 2016, doi:10.5194/amt-9-3491-2016.

567 Karion, A., Sweeney, C., Tans, P., and Newberger, T.: AirCore: An Innovative Atmospheric Sampling System, *J.*
568 *Atmos. Ocean. Tech.*, 27, 1839–1853, doi:10.1175/2010JTECHA1448.1, 2010.

569 Karion, A., C. Sweeney, S. Wolter, T. Newberger, H. Chen, A. Andrews, J. Kofler, D. Neff, and P. Tans (2013),
570 Long-term greenhouse gas measurements from aircraft, *Atmos. Meas. Tech.*, 6(3), 511–526, doi:10.5194/amt-6-
571 511-2013.

572 Keeling, C. D. and Rakestraw, N. W.: The concentration of carbon dioxide in the atmosphere, *J. Geophys. Res.*, 65,
573 2502-2502, 1960.

574 Keppel-Aleks, G., Wennberg, P. O., Washenfelder, R. A., Wunch, D., Schneider, T., Toon, G. C., Andres, R. J.,
575 Blavier, J.-F., Connor, B., Davis, K. J., Desai, A. R., Messerschmidt, J., Notholt, J., Roehl, C. M., Sherlock, V.,
576 Stephens, B. B., Vay, S. A., and Wofsy, S. C.: The imprint of surface fluxes and transport on variations in total
577 column carbon dioxide, *Biogeosci.*, 9, 875– 891, doi:10.5194/bg-9-875-2012, 2012.

578 Kulawik, S. S., Worden, J. R., Wofsy, S. C., Biraud, S. C., Nassar, R., Jones, D. B. A., Olsen, E. T., Jimenez, R.,
579 Park, S., Santoni, G. W., Daube, B. C., Pittman, J. V., Stephens, B. B., Kort, E. A., Osterman, G. B. and Team,
580 T. E. S.: Comparison of improved Aura Tropospheric Emission Spectrometer CO₂ with HIPPO and SGP
581 aircraft profile measurements, *Atmos. Chem. Phys.*, 13, 3205-3225, 2013.

582 Lauvaux, T., Schuh, A. E., Uliasz, M., Richardson, S., Miles, N., Andrews, A. E., Sweeney, C., Diaz, L. I., Martins,
583 D., Shepson, P. B. and Davis, K. J.: Constraining the CO₂ budget of the corn belt: exploring uncertainties from
584 the assumptions in a mesoscale inverse system, *Atmos. Chem. Phys.*, 12, 337-354, 2012.

585 Machida, T., Kita, K., Kondo, Y., Blake, D., Kawakami, S., Inoue, G. and Ogawa, T. : Vertical and meridional
586 distributions of the atmospheric CO₂ mixing ratio between northern midlatitudes and southern subtropics, *J.*
587 *Geophys. Res. Atmos.*, 108(D3), 8401, doi:10.1029/2001JD000910, 2002.

588 Machida, T., Matsueda, H., Sawa, Y., Nakagawa, Y., Hirokuni, K., Kondo, N., Goto, K., Nakazawa, T., Ishikawa, K.
589 and Ogawa, T.: Worldwide Measurements of Atmospheric CO₂ and Other Trace Gas Species Using
590 Commercial Airlines, *J. Atmos. Ocean. Tech.*, 25(10), 1744-1754, 2008.

591 Mesinger, F., DiMego, G., Kalnay, E., Mitchell, K., Shafran, P. C., Ebisuzaki, W., Jovic, D., Woollen, J., Rogers,
592 E., Berbery, E. H., Ek, M. B., Fan, Y., Grumbine, R., Higgins, W., Li, H., Lin, Y., Manikin, G., Parrish, D., and
593 Shi, W.: North American regional reanalysis, *B. Am. Meteorol. Soc.*, 87, 343–360, doi:10.1175/BAMS-87-3-
594 343, 2006.

595 Messerschmidt, J., Geibel, M. C., Blumenstock, T., Chen, H., Deutscher, N. M., Engel, A., Feist, D. G., Gerbig, C.,
596 Gisi, M., Hase, F., Katrynski, K., Kolle, O., Lavric, J. V., Notholt, J., Palm, M., Ramonet, M., Rettinger, M.,
597 Schmidt, M., Sussmann, R., Toon, G. C., Truong, F., Warneke, T., Wennberg, P. O., Wunch, D. and Xueref-
598 Remy, I.: Calibration of TCCON column-averaged CO₂: the first aircraft campaign over European TCCON
599 sites, *Atmos. Chem. Phys.*, 11(21), 10765-10777, 2011.

600 Miller, C. E., Crisp, D., DeCola, P. L., Olsen, S. C., Randerson, J. T., Michalak, A. M., Alkhaled, A., Rayner, P.,
601 Jacob, D. J., Suntharalingam, P., Jones, D. B. A., Denning, A. S., Nicholls, M. E., Doney, S. C., Pawson, S.,
602 Boesch, H., Connor, B. J., Fung, I. Y., O'Brien, D., Salawitch, R. J., Sander, S. P., Sen, B., Tans, P., Toon, G.
603 C., Wennberg, P. O., Wofsy, S. C., Yung, Y. L. and Law, R. M.: Precision requirements for space-based X-CO₂
604 data, *J. Geophys. Res. Atmos.*, 112, D10314, doi:10.1029/2006JD007659, 2007.

605 Miller, J. B., Lehman, S. J., Montzka, S. A., Sweeney, C., Miller, B. R., Karion, A., Wolak, C., Dlugokencky, J.,
606 Southon, J., Turnbull, J. C., and Tans, P. P.: Linking emissions of fossil fuel CO₂ and other anthropogenic trace
607 gases using atmospheric ¹⁴C, *J. Geophys. Res.*, 117, D08302, doi:10.1029/2011JD017048, 2012.

608 Miller, S. T. K., Keim, B. D., Talbot, R. W. and Mao, H.: Sea breeze: Structure, forecasting, and impacts, *Rev.*
609 *Geophys.*, 41, 1011, doi:10.1029/2003RG000124, 2003.

610 Miyamoto, Y., Inoue, M., Morino, I., Uchino, O., Yokota, T., Machida, T., Sawa, Y., Matsueda, H., Sweeney, C.,
611 Tans, P. P., Andrews, A. E. and Patra, P. K.: Atmospheric column-averaged mole fractions of carbon dioxide at
612 53 aircraft measurement sites, *Atmos. Chem. Phys.*, 13(10), 5265-5275, 2013.

613 Peters, W., Jacobson, A. R., Sweeney, C., Andrews, A. E., Conway, T. J., Masarie, K., Miller, J. B., Bruhwiler, L.
614 M. P., Petron, G., Hirsch, A. I., Worthy, D. E. J., van der Werf, G. R., Randerson, J. T., Wennberg, P. O., Krol,
615 M. C. and Tans, P. P.: An atmospheric perspective on North American carbon dioxide exchange:
616 CarbonTracker, *Proc. Natl. Acad. Sci. U. S. A.*, 104(48), 18925-18930, 2007.

617 Ramonet, M., Ciais, P., Nepomniachii, I., Sidorov, K., Neubert, R. E. M., Langendorfer, U., Picard, D., Kazan, V.,
618 Biraud, S., Gusti, M., Kolle, O., Schulze, E. D. and Lloyd, J.: Three years of aircraft-based trace gas
619 measurements over the Fyodorovskoye southern taiga forest, 300 km north-west of Moscow, *Tellus B-Chem.*
620 *Phys. Meteor.*, 54(5), 713-734, 2002.

621 Reuter, M., Bovensmann, H., Buchwitz, M., Burrows, J. P., Connor, B. J., Deutscher, N. M., Griffith, D. W. T.,
622 Heymann, J., Keppel-Aleks, G., Messerschmidt, J., Notholt, J., Petri, C., Robinson, J., Schneising, O., Sherlock,
623 V., Velasco, V., Warneke, T., Wennberg, P. O. and Wunch, D.: Retrieval of atmospheric CO₂ with enhanced
624 accuracy and precision from SCIAMACHY: Validation with FTS measurements and comparison with model
625 results, *J. Geophys. Res. Atmos.*, 116, 2011.

626 Reuter, M., Buchwitz, M., Hilker, M., Heymann, J., Schneising, O., Pillai, D., Bovensmann, H., Burrows, J. P.,
627 Bosch, H., Parker, R., Butz, A., Hasekamp, O., O'Dell, C. W., Yoshida, Y., Gerbig, C., Nehr Korn, T.,
628 Deutscher, N. M., Warneke, T., Notholt, J., Hase, F., Kivi, R., Sussmann, R., Machida, T., Matsueda, H. and
629 Sawa, Y.: Satellite-inferred European carbon sink larger than expected, *Atmos. Chem. Phys.*, 14(24), 13739-
630 13753, 2014.

631 Reuter, M., Buchwitz, M., Hilker, M., Heymann, J., Bovensmann, H., Burrows, J. P., Houweling, S., Liu, Y. Y.,
632 Nassar, F., Chevallier, F., Cias, P., Marshall, J., and Reichstein, M.: How much CO₂ is taken up by the
633 European terrestrial biosphere? *Bull. Am. Meteor. Soc.*, 98, 665–671, doi:10.1175/BAMS-D-15-00310.1, 2017.

634 Saitoh, N., Kimoto, S., Sugimura, R., Imasu, R., Kawakami, S., Shiomi, K., Kuze, A., Machida, T., Sawa, Y. and
635 Matsueda, H.: Algorithm update of the GOSAT/TANSO-FTS thermal infrared CO₂ product (version 1) and
636 validation of the UTLS CO₂ data using CONTRAIL measurements, *Atmos. Meas. Tech.*, 9(5), 2119-2134,
637 2016.

638 Stephens, B. B., Gurney, K. R., Tans, P. P., Sweeney, C., Peters, W., Bruhwiler, L., Ciais, P., Ramonet, M.,
639 Bousquet, P., Nakazawa, T., Aoki, S., Machida, T., Inoue, G., Vinnichenko, N., Lloyd, J., Jordan, A., Heimann,
640 M., Shibistova, O., Langenfelds, R. L., Steele, L. P., Francey, R. J. and Denning, A. S.: Weak northern and
641 strong tropical land carbon uptake from vertical profiles of atmospheric CO₂, *Science*, 316(5832), 1732-1735,
642 2007.

643 Sweeney, C., Karion, A., Wolter, S., Newberger, T., Guenther, D., Higgs, J. A., Andrews, A. E., Lang, P. M., Neff,
644 D., Dlugokencky, E., Miller, J. B., Montzka, S. A., Miller, B. R., Masarie, K. A., Biraud, S. C., Novelli, P. C.,
645 Crotwell, M., Crotwell, A. M., Thoning, K. and Tans, P. P.: Seasonal climatology of CO₂ across North America
646 from aircraft measurements in the NOAA/ESRL Global Greenhouse Gas Reference Network, *J. Geophys. Res.*
647 *Atmos.*, 120(10), 5155-5190, 2015.

648 Tanaka, M., Nakazawa, T. and Aoki, S.: Concentration of atmospheric carbon-dioxide over Japan, *J. Geophys. Res.*
649 *Ocean.*, 88(C2), 1339-1344, DOI: 10.1029/JC088iC02p01339, 1983.

650 Tanaka, T., Miyamoto, Y., Morino, I., Machida, T., Nagahama, T., Sawa, Y., Matsueda, H., Wunch, D., Kawakami,
651 S. and Uchino, O.: Aircraft measurements of carbon dioxide and methane for the calibration of ground-based
652 high-resolution Fourier Transform Spectrometers and a comparison to GOSAT data measured over Tsukuba
653 and Moshiri, *Atmos. Meas. Tech.*, 5(8), 2003-2012, 2012.

654 Tans, P. P., Conway, T. J. and Nakazawa, T.: Latitudinal distribution of the sources and sinks of atmospheric
655 carbon-dioxide derived from surface observations and an atmospheric transport model, *J. Geophys. Res.*
656 *Atmos.*, 94(D4), 5151-5172, 1989.

657 Tans, P. P., Fung, I. Y. and Takahashi, T.: Observational constraints on the global atmospheric co2 budget, *Science*,
658 247(4949), 1431-1438, 1990.

659 Thoning, K. W., Tans, P. P. and Komhyr, W. D.: Atmospheric carbon-dioxide at Mauna Loa observatory. 2.
660 Analysis of the NOAA GMCC data, 1974-1985, *J. Geophys. Res. Atmos.*, 94(D6), 8549-8565, 1989.

661 Washenfelder, R. A., Toon, G. C., Blavier, J. F., Yang, Z., Allen, N. T., Wennberg, P. O., Vay, S. A., Matross, D.
662 M. and Daube, B. C.: Carbon dioxide column abundances at the Wisconsin Tall Tower site, *J. Geophys. Res.*
663 *Atmos.*, 111, D22305, doi:10.1029/2006JD007154, 2006.

664 Wofsy, S. C.: HIAPER Pole-to-Pole Observations (HIPPO): finegrained, global-scale measurements of climatically
665 important atmospheric gases and aerosols, *Philos. T. R. Soc. A*, 369, 2073–2086, doi:10.1098/rsta.2010.0313,
666 2011.

667 Wunch, D., Toon, G. C., Wennberg, P. O., Wofsy, S. C., Stephens, B. B., Fischer, M. L., Uchino, O., Abshire, J. B.,
668 Bernath, P., Biraud, S. C., Blavier, J. F. L., Boone, C., Bowman, K. P., Browell, E. V., Campos, T., Connor, B.
669 J., Daube, B. C., Deutscher, N. M., Diao, M., Elkins, J. W., Gerbig, C., Gottlieb, E., Griffith, D. W. T., Hurst,
670 D. F., Jimenez, R., Keppel-Aleks, G., Kort, E. A., Macatangay, R., Machida, T., Matsueda, H., Moore, F.,
671 Morino, I., Park, S., Robinson, J., Roehl, C. M., Sawa, Y., Sherlock, V., Sweeney, C., Tanaka, T. and Zondlo,
672 M. A.: Calibration of the Total Carbon Column Observing Network using aircraft profile data', *Atmos. Meas.*
673 *Tech.*, 3(5), 1351-1362, 2010.

674 Wunch, D., Wennberg, P. O., Messerschmidt, J., Parazoo, N. C., Toon, G. C., Deutscher, N. M., Keppel-Aleks, G.,
675 Roehl, C. M., Randerson, J. T., Warneke, T., and Notholt, J.: The covariation of Northern Hemisphere
676 summertime CO2 with surface temperature in boreal regions, *Atmos. Chem. Phys.*, 13, 9447–9459,
677 doi:10.5194/acp-13-9447-2013, 2013.