# Observations of atmospheric <sup>14</sup>CO<sub>2</sub> at Anmyeondo GAW station, Korea: Implications for fossil fuel CO<sub>2</sub> and emission ratios

Haeyoung Lee<sup>1,2</sup>, Edward J. Dlugokencky<sup>3</sup>, Jocelyn C Turnbull<sup>4,5</sup>, Sepyo Lee<sup>1</sup>, Scott J. Lehman<sup>6</sup>,
 John B Miller<sup>3</sup>, Gabrielle Petron<sup>3,5</sup>, Jeongsik Lim<sup>7,8</sup>, and Gang-Woong Lee<sup>2</sup>, Sang-Sam Lee<sup>1</sup> and
 Young-San Park<sup>1</sup>

6 7

8 Correspondence to Haeyoung Lee (leehy80@korea.kr)

9

<sup>1</sup>National Institute of Meteorological Sciences, Jeju, 63568, Republic of Korea

- 11 <sup>2</sup>Atmospheric Chemistry Laboratory, Hankuk University of Foreign Studies, Gyeonggi-do, 17035, Republic of 12 Korea
- 13 <sup>3</sup>NOAA, Global Monitoring Laboratory, Boulder, Colorado, USA
- <sup>4</sup> National Isotope Center, GNS Science, Lower Hutt, New Zealand
- 15 <sup>5</sup>CIRES, University of Colorado, Boulder, Colorado, USA
- 16 <sup>6</sup>INSTAAR, University of Colorado, Boulder, Colorado, USA
- <sup>7</sup>Korea Research Institute of Standard and Science, Daejeon, 34113, Republic of Korea
- 18 <sup>8</sup> University of Science and Technology, Daejeon, 34113, Republic of Korea
- 19 Abstract. To understand Korea's carbon dioxide  $(CO_2)$  emissions and sinks as well as those of
- 20 the surrounding region, we used 70 flask-air samples collected during May 2014 to August 2016
- 21 at Anmyeondo (AMY, 36.53° N, 126.32° E; 46 m a.s.l) World Meteorological Organization
- 22 (WMO) Global Atmosphere Watch (GAW) station, located on the west coast of South Korea, for
- 23 analysis of observed <sup>14</sup>C in atmospheric CO<sub>2</sub> as a tracer of fossil fuel CO<sub>2</sub> contribution ( $C_{ff}$ ).
- 24 Observed <sup>14</sup>C/C ratios in CO<sub>2</sub> (reported as  $\Delta$  values) at AMY varied from -59.5 to 23.1 ‰ with a
- 25 measurement uncertainty of  $\pm 1.8$ %. The derived mean value  $C_{\rm ff}$  of (9.7 $\pm$ 7.8)  $\mu$ mol mol<sup>-1</sup> (1 $\sigma$ ) is
- 26 greater than that found in earlier observations from Tae-Ahn Peninsula (TAP, 36.73° N, 126.13°
- 27 E, 20 m a.s.l., 28 km away from AMY) of  $(4.4\pm5.7)$  µmol mol<sup>-1</sup> from 2004 to 2010. The
- 28 enhancement above background mole fraction of sulfur hexafluoride ( $\Delta x(SF_6)$ ) and carbon
- 29 monoxide ( $\Delta x(CO)$ ) correlate strongly with  $C_{ff}$  (r > 0.7) and appear to be good proxies for fossil

30 fuel CO<sub>2</sub> at regional and continental scales. Samples originating from the Asian continent had

31 greater  $\Delta x(CO)$ :  $C_{ff}(R_{CO})$  values, (29±8) to (36±2) nmol  $\mu$ mol<sup>-1</sup>, than in Korean local air ((8±2))

32  $nmol \ \mu mol^{-1}$ ). Air masses originating in China showed  $(1.6 \pm 0.4)$  to  $(2.0 \pm 0.1)$  times greater  $R_{CO}$ 33 than a bottom-up inventory suggesting that China's CO emissions are underestimated in the 34 inventory while observed  $R_{SF6}$  values are 2-3 times greater than inventories for both China and 35 Korea. However, both  $R_{CO}$  derived from inventories and observations have decreased relative to

36 previous studies, indicating that combustion efficiency is increasing in both China and South
37 Korea.

#### 38 1 Introduction

Carbon Dioxide (CO<sub>2</sub>) is the principle cause of climate change in the industrial era, and is 39 increasing in the atmosphere at  $(2.4\pm0.4) \mu mol mol^{-1} a^{-1}$  in a recent decade globally (where 0.4 40 41 is the standard deviation of annual growth rates; www.esrl.noaa.gov/gmd/ccgg/trends/, last access: 6 December 2019). This increase is by release of CO<sub>2</sub> from fossil fuel combustion that 42 has been demonstrated through <sup>14</sup>C analysis of tree rings from the last two centuries (Stuiver and 43 44 Quay, 1981; Suess, 1955; Tans et al., 1979). Atmospheric measurement program for the ratio <sup>14</sup>C/C in CO<sub>2</sub> was initiated in the 1950s and 1960s (Rafter and Fergusson, 1957; Nydal, 1996). 45 Observed <sup>14</sup>C/C ratios are reported in Delta notation ( $\Delta$ (<sup>14</sup>CO<sub>2</sub>)) as fractionation-corrected permil 46 47 (or ‰) deviations from the absolute radiocarbon standard (Stuiver and Polach, 1977). Many studies show that the variation of  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) is an unbiased and now widely used tracer for CO<sub>2</sub> 48 49 emitted from fossil-fuel combustion (Levin et al., 2003; Turnbull et al., 2006; Graven et al., 2009; 50 Van der Laan et al., 2010; Miller et al., 2012). Therefore measurements of  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) are 51 important to test the effectiveness of emission reduction strategies to mitigate the rapid 52 atmospheric CO<sub>2</sub> increase, since they can partition observed CO<sub>2</sub> enhancements,  $\Delta x$ (CO<sub>2</sub>), into 53 fossil fuel CO<sub>2</sub> ( $C_{\rm ff}$ ) and biological CO<sub>2</sub> ( $C_{\rm bio}$ ) components with high confidence (Turnbull et al., 54 2006).

55 When trace gases are co-emitted with  $C_{\rm ff}$ , correlations of their enhancements with  $C_{\rm ff}$  improve 56 understanding of the emission sources of both  $C_{\rm ff}$  and the co-emitted tracers. For example, CO 57 and CH<sub>4</sub> emission inventories are typically more uncertain than the fossil fuel CO<sub>2</sub> emission inventory, since fossil fuel CO2 emissions related to complete combustion are generally well 58 59 estimated while emissions related to incomplete combustion and agricultural activities are poorly 60 constrained (Kurokawa et al., 2013). Temporal changes in the observed emission ratio of a trace 61 gas to  $C_{\rm ff}$  can be used to examine emission trends in the trace gas (Tohijima et al., 2014). 62 Therefore the observed emission ratios of trace gases to  $C_{\rm ff}$  can be used to evaluate bottom-up 63 inventories of various trace gases (e.g., Miller et al., 2012). Here, we used two trace gases, 64 carbon monoxide (CO) and sulfur hexafluoride (SF<sub>6</sub>) for this analysis. CO is produced along 65 with CO<sub>2</sub> during incomplete combustion of fossil fuels and biomass. CO enhancements above 66 background ( $\Delta x(CO_2)$ ) correlate well with  $C_{\rm ff}$  and have been used as a fossil fuel tracer 67 (Zondervan and Meijer, 1996; Gamnitzer et al., 2006; Turnbull et al., 2011a; Turnbull et al., 68 2011b; Tohijima et al., 2014). SF<sub>6</sub> is an entirely anthropogenic gas and is widely used as an arc 69 quencher in high-voltage electrical equipment (Geller et al., 1997). At regional to continental 70 scales, persistent small leaks to the atmosphere of SF<sub>6</sub> are typically co-located with fossil fuel 71  $CO_2$  sources and allow SF<sub>6</sub> to be used as an indirect  $C_{\rm ff}$  tracer, if the leaks are co-located with  $C_{\rm ff}$ 72 emissions at the location and scale of interest (Turnbull et al., 2006; Rivier et al., 2006).

73 South Korea is a rapidly developing country with fast economic growth, and it is located next to 74 China, which is the world's largest emitter of anthropogenic CO<sub>2</sub> (Boden et al., 2017; Janssens-Maenhout et al., 2017). The first  $\Delta(^{14}CO_2)$  measurements in South Korea were reported by 75 76 Turnbull et al. (2011a) based on air samples collected during October 2004 to March 2010 at Tae-Ahn Peninsula (TAP, 36.73° N, 126.13° E, 20 m a.s.l.). This study showed that observed 77  $CO_2$  at this site was often influenced by Chinese emissions and the observed ratio of  $\Delta x(CO)$ :  $C_{\rm ff}$ 78  $(R_{\rm CO})$  was greater than expected from bottom-up inventories. However South Korean  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) 79 80 data are still limited and the ratio of the other trace gases to  $C_{\rm ff}$  barely discussed.

81 Here we use whole-air samples collected in glass flasks during May 2014 to August 2016 at Anmyeondo (AMY, 36.53° N, 126.32° E; 46 m a.s.l.) World Meteorological Organization 82 83 (WMO) Global Atmosphere Watch (GAW) station, located on the west coast of South Korea and 84 about 28 km SSE of TAP, where the first study was conducted. We decompose observed CO<sub>2</sub> enhancements into their fossil fuel and biological components at AMY to understand sources and 85 86 sinks of CO<sub>2</sub>. We also implemented cluster analysis using the NOAA Hybrid Single Particle 87 Lagrangian Integrated Trajectory Model (HYSPLIT) to calculate back-trajectories for sample 88 times and dates. Based on clusters of trajectories from specific regions, trace gas enhancement:  $C_{\rm ff}$  ratios and correlation coefficients were analyzed, especially focused on SF<sub>6</sub> and CO, to 89 determine the potential of alternative proxies to  $\Delta$ (<sup>14</sup>CO<sub>2</sub>). Finally we compared our  $\Delta x$ (CO): $C_{\rm ff}$ 90 91 ratio with ratios determined from bottom-up inventories (EDGARv4.3.2 and Korea's National 92 Inventory Report in 2018) to evaluate reported CO emissions and how they've changed since 93 2010.

#### 95 2. Materials and Methods

#### 96 **2.1 Sampling site and methods**

97 The AMY GAW station is managed by the National Institute of Meteorological Sciences (NIMS) 98 in the Korea Meteorological Administration (KMA). It has the longest record of continuous CO<sub>2</sub> 99 measurement in South Korea, beginning in 1999. It is located on the west coast of Korea about 100 130 km southwest of the megacity of Seoul, whose population was 9.8 million in 2017. 101 Semiconductor and other industries exist within a 100 km radius of the station. Also, the largest 102 thermal power plants fired by coal and heavy oil in South Korea are within 35 km to the 103 northeast and southeast of the station. The closest town, around 30 km to the east of AMY, is 104 well known for its livestock industries. Local economic activities are related to agriculture, e.g., 105 production of rice paddies, sweet potatoes, and onions, and the area is also known for its leisure 106 opportunities that increase traffic and tourists in summer, indicating the complexity of 107 greenhouse gas sources around AMY. On the other hand, air masses often arrive at AMY from 108 the west and south, which is open to the Yellow Sea. Therefore AMY observes enhanced CO<sub>2</sub> 109 compared to many other East Asian stations due not only to numerous local sources but also 110 long-range transport of air-masses from the Asian continent (Lee et al., 2019).

Two pairs of flask-air samples (4 flasks total, 2 L, borosilicate glass with Teflon O-ring sealed stopcocks) were collected about weekly from a 40 m tall tower at AMY, regardless of wind direction and speed from May 2014 to August 2016, generally between 1400 to 1600 local time (Table S1) using a semi-automated portable sampler. A pair of flasks was flushed for 10 min at 5-6 L min<sup>-1</sup> then pressurized to 0.38 bar in less than 1 min. A second pair is collected shortly after the first (within 20 min). The portable sampler was checked for leaks after pressurizing by observing the pressure gauge before closing the stopcocks. Batches of sampled flasks wereshipped to Boulder, CO, USA every two months.

119 A total of 70 sets were collected and analyzed at the National Oceanic and Atmospheric 120 System Research Laboratory/Global Administration/Earth Monitoring Division (NOAA/ESRL/GMD) for CO<sub>2</sub>, CO, and SF<sub>6</sub> and for  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) by University of Colorado 121 122 Boulder, Institute of Arctic and Alpine Research (INSTAAR). NOAA/ESRL/GMD analyzed 123  $CO_2$  using a non-dispersive infrared analyzer,  $SF_6$  using gas chromatography (GC) with electron 124 capture detection, and CO by vacuum UV, resonance fluorescence. All analyzers were calibrated 125 with the appropriate WMO mole fraction scales (WMO-X2007 scale for CO<sub>2</sub>, WMO-X2014A 126 scale for CO, and WMO-X2014 for SF<sub>6</sub>; https://www.esrl.noaa.gov/gmd/ccl/, last access: 4 127 December 2019). The measurement and analysis methods for those gases are described in detail 128 (http://www.esrl.noaa.gov/gmd/ccgg/behind the scenes/measurementlab.html, last access: 4 129 December 2019). Measurement uncertainties for CO<sub>2</sub> and SF<sub>6</sub> are reported as 68% confidential intervals. For CO<sub>2</sub>, it is 0.07  $\mu$ mol mol<sup>-1</sup> for all measurements used here. For SF<sub>6</sub>, it is 0.04 pmol 130 131 mol<sup>-1</sup>. For CO, measurement uncertainty has not yet been formally evaluated, but is estimated at 1 nmol mol<sup>-1</sup> (68% confidence interval). All CO<sub>2</sub>, SF<sub>6</sub> and CO data at AMY can be downloaded 132 133 through ftp://aftp.cmdl.noaa.gov/data/trace gases/. When we compare NOAA's CO<sub>2</sub> 134 measurements from flask-air with quasi-continuous measurements by KMA at AMY, the difference was -0.11±2.32  $\mu$ mol mol<sup>-1</sup> (mean±1  $\sigma$ ), close to GAW's compatibility goal for CO<sub>2</sub> 135 136  $(\pm 0.1 \text{ ppm for Northern Hemisphere measurements, Lee et al., 2019}).$ 

137 The analysis methods for  $\Delta({}^{14}\text{CO}_2)$  are described by Lehman et al.(2013). Measurement 138 repeatability of  $\Delta({}^{14}\text{CO}_2)$  in aliquots of whole air extracted from surveillance cylinders is 1.8‰ 139 (1  $\sigma$ ), roughly equating to 1 µmol mol<sup>-1</sup>  $C_{\rm ff}$  detection capability from the measurement 140 uncertainty alone. The  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) data at AMY was tabulated in Table S1. Among four flasks, the 141 air from two flasks, after analysis for greenhouse gas mole fractions, was combined and analyzed 142 for  $\Delta$ (<sup>14</sup>CO<sub>2</sub>).

143

# 144 2.2 Data analysis method using $\Delta$ (<sup>14</sup>CO<sub>2</sub>) data

### 145 **2.2.1 Calculation of** *C*<sub>ff</sub> and *C*<sub>bio</sub>

146 As Turnbull et al. (2009) suggested the observed  $CO_2(C_{obs})$  at AMY can be defined as:

147 
$$C_{\text{obs}} = C_{\text{bg}} + C_{\text{ff}} + C_{\text{other}}(1)$$

148 where  $C_{bg}$ ,  $C_{ff}$  and  $C_{other}$  are the background, recently added fossil fuel  $CO_2$  and the  $CO_2$  derived 149 from the other sources.

- 150 According to Tans et al. (1993), the product of  $CO_2$  abundance and its isotopic ratio is conserved;
- 151 the isotopic mass balance can be described as below:
- 152  $\Delta_{obs}C_{obs} = \Delta_{bg}C_{bg} + \Delta_{ff}C_{ff} + \Delta_{other}C_{other}$  (2)
- 153 where  $\Delta$  is the  $\Delta$ (<sup>14</sup>C) of each CO<sub>2</sub> component of Equ. (1).
- 154 Therefore we can calculate fossil fuel  $CO_2$  by combining equations (1) and (2) as:

155 
$$C_{\rm ff} = \frac{C_{\rm bg}(\Delta_{\rm obs} - \Delta_{\rm bg})}{\Delta_{\rm ff} - \Delta_{\rm bg}} - \frac{C_{\rm other}(\Delta_{\rm other} - \Delta_{\rm bg})}{\Delta_{\rm ff} - \Delta_{\rm bg}} \quad (3)$$

156 Fossil fuel derived CO<sub>2</sub> contains no <sup>14</sup>C because the half–life of <sup>14</sup>C is (5700 $\pm$ 30) years (Godwin,

157 1962) while these fuels are hundreds of millions of years old. As we mentioned in the section 1,  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) is reported as a per mil (‰) deviation from the absolute radiocarbon reference standard 158 159 corrected for fractionation and decay with simplified а form;  $\Delta({}^{14}C) \approx [R_{sample}({}^{14}C/C)/R_{standard}({}^{14}C/C)-1]1000\%$ , where  $R({}^{14}C/C)$  is the  ${}^{14}C/C$  amount ratio. 160 Therefore  $\Delta_{\rm ff}$  is set at -1000‰ (Stuiver and Pollach, 1977). Background values ( $\Delta_{\rm bg}$ ) in equations 161 162 (1) to (3) are determined from measurements from background air collected at Niwot Ridge, 163 Colorado, a high altitude site at a similar latitude as AMY (NWR, 40.05° N, 105.58° W, 3,526 m 164 a.s.l.). Turnbull et al. (2011a) showed that the choice of background values did not significantly 165 influence derived enhancements due to the large regional and local signal at TAP, 28 km from AMY. NWR  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) and other trace gas background values are selected using a flagging 166 167 system to exclude polluted samples (Turnbull et al., 2007), and then fitted with a smooth curve 168 following Thoning et al. (1989).

169 The second term of equation (3) is typically a small correction for the effect of other sources of 170  $CO_2$  that have a  $\Delta$ (<sup>14</sup>C) differing by a small amount that of the atmospheric background, such as 171  $CO_2$  from the 1) nuclear power industry, 2) oceans, 3) photosynthesis and 4) heterotrophic 172 respiration.

173 1) The nuclear power industry produces <sup>14</sup>C that can influence the  $C_{\rm ff}$  calculation. South Korea 174 has nuclear power plants along the east coast that may influence AMY air samples when air-175 masses originated from the eastern part of Korea (Figure 1). It is also possible that Chinese 176 nuclear plants could influence some samples. Here we did not make any correction for this since 177 most nuclear installations in this region are pressurized water reactors, which produce mainly <sup>14</sup>C 178 in  $CH_4$  rather than  $CO_2$  (Graven and Gruber, 2011). 2) For the ocean, although there may also be 179 a small contribution from oceanic carbon exchange across the Yellow Sea, we consider this 180 effect small enough to ignore (Turnbull et al., 2011a). It was also demonstrated there is no 181 significant bias from the oceans including East China Sea (Song et al., 2018), even at coastal 182 sites in the Northern Hemisphere (Turnbull et al., 2009). Larger scale ocean exchange and also 183 stratospheric exchange affect both background and observed samples equally, so they can be ignored in the calculations. 3) For the photosynthetic terms, <sup>14</sup>C in CO<sub>2</sub> accounts for natural 184 185 fractionation during uptake, so we also set this observed value the same as the background value. 186 4) Therefore we only consider heterotrophic respiration. For land regions, where most fossil fuel 187 emissions occur, heterotrophic respiration could be a main contributor to the second term of equation (3) due to <sup>14</sup>C disequilibrium potentially. When this value is ignored,  $C_{\rm ff}$  would be 188 189 consistently underestimated (Palstra et al., 2008; Riley et al., 2008; Hsueh et al., 2007; Turnbull et al., 2006). For this, corrections were estimated as (-0.2 $\pm$ 0.1) µmol mol<sup>-1</sup> during winter and (-190

191  $0.5\pm0.2$ ) µmol mol<sup>-1</sup> during summer (Turnbull et al., 2009; Turnbull et al., 2006).

192 CO<sub>2</sub> enhancements relative to baseline CO<sub>2</sub> are defined as  $\Delta x(CO_2)$ , with the excess signal of 193  $C_{obs}$  minus  $C_{bg}$  in Equ.(1). Partitioning of  $\Delta x(CO_2)$  into  $C_{ff}$  and  $C_{bio}$  is calculated simply from the 194 residual of the difference between observed  $\Delta x(CO_2)$  and  $C_{ff}$ .

#### 195 **2.2.2** The ratio of trace gas enhancement to C<sub>ff</sub> and its correlation

196 To obtain the correlation coefficient (r) between  $C_{\rm ff}$  and other trace gas enhancements ( $\Delta x(x) = x_{\rm obs} - x_{\rm bg}$ ) and the ratio of any trace gas to  $C_{\rm ff}(R_{\rm gas})$ , we use reduced major axis (RMA) regression 198 analysis (Sokal and Rohlf, 1981). The distributions of  $R_{\rm gas}$  are normally broad and non-Gaussian and RMA analysis is a relatively robust method of calculating the slope of two variables that show some causative relationship. Here,  $x_{bg}$  was derived from NWR with the same method described in section 2.2.1. The relevant equations are presented from Equ. S1 to Equ. S3. Results for each species are given in Table 1.

203

#### 204 **2.3 HYSPLIT cluster analysis**

205 HYSPLIT trajectories were run using Unified Model-Global Data Assimilation and Prediction 206 System (UM-GDAPS) weather data at 25 km by 25 km horizontal resolution to determine the 207 regions that influence air mass transport to AMY. A total of 70 air-parcel back-trajectories were 208 calculated for 72-h periods at 3-h intervals matching the time of each flask-air sample taken at 209 AMY from May 2014 to August 2016. We assign the sampling altitude as 500 m, since it was 210 demonstrated that HYSPLIT and other particle dispersion back-trajectory models (e.g., 211 FLEXPART) are consistent at 500 m altitude (Li et al., 2014). Cluster analysis of the resulting 212 70 back-trajectories categorized six pathways through which air parcels arrive at AMY during 213 the time period of interest.

Among the calculated back-trajectories, 67% indicate air masses originating from the Asian continent. Back-trajectories of continental background air (CB) originating in Russia and Mongolia occurred 13% of the time. 23% of the trajectories originated and travelled through northeast China (CN). The CN region includes Inner Mongolia and Liaoning, one of the most populated regions in China with 43.9 million people in 2012. These CN air masses arrive in South Korea after crossing through western North Korea. 17% of the trajectories are derived from central eastern China around the Shandong area (CE). The CE region contains Shandianzi (SDZ, 40.65° N, 117.12° E, 287 m a.s.l.) located next to the megacities of Beijing and Tianjin, which are some of China's highest CO<sub>2</sub> emitting regions (Gregg et al., 2008). 14% are Ocean Background (OB) derived from the East China Sea. Among them, a few of the trajectories passed over the eastern part of China (e.g., over Shanghai) with high altitude (1000 m). Flow from South Korea also travels through heavily industrialized and/or metropolitan regions in South Korea (Korea Local, KL, 19%) and under stagnant conditions (Polluted Local region, PL, 14%). Some of the KL air-masses have also passed over the East Sea and Japan.

228

#### 229 **3. Results and discussions**

## 230 **3.1 Observed** $\Delta$ (<sup>14</sup>CO<sub>2</sub>) and portioning of CO<sub>2</sub> into C<sub>ff</sub> and C<sub>bio</sub>

AMY  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) values are almost always lower than those observed at NWR, which we consider 231 232 to be broadly representative of background values for the mid-latitude Northern Hemisphere (Figure 2). NWR  $\Delta$ (<sup>14</sup>CO<sub>2</sub>), which is based on weekly air samples, was in the range 10.0 to 21.2 233 234 %, with an average  $(16.6\pm3)$ %  $(1\sigma, \text{ standard deviation})$  from May 2014 to August 2016. Waliguan (WLG, 36.28° N, 100.9° E, 3816 m a.s.l.), an Asian background GAW station in 235 China, also showed similar  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) levels to NWR with an average of (17.1±6.8)‰ in 2015 236 (Niu et al., 2016, measurement uncertainty  $\pm 3\%$ , n=20).  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) at AMY varied from -59.5 to 237 23.1‰ and had a mean value of  $(-6.2\pm18.8)$ ‰  $(1\sigma, n=70)$  during the measurement period 238 (Table S1). This was similar to results from observations at SDZ, which is located about 100 km 239

northeast of Beijing, in the range of -53.0 to 32.6% with an average  $(-6.8\pm21.1)$ %  $(1\sigma, n=32)$ 

241 during Sep 2014 to Dec 2015 (Niu et al., 2016).

243

242 Calculated  $C_{\rm ff}$  at AMY ranges between -0.05 and 32.7  $\mu$ mol mol<sup>-1</sup> with an average of (9.7±7.8)

 $\mu$ mol mol<sup>-1</sup> (1 $\sigma$ , n=70); high C<sub>ff</sub> was observed regardless of season (Figure 2 (a)). One negative

 $C_{\rm ff}$  value of -0.05 µmol mol<sup>-1</sup> was estimated due to greater AMY  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) than NWR on July 30, 244 2014. Although negative  $C_{\rm ff}$  values are non-physical, this value is not significantly different from 245 zero, and is reasonable given that this air originated from the OB sector. The range of  $C_{\rm ff}$  in the 246 AMY samples is similar to that observed at TAP from 2004 to 2010 (-1.6 to 42.9  $\mu$ mol mol<sup>-1</sup> 247  $C_{\rm ff}$ ), but  $C_{\rm ff}$  is on average about twice as high at AMY as in the 2004 to 2010 TAP samples 248 (mean (4.4±5.7) µmol mol<sup>-1</sup>, n=202) (Turnbull et al., 2011a). A more detailed comparison of 249 250 results based on differences between samples derived from the Asian continent and Korea local 251 air is provided in section 3.2.

Estimated  $C_{\text{bio}}$ , as defined in section 2.2.1, varied from -18.1 to 15.7  $\mu$ mol mol<sup>-1</sup> (mean (0.9\pm5.8)

 $\mu$ mol mol<sup>-1</sup>) at AMY (Figure 2 (c)).  $C_{bio}$  showed a strong seasonal cycle with the lowest values from July to September when photosynthetic drawdown is expected to be strongest, in good agreement with the previous TAP study (Turnbull et al., 2011a). Even though  $C_{bio}$  was at times negative, mainly due to photosynthesis during summer, the largest positive  $C_{bio}$  was also observed in summer.

258 The largest  $C_{\rm ff}$  by season was observed in order of winter (DJF, (11.3±7.6), n=14) > summer 259 (JJA, (10.7±9.2), n=11) > spring (MAM, (8.6±8.0), n=22) > autumn (SON, (7.6±5.6), n=17) with a unit of  $\mu$ mol mol<sup>-1</sup>. When we consider only positive contributions of  $C_{\text{bio}}$  samples, the order was summer ((4.6±4.0), n=14) > autumn ((4.1±2.5), n=9) > spring ((3.8±2.6), n=13) > winter ((3.4±2.5), n=11) with a unit of  $\mu$ mol mol<sup>-1</sup>.

 $C_{\rm ff}$  in summer was nearly as high as in winter. This is because lower wind speeds are observed at AMY during summer (Lee et al., 2019). When we analyzed seasonal boundary layer height for each sample by UM-GDAPS, it also showed similar result that it was highest in winter (with a range from 150 m to 1100 m) and lowest in summer (with a range from 100 m to 500 m). This suggests that these high summer  $C_{\rm ff}$  values may reflect emission from local activities, which were described in section 2.1, more than in other seasons.

The highest  $C_{bio}$  value was also observed in the summer, PL sector. PL sector showed that positive  $C_{bio}$  correlates with CH<sub>4</sub>, which is a tracer for agriculture when observed in TAP local air masses. Turnbull et al.(2011a) also showed similar results.

In winter,  $C_{\text{bio}}$  was relatively lower than in other seasons while  $C_{\text{ff}}$  was highest. During winter, AMY is mainly affected by long-range transport of air-masses from China due to the Siberian high (Lee et al., 2019). Therefore air samples were less affected by local activities in winter but  $C_{\text{bio}}$  still contributed almost 23% to  $\Delta x(\text{CO}_2)$ . In the dry season (from October to March), forest fires, which contribute the largest portion of total CO<sub>2</sub> emissions from open fires at the national scale, are concentrated in northeastern and southern China (Yin et al., 2019). The highest CO was observed in winter ((449.1±244.1) nmol mol<sup>-1</sup> (1 $\sigma$ ) in winter while (236.8±124.4) nmol

279  $mol^{-1}(1\sigma)$  in summer), which also supports biomass burning and bio fuels as large contributors

280 to observed  $CO_2$  enhancements in winter. Turnbull et al. (2011a) also showed that 20-30% of

winter CO<sub>2</sub> enhancements at TAP were likely contributed by biofuel combustion, along with
 plant, soil, human, and animal respiration.

Regardless of the source, we find that  $C_{\text{bio}}$  contributes substantially to atmospheric CO<sub>2</sub> enhancements at AMY in air masses affected by local and long-range transport, so when only CO<sub>2</sub> enhancements above background are compared to bottom-up inventories, it can make a bias due to  $C_{\text{bio}}$  contributions.

287

#### 288 3.2 C<sub>ff</sub> comparison between Korea Local and Asian Continent samples

To more clearly identify samples originating from the Asian continent (trajectory clusters CB, CN, CE, and OB) and Korea Local (trajectory cluster KL) after cluster analysis of the 70 sets of measurements, we use wind speed data from the Automatic Weather System (AWS) installed at the same level as the air sample inlet at AMY. Among the data from CB, CN, CE, OB, and KL, when wind speed was less than 3 m/s, we assumed that those samples could be affected by local pollution. PL was also ruled out since it was affected by local pollutions under the stagnant condition. Therefore we use only 41 sets of observations for this analysis (Table 1).

296  $C_{\rm ff}$  is highest in the order CE > CN > KL > CB > OB (Table 1). During the measurement period, 297 the averages from Asian continent (sectors CE and CN) were higher than KL without the 298 baseline sector (CB and OB). The calculated mean  $C_{\rm ff}$  using only CE, CN, CB and OB, which 299 sample substantial outflow from the Asian Continent, was (7.6±3.9) µmol mol<sup>-1</sup>.

When we compared the KL samples ( $(8.6\pm5.3)$  µmol mol<sup>-1</sup>) with those from Korea Local air-300 masses observed at TAP ((8.5±8.6)  $\mu$ mol mol<sup>-1</sup>, n=58, Turnbull et al., 2011a), mean C<sub>ff</sub> was 301 quite similar (Figure 3). However, when comparing the  $C_{\rm ff}$  values from CB air masses in this 302 study and TAP far-field (from China) samples (n=144, Turnbull et al., 2011a), C<sub>ff</sub> almost 303 doubled from (2.6±2.4) to (4.3±2.1)  $\mu$ mol mol<sup>-1</sup>, even though they might be expected to have 304 305 had similar air mass back-trajectories. We also compared the values at SDZ from 2009 to 2010 306 (Turnbull et al., 2011a) and in 2015 (Niu et al., 2016); they also increased, not only in the 307 samples that were affected by Beijing and North China Plain (SDZ-BN), which are comparably 308 polluted, but also in the samples that were affected by northeast China (SDZ-NE). For SDZ-BN samples,  $C_{\rm ff}$  increased from (10±1) to (16±7.6) µmol mol<sup>-1</sup> from 2009/2010 (n=32) to 2015 309 (n=32). The AMY samples from CE, which flow over Beijing, showed (11.2 $\pm$ 8.3) µmol mol<sup>-1</sup> of 310  $C_{\rm ff}$  and were also slightly greater than the 2009 – 2010 SDZ-BN samples (Turnbull et al., 2011a). 311 For SDZ-NE samples,  $C_{\rm ff}$  was (3±7) µmol mol<sup>-1</sup> in 2009 to 2010 and increased to (7.6±6.8) 312 µmol mol<sup>-1</sup> in 2015. Since the SDZ-NE samples are affected by northeast China according to 313 314 Turnbull et al. (2011a) and Niu et al. (2016), we also see CN that originated from northeast china (NE) and its mean value of  $C_{\rm ff}$  had increased around (10.6±6.9) µmol mol<sup>-1</sup> compared to those 315 316 values in 2009 to 2010.

317 It has been suggested that inter-annual variability in observed mean  $C_{\rm ff}$  in South Korea could 318 reflect changing fossil fuel CO<sub>2</sub> emissions, or could indicate inter-annual variability in the air 319 mass trajectories of the (small) dataset of flask-air samples (Turnbull et al., 2011a). Even though the growth rate of  $C_{\rm ff}$  emission has been decreasing slowly in East Asia since 2010 due to emission reduction policies (Labzovskii et al., 2019), reported emissions increased 16.7% in China and 1.8% in South Korea from 2010 to 2016 (Janssens-Maenhout et al., 2017). This is broadly consistent with the flat trend in observed  $C_{\rm ff}$  in KL air masses, and in the upward trend in  $C_{\rm ff}$  observed in air-masses flowing out from Asia. Therefore it is possible that AMY mean  $C_{\rm ff}$ increased relative to the earlier TAP observations due to increased fossil fuel emissions from the Asian continent.

On the other hand, those values from this study showed large variability with small sample numbers due to different sampling strategy, environment, and synoptic conditions such as boundary layer height at the sampling time from reference studies. Further study will be necessary to understand those increased values.

331

#### 332 **3.3.** Correlation of $C_{\rm ff}$ with SF<sub>6</sub> and its emission ratios

We calculated correlation coefficients (r from Equ. (S3)) between SF<sub>6</sub> and CO enhancements with  $C_{\rm ff}$  and their ratios from Equ. (S1) with the 50 samples that were described in section 3.2 including PL sector (n=9) and whose values are tabulated in Table 1.

The correlations of CO enhancements ( $\Delta x$ (CO)) with  $C_{\rm ff}$  were strong (r > 0.7) in all sectors except PL, while SF<sub>6</sub> enhancements ( $\Delta x$ (SF<sub>6</sub>)) correlated strongly with  $C_{\rm ff}$  (r > 0.8) for CE and OB in outflow from the Asian Continent and KL.  $R_{\rm CO}$  and  $R_{\rm SF6}$  were different between Korea Local and outflows from the Asian Continent. Here we discuss  $R_{\rm SF6}$  and section 3.4 discuss  $R_{\rm CO}$ more detail. 341 For SF<sub>6</sub>, observed mean levels were high in order of (KL, PL) > (CN, CE) > (OB, CB) (Table 1). 342  $SF_6$  in KL and PL were higher than from the Asian Continent, since South Korea has larger  $SF_6$ emissions than most countries (ranked at 4<sup>th</sup> as of 2010 according to the EDGAR4.2.) because of 343 344 liquid-crystal display (LCD) and electrical equipment production (Fang et al., 2014). Even 345 though both KL and PL showed higher  $SF_6$  mole fraction than outflows of Asian Continent, the 346 correlation is different between KL and PL (Table 1). Under stagnant conditions, emitted SF<sub>6</sub> is 347 less diluted by mixing, so that in PL,  $\Delta x(SF_6)$  correlated weakly with C<sub>ff</sub>. On the other hand, KL, CE and OB showed strong correlations (r > 0.8). Those three sectors are also larger SF<sub>6</sub> sources 348 349 compared to other regions, according to SF<sub>6</sub> emission estimates for Asia (Fang et al., 2014). 350 Because long-range transport allows time for mixing, SF<sub>6</sub> and C<sub>ff</sub> emissions are effectively co-351 located at not only continental scales but also regional scales. Thus SF<sub>6</sub> can be a good tracer of 352 fossil fuel CO<sub>2</sub> for those regions.

The correlation between  $\Delta x(SF_6)$  and  $C_{\rm ff}$  was strong in CE, OB and KL, however,  $R_{\rm SF6}$  is different between South Korea and outflow from the Asian continent (Figure S2). In a previous study, observed  $R_{\rm SF6}$  was 0.02 to 0.03 pmol µmol<sup>-1</sup> at NWR in 2004 (Turnbull et al., 2006). Here, the ratio was at (0.19±0.03) and (0.17±0.03) pmol µmol<sup>-1</sup> for CE and OB respectively. For KL,

357 it was (0.66±0.16) pmol  $\mu$ mol<sup>-1</sup> indicating much larger ratios than in outflow from the Asian

358 continent. Further, observed  $R_{SF6}$  is 2 to 3 times greater for all air masses than predicted from 359 bottom-up inventories based on national scale roughly. For this calculation, we use EDGAR4.3.2 360 for CO<sub>2</sub> and EDGAR4.2 for SF<sub>6</sub>. We repeat the calculations for both CO<sub>2</sub> and SF<sub>6</sub> with Korea's 361 National Inventory Report (KNIR, Greenhouse Gas Inventory and Research Center, 2018). 362 Using SF<sub>6</sub> for 2010 from EDGAR4.2, we obtain  $R_{SF6}$  of 0.08 pmol µmol<sup>-1</sup> for China while for

South Korea it was 0.14 pmol µmol<sup>-1</sup>. Especially for South Korea, this is much lower than the 363 observed R<sub>SF6</sub>. When KL R<sub>SF6</sub> was compared to ratios calculated from the KNIR inventory (0.27 364 pmol  $\mu$ mol<sup>-1</sup> for 2010 and 0.22 pmol  $\mu$ mol<sup>-1</sup> for 2014), it was closer to observed  $R_{SF6}$  than 365 366 EDGAR, but still underestimated (Figure S3 and S2). This result suggests that the observed ratio 367 could be used to re-evaluate the bottom-up inventories (Rivier et al., 2006), especially targeting 368 the Asian continent. Even though KL  $R_{SF6}$  showed greater uncertainty than CE and OB, it is still 369 greater than bottom-up inventories, such as KNIR and EDGAR. Therefore it would be useful to 370 get more data to try and derive a more robust estimate to evaluate SF<sub>6</sub> emission inventories for 371 Korea.

372

#### 373 **3.4 Correlation of C**<sub>ff</sub> with CO and its emission ratios

High CO was mainly observed in outflow from the Asian continent in order of CE > CN > PL > 374 (CB, KL) > OB (Table 1). The order of CO is quite different to that of SF<sub>6</sub>. CO from KL and PL 375 376 is lower than from outflow from the Asian continent, except for the OB sector, indicating that 377 high CO can be a tracer of outflow from the Asian continent. Since CO is produced during 378 incomplete combustion of fossil fuel and biomass, it is more closely related to fossil fuel CO<sub>2</sub> emissions than the other trace gases. Therefore in most cases the correlation between CO and  $C_{\rm ff}$ 379 was strong. R<sub>CO</sub> was very different between air masses originating from South Korea Local 380  $((8\pm2) \text{ nmol } \mu\text{mol}^{-1})$  and the Asian continent  $((29\pm8) \text{ to } (36\pm2) \text{ nmol } \mu\text{mol}^{-1})$ , due to differences 381 382 in combustion efficiencies and the use of catalytic converters. The higher continental emission 383 ratios may also result from some contribution of biofuel combustion and agricultural burning in 384 the Asian continent, which have significantly higher CO emission than fossil-fuel combustion

385 (Akagi et al., 2011). For example, for CB the CO level is similar to KL while  $R_{CO}$  is higher than 386 KL with low  $C_{\rm ff}$ .

387 Typically CO shows seasonal variations with lower values in summer due to the atmospheric 388 chemical sink, OH. Among the samples, the samples collected in summer were mainly rejected 389 through wind speed cut-off (less than 3 m/s) since AMY has lower wind speed in summer (Lee 390 et al., 2019). Only OB sector includes 4 summer samples (of 7), because summer air masses are mainly from the southern part of the Yellow Sea (Lee et al., 2019). However, we assumed  $R_{CO}$  is 391 392 less affected by the summer sink, since only two  $\Delta x(CO)$  samples were negative for OB (Figure 393 S1) and  $R_{\rm CO}$  was consistent whether or not the negative  $\Delta x({\rm CO})$  values were considered. To 394 compare emission ratios derived from atmospheric observations with those from inventories for 395 2000 to 2012, we calculated inventory emission ratio ( $I_{CO/CO2}$ ) as:

$$396 \qquad I_{\rm CO/CO2} = E_{\rm CO}/E_{\rm CO2} \,\, {\rm X} \,\, M_{\rm CO2}/M_{\rm CO}$$

397 Where,  $E_{CO}$  and  $E_{CO2}$  are total CO and fossil fuel CO<sub>2</sub> emissions in gigagrams (Gg a<sup>-1</sup>, 10<sup>9</sup> g a<sup>-1</sup>)

from the bottom-up national inventory.  $M_X$  is the molar masses of CO and CO<sub>2</sub> in g mol<sup>-1</sup>.

We use EDGAR4.3.2 (Janssens-Maenhout et al., 2017) and KNIR (Greenhouse Gas Inventory
and Research Center, 2018) for inventory information for both CO and CO<sub>2</sub>.

401 The uncertainty of EDGAR4.3.2 fossil fuel CO<sub>2</sub> emissions was reported as a 95% confidence

402 interval (Janssens-Maenhout et al., 2019), ±5.4% for China and ±3.6% for South Korea

403 (personal communication with EDGAR team). The uncertainties of CO and SF<sub>6</sub> emissions were

404 not reported by EDGAR. For KNIR, the CO<sub>2</sub> 2016 emission uncertainty in the energy sector was

405 ±3% (Greenhouse Gas Inventory and Research Center, 2018). KNIR does not provide
406 uncertainties for other emission sectors of CO<sub>2</sub>, nor from emissions of CO and SF<sub>6</sub>.

407 In Fig. 4 we confirm that the CO to  $C_{\rm ff}$  emission ratios ( $R_{\rm CO}$ ) derived from both observations and 408 inventories for China and South Korea are decreasing. Since  $C_{\rm ff}$  emissions appear to be flat 409 (South Korea) or slightly increasing (China), this indicates that combustion efficiency and/or 410 scrubbing of CO is improving.

411 For South Korea, EDGAR4.3.2 indicated that CO emissions from the energy sector (98% to 99% 412 of total emission) decreased by 47% between the 1997 and 2012. South Korean fossil fuel CO<sub>2</sub> 413 emissions increased until 2011 and remained mostly constant from 2011 to 2016 ((603,901±4,315) Gg  $a^{-1}$  CO<sub>2</sub>) (Figure S4). Therefore the decreased trend in the emission ratio 414 415 seems to reflect recent decreases in CO emissions in South Korea. Turnbull et al. (2011a) determined an observed mean  $R_{\rm CO}$  of (13±3) nmol µmol<sup>-1</sup> during 2004 to 2010. Suntharalingam 416 et al. (2004) estimated  $R_{\rm CO}$  15.4 nmol  $\mu$ mol<sup>-1</sup> for South Korea in 2001 from CO<sub>2</sub> and CO airborne 417 observations (C<sub>ff</sub> was not determined). Recently, the KORUS-AQ campaign, which was 418 conducted over Seoul from May to June in 2016, estimated  $R_{CO}$  as 9 nmol  $\mu$ mol<sup>-1</sup> (Tang et al., 419 2018) based on CO<sub>2</sub> and CO observations ( $C_{\rm ff}$  was not determined). Our study gives  $R_{\rm CO}$  of (8±2) 420 nmol  $\mu$ mol<sup>-1</sup> for South Korea, slightly but not significantly lower than the KORUS-AQ result for 421 422 Seoul. Different contributions of  $C_{\text{bio}}$  and  $C_{\text{ff}}$  to total CO<sub>2</sub> may bias the  $R_{\text{CO}}$  calculation when total CO<sub>2</sub> was used in the KORUS-AQ study (e.g., Miller et al., 2012). The South Korean 423 national  $R_{CO}$  from EDGAR4.3.2 in 2012 was 6.7 nmol µmol<sup>-1</sup>, consistent with our observations. 424 Using KNIR for 2016, we obtain  $R_{CO}$  of 2.1 nmol  $\mu$ mol<sup>-1</sup>. KNIR suffers from a large number of 425

missing CO emission sources compared to the EDGAR, as indicated by their reported emissions,
638.3 and 2580.8 Gg a<sup>-1</sup> in 2012, respectively (Figure S5). For example, CO emissions recently
derived from fugitive emissions and residential/other sectors increased to 14% and 11.5% of total
emission respectively in EDGAR but were not reported in KNIR.

430 For China the inventories estimate that CO emissions from the energy sector,  $(96.5\pm0.2)$ %, were 431 almost constant through the 1990s, and then increased during the early-2000s from industrial 432 processes (8.8% of total emissions in 2012). Fossil fuel CO<sub>2</sub> emission in China also increased until 2013 and then stayed roughly constant at (10,461,890±60,571) Gg a<sup>-1</sup> according to 433 434 EDGAR4.3.2. Thus even though both emissions show an increase from 2000 to 2016 for fossil fuel CO<sub>2</sub> and to 2012 for CO, the emission ratio decreased (Figure S4 and Figure 4) seeming to 435 436 indicate that combustion efficiency is improving. Many studies observed decreasing  $R_{CO}$  in China from 2000 to 2010 (Turnbull et al., 2011a; Wang et al., 2010). Suntharalingam et al. (2004) 437 reported  $R_{\rm CO}$  was 55 nmol µmol<sup>-1</sup> in 2001 ( $C_{\rm ff}$  was not determined). In the Beijing region,  $R_{\rm CO}$ 438 decreased from 57.80 to 37.59 nmol µmol<sup>-1</sup> during 2004 to 2008 (Wang et al., 2010). The overall 439  $R_{CO}$  was (47±2) nmol µmol<sup>-1</sup> at SDZ for 2009-2010 and (44±3) nmol µmol<sup>-1</sup> in air-masses that 440 originated from the Asian continent from 2005 to 2009 (Turnbull et al., 2011a). Tohjima et al. 441 (2014) explained that surface based  $R_{\rm CO}$  decreased from 45 to 30 nmol  $\mu$ mol<sup>-1</sup> in outflow air 442 masses from China from 1998 to 2010. Fu et al. (2015) also observed  $R_{CO}$  of 29 nmol  $\mu$ mol<sup>-1</sup> 443

- 444 over mainland China in 2009. In Beijing, which is located along the path of CE, it was (30.4±1.6)
- 445 nmol  $\mu$ mol<sup>-1</sup> and (29.6±3.2) nmol  $\mu$ mol<sup>-1</sup> for Xiamen in 2016, which is in the OB sector (Niu et
- 446 al., 2018). During KORUS-AQ in 2016,  $R_{CO}$  of 28 nmol  $\mu$ mol<sup>-1</sup> was observed over the Yellow

447 Sea. Some of those studies did not differentiate  $C_{\rm ff}$  from the total CO<sub>2</sub> enhancement, so, although 448  $R_{\rm CO}$  still includes uncertainties, it is continually decreasing.

449 In this study  $R_{CO}$  is (29±8), (31±8), (36±2), and (31±4) nmol µmol<sup>-1</sup> for CB, CN, CE and OB,

450 consistent with Tang et al.(2018) and Liu et al.(2018). On the other hand,  $R_{CO}$  in CE is higher 451 than in other sectors in this study. The Shandong area, which is located in the path of CE, has 452 been plagued with problems of combustion inefficiency and ranked as the largest consumer of 453 fossil fuels in all of China (Chen and Li, 2009). The uncertainties in our observed  $R_{CO}$  for this 454 region overlap with other sectors such as CB, CN and OB, so further monitoring of the ratios 455 will help to get more detailed information.

456 In South Korea and China, atmosphere-based  $R_{CO}$  values calculated by this study are (1.2±0.3)

457 times (with KL),  $(1.6\pm0.4)$ ,  $(1.7\pm0.4)$ ,  $(2\pm0.1)$  and  $(1.7\pm0.2)$  times greater (with CB, CN, CE

and OB) than in the inventory, respectively (Figure 4). This is in agreement with previous studies
(Turnbull et al., 2011a; Kurokawa et al., 2013; Tohjima et al., 2014). One explanation is that
EDGAR does not reflect secondary CO production, which can be a significant contributor to CO
(Kurokawa et al., 2013). Also, CO derived from biomass burning and biofuels was not included
in this inventory. Therefore, this indicates that top-down observations are necessary to evaluate
and improve bottom-up emission products.

464

#### 465 **4. Summary and Conclusions**

466 To understand  $CO_2$  sources and sinks in Korea as well as those of the surrounded region, we 467 collected  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) with 70 flask samples from May 2014 to August 2016. We summarized our 468 results below.

- 469 1) Observed  $\Delta(^{14}CO_2)$  values at AMY ranged from -59.5 to 23.1‰ (a mean value of
- 470

 $(-6.2\pm18.8)$ % (1 $\sigma$ )) during the study period, almost always lower than those observed at

- 471 NWR, which we consider to be broadly representative of background values for the mid472 latitude Northern Hemisphere. This reflects the strong imprint of fossil fuel-CO<sub>2</sub>
  473 emissions recorded in AMY air samples.
- 474 2) Calculated  $C_{\rm ff}$  using  $\Delta(^{14}{\rm CO}_2)$  at AMY ranges between -0.05 and 32.7 µmol mol<sup>-1</sup> with 475 an average of (9.7±7.8) µmol mol<sup>-1</sup> (1 $\sigma$ ); this average is twice as high as in the 2004 to
- 2010 TAP samples (mean (4.4 $\pm$ 5.7) µmol mol<sup>-1</sup>) (Turnbull et al., 2011a). We also 476 477 observed high C<sub>ff</sub> regardless of the season or source region. After separately identifying 478 samples originating from the Asian continent and the Korean peninsula, we determined 479 that the mean  $C_{\rm ff}$  increased relative to the earlier observations due to increased fossil fuel 480 emissions from the Asian continent as showing by the consistent growth in reported 481 emissions, which increased 16.7% in China and only 1.8% in South Korea from 2010 to 482 2016. Note, however, that our data span a relatively limited time period and are subject 483 to different synoptic conditions during the sampling time from previous studies, so a 484 longer time-series would increase confidence in tracking this change.
- 485 3) Because  $\Delta x(CO)$  and  $\Delta x(SF_6)$  agreed well with  $C_{\rm ff}$ , but showed different slopes for Korea 486 and the Asian continent, those  $R_{\rm gas}$  values can be indicators of air mass origin and those

487		gases can be proxies for $C_{\rm ff}$ . Overall, we have confirmed that both $R_{\rm CO}$ derived from
488		inventory and observation have decreased relative to previous studies, indicating that
489		combustion efficiency is increasing in both China and South Korea.
490	4)	However, atmosphere-based $R_{gas}$ values are greater than bottom-up inventories. For CO,
491		our values are $(1.2\pm0.3)$ times and $(1.6\pm0.4)$ to $(2.0\pm0.1)$ times greater than in inventory
492		values for South Korea and China, respectively. This discrepancy may arise from several
493		sources including the no contribution of atmospheric chemical CO production such as
494		oxidation of CH <sub>4</sub> and non-methane VOCs. Observed $R_{SF6}$ is 2 to 3 times greater than in
495		inventories. Therefore those values in our study can be used for improving bottom-up
496		inventories in the future.
497	5)	Finally, we stress that because $C_{\text{bio}}$ contributes substantially to $\Delta x(\text{CO}_2)$ , even in winter,
498		$\Delta^{14}$ C-based $C_{\rm ff}$ (and not $\Delta x(\rm CO_2)$ ) is required for accurate calculation of both $R_{\rm CO}$ and

 $R_{\rm SF6}$ .



Figure 1. A total of 70 air-parcel back-trajectories were calculated for 72-h periods at 3-h
intervals from May 2014 to August 2016 using the HYSPLIT model in conjunction with KMA
UM GDAPS data at 25 km by 25 km resolution. Station locations are: WLG (Waliguan, 36.28°
N, 100.9° E, 3816 m a.s.l.), SDZ (Shandianzi, 40.65° N, 117.12° E, 287 m a.s.l.), and AMY
(Anmyeondo, 36.53° N, 126.32° E, 86 m a.s.l.). TAP (Tae-Ahn Peninsula, 36.73° N, 126.13° E,
20 m a.s.l.) is around 28 km northeast from AMY.



Figure 2. Time series of (a) observed CO<sub>2</sub> dry air mole fraction (open circles) and observed CO<sub>2</sub> ( $C_{obs}$ ) minus  $C_{ff}$  calculated from  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) (closed circles). (b)  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) at AMY (black circles) and at NWR (Niwot Ridge, line), baseline data. (c) Time series of  $C_{ff}$  and  $C_{bio}$  calculated from  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) (left) and the frequency distribution at AMY (right).



Figure 3. Calculated  $C_{\rm ff}$  (µmol mol<sup>-1</sup>). Red bars are for KL and blue bars are for Korea far-field (China) (2004-2010 from Turnbull et al. (2011a)). Black bars are for SDZ-BN samples that were affected by Beijing and North China plain. Gray bars for SDZ-NE indicate samples that were affected by regions northeast of SDZ. SDZ (2009-2010) is from Turnbull et al. (2011a) and SDZ (2015) is from Niu et al. (2016). Hatched red, blue, black and grey bars are derived from this study during 2014 to 2016.

![](_page_27_Figure_0.jpeg)

Figure 4.  $R_{CO}$  for China (a) and for South Korea (b). Black circles: EDGARv.4.3.2 emission inventory. Black squares: National Inventory Report, Korea (2018). Blue symbols are from other studies (Suntharalingam et al., 2004; Wang et al., 2010; Turnbull et al., 2011a; Tohjima et al., 2014; Liu et al., 2018; Tang et al., 2018). Red symbols: This study. Y-error bars: uncertainty in the slope according to equation (S2). X-error bars: the period for the mean value.

529	Table 1. Means and standard deviations of $C_{\rm ff}$ (µmol mol <sup>-1</sup> ), CO (nmol mol <sup>-1</sup> ) and SF <sub>6</sub> (pmol
530	mol <sup>-1</sup> ) (total N=50, without PL N=41). The correlations (r) and the ratio ( $R_{gas}$ ) of enhancement
531	between $C_{\rm ff}$ were determined by Reduced Major Axis (RMA) regression analysis on each scatter
532	plot to obtain regression slopes. The uncertainty of $R_{gas}$ refers to equation (S2). When r is less
533	than 0.7, $R_{gas}$ was not included here. N is the number of data. The unit of $R_{CO}$ is nmol $\mu$ mol <sup>-1</sup> and
534	for $R_{SF6}$ it is pmol $\mu$ mol <sup>-1</sup> . A plot of $R_{CO}$ and $R_{SF6}$ is shown in Figure S1. CB represents
535	continental background, CN north east China, CE central eastern China, OB ocean background,
536	KL Korea local and PL polluted local air-mass

	(	Outflow from t	South Korea			
	CB (N=7)	CN (N=9)	CE (N =9)	OB (N =7)	KL (N =9)	PL (N =9)
$C_{ m ff}$	4.3±2.1	10.6±6.9	11.2±8.3	4.1±2.7	8.6±5.3	15.6±11.6
СО	233±59	353±219	473±293	169±90	228±40	259±100
SF <sub>6</sub>	9.0±0.4	10.1±1.2	10.1±1.5	9.2±0.5	13.0±3.3	12.7±6.2
R <sub>CO</sub> (r)	29±8 (0.80)	31±8 (0.76)	36±2 (0.98)	31±4 (0.96)	8±2 (0.74)	- (0.44)
R <sub>SF6</sub> (r)	(0.63)	- (0.48)	0.19±0.03 (0.91)	0.17±0.03 (0.94)	0.66±0.16 (0.76)	(0.38)

# 539 Data availability540

541 Our CO<sub>2</sub>, CO, SF<sub>6</sub> data from AMY and NWR can be downloaded from 542 ftp://aftp.cmdl.noaa.gov/data/trace\_gases.  $\Delta$ (<sup>14</sup>CO<sub>2</sub>) data are provided in the supplementary 543 material of this paper.

544

#### 545 Author contributions

546 HL wrote this paper and analyzed all data. HL and GWL designed this study. EJD and JCT 547 guided and reviewed this paper. SL collected samples and gave the information of the data at 548 AMY. EJD, JCT, SJL, JBM, GP, and JL provided data and reviewed the manuscript. All authors 549 contributed this work.

550

#### 551 ACKNOWLEDGMENT

This work was funded by the Korea Meteorological Administration Research and Development
Program "Research and Development for KMA Weather, Climate, and Earth system Services–
Development of Monitoring and Analysis Techniques for Atmospheric Composition in Korea"
under Grant (KMA2018-00522).

556

#### 557 **REFERENCES**

558 Akagi, S. K., R. J. Yokelson, C. Wiedinmyer, M. J. Alvarado, J. S. Reid, T. Karl, J. D. Crounse,

559 P. O. Wennberg: Emission factors for open and domestic biomass burning for use in atmospheric

560 models, Atmos. Chem. Phys. 11, 4039-4027, doi:10.5194/acp-11-4039-2011, **2011** 

- Boden, T.A., G. Marland, and R.J. Andres: National CO<sub>2</sub> Emissions from Fossil-Fuel Burning,
  Cement Manufacture, and Gas Flaring: 1751-2014, Carbon Dioxide Information Analysis Center,
  Oak Ridge National Laboratory, U.S. Department of Energy, doi 10.3334/CDIAC/00001\_V2017,
  2017
- 565 Chen, Y. Y. Li: Low-carbon economy and China's regional energy use research. *Jilin Univ. J.*566 Soc. Sci. Ed. 49(2), 66-73, 2009
- 567 Fang, X., R. L. Thompson, T. Saito, Y. Yokouchi, J. Kim, S. Li, K. R. Kim, S. Park, F. Graziosi,
- A. Stohl: Sulfur hexafluoride (SF<sub>6</sub>) emissions in East Asia determined by inverse modeling. *Atmos. Chem. Phys.* 14, 4779–4791, doi:10.5194/acp-14-4779-2014, 2014
- 570 Fu, X. W., H. Zhang, C.-J. Lin, X. B. Feng, L. X. Zhou, S. X. Fang: Correlation slopes of 571 GEM/CO, GEM/CO<sub>2</sub>, and GEM/CH<sub>4</sub> and estimated mercury emissions in China, South Asia, the 572 Indochinese Peninsula, and Central Asia derived from observations in northwestern and 573 southwestern China. *Atmos. Chem. Phys.* 15, 1013-1028, doi:10.5194/acp-15-1013-2015, **2015**
- Gamnitzer, U., U. Karstens, B. Kromer, R. E. M. Neubert, H. Schroeder, I. Levin: Carbon
  monoxide: A quantitative tracer for fossil fuel CO<sub>2</sub>?. *J. Geohys. Res.*, 111, D22302,
  doi:10.1029/2005JD006966, 2006
- Geller, L. S., J. W. Elkins, J. M. Lobert, A. D. Clarke, D. F. Hurst, J. H. Butler, R. C. Myers:
  Tropospheric SF<sub>6</sub>: Observed latitudinal distribution and trends, derived emissions and
  interhemispheric exchange time. *Geophys. Res. Lett.*, 24(6), 675–678, doi:10.1029/97GL00523,
  1997

Graven, H. D. N. Gruber: Continental-scale enrichment of atmospheric <sup>14</sup>CO<sub>2</sub> from the nuclear
power industry: Potential impact on the estimation of fossil fuel-derived CO<sub>2</sub>. *Atmos. Chem. Phys. Discuss. 11*, 14,583–14,605, doi:10.5194/acpd-11-14583-2011, 2011

584 Graven, H. D., B. B. Stephens, T. P. Guilderson, T. L. Campos, D. S. Schimel, J. E. Campbell, R.

- 585 F. Keeling: Vertical profiles of biospheric and fossil fuel-derived CO<sub>2</sub> and fossil fuel CO<sub>2</sub>:CO
- 586 ratios from airborne measurements of <sup>14</sup>C, CO<sub>2</sub> and CO above Colorado, USA, *Tellus*, 61, 536-
- 587 546, DOI:10.1111/j.1600-0889.2009.00421.x, **2009**
- Gregg, J. S. R. J. Andres, G. Marland: China: Emissions pattern of the world leader in CO<sub>2</sub>
  emissions from fossil fuel consumption and cement production, *Geophys. Res. Lett.* 35, L08806,
  doi:10.1029/2007GL032887, 2008
- Greenhouse Gas Inventory and Research Center: National Greenhouse Gas Inventory Report of
  Korea; National statistics-115018, 11-1480906-000002-10,
  www.gir.go.kr/home/index.do?menuld=36 (in Korean), 2018
- 594 Hsueh, D. Y., N. Y. Krakauer, J. T. Randerson, X. Xu, S. E. Trumbore, J. R. Southon: Regional
- patterns of radiocarbon and fossil fuel derived CO<sub>2</sub> in surface air across North America, *Geophys. Res. Lett.*, *34*, L02816, doi:10.1029/2006GL027032, **2007**
- 597 Janssens-Maenhout, G., M. Crippa, D. Guizzardi, M. Muntean, E. Schaaf, J.G.J. Olivier,
- 598 J.A.H.W. Peters, K.M. Schure: Fossil CO<sub>2</sub> and GHG emissions of all world countries, EUR
- 599 28766 EN, Publications Office of the European Union, Luxembourg, ISBN 978-92-79-73207-2,
- 600 doi:10.2760/709792, JRC107877, 2017

- 601 Janssens-Maenhout, G.; M. Crippa, D. Guizzardi, M. Muntean, E. Schaaf, F. Dentener, P.
- 602 Bergamaschi, V. Pagliari, J. G. J. Olivier, J. A. H. W. Peters, J. A. van Aardenne, S. Monni, U.
- 603 Doering, A. M. R. Petrescu, E. Solazzo, G. D. Oreggioni: EDGAR v4.3.2 Global Atlas of the
- 604 three major greenhouse gas emissions for the period 1970–2012, *Earth Syst. Sci. Data*, 11, 959–
- 605 1002, https://doi.org/10.5194/essd-11-959-2019, 2019
- Kurokawa, J., T. Ohara, T. Morikawa, S. Hanayama, G. Janssens-Maenhout, T. Fukui, K.
  Kawashima, H. Akimoto: Emissions of air pollutants and greenhouse gases over Asian
  regionsduring 2000–2008: Regional Emission inventory in ASia (REAS) version 2, *Atmos. Chem. Phys. 13*, 11019–11058, doi:10.5194/acp-13-11019-2013, 2013
- 610 Labzovskiia, L.D., H. W. L. Mak, S. T. Keneaa, J.-S. Rhee, A. Lashkari, S. Li, T.-Y. Goo, Y.-S.
- 611 Oh, Y.-H. Byun: What can we learn about effectiveness of carbon reduction policies from
- 612 interannual variability of fossil fuel CO<sub>2</sub> emissions in East Asia? *Environ. Sci. Policy.* 96, 132-
- 613 140, https://doi.org/10.1016/j.envsci.2019.03.011, 2019
- 614 Lee, H., S.-O. Han, S.-B. Ryoo, J.-S. Lee, G.-W. Lee: The measurement of atmospheric CO<sub>2</sub> at
- 615 KMA GAW regional stations, its characteristics, and comparisons with other East Asian sites.
- 616 Atmos. Chem. Phys. 19, 2149–2163, doi.org/10.5194/acp-19-2149-2019, 2019
- 617 Lehman, S.J., J. B. Miller, C. Wolak, J.R. Southon, P.P. Tans, S.A. Montzka, C. Sweeney, A. E.
- 618 Andrews, B.W. LaFranchi, T. P. Guilderson: Allocation of terrestrial carbon sources using <sup>14</sup>CO<sub>2</sub>:
- 619 methods, measurement, and modelling. *Radiocarbon*. 55(2–3):1484–95, 2013
- 620 Le Quéré, C., R. M. Andrew, P. Friedlingstein, S. Sitch, J. Hauck, J. Pongratz, P. A. Pickers, J. I.
- 621 Korsbakken, G. P. Peters, J. G. Canadell, A. Arneth, V. K. Arora, L. Barbero, A. Bastos, L. Bopp,

- 622 F. Chevallier, L. P. Chini, P. Ciais, S. C. Doney, T. Gkritzalis, D. S. Goll, I. Harris, V. Haverd, F.
- 623 M. Hoffman, M. Hoppema, R. A. Houghton, G. Hurtt, T. Ilyina, A. K. Jain, T. Johannessen, C. D.
- 624 Jones, E. Kato, R. F. Keeling, K. K. Goldewijk, P. Landschützer, N. Lefèvre, S. Lienert, Z. Liu,
- D. Lombardozzi, N. Metzl, D. R. Munro, J. E. M. S. Nabel, S. Nakaoka, C. Neill, A. Olsen, T.
- 626 Ono, P. Patra, A. Peregon, W. Peters, P. Peylin, B. Pfeil, D. Pierrot, B. Poulter, G. Rehder, L.
- 627 Robertson, E.M. Rocher, C. Rödenbeck, U. Schuster, J. Schwinger, R. Séférian, I. Skjelvan, T.
- 628 Steinhoff, A. Sutton, P. P. Tans, H. Tian, B. Tilbrook, F. N. Tubiello, I. T. vander Laan-Luijkx,
- 629 G. R. vander Werf, N. Viovy, A. P. Walker, A.J. Wiltshire, R. Wright, S. Zaehle, Bo. Zheng:
- 630 Global Carbon Budget 2018. Earth Syst. Sci. Data. 10, 2141–2194, https://doi.org/10.5194/essd-

#### 631 <u>10-2141-2018</u>, **2018**

- Levin, I., B., M. S. Kromer, H. Sartorius: A novel approach for independent budgeting of fossil
  fuel CO<sub>2</sub> over Europe by <sup>14</sup>CO<sub>2</sub> observations, *Geophys. Res. Lett.* 30(23), 2194,
  doi:10.1029/2003GL018477, 2003
- 635 Li, S., J. Kim, S. Park, S.-K. Kim, M.-K. Park, J. Mühle, G.-W. Lee, M. Lee, C. O. Jo, K.-R.
- 636 Kim: Source identification and apportionment of halogenated compounds observed at a remote
- 637 site in East Asia. *Eviron. Sci. Technol.* 48, 491–498, doi.org/10.1021/es402776w, **2014**
- 638 Miller, J.B., S. J. Lehman, S. A. Montzka, C. Sweeney, B. R. Miller, A. Karion, C. Wolak, E. J.
- Dlugokencky, J. Southon, J. C. Turnbull, P.P. Tans: Linking emissions of fossil fuel CO<sub>2</sub> and
   other anthropogenic trace gases using atmospheric <sup>14</sup>CO<sub>2</sub>. *J.Geophys.Res.* 117, D08302,
   doi:10.1029/2011JD017048, 2012
- 642 Niu, Z., W. Zhou, X. Feng, T. Feng, S. Wu, P. Cheng, X. Lu, H. Du, X. Xiong, Y. Fu: 643 Atmospheric fossil fuel  $CO_2$  traced by <sup>14</sup> $CO_2$  and air quality index pollutant observations in

- 644 Beijing and Xiamen, China. *Environ. Sci. Pollut. Res.* 25, 17109–17117,
  645 doi.org/10.1007/s11356-018-1616-z, 2018
- 646 Niu, Z., W. Zhou, P. Cheng, S. Wu, X. Lu, X. Xiong, H. Du, Y. Fu: Observations of atmospheric
- $\Delta^{14}$ CO<sub>2</sub> at the global and regional background sites in China: Implication for fossil fuel CO<sub>2</sub>
- 648 inputs. Eviron. Sci. Technol. 50, 12122–12128 DOI: 10.1021/acs.est.6b02814, 2016
- 649 Nydal, R., and K. Lövseth, Carbon-14 measurements in atmospheric CO<sub>2</sub> from Northern and
- 650 Southern Hemisphere sites, 1962–1993, technical report, Carbon Dioxide Inf. Anal. Cent., Oak
- 651 Ridge Natl. Lab., U.S. Dep. of Energy, Oak Ridge, Tenn, 1996
- 652 Rafter, T. A., and G. J. Fergusson, "Atom Bomb Effect"—Recent increase of Carbon-14 content
- 653 of the atmosphere and biosphere, *Science*, 126(3273), 557–558, **1957**
- 654 Palstra, S. W., U. Karstens, H.-J. Streurman, H. A. J. Meijer: Wine ethanol <sup>14</sup>C as a tracer for
- 655 fossil fuel CO<sub>2</sub> emissions in Europe: Measurements and model comparison, J. Geophys. Res.,
- 656 *113*, D21305, doi:10.1029/2008JD010282, **2008**
- Riley, W. G., D. Y. Hsueh, J. T. Randerson, M. L. Fischer, J. Hatch, D. E. Pataki, W. Wang, M.
  L. Goulden: Where do fossil fuel carbon dioxide emissions from California go? An analysis
  based on radiocarbon observations and an atmospheric transport model, *J. Geophys. Res.*, *113*,
  G04002, doi:10.1029/2007JG000625, **2008**
- 661 Rivier, L., P. Ciais, D. A. Hauglustaine, P. Bakwin, P. Bousquet, P. Peylin, A. Klonecki: 662 Evaluation of  $SF_6$ ,  $C_2Cl_4$ , and CO to approximate fossil fuel  $CO_2$  in the Northern Hemisphere 663 using a chemistry transport model. *J. Geophys. Res.* 111, D16311, doi:10.1029/2005JD006725, 664 **2006**

- 665 Suntharalingam, P., D. J. Jacob, P. I. Palmer, J. A. Logan, R.M. Yantosca, Y. Xiao, M. J. Evans:
- 666 Improved quantification of Chinese carbon fluxes using CO<sub>2</sub>/CO correlations in Asian outflow, J.

667 Geophys. Res. 109, D18S18, doi:10.1029/2003JD004362, 2004

- 668 Suess, H. E. Radiocarbon concentration in modern wood, *Science*, 122,415, 1955
- 669 Stuiver, M., P. Quay: Atmospheric <sup>14</sup>C changes resulting from fossil fuel CO<sub>2</sub> release and cosmic
- 670 ray flux variability, Earth Planet. Sci. Lett. 53, 349-362, 1981
- 671 Tang, W., A. F. Arellano, J. P. DiGangi, Y. Choi, G. S. Diskin, A. Agustí-Panareda, M.
- 672 Parrington, S. Massart, B. Gaubert, Y. Lee, D. Kim, J. Jung, J. Hong, J.-W. Hong, Y. Kanaya, M.
- 673 Lee, R. M. Stauffer, A. M. Thompson, J. H. Flynn, J.-H. Woo: Evaluating high-resolution
- 674 forecasts of atmospheric CO and CO<sub>2</sub> from a global prediction system during KORUS-AQ field
- 675 campaign. Atmos. Chem. Phys. 18, 11007–11030, doi.org/10.5194/acp-18-11007-2018, 2018
- 676 Tans, P. P.; J. A. Berry, R. F. Keeling: Oceanic<sup>13</sup>C/<sup>12</sup>C observations: A new window on ocean
- 677 CO<sub>2</sub> uptake. *Global Biogeochem. Cycles.* 7(2), 353–368, doi:10.1029/93GB00053, **1993**
- 678 Sokal, R. R., and F. J. Rohlf. 1981. Biometry. 2nd edition. Freeman, NY.
- Song Jinming, Baoxiao Qu, Xuegang Li, Huamao Yuan, Ning Li, Liqin Duan: Carbon
  sinks/sources in the Yellow and East China Seas-Air-sea interface exchange, dissolution in
  seawater, and burial in sediments. *Science China Earth Sciences*. 61, 1583-1593, 2018
- 682 Stuiver, M., Polach H. A. Discussion: Reporting of <sup>14</sup>C data, *Radiocarbon*, 19(3), 355–363, **1977**
- Tans, P.P., A.F.M. de Jong, W.G. Mook: Natural atmospheric <sup>14</sup>C variation and the Suess effect,
- 684 Science, 280, 826-828, **1979**

- Thoning, K. W., P. P. Tans, W. D. Komhyr: Atmospheric Carbon dioxide at Mauna Loa
  Observatory 2. Analysis of the NOAA GMCC Data, 1984–1985, *J. Geophys. Res. 94*, 8549–
  8565, 1989
- 688 Tohjima, Y., M. Kubo, C. Minejima, H. Mukai, H. Tanimoto, A. Ganshin, S. Maksyutov, K.
- 689 Katsumata, T. Machida, K. Kita: Temporal changes in the emissions of CH<sub>4</sub> and CO from China
- 690 estimated from CH<sub>4</sub>/CO<sub>2</sub> and CO/CO<sub>2</sub> correlations observed at Hateruma Island. *Atmos. Chem.*
- 691 Phys. 14, 1663–1677, doi:10.5194/acp-14-1663-2014, 2014
- Turnbull, J., P. Rayner, J. Miller, T. Naegler, P. Ciais, A. Cozic: On the use of <sup>14</sup>CO<sub>2</sub> as a tracer
- 693 for fossil fuel CO<sub>2</sub>: Quantifying uncertainties using an atmospheric transport model, J. Geophys.
- 694 Res. 114, D22302, doi:10.1029/2009JD012308, 2009
- Turnbull, J. C., S. J. Lehman, J. B. Miller, R. J. Sparks, J. R. Southon, P. P. Tans: A new high
  precision <sup>14</sup>CO<sub>2</sub> time series for North American continental air. *J. Geophys. Res. 112*, D11310,
  doi:10.1029/2006JD008184, 2007
- 698 Turnbull, J. C., P. P. Tans, S. J. Lehman, D. Baker, T. J. Conway, Y. S. Chung, J. Gregg, J. B.
- 699 Miller, J. R. Southon, L.-X. Zhou: Atmospheric observations of carbon monoxide and fossil fuel
- 700 CO<sub>2</sub> emissions from East Asia. J. Geophys. Res., 116, D24306,doi:10.1029/2011JD016691,
- 701 **2011a**
- 702 Turnbull, J. C., A. Karion, M. L. Fischer, I. Faloona, T. Guilderson, S. J. Lehman, B. R. Miller, J.
- 703 B. Miller, S. Montzka, T. Sherwood, S. Saripalli, C. Sweeney, P. P. Tans: Assessment of fossil
- fuel carbon dioxide and other anthropogenic trace gas emissions from airborne measurements

705 over Sacramento, California in spring 2009, Atmos. Chem. Phys. 11(2), 705–721,
706 doi:10.5194/acp-11-705-2011, 2011b

707 Turnbull, J. C. J. B. Miller, S. J. Lehman, P. P. Tans, R. J. Sparks, J. Southon: Comparison of 708 <sup>14</sup>CO<sub>2</sub>, CO, and SF<sub>6</sub> as tracers for recently added fossil fuel CO<sub>2</sub> in the atmosphere and 709 implications for biological  $CO_2$ exchange, Geophys. Res. Lett., 33. L01817. 710 doi:10.1029/2005GL024213, 2006

711 Van Der Laan, S, U. Karstens, R.E.M. Neubert, I.T. Van Der Laan-Luijkx and H.A.J. Meijer:

712 Observation-based estimates of fossil fuel-derived CO<sub>2</sub> emissions in the Netherlands using  $\Delta^{14}$ C,

713 CO and <sup>222</sup>Radon, *Tellus B: Chemical and Physical Meteorology*, 62:5, 389-402,
714 DOI:10.1111/j.1600-0889.2010.00493.x. 2010

- Wang, Y. J. W. Munger, S. Xu, M. B. McElroy, J. Hao, C. Nielsen, H. Ma: CO<sub>2</sub> and its
  correlation with CO at a rural site near Beijing: Implications for combustion efficiency in China, *Atmos. Chem. Phys.* 10, 8881–8897, doi:10.5194/acp-10-8881-2010, 2010
- Yin, L., P. Du, M. Zhang, M. Liu, T. Xu, Y. Song: Estimation of emissions from biomass
  burning in China (2003–2017) based on MODIS fire radiative energy data, *Biogeosciences*, 16,
  1629–1640. 2019
- Zondervan, A., and Meijer, H. A. J: Isotopic characterization of CO<sub>2</sub> sources during regional
  pollution events using isotopic and radiocarbon analysis, *Tellus B: Chemical and Physical Meteorology*, 48(4), 601–612, doi:10.1034/j.1600-0889.1996.00013.x, 1996