

# FTIR time series of tropospheric HCN in eastern China: seasonality, interannual variability and source attribution

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## Abstract:

We analyzed seasonality and interannual variability of tropospheric HCN columns in densely populated eastern China for the first time. The results were derived from solar absorption spectra recorded with ground-based high spectral resolution Fourier transform infrared (FTIR) spectrometer at Hefei (117°10'E, 31°54'N) between 2015 and 2018. The tropospheric HCN columns over Hefei, China showed significant seasonal variations with three monthly mean peaks throughout the year. The magnitude of the tropospheric HCN column peak in May > September > December. The tropospheric HCN column reached a maximum monthly mean of  $(9.8 \pm 0.78) \times 10^{15}$  molecules/cm<sup>2</sup> in May and a minimum monthly mean of  $(7.16 \pm 0.75) \times 10^{15}$  molecules/cm<sup>2</sup> in November. In most cases, the tropospheric HCN columns at Hefei (32°N) are higher than the FTIR observations at Ny Alesund (79°N), Kiruna (68°N), Bremen (53°N), Jungfraujoch (47°N), Toronto (44°N), Rikubetsu (43°N), Izana (28°N), Mauna Loa (20°N), La Reunion Maito (21°S), Lauder (45°S), and Arrival Heights (78°S) that are affiliated with the Network for Detection of Atmospheric Composition Change (NDACC). Enhancements of tropospheric HCN column were observed between September 2015 and July 2016 compared to the same period of measurements in other years. The magnitude of the enhancement ranges from 5 to 46% with an average of 22%. Enhancement of tropospheric HCN ( $\Delta$ HCN) is correlated with the concurrent enhancement of tropospheric CO ( $\Delta$ CO), indicating that enhancements of tropospheric CO and HCN were due to the same sources. The GEOS-Chem tagged CO simulation, the global fire maps and the Potential Source Contribution Function values

1 (PSCFs) calculated using back trajectories revealed that the seasonal maxima in May is largely due  
2 to the influence of biomass burning in South Eastern Asia (SEAS) ( $41 \pm 13.1\%$ ), Europe and Boreal  
3 Asia (EUBA) ( $21 \pm 9.3\%$ ) and Africa (AF) ( $22 \pm 4.7\%$ ). The seasonal maxima in September is  
4 largely due to the influence of biomass burnings in EUBA ( $38 \pm 11.3\%$ ), AF ( $26 \pm 6.7\%$ ), SEAS ( $14$   
5  $\pm 3.3\%$ ), and Northern America (NA) ( $13.8 \pm 8.4\%$ ). For the seasonal maxima in December,  
6 dominant contributions are from AF ( $36 \pm 7.1\%$ ), EUBA ( $21 \pm 5.2\%$ ), and NA ( $18.7 \pm 5.2\%$ ). The  
7 tropospheric HCN enhancement between September 2015 and July 2016 at Hefei ( $32^\circ\text{N}$ ) were  
8 attributed to an elevated influence of biomass burnings in SEAS, EUBA, and Oceania (OCE) in this  
9 period. In particular, an elevated fire number in OCE in the second half of 2015 dominated the  
10 tropospheric HCN enhancement in September – December 2015. An elevated fire number in SEAS  
11 in the first half of 2016 dominated the tropospheric HCN enhancement in January – July 2016.

## 12 13 **1 Introduction**

14 Atmospheric hydrogen cyanide (HCN) is an extremely hazardous gas that threaten human  
15 health and terrestrial ecosystems (Andreae and Merlet, 2001; Akagi et al., 2011; Rinsland et al.,  
16 2002). Improved knowledge of the physical and chemical mechanisms which drive the observed  
17 HCN variability is of great significance because HCN plays an important role in the global nitrogen  
18 cycle (Andreae and Merlet, 2001; Li et al., 2003). It is well established that biomass burning is the  
19 major source of tropospheric HCN and industrial emissions contribute additional minor sources of  
20 HCN (Bange and Williams, 2000; Holzinger et al., 1999; Lobert et al., 1990). Li et al. (2009)  
21 estimates a global source of HCN from biomass burning of  $0.4 - 3.2 \text{ Tg N yr}^{-1}$  and from burning  
22 domestic biofuel of  $0.2 \text{ Tg N yr}^{-1}$  (Li et al., 2009). Bertschi et al. (2003) estimates a global fossil  
23 fuel combustion source of  $0.04 \text{ Tg N yr}^{-1}$ , negligibly small in comparison (Bertschi et al., 2003).  
24 The principle pathway for HCN sink is ocean uptake which accounts for 0.73 to  $1.0 \text{ Tg N/year}$  (Li  
25 et al., 2009). Additional minor sinks of HCN are attributed to atmospheric reaction with hydroxyl  
26 radical (OH) and  $\text{O}(^1\text{D})$ , and photolysis (Li et al., 2000; Nagahama and Suzuki, 2007). The life time  
27 of HCN is 2 – 5 months in the troposphere and several years in the stratosphere. Li et al. (2003,  
28 2009), Lupu et al. (2009), Vigouroux et al. (2012), and Zeng et al. (2012) showed that the observed  
29 variability of HCN can be reproduced by the chemical model simulations where biomass burning  
30 and ocean uptake provide the main source and sink, respectively (Li et al., 2009; Li et al., 2003;  
31 Lupu et al., 2009; Vigouroux et al., 2012; Zeng et al., 2012).

32 With the rapid economic growth in China over the past three decades, the anthropogenic  
33 emissions have increased dramatically, raising concerns about worsening air quality in China (Tang  
34 et al., 2012; Chan, 2017; Xing et al., 2017; Wang et al., 2017). These emissions are from automobile  
35 exhaust, industrial processes and biomass burning. Many researchers have evaluated regional  
36 emissions in various pollution regions (e.g., the Jing-Jin-Ji region, the Yangtze River Delta region,  
37 and the Pearl River Delta region, Fig. S1), but the relative contribution of the biomass burning,  
38 automobile exhaust, and industrial processes is seldom mentioned in the literature (Tang et al., 2012;  
39 Chan, 2017; Wang et al., 2017; Sun et al., 2018a; Xing et al., 2017). This is because both industrial  
40 emissions and biomass burning are major sources of the trace gases (e.g. carbon monoxide (CO),  
41 formaldehyde (HCHO) and carbon dioxide ( $\text{CO}_2$ )) that were used to evaluate regional emissions in  
42 the literature, and it is hard to quantify their relative contribution under the complex pollution  
43 condition in China (Chan et al., 2018; Tang et al., 2012; Wang et al., 2017; Xiaoyan et al., 2010;  
44 Xing et al., 2017). It has been proved that HCN is an unambiguous tracer of biomass burning  
45 emission due to its inactive chemical feature and long lifetime (Rinsland et al., 2002; Zhao et al.,  
46 2002). Therefore, measurements of HCN made in the polluted troposphere over eastern China at  
47 middle latitudes are particularly useful in determining the potential biomass burning sources that  
48 drive the observed tropospheric HCN seasonality and interannual variability in China.

49 Ground based high-resolution Fourier Transform Spectroscopy (FTIR) measurements of trace  
50 gases made by Anhui Institute of Optics and Fine Mechanics, Chinese Academy of Sciences  
51 (AIOFM-CAS) at Hefei ( $117^\circ 10'\text{E}$ ,  $31^\circ 54'\text{N}$ , 30 m a.s.l. (above sea level)) is one of few multiyear  
52 time series of trace gases on Asian continent (Sun et al., 2018a; Sun et al., 2018b). These  
53 measurements are crucial to understanding global warming, regional pollution, and long term  
54 transport. Both HCN and CO are regularly measured at Hefei ( $32^\circ\text{N}$ ) using the FTIR observations,  
55 where influences from biomass burning occurring at long distances or locally can be assessed.

56 In this study, we analyze the first multiyear measurements of tropospheric HCN in densely  
57 populated eastern China. In section 2 the retrieval strategy to derive HCN from high resolution FTIR

1 spectrometry and the methods for a GEOS-Chem tagged CO simulation and potential source  
2 contribution function (PSCF) calculation are summarized. In section 3 we present the seasonal and  
3 interannual variability of tropospheric HCN columns measured at Hefei (32° N), China and  
4 comparisons with the measurements affiliated with Network for Detection of Atmospheric  
5 Composition Change (NDACC, <http://www.ndacc.org/>, last accessed on 3 June 2019). The potential  
6 sources that drive the observed HCN variability are determined by using the GEOS-Chem tagged  
7 CO simulation, the global fire maps and the PSCFs analysis in section 4. The work concludes with  
8 a summary in section 5. This study aims to improve our understanding of regional biomass burning  
9 characteristic and transport, and contribute to the evaluation of the global nitrogen cycle.

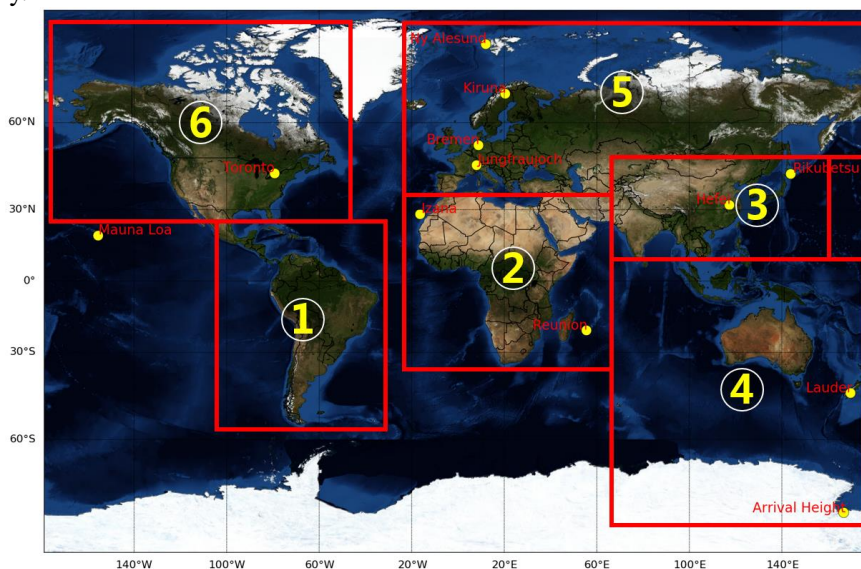
## 10 2 Methods

### 11 2 FTIR observations

#### 12 2.1.1 Site description and instrumentation

13 The routine observations of atmospheric trace gases using ground based high-resolution FTIR  
14 spectrometer at Hefei (117°10'E, 31°54'N, 30 m a.s.l.) started in July 2014. Location of Hefei site  
15 alongside those of the NDACC FTIR stations selected for comparison are shown in Fig.1.  
16 Geographical source regions used in the standard GEOS-Chem tagged CO simulation are also  
17 marked in Fig.1. Detailed description of Hefei site can be found in Tian et al., 2017. We follow the  
18 NDACC requirements, and plan to apply for acceptance within the NDACC in the future.

19 A Bruker IFS 125 HR with maximum optical path difference (OPD) of 900 cm is used to take  
20 the solar spectra (Tian et al., 2017). Defined as 0.9/OPD, this instrument can reach the highest  
21 spectral resolution of 0.001 cm<sup>-1</sup>. However, all mid-infrared (MIR) spectra are recorded with a  
22 spectral resolution of 0.005 cm<sup>-1</sup> to follow NDACC convention. This spectral resolution is sufficient  
23 to resolve the optical absorption structure of all gases in the atmosphere. The FTIR spectrometer  
24 covered a wide spectral range (about 600 – 4500 cm<sup>-1</sup>) but, depending on the species, specific  
25 detectors and band-pass filters are applied (Sun et al. 2018a). In this study, the instrument is  
26 equipped with a KBr beam splitter, an InSb detector, and a filter centered at 2900 cm<sup>-1</sup> for HCN  
27 measurements, and a KBr beam splitter, an InSb detector, and a filter centered at 2400 cm<sup>-1</sup> for CO  
28 measurements. The entrance field stop size ranging from 0.80 to 1.5 mm was employed to maximise  
29 the signal to noise ratio (SNR) consistent with the maximum frequency possible for the selected  
30 wavenumber range. The number of measurements within a day varies from 1 to 20. In total, there  
31 were 651 and 649 days of qualified measurements between 2015 and 2018 for CO and HCN,  
32 respectively.



33 Fig. 1 Location of Hefei site alongside those of the NDACC FTIR stations (yellow dots) that are selected for  
34 comparison. Geographical source regions used in the standard GEOS-Chem tagged CO simulation are also shown.  
35 See Table 3 for latitude and longitude definitions  
36

#### 37 2.1.2 Retrieval strategy

38 The SFIT4 (version 0.9.4.4) algorithm is used to retrieve the vertical profiles of CO and HCN  
39 (Viatte et al., 2014). Both CO and HCN are standard NDACC species, and we follow the NDACC  
40 recommendation for micro windows (MWs) selection and the interfering gases consideration

(<http://www.ndaccdemo.org/>, last accessed on 23 May 2019). The retrieval inputs for CO and HCN are summarized in Table 1. Time series of tropospheric CO columns between 2014 and 2017 at Hefei (32°N) measured from the FTIR have been reported in Sun et al. (2018a) and the detailed description of CO profile retrieval can be found therein. Time series of tropospheric HCN columns at Hefei (32°N) are presented for the first time. Temperature and pressure profiles are extracted from National Centers for Environmental Protection (NCEP) 6-hourly reanalysis data (De Maziere et al., 2018) and all spectroscopic absorption parameters are prescribed from HITRAN 2008 database (Rothman et al., 2009). The water vapor (H<sub>2</sub>O) *a priori* profile is interpolated from the NCEP 6-hourly reanalysis data and *a priori* profiles of other gases are from the Whole-Atmosphere Community Climate Model (WACCM) v6 special run for NDACC.

Three MWs were used for CO: a strong line at 2057.7–2058 cm<sup>-1</sup> and two weak lines at 2069.56–2069.76 cm<sup>-1</sup> and 2157.5–2159.15 cm<sup>-1</sup> (Sun et al., 2018a). For HCN, two MWs were used: 3268.00–3268.38 cm<sup>-1</sup> and 3287.00–3287.48 cm<sup>-1</sup> (Mahieu et al., 1997; Lutsch et al., 2016; Notholt et al., 2000). In order to minimize the cross absorption interference, profiles of ozone (O<sub>3</sub>) and nitrous oxide (N<sub>2</sub>O) and columns of H<sub>2</sub>O, carbonyl sulfide (OCS) and CO<sub>2</sub> are simultaneously retrieved in addition to the CO profile. Profile of H<sub>2</sub>O and columns of O<sub>3</sub>, C<sub>2</sub>H<sub>2</sub>, and CH<sub>4</sub> are simultaneously retrieved in addition to the HCN profile. No de-weighting SNR is used for HCN and a de-weighting SNR of 500 is used in the three MWs for CO.

The diagonal elements of *a priori* profile covariance matrices  $S_a$  are set to standard deviation of the WACCM v6 special run for NDACC, and its non-diagonal elements are set to zero. The diagonal elements of the measurement noise covariance matrices  $S_e$  are set to the inverse square of the SNR calculated from each individual spectrum and its non-diagonal elements are set to zero. The measured instrument line shape (ILS) is included in the retrieval (Hase, 2012; Sun et al., 2018a).

Table 1. Retrieval inputs used for CO and HCN.

Gases	CO	HCN
Code	SFIT4 v 0.9.4.4	SFIT4 v 0.9.4.4
Spectroscopic parameters	HITRAN 2008	HITRAN 2008
P, T, H <sub>2</sub> O profiles	NCEP reanalysis data	NCEP reanalysis data
A priori profiles of all gases except H <sub>2</sub> O	WACCM v6	WACCM v6
Micro windows for profile retrievals (cm <sup>-1</sup> )	2057.7 - 2058 2069.56–2069.76 2157.5–2159.15	3268.00 – 3268.38 3287.00–3287.48
Retrieved interfering gases	O <sub>3</sub> , N <sub>2</sub> O, CO <sub>2</sub> , OCS, H <sub>2</sub> O	H <sub>2</sub> O, O <sub>3</sub> , C <sub>2</sub> H <sub>2</sub> , CH <sub>4</sub>
SNR for de-weighting	500	None
$S_a$	WACCM v6 standard deviation	WACCM v6 standard deviation
$S_e$	SNR calculated from each individual spectrum within 2526.23 – 2526.62	SNR calculated from each individual spectrum within 3381.16 – 3381.54
ILS	LINEFIT145 analysis	LINEFIT145 analysis
Error analysis	Systematic error: line intensity, line pressure broadening, line temperature broadening, solar zenith angle, background curvature, solar line strength, optical path difference, field of view, phase	
	Random error: -Measurement error -Smoothing error -Interference errors: interfering species, retrieval parameters - Other errors: zero level, temperature	

### 2.1.3 Averaging kernels and error budget

The partial column averaging kernels of CO and HCN at selected layers are shown in Fig. 2. The CO averaging kernels have three maxima at the surface, 7 km, and 14 km, respectively. The HCN averaging kernels have only one maxima at 10 km. Both CO and HCN retrievals show good vertical sensitivity in the whole troposphere where CO exhibits the best sensitivity with two maxima in the troposphere (Sun et al., 2018a). Typical degrees of freedom (DOFS) obtained at Hefei (32°N) over the total atmosphere for CO and HCN are about  $2.8 \pm 0.3$  ( $1\sigma$ ) and  $1.3 \pm 0.2$  ( $1\sigma$ ), respectively (Table 2). In this study, only partial columns of CO and HCN within a broad layer between surface and 15 km are considered. The selected layer corresponds roughly to the total troposphere over eastern China, as the mean tropopause height calculated from NCEP reanalysis data is around 15 km over four seasons. The selected layer corresponds to  $2.3 \pm 0.2$  ( $1\sigma$ ) and  $1.0 \pm 0.1$  ( $1\sigma$ ) of DOFS for CO and HCN, respectively.

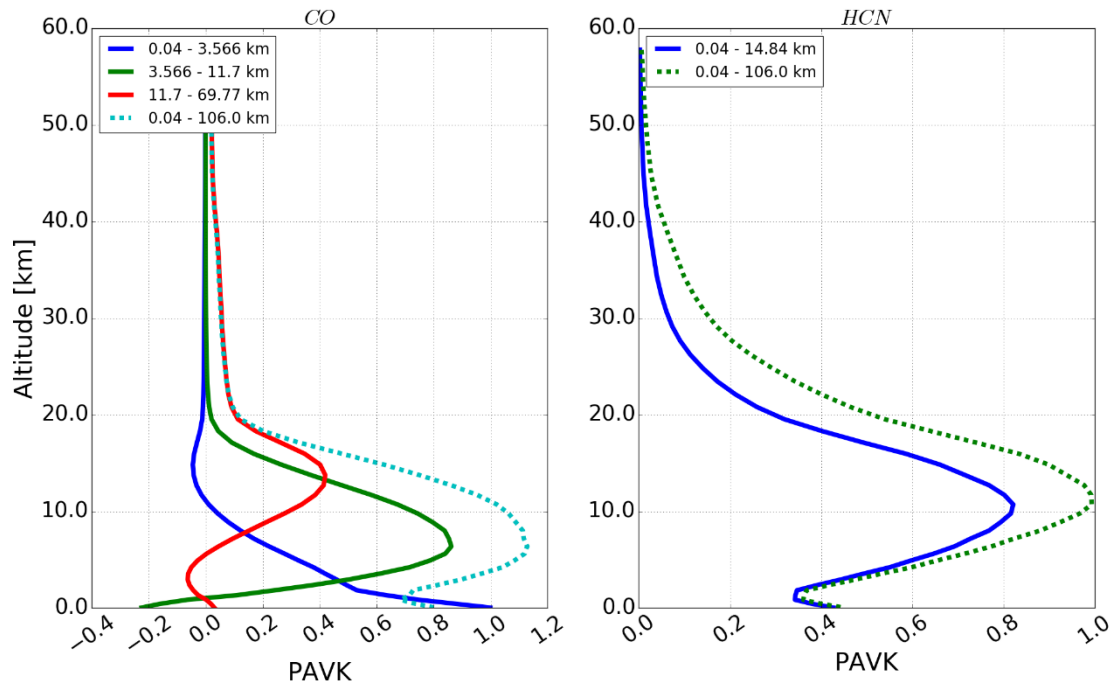


Fig. 2 Partial column averaging kernels (PAVKs) (ppmv / ppmv) for CO and HCN retrievals.

We calculated the error budget following the formalism of Rodgers, 2000, and separated all error items into systematic error or random error depending on whether they are constant over consecutive measurements, or vary randomly. Table 2 summarizes the random, the systematic, and the combined error budget of tropospheric CO and HCN columns. The error items included in the error budget are listed in Table 1. For CO, the major systematic error is line intensity uncertainty, and the major random error are zero level uncertainty and temperature uncertainty. For HCN, the major systematic error are line intensity uncertainty and line pressure broadening uncertainty, the major random error are smoothing error and measurement error. Total retrieval errors for tropospheric CO and HCN columns between surface and 15 km are estimated to be 8.3 and 14.2%, respectively.

Table 2. Retrieval error budgets and DOFs for tropospheric CO and HCN.

Gases	CO	HCN
Temperature uncertainty	2.5%	0.2%
Zero level uncertainty	5.2%	1.5%
Retrieval parameters uncertainty	< 0.1%	2.0%
Interfering species uncertainty	< 0.1%	1.3%
Measurement Error	< 0.1%	6.8%
Smooth Error	0.1%	11.0%
<b>Total Random Error</b>	<b>5.7%</b>	<b>13.2%</b>
Background curvature uncertainty	< 0.1%	*
Optical path difference uncertainty	< 0.1%	< 0.1%
Field of view uncertainty	< 0.1%	< 0.1%
Solar line strength uncertainty	< 0.1%	< 0.1%
Phase uncertainty	*	< 0.1%
Solar zenith angle uncertainty	0.1%	< 0.1%
Line temperature broadening uncertainty	0.13%	0.3%
Line pressure broadening uncertainty	0.87%	3.5%
Line intensity uncertainty	6.0%	3.7%
<b>Total Systematic Error</b>	<b>6.1%</b>	<b>5.1%</b>
<b>Total Errors</b>	<b>8.3%</b>	<b>14.2%</b>
<b>DOFS (-)</b>	<b>2.2</b>	<b>1.0</b>

\* Not included into error budget since they are retrieved together with the target gas

## 2.2 GEOS-Chem tagged simulation

A GEOS-Chem tagged CO simulation is used to interpret the influence of biomass burning sources on HCN tropospheric columns at Hefei (32°N) (Bey et al., 2001). The GEOS-Chem simulation was designed following Lutsch et al., 2019 and is described here. GEOS-Chem version 12.2.1 and the Goddard Earth Observing System-Forward Processing (GEOS-FP) product with

1 assimilated meteorological data observations from the NASA Global Modeling and Assimilation  
 2 Office (GMAO) was used. For driving the GEOS-Chem model, the GEOS-FP meteorological data  
 3 with a native horizontal resolution of  $0.25^\circ$  latitude  $\times$   $0.3125^\circ$  longitude were downgraded to  $2^\circ$   
 4 latitude  $\times$   $2.5^\circ$  longitude and a vertical resolution of 72 hybrid levels (extending from surface to  
 5 0.01 hPa). The temporal resolution of surface variables and boundary layer height are 1hr and other  
 6 variables are 3 hr.

7 A 1-year spin-up from July 2014 to July 2015 was used to initialize the simulation. Time-steps  
 8 of 1 hr and 10 min for the chemical and transport operators, respectively, were used. Biomass  
 9 burning emissions are from Global Fire Assimilation System (GFAS) v1.2 (Kaiser et al., 2012;  
 10 Giuseppe et al., 2018). GFASv1.2 emissions have a  $0.1^\circ \times 0.1^\circ$  horizontal resolution with 1-hourly  
 11 temporal resolution. Global anthropogenic and biofuel emissions are from the Community  
 12 Emissions Data System (CEDS) inventory (Hoesly et al., 2018). In particular, the latest Multi-  
 13 resolution Emission Inventory for China (MEIC) is used to provide Chinese anthropogenic  
 14 emissions (Li et al., 2017). Biogenic emissions of precursor VOCs are from the Model of Emissions  
 15 of Gases and Aerosols from Nature (MEGANv2.1; Guenther et al., 2012) and biofuel emissions are  
 16 from Yevich and Logan (2003). The OH fields were obtained from monthly mean OH  
 17 concentrations archived from a previous full-chemistry simulation. GEOS-Chem releases surface  
 18 emissions assuming a uniform distribution in the boundary layer, and boundary layer mixing is  
 19 implemented using the non-local mixing scheme of Holtslag and Boville (1993). Biomass emissions  
 20 are released by uniformly distributing emissions from the surface to the mean altitude of maximum  
 21 injection based on the injection height information in GFASv1.2 as described in Rémy et al. (2017).

22 GEOS-Chem version 12.2.1 tagged CO simulation includes the improved secondary CO  
 23 production scheme of Fisher et al. (2017), which assumes production rates of CO from CH<sub>4</sub> and  
 24 NMVOC (non-methane volatile organic compounds) oxidation from a GEOS-Chem full-chemistry  
 25 simulation. The tracers of anthropogenic, biomass burning, CH<sub>4</sub> and NMVOC oxidations are  
 26 implemented following the standard GEOS-Chem tagged CO simulation (<http://geos-chem.org/>, last  
 27 accessed on 8 April 2020). In this study, we only investigate the influence from the biomass burning  
 28 sources. The regional definition of all biomass burning tracers are shown in Fig. 1 and tabulated in  
 29 Table 3.

30 Table 3. Regional definitions of all biomass burning tracers implemented in the standard GEOS-Chem tagged CO  
 31 simulation

No.	Tracer	Description	Region
1	SA	Biomass burning CO emitted over South America	112.5°W - 32.5°W; 56°S - 24°N
2	AF	Biomass burning CO emitted over Africa	17.5°W - 70.0°E; 48.0°S - 36.0°N
3	SEAS	Biomass burning CO emitted over Southeast Asia	70.0°E - 152.5°E; 8.0°N - 45.0°N
4	OCE	Biomass burning CO emitted over Oceania	70.0°E - 170.0°E; 90.0°S - 8.0°N
5	EUBA	Biomass burning CO emitted over Europe and Boreal Asia	17.5°W - 72.5°E; 36.0°N - 45.0°N and 17.5°W - 172.5°E; 45.0°N - 88.0°N
6	NA	Biomass burning CO emitted over North America	173°W - 50°W; 24.0°N - 88.0°N

### 32 2.3 Potential source contribution function

33 We used the potential source contribution function (PSCF) analysis method to identify air  
 34 masses associated with high levels of air pollutants. The PSCF assumes that back trajectories  
 35 arriving at times of higher concentrations likely point to the more significant pollution directions  
 36 (Ashbaugh et al., 1985). PSCF has been applied in many studies to locate air masses associated with  
 37 high levels of air pollutants (Kaiser et al., 2007; Dimitriou and Kassomenos, 2015; Yin et al., 2017).  
 38 In this study, PSCF values were calculated using back trajectories that were calculated by HYSPLIT.  
 39 The top of the model was set to 10 km. The PSCF values for the grid cells in the study domain were  
 40 based on a count of the trajectory segment that terminated within each cell (Ashbaugh et al., 1985).  
 41 The number of endpoints that fall in the  $i^{th}$  cell is designated  $n_i$ . The number of endpoints for the  
 42 same cell having arrival times at the sampling site corresponding to concentrations higher than an  
 43 arbitrarily set criterion is defined to be  $m_i$ . In this study, we calculated the PSCF values based on  
 44 trajectories corresponding to concentrations that exceeded the monthly mean level of tropospheric

1 HCN column during measurement. The PSCF value for the  $ij^{\text{th}}$  cell is then defined as:

$$2 \quad PSCF_{ij} = m_{ij}/n_{ij} \quad (1)$$

3 The unitless PSCF value can be interpreted as the conditional probability that the  
4 concentrations of a given analyte greater than the criterion level are related to the passage of air  
5 parcels through the  $ij^{\text{th}}$  cell during transport to the receptor site. That is, cells with high PSCF values  
6 are associated with the arrival of air parcels at the receptor site that have concentrations of the  
7 analyte higher than the criterion value. These cells are indicative of areas of ‘high potential’  
8 contributions for the constituent.

9 Identical  $PSCF_{ij}$  values can be obtained from cells with very different counts of back-trajectory  
10 points (e.g., grid cell A with  $m_{ij} = 400$  and  $n_{ij} = 800$  and grid cell B with  $m_{ij} = 4$  and  $n_{ij} = 8$ ). In this  
11 extreme situation grid cell A has 100 times more air parcels passing through than grid cell B.  
12 Because of the sparse particle count in grid cell B, the PSCF values are more uncertain. To account  
13 for the uncertainty due to low values of  $n_{ij}$ , the PSCF values were scaled by a weighting function  
14  $W_{ij}$  (Polissar et al., 1999). The weighting function reduced the PSCF values when the total number  
15 of endpoints in a cell was less than approximately 3 times the average value of the end points per  
16 cell. In this case,  $W_{ij}$  was set as follows:

$$17 \quad W_{ij} = \begin{cases} 1.00 & n_{ij} > 3N_{ave} \\ 0.70 & 3N_{ave} > n_{ij} > 1.5N_{ave} \\ 0.42 & 1.5N_{ave} > n_{ij} > N_{ave} \\ 0.05 & N_{ave} > n_{ij} \end{cases} \quad (2)$$

18 where  $N_{ave}$  represents the mean  $n_{ij}$  of all grid cells. The weighted PSCF values were  
19 obtained by multiplying the original PSCF values by the weighting factor.

### 20 3 FTIR time series and comparisons with NDACC counterparts

21 The new HCN data are compared with the concurrent measurements regularly measured at  
22 eleven NDACC stations to investigate the representativeness of the observation site at Hefei (32°N)  
23 in polluted eastern China. These NDACC stations cover over a wide latitude range from 77.8°S to  
24 78.9°N and a wide longitude range from 79°W to 170°E (<http://www.ndaccdemo.org/>, last access  
25 on 19 July 2019). Most of these NDACC stations use the same instrument and retrieval algorithm  
26 as those of Hefei (32°N). Alternatively, the high resolution spectrometers Bruker 125M, 120HR, or  
27 Bomem DA8 and the retrieval algorithm PROFFIT are used in other stations. It has been  
28 demonstrated that the profiles derived from these different instruments and algorithms are in  
29 excellent agreement (Hase et al., 2004; De Maziere et al., 2018). In addition, we show the time  
30 series of tropospheric CO columns, also measured with FTIR spectrometer, because we will discuss  
31 the correlation between HCN and CO, and quantify the influence of biomass burning sources on  
32 HCN columns at Hefei (32°N) by using a tagged CO simulation. The upper limit of 15 km is above  
33 the tropopause at most of the NDACC stations. For most NDACC stations, the surface – 15 km  
34 layer is a mixture of troposphere and a part of stratosphere. However, we did not find major changes  
35 in the results of this study when choosing a lower upper limit such as 12 km. Thus we have chosen  
36 the same upper limits for all stations. The geolocations of all FTIR stations and their seasonal  
37 maximum, minimum and variabilities are summarized in Table 4.

38 Table 4. Tropospheric HCN and CO columns at Hefei (32°N), China from 2015 to 2018 alongside those of the  
39 NDACC FTIR stations. All stations are organised as a function of decreasing latitude.

Station	Location (Lon., Lat., Alt. in km)	Instrument	Algorithm	Maximum (molecules cm <sup>-2</sup> )		Minimum (molecules cm <sup>-2</sup> )	
				HCN (10 <sup>15</sup> )	CO (10 <sup>18</sup> )	HCN (10 <sup>15</sup> )	CO (10 <sup>18</sup> )
Ny Alesund	(12°E, 79°N, 0.02)	125HR	SFIT4	5.94 ± 1.20 (August)	2.11 ± 0.11 (March)	3.75 ± 0.37 (March)	1.56 ± 0.12 (July)
Kiruna	(20°E, 68°N, 0.42)	125HR	PROFFIT	5.81 ± 0.58 (August)	2.1 ± 0.01 (January)	2.43 ± 0.27 (January)	1.45 ± 0.09 (July)
Bremen	(9°E, 53°N, 0.03)	125HR	SFIT4	6.11 ± 0.87 (August)	2.32 ± 0.13 (March)	2.85 ± 0.25 (January)	1.63 ± 0.19 (July)
Jungfraujoch	(8°E, 46.5°N, 3.58)	125HR	SFIT4	4.68 ± 0.63 (May)	1.14 ± 0.08 (March)	2.1 ± 0.29 (February)	0.88 ± 0.08 (July)
Toronto	(79°W, 44°N, 0.17)	Bomem DA8	SFIT4	5.92 ± 1.13 (May)	2.19 ± 0.15 (April)	3.12 ± 1.02 (November)	1.74 ± 0.1 (October)
Rikubetsu	(144°E, 43°N, 0.38)	125HR	SFIT4	7.0 ± 1.92 (May)	2.32 ± 0.31 (March)	2.86 ± 0.44 (February)	1.79 ± 0.14 (October)
<b>Hefei</b>	<b>(117°E, 32°N, 0.03)</b>	<b>125HR</b>	<b>SFIT4</b>	<b>9.8 ± 0.78 (May)</b>	<b>3.38 ± 0.43 (February)</b>	<b>7.16 ± 0.75 (November)</b>	<b>2.29 ± 0.48 (July)</b>



Izana	(16°W, 28°N, 2.37)	125HR	PROFFIT	$5.33 \pm 1.2$ (May)	$1.41 \pm 0.14$ (April)	$2.59 \pm 0.28$ (October)	$1.1 \pm 0.08$ (October)
Mauna Loa	(24°W, 20°N, 3.40)	125M	SFIT4	$4.49 \pm 1.8$ (April)	$1.36 \pm 0.31$ (April)	$2.07 \pm 0.43$ (August)	$0.8 \pm 0.04$ (August)
La Reunion Maido	(55°E, 21°S, 2.16)	125HR	SFIT4	$6.91 \pm 2.45$ (November)	$1.46 \pm 0.17$ (October)	$2.56 \pm 0.48$ (May)	$1.0 \pm 0.1$ (April)
Lauder	(170°E, 45°S, 0.37)	120HR	SFIT4	$5.29 \pm 1.18$ (November)	$1.28 \pm 0.19$ (October)	$1.94 \pm 0.28$ (July)	$0.89 \pm 0.09$ (February)
Arrival Heights	(167°E, 78°S, 0.2)	120HR	SFIT4	$3.22 \pm 0.51$ (February)	$1.0 \pm 0.04$ (October)	$1.78 \pm 0.21$ (September)	$0.67 \pm 0.03$ (April)

### 3.1 Seasonal variation

The monthly means of the tropospheric CO and HCN columns at the twelve FTIR stations are shown in Fig. 3. As commonly observed at Hefei (32°N), three monthly mean peaks are evident for tropospheric HCN and CO columns. The magnitude of the tropospheric HCN peak at Hefei (32°N) in May > September > December, while for tropospheric CO column, the magnitude of the peak at Hefei (32°N) in February > September > December. Note that the largest seasonal peak of HCN occurs in May which is 3 months later than that of CO which occurs in February, but the other two seasonal peaks for both species occur in the same months, i.e., in September and December respectively. Otherwise, their seasonal cycles show similarities.

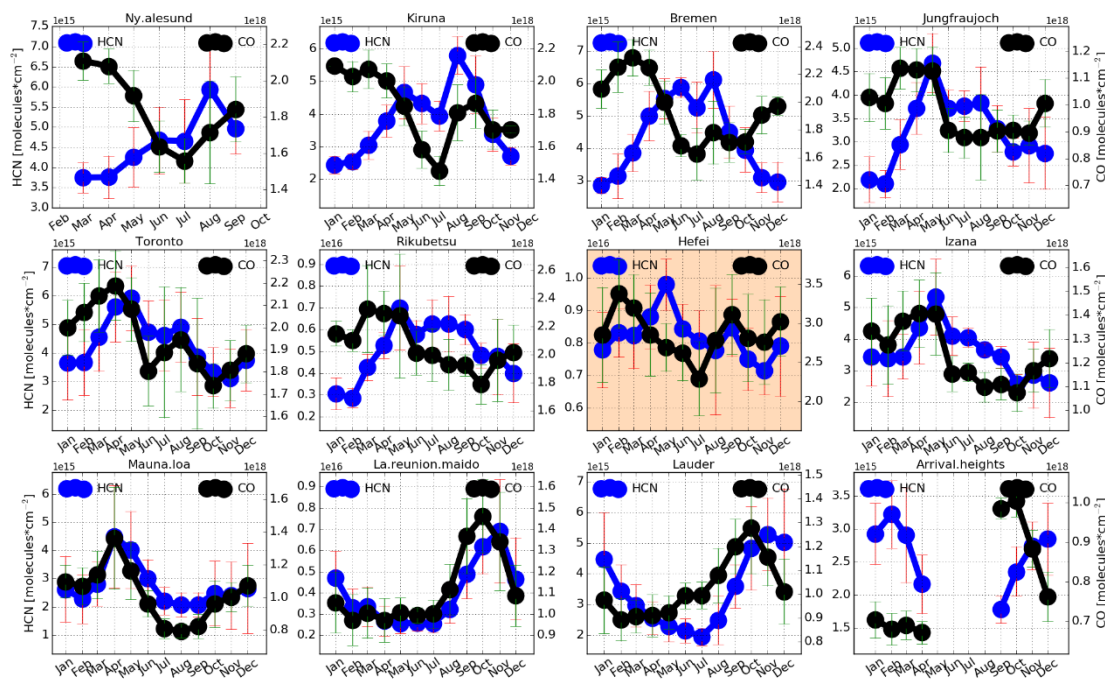


Fig. 3. Monthly means of the tropospheric CO and HCN columns at Ny Alesund, Kiruna, Bremen, Jungfrauoch, Toronto, Rikubetsu, Hefei, Izana, Mauna Loa, La Reunion Maido, Lauder, and Arrival Heights from 2015 to 2018. Vertical error bars represent  $1\sigma$  within that month. All stations are organised as a function of decreasing latitude.

The tropospheric HCN and CO columns at Hefei (32°N) are higher than the NDACC FTIR observations (see Fig. S2). The tropospheric HCN column reached a maximum of  $(9.8 \pm 0.78) \times 10^{15}$  molecules/cm<sup>2</sup> in May and a minimum of  $(7.16 \pm 0.75) \times 10^{15}$  molecules/cm<sup>2</sup> in November. The tropospheric CO column reached a maximum of  $(3.38 \pm 0.43) \times 10^{18}$  molecules/cm<sup>2</sup> in February and a minimum of  $(2.29 \pm 0.48) \times 10^{18}$  molecules/cm<sup>2</sup> in July (Table 4). In comparison, the seasonal maxima and minima of tropospheric HCN columns at the selected NDACC FTIR stations varied over  $(3.22 \pm 0.51)$  to  $(7.0 \pm 1.92) \times 10^{15}$  molecules/cm<sup>2</sup> and  $(1.78 \pm 0.21)$  to  $(3.75 \pm 0.37) \times 10^{15}$  molecules/cm<sup>2</sup>, respectively. The seasonal maxima and minima of tropospheric CO columns at the selected NDACC FTIR stations varied over  $(1.0 \pm 0.04)$  to  $(2.32 \pm 0.31) \times 10^{18}$  molecules/cm<sup>2</sup> and  $(0.67 \pm 0.03)$  to  $(1.79 \pm 0.14) \times 10^{18}$  molecules/cm<sup>2</sup>, respectively (Table 4).

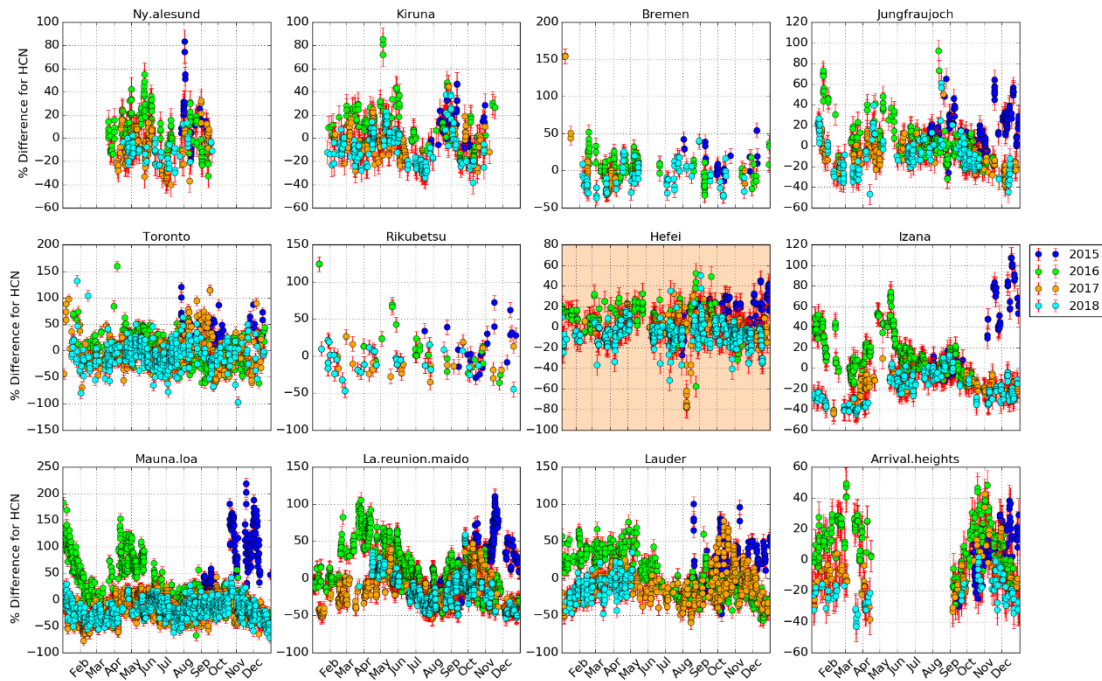
In the northern hemisphere, the phase of the seasonal maxima for tropospheric HCN columns generally occur in spring or summer, and for CO occur in winter or spring. While in the southern hemisphere, the phase of the seasonal maxima for both tropospheric HCN and CO columns occur in autumn or winter.

### 3.2 Interannual variability and enhancement



1 In order to study the interannual variability of HCN and CO, fractional differences in the  
 2 tropospheric HCN and CO columns relative to their seasonal mean values represented by the cosine  
 3 fitting at the twelve FTIR stations are shown in Fig.4 and Fig.5, respectively. Enhancements of both  
 4 tropospheric HCN and CO columns between September 2015 and July 2016 at Hefei (32°N) were  
 5 observed compared to the same period of measurements in other years. For HCN, the magnitude of  
 6 the enhancement ranges from 5 to 46% with an average of 26%. The significant enhancements  
 7 occurred in December 2015 and May 2016 with peaks of 46% and 38%, respectively. By contrast,  
 8 the magnitude of the enhancement in tropospheric CO column at Hefei (32°N) between September  
 9 2015 and July 2016 ranges from 4 to 59% with an average of 27%. The tropospheric CO columns  
 10 were elevated over its seasonal means by more than 20% from March to April 2016. In addition, an  
 11 enhancement magnitude of more than 40% were occasionally observed in August and September  
 12 for both HCN and CO at Hefei (32°N).

13 The enhancements of both tropospheric HCN and CO columns within the same period were  
 14 also observed at the selected NDACC stations except Ny Alesund (79°N) and Kiruna (68°N). The  
 15 winter enhancements were not shown over Ny Alesund (79°N) and Kiruna (68°N) because of the  
 16 polar night in the Arctic which interrupted the observations in winter. The magnitude of the  
 17 enhancement in tropospheric HCN column at the selected NDACC stations between September  
 18 2015 and July 2016 ranges from 3 to 213%, and for CO ranges from 4 to 62%.



19  
 20 Fig.4. Fractional difference in the partial columns (surface - 15 km) of HCN from 2015 to 2018 at Ny Alesund,  
 21 Kiruna, Bremen, Jungfraujoch, Toronto, Rikubetsu, Hefei, Izana, Mauna Loa, La Reunion Maito, Lauder, and  
 22 Arrival Heights relative to their seasonal mean values. Vertical error bars represent the estimated retrieval errors.  
 23 All stations are organised as a function of decreasing latitude.

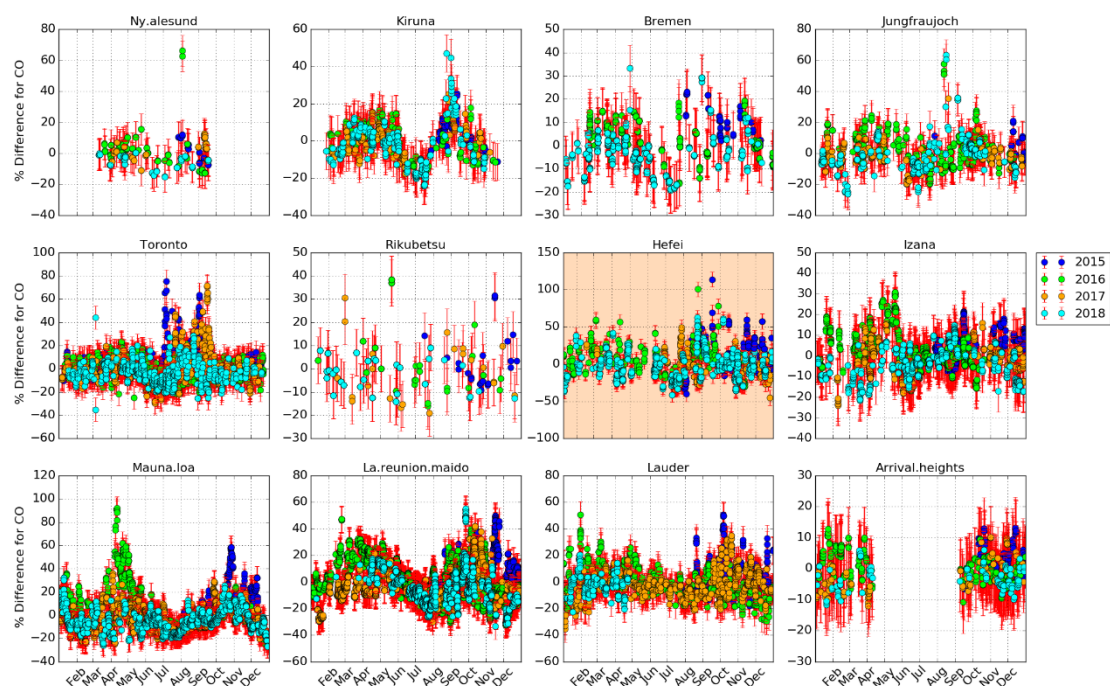


Fig.5. The same as Fig.4 but for CO.

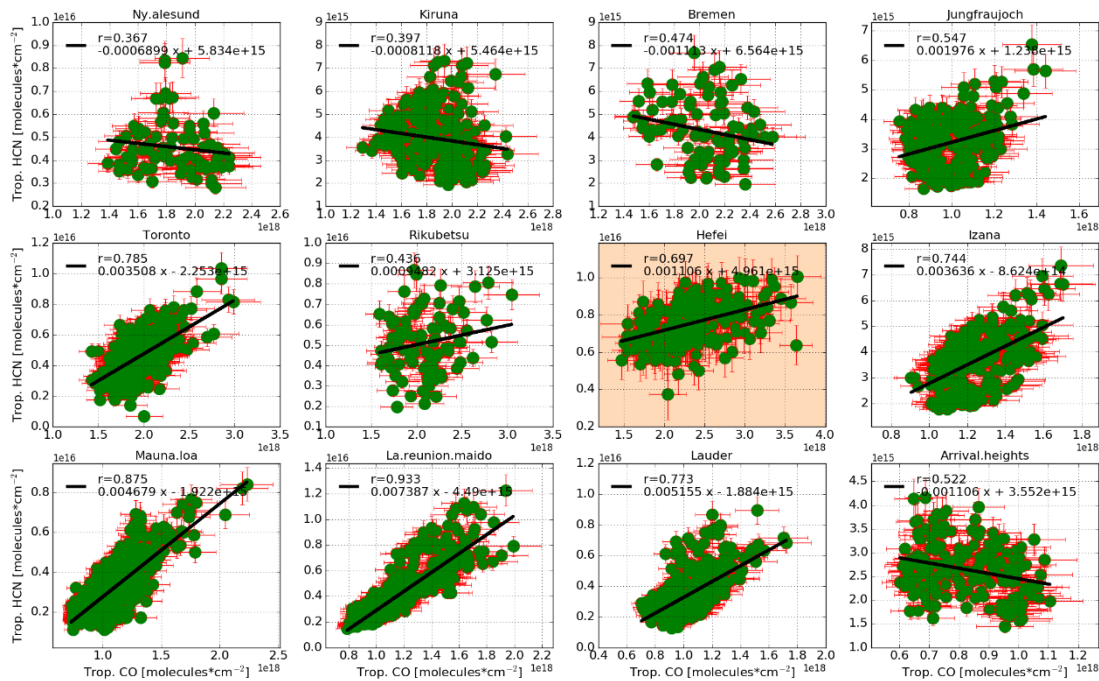
### 3.3 Correlation with CO and enhancement ratios

The tropospheric HCN columns at the twelve FTIR stations from 2015 to 2018 have been plotted against the coincident CO partial columns (Fig.6). In Fig.7, the correlations between the tropospheric HCN and CO columns at Hefei (32°N) for all spectra recorded throughout the year (gray dots) and those recorded within the selected periods (green dots) are compared. We followed the least squares procedure of York et al., 2004 to fit the coincident measurements using a linear regression, and incorporated the errors in both ordinal and abscissa coordinates into the uncertainty estimation.

Biomass burning is the dominant source of HCN and industrial emissions only contribute additional minor sources (Bange and Williams, 2000; Holzinger et al., 1999; Lobert et al., 1990). In contrast, anthropogenic, biomass burning, CH<sub>4</sub> and NMVOC oxidations are major sources of CO and their contributions are season and location dependent. Therefore, the correlation between HCN and CO tropospheric column is also season and site dependent. High correlation of these two species is supposed to be observed if biomass burning dominates the CO variability, and vice versa. For the period of 2015 to 2018 in this study, moderate overall correlations between HCN and CO tropospheric columns were present at Jungfrauoch (47°N) and Rikubetsu (43°N), and negative overall correlations were present at Ny Alesund (79°N), Kiruna (68°N), Bremen (53°N), and Arrival Heights (78°S). However, high correlation of these two species were seen at Toronto (44°N), Hefei (32°N), Izana (28°N), Mauna Loa (20°N), La Reunion Maido (21°S), and Lauder (45°S) throughout the year probably because the portion of the fire-affected seasonal measurements at these stations are larger than those at other stations (Fig.6). For the measurements at Hefei (32°N), the high correlations between HCN and CO tropospheric columns deduced from the measurements without March and April ( $R=0.67$ , Fig.7 (a)), in May ( $R=0.69$ , Fig.7 (b)), in September ( $R=0.77$ , Fig.7 (c)), and in December ( $R=0.65$ , Fig.7 (d)) are consistent with that deduced from all measurements ( $R=0.70$ ) (Table 5). However, the correlation slope for the May, September, and December tropospheric columns differ from the annual one, indicating different biomass burning sources in different periods.

For fire-affected measurements, the slope  $\Delta\text{HCN}/\Delta\text{CO}$  defined as enhancement ratio ( $\text{EnhR}_{\text{HCN}}$ ) is an effective quantity to identify biomass burning emissions (Holzinger et al., 1999; Lutsch et al., 2016; Rinsland et al., 2002; Viatte et al., 2015; Vigouroux et al., 2012; Zhao et al., 2000). Depending on the burnt biomaterials, fire type, the phase of the fire, and the travel time of the plumes, the reported  $\text{EnhR}_{\text{HCN}}$  varied by 2 orders of magnitude. The mean  $\text{EnhR}_{\text{HCN}}$  of  $1.34 \times 10^{-3}$  at Hefei (32°N) falls between the wide range of the HCN/CO ratios measured in laboratory ( $0.4 - 7.1 \times 10^{-3}$  in the work of (Yokelson et al., 1997) and  $0.4 - 2.6 \times 10^{-3}$  in the work of (Holzinger et al., 1999), and  $0.94 - 7.4 \times 10^{-3}$  in the NDACC FTIR measurements (Fig. 6). The mean  $\text{EnhR}_{\text{HCN}}$  at Hefei (32°N) is close

1 to that at Rikubetsu (43°N) indicating that these two Asian stations share similar biomass burning  
 2 sources throughout the year. The mean  $\text{EnhR}_{\text{HCN}}$  at Hefei (32°N) is lower than those measured at  
 3 Jungfraujoch (47°N), Toronto (44°N), Izana (28°N), Mauna Loa (20°N), Lauder (45°S), and La  
 4 Reunion Maito (21°S) because the emissions of crop residue burning which dominates the HCN  
 5 enhancements at Hefei (32°N) is lower than those of the boreal or tropical forest burning, which  
 6 account for the HCN enhancements at aforementioned NDACC stations (Akagi et al., 2011; Akagi  
 7 et al., 2012; Rinsland et al., 2007; Vigouroux et al., 2012). On the other hand, the Hefei (32°N)  
 8 site is located in the densely populated part of China, therefore emissions of fossil fuel combustion such  
 9 as automobile exhaust and industrial processes could elevate the CO background level and hence  
 10 lessen the  $\text{EnhR}_{\text{HCN}}$ .

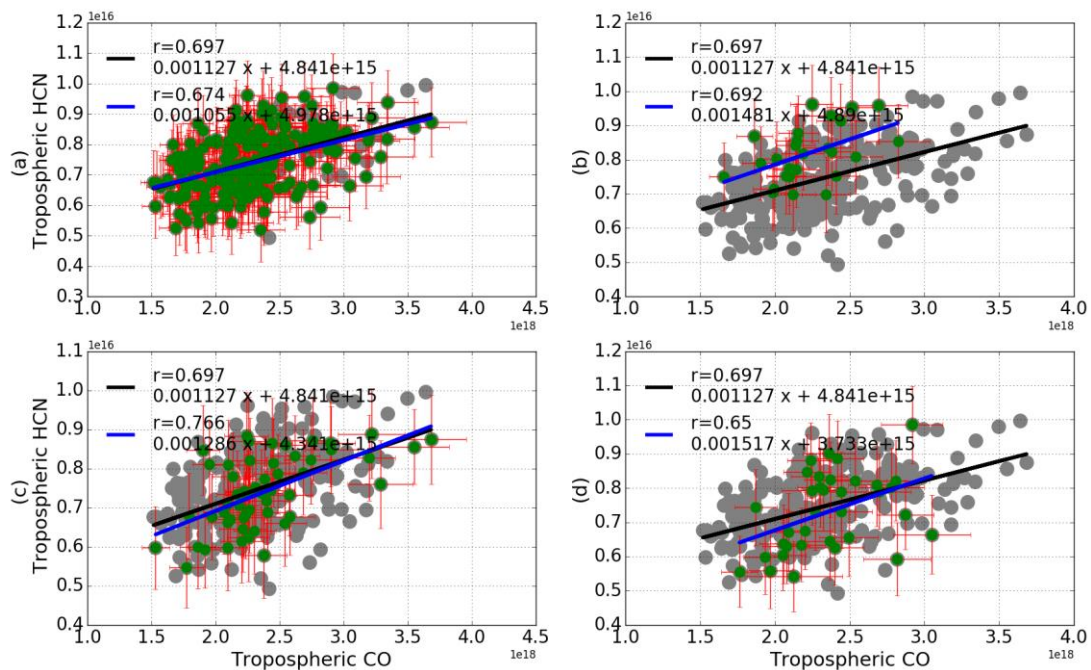


11  
 12 Fig. 6. Correlation plots of daily mean partial columns (surface - 15 km) of HCN versus CO (molecules/cm<sup>2</sup>). The  
 13 linear equation of the fit and the resulting correlation coefficient  $r$  are shown. The black line is a linear least-squares  
 14 fit of respective data. All stations are organised as a function of decreasing latitude. Error bars represent the retrieval  
 15 uncertainties.

16 Table 5. Correlation between HCN and CO tropospheric columns within each selected period at Hefei (32°N),  
 17 China.  $N$  is the number of points,  $R$  is the correlation coefficient and  $\text{EnhR}_{\text{HCN}}$  is the enhancement ratio.

Gas	Period	without March and April	May	September	December	Mean
HCN	$N$	239	26	56	35	-
	$R$	0.67	0.69	0.77	0.65	0.7
	$\text{EnhR} \times 10^{-3}$	1.06	1.48	1.29	1.52	1.34





1  
2 Fig. 7. Correlation plots of daily mean tropospheric columns of HCN versus CO (molecules/cm<sup>2</sup>) at Hefei (32°N).  
3 The gray dots represent all measurements and the green dots represent the measurements within the selected period:  
4 (a) measurements without March and April; (b) measurements in May; (c) measurements in September; (d)  
5 measurements in December. The linear equation of the fit and the resulting correlation coefficient  $r$  are shown. The  
6 black line is a linear least-squares fit of the gray data and the blue line is for the green data. Error bars represent the  
7 retrieval uncertainties.

#### 8 4 Source attribution

9 In order to determine what drives the seasonality and interannual variability of tropospheric  
10 HCN in eastern China, it is necessary to match the observed time series with actual biomass burning  
11 events, and show that the generated plumes are capable of travelling to the observation site. We did  
12 this by using various independent data sets.

13 1. The 1-hourly instantaneous CO VMR (volume mixing ratio) profiles of the tracers listed in  
14 Table 3 provided by a GEOS-Chem tagged CO simulation performed as described in Section 2.2.

15 2. The global fire atlas data archived by the Fire Information for Resource Management System  
16 (FIRMS) which generates fire information from NASA's Moderate Resolution Imaging  
17 Spectroradiometer (MODIS) and NASA's Visible Infrared Imaging Radiometer Suite (VIIRS)  
18 (<https://firms.modaps.eosdis.nasa.gov/download/>, last access on 23 May 2019). We have only taken  
19 the fire number with a retrieval confidence value of larger than 60% into account.

20 3. Three dimensional kinematic back trajectories at designated elevations calculated by the Air  
21 Resources Laboratory (ARL, <http://ready.arl.noaa.gov/HYSPLIT.php>, last accessed on 23 May  
22 2019) Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model using Global  
23 Data Assimilation System (GDAS) meteorological fields (<https://ready.arl.noaa.gov/gdas1.php>, last  
24 accessed on 23 May 2019).

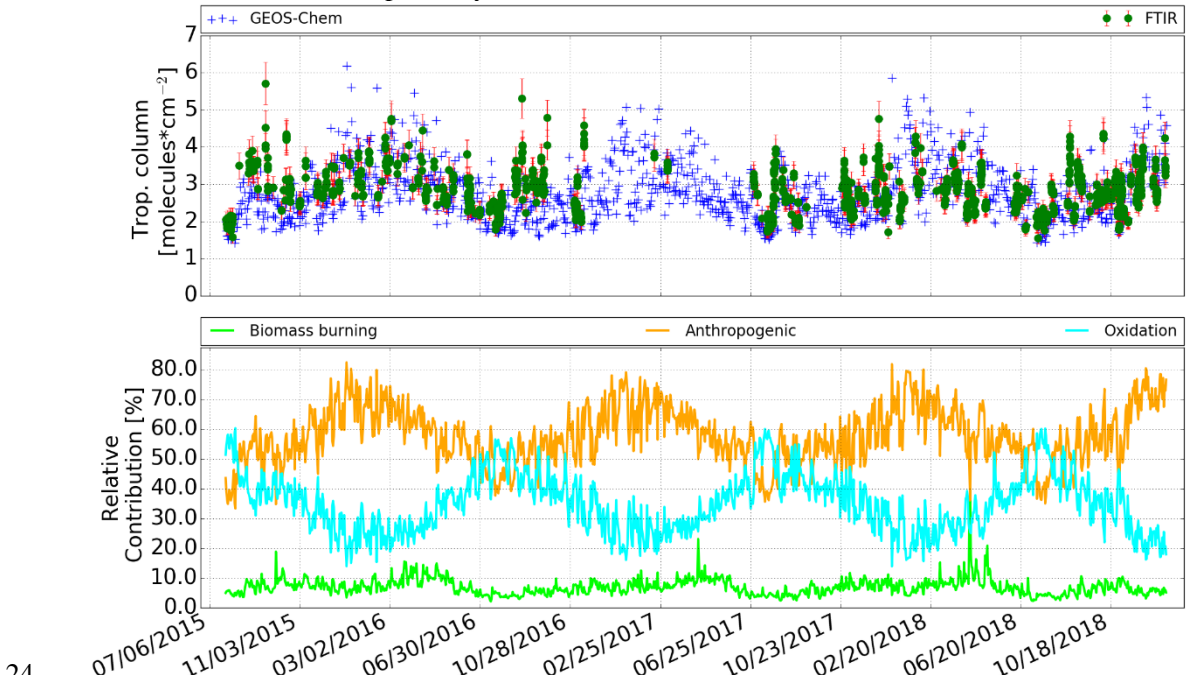
25 4. The PSCF values calculated by MeteInfo as described in Section 2.3 using HYSPLIT back  
26 trajectories (<http://meteothink.org/index.html>, last accessed on 17 December 2019).

#### 27 4.1 Attribution for the seasonality

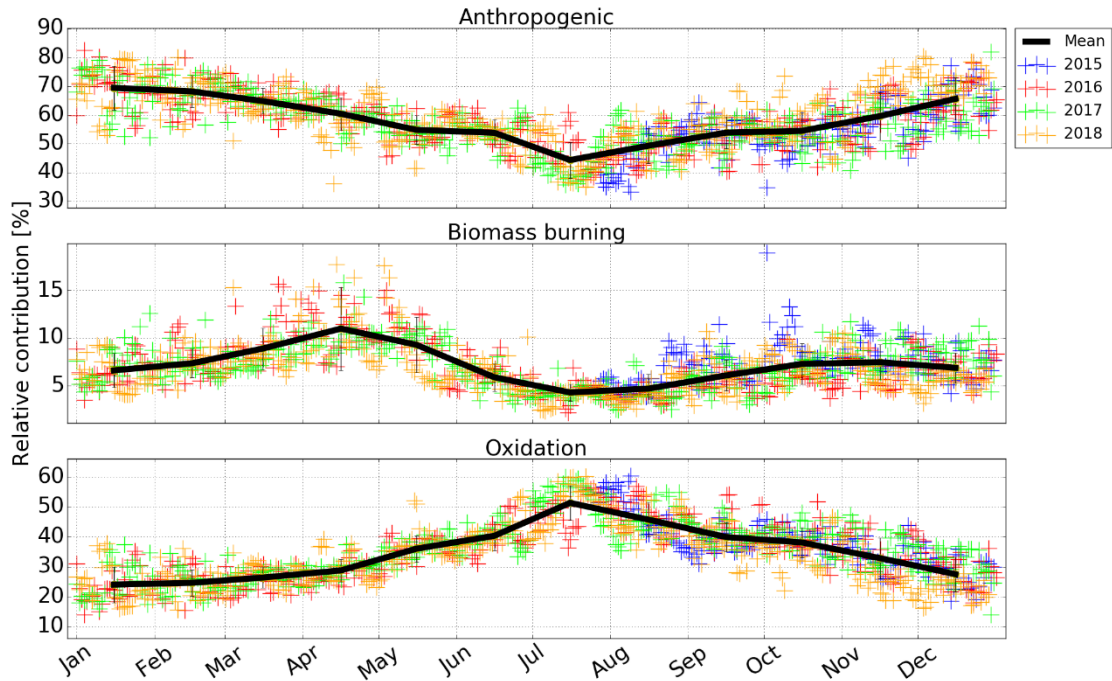
28 The GEOS-Chem tagged CO simulation provides a means of evaluating the contribution of  
29 CO from anthropogenic, biomass burning and oxidation sources to the measured CO columns at  
30 Hefei (32°N). Source attribution is performed as follows. First, the GEOS-Chem CO VMR profiles  
31 of all tracers in the grid box containing the Hefei (32°N) site were linearly interpolated and regridded  
32 onto the FTIR vertical retrieval grid. This was necessary in order to account for the differences in  
33 the vertical levels of the model and the FTIR (Barret et al., 2003). Then, the interpolated GEOS-  
34 Chem CO profiles were smoothed by the FTIR CO averaging kernel following Rodgers and Connor  
35 (2003). Finally, we compared the partial columns calculated from the smoothed GEOS-Chem CO  
36 profiles with the FTIR ones. Fig.8 shows the daily-averaged GEOS-Chem and FTIR CO  
37 tropospheric columns (surface-15 km) for the simulation period from 2015 - 2018. The relative  
38 contribution of anthropogenic, biomass burning and oxidation tracers are also shown. The GEOS-

1 Chem and FTIR CO tropospheric columns are in good agreement.  
 2 The combination of the anthropogenic source and the oxidations of CH<sub>4</sub> and NMVOCs is the  
 3 greatest contribution to the tropospheric CO column at Hefei (32°N). The magnitude of this  
 4 combination source varies over 80 to 95% throughout the year. In contrast, the magnitude of biomass  
 5 burning source varies over 5 to 20%. As shown in Fig.9, the anthropogenic, biomass burning and  
 6 oxidation sources are all seasonal dependent due to the magnitude of the emissions and the influence  
 7 of seasonally variable transport. The onset of the anthropogenic contribution begins in July with a  
 8 maximum in December. In contrast to the anthropogenic influence, the onset of the oxidation  
 9 contribution begins in January with a maximum in July, as a result of maximum NMVOC emissions  
 10 in Summer (Sun et al., 2018b). For biomass burning contribution, two onsets were observed. One  
 11 begins in January with a maximum in April and the other one begins in July with a maximum in  
 12 October.

13 After normalizing each biomass burning tracer listed in Table 3 to the total biomass burning  
 14 contribution, the normalized relative contribution of each individual biomass burning tracer to the  
 15 total biomass burning associated CO tropospheric column was obtained in Fig.10. The results show  
 16 that the seasonal maxima in May is largely due to the influence of SEAS biomass burning ( $41 \pm$   
 17  $13.1\%$ ). Moderate contributions from EUBA ( $21 \pm 9.3\%$ ) and AF ( $22 \pm 4.7\%$ ), and small  
 18 contributions from SA ( $7.8 \pm 2.9\%$ ), OCE ( $1.5 \pm 0.8\%$ ), and NA ( $7.7 \pm 1.9\%$ ) are also observed. The  
 19 seasonal maxima in September is largely due to the influence of EUBA ( $38 \pm 11.3\%$ ) and AF ( $26 \pm$   
 20  $6.7\%$ ) biomass burnings. Remaining contributions are from SA ( $5.1 \pm 2.7\%$ ), SEAS ( $14 \pm 3.3\%$ ),  
 21 OCE ( $8.9 \pm 7.4\%$ ), and NA ( $13.8 \pm 8.4\%$ ). For the seasonal maxima in December, contributions  
 22 from AF, SA, SEAS, EUBA, OCE, and NA are  $36 \pm 7.1\%$ ,  $11 \pm 1.9\%$ ,  $11 \pm 3.6\%$ ,  $21 \pm 5.2\%$ ,  $4.8 \pm$   
 23  $2.7\%$ , and  $18.7 \pm 5.2\%$ , respectively.

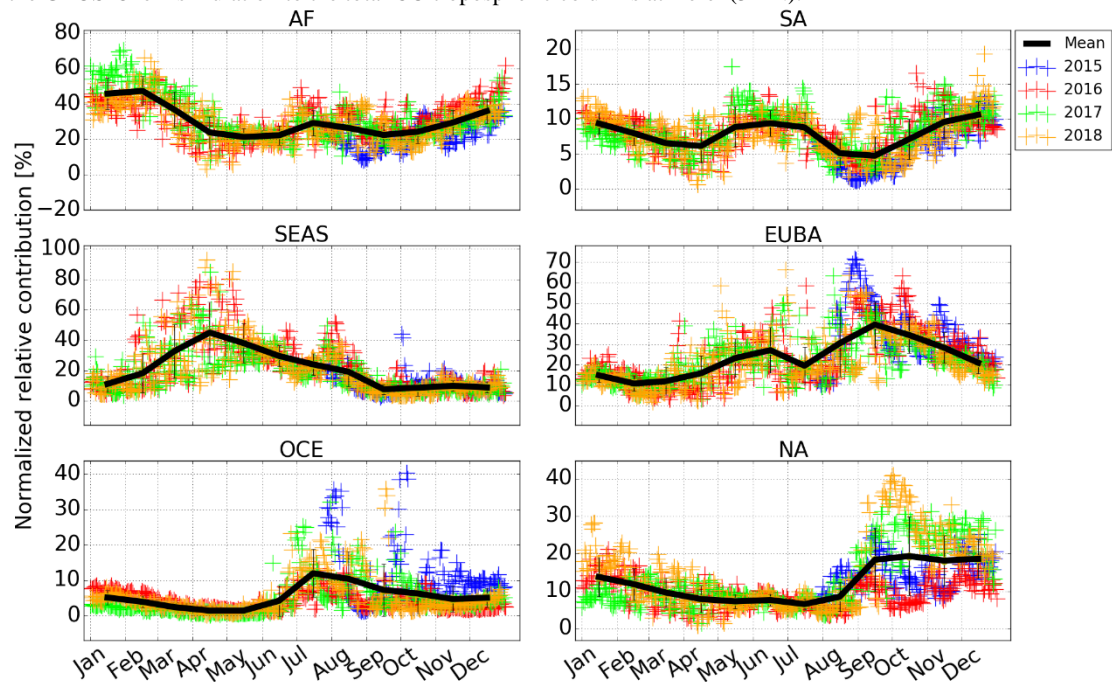


24 Fig. 8. Daily-mean CO tropospheric column time series of FTIR and GEOS-Chem (top panel) from 2015-2018 at  
 25 Hefei (32°N). The bottom panel shows the relative contribution (%) of the anthropogenic, biomass burning, and  
 26 oxidation tracers in the GEOS-Chem simulation to the total CO tropospheric columns at Hefei (32°N).  
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Fig. 9. Seasonality of the relative contribution (%) of the anthropogenic, biomass burning, and oxidation tracers in the GEOS-Chem simulation to the total CO tropospheric columns at Hefei (32°N).



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Fig. 10. Seasonality of the normalized relative contribution (%) of the AF, SA, SEAS, EUBA, OCE, and NA biomass burning tracers in the GEOS-Chem simulation to the total biomass burning associated CO tropospheric column at Hefei (32°N).

#### 4.2 Attribution for transport pathway

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For each seasonal enhancement of the tropospheric HCN, transport pathway is determined as follows. First, the GEOS-Chem tagged CO simulation is used to calculate the relative contribution of each biomass burning tracer (Fig. 10). For the tracer with a high contribution, the FIRMS global fire map is used to search for potential fire events occurring before the phase of tropospheric HCN enhancement within a one month period. Then, we generated an ensemble of HYSPLIT back trajectories with different travel times and arrival altitudes to judge whether these plumes are capable of travelling to the observation site. For example, for each intensive biomass burning event detected at a specific period, we generated ten back trajectories at different arrival altitudes ranging from 1.5 to 12 km, and modified the end time of these back-trajectories within one day of the

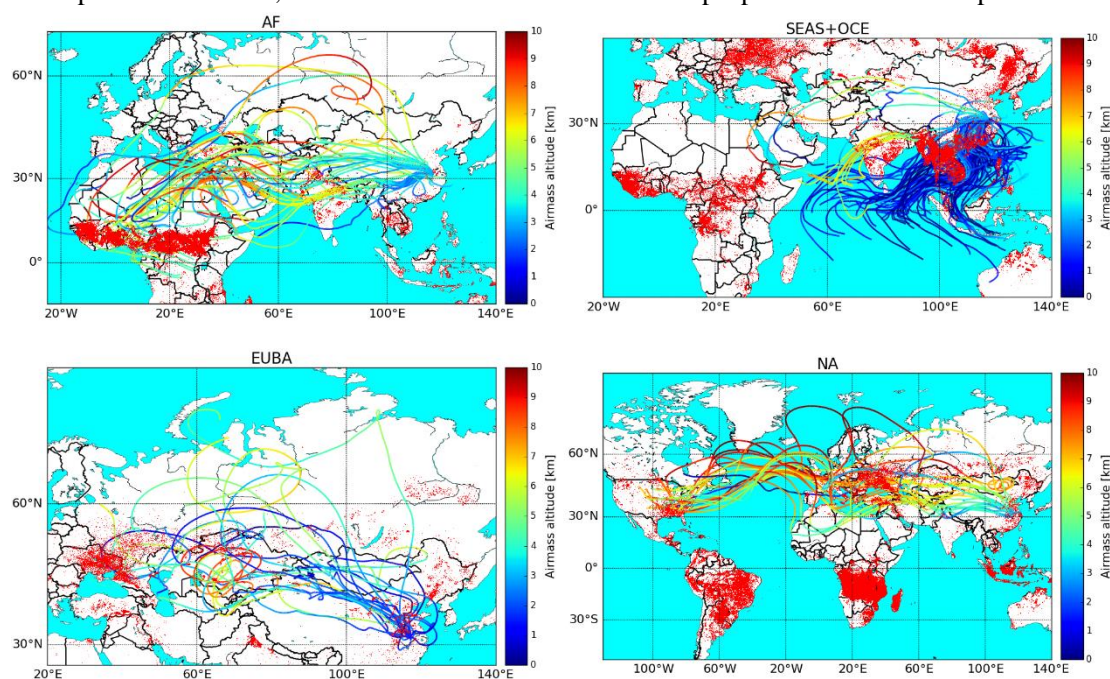


1 observed enhancement. If the back-trajectories intersect a region where the FIRMS fire data  
 2 indicates an intensive fire event and the travel duration is within a reasonable range, then this  
 3 specific fire event could contribute to the observed enhancements at Hefei (32°N) in eastern China.  
 4 The transport pathway for this enhancement is finally determined.

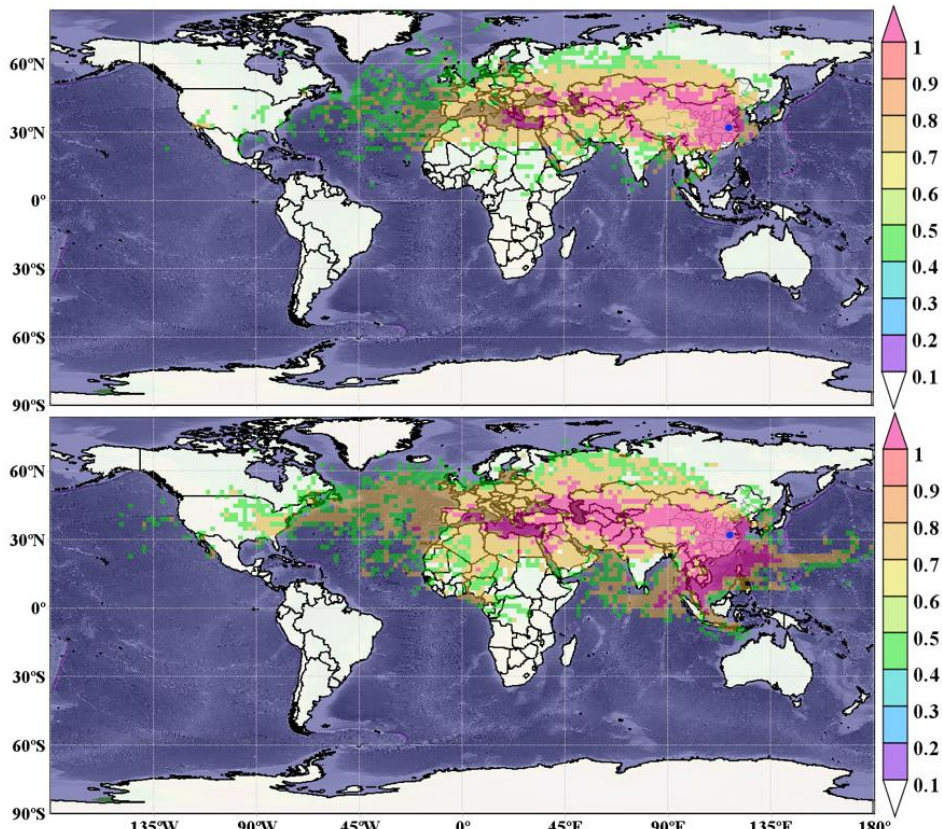
5 Fig. 11 demonstrates travel trajectories of the plumes occurred in AF, SEAS & OCE, EUBA,  
 6 and NA that reached Hefei (32°N) through long range transport. Fig. 12 shows the PSCFs calculated  
 7 using 13-day HYSPLIT back trajectories that are coincident with the FTIR measurement time. The  
 8 eastern China, South Asia, Central Asia, Eastern Europe, and Northern Africa had high PSCF  
 9 weight values in both the first and the second half of the year. The large areas of Southeastern Asian  
 10 countries including Philippines, Malaysia, and Indonesia, and the Eastern North America were the  
 11 additional regions with potentially high PSCF weight values in the second half of the year. Generally,  
 12 trajectories with the same travel time in the second half of the year are longer than those in the first  
 13 half year, resulting in broader areas with potentially high PSCF weight values.

14 As shown in Figs.13 and 14, the seasonal biomass burning typically occurs in July – September  
 15 in southern Africa and in November – February in central Africa. These AF emissions can be  
 16 transported to eastern China along with the southwestern wind which contributed 25 – 45% of the  
 17 tropospheric HCN in these periods. The seasonal biomass burning typically occurs in March – May  
 18 and July – November in central Europe, and in June – September in Siberia. These EUBA emissions  
 19 can be transported to eastern China along with the northwestern or northern wind which contributed  
 20 27 – 40% of the tropospheric HCN in these periods. The seasonal biomass burning typically occurs  
 21 in March – May in India and South Asia peninsula. Largely driven by deep convection followed by  
 22 northward transport into the mid-latitude westerlies (Liu et al., 2003), these emissions can be  
 23 transported to eastern China which contributed to the tropospheric HCN peak in May. The seasonal  
 24 biomass burning typically occurs in March – May, July – September, and November – December in  
 25 the eastern part of China. All these emissions can be transported to the observation site at Hefei  
 26 (32°N) under favorable meteorological conditions and thus contribute to all the seasonal  
 27 tropospheric HCN peaks. The SEAS contribution (mainly China, India and South Asia peninsula)  
 28 varies over 25 to 80% in March to August.

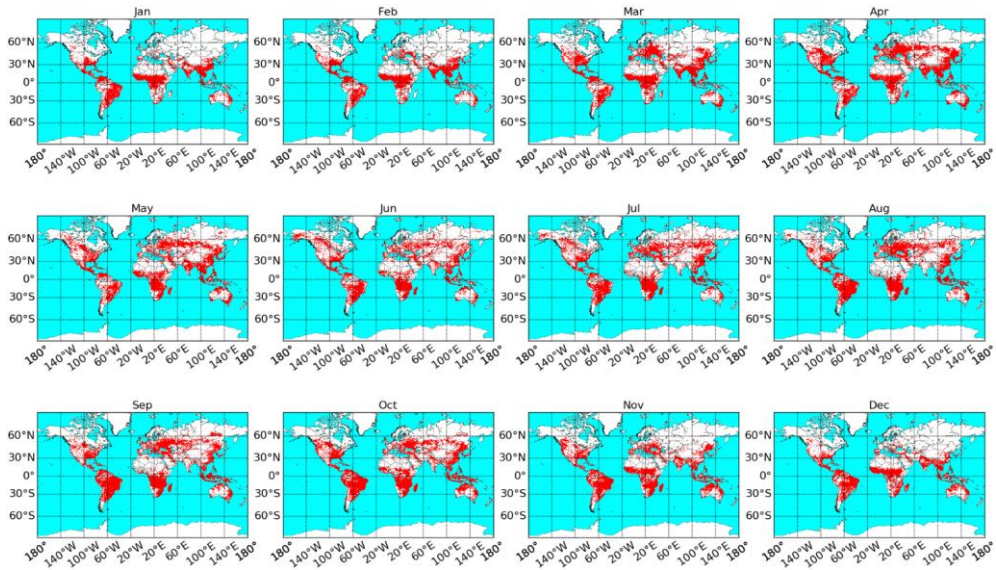
29 Additionally, a small to moderate portion of wildfire events in central SA, eastern NA, and  
 30 Northern OCE in autumn or winter could transport to the observation site through large-scale  
 31 atmospheric circulation, which contributed 5 – 20% of the tropospheric HCN in these periods.



32 Fig.11. Travel trajectories of the plumes occurred in AF, SEAS & OCE, EUBA, and NA that reached Hefei (32°N)  
 33 through long range transport. Travel times are 13, 7, 10, and 14 days, respectively. For clarity, only few trajectories  
 34 are selected for demonstration. FIRMS fire numbers are shown in red dots for 13, 7, 10, and 14 days prior to the  
 35 arrival time, respectively.  
 36

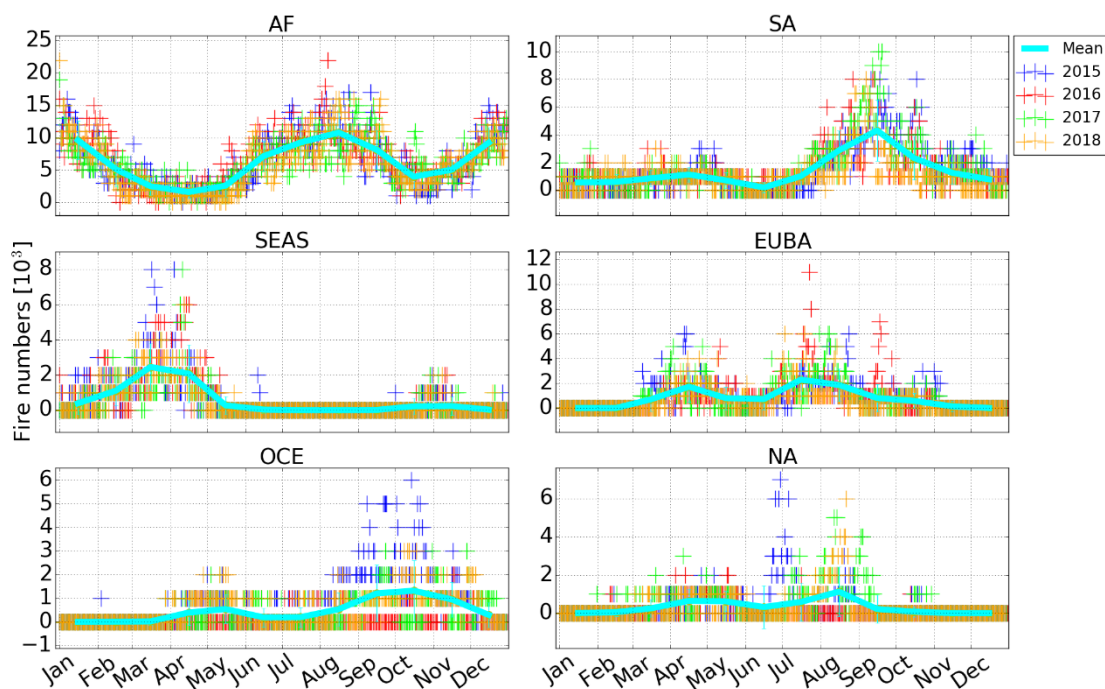


1  
2 Fig. 12. Likely source areas of air mass associated with higher HCN concentrations at Hefei (32°N) in the first half  
3 year (top panel) and the second half year (bottom panel) identified using PSCF.



4  
5 Fig. 13. Global fire map in January to December 2015 accumulated from the FIRMS fire atlas.





1  
2 Fig.14. Seasonality of total fire numbers within the AF, SA, SEAS, EUBA, OCE, and NA tracers. All data are  
3 accumulated from the FIRMS fire atlas.

#### 4 **4.3 Attribution for interannual variability**

5 In Fig. 9, the biomass burning contribution was elevated by 5 – 15% between September 2015  
6 and July 2016, while no elevations were observed for anthropogenic and oxidation influence. As a  
7 result, enhancements of both tropospheric HCN and CO columns between September 2015 and July  
8 2016 at Hefei (32°N) were attributed to an elevated influence of biomass burning. In Fig.10, the  
9 relative contribution (%) of the SEAS, EUBA, and OCE biomass burning tracers to the total biomass  
10 burning associated CO tropospheric column were elevated by 5 – 20%, 8 – 27%, 8 – 31%,  
11 respectively, in the second half of 2015 compared to the same period in other years. The relative  
12 contribution (%) of the SEAS and OCE biomass burning tracers to the total biomass burning  
13 associated CO tropospheric column were elevated by 8 – 39% and 2 – 7%, respectively, in the first  
14 half of 2016 compared to the same period in other years.

15 The statistical results of the FIRMS fire atlas data in Fig.14 show that, the fire numbers in the  
16 SEAS, EUBA, and OCE regions elevated by 21.89%, 15.72%, and 32.68% between September  
17 2015 and July 2016 compared to the same period in other years. These elevated fire numbers in  
18 EUBA, SEAS and OCE drove the enhancements of tropospheric HCN and CO columns between  
19 September 2015 and July 2016 at Hefei (32°N). In particular, the number of fires in OCE in the  
20 second half of 2015 was greatly elevated in comparison with the other years, acting as a dominant  
21 source of tropospheric HCN enhancement in September – December 2015. The fire numbers  
22 elevated significantly in the SEAS region in the first half of 2016, which dominated the tropospheric  
23 HCN enhancement in January – July 2016.

24 Many studies have revealed that the El Niño Southern Oscillation (ENSO) can cause large  
25 scale variations in the convection, circulation, and air temperature of the global atmosphere-ocean  
26 system (Liu et al., 2017; Zhao et al., 2002), which could affect the distribution, frequency, and  
27 intensity of biomass burning emissions (Schaefer et al., 2018). Furthermore, ENSO could also alter  
28 the destruction processes of tropospheric species through their photochemical reactions with  
29 tropospheric OH (Zhao et al., 2002). Zhao et al. (2002) found that the abnormal enhancement of  
30 tropospheric CO and HCN observed in northern Japan in 1998 were associated with the 1997–1998  
31 ENSO events (Zhao et al., 2002). There is a close correlation between ENSO and HCN columns at  
32 Lauder (45°S) (Zeng et al., 2012; Schaefer et al., 2018), and Schaefer et al. (2018) quantified a  
33 detectable ENSO influence on biomass burning of up to 51-55% (Schaefer et al., 2018; Zeng et al.,  
34 2012). Very likely, the elevated fire number which caused significant enhancements between  
35 September 2015 and July 2016 for tropospheric CO and HCN columns at Hefei and most selected  
36 NDACC stations were related to the 2015 – 2016 ENSO events.

1 Compared to the northwestern part of China such as the Xinjiang province and the Tibet plateau,  
2 the densely populated eastern parts of China are more suitable for crop planting because of fertile  
3 soil and adequate water resources. Historically, Chinese farmers burned their crop residue (such as  
4 rice, corn, and wheat straws) after harvest to fertilize the soil for the coming farming season. Post-  
5 harvest crop residue is a fine fuel that burns directly in the field and mostly by flaming in many  
6 mechanized agricultural systems. In contrast, when crops are harvested by hand the residue is often  
7 burned in large piles that may smolder for weeks.

8 This seasonal crop residue burning season typically occurs in the spring and summer seasons  
9 and also occasionally occurs in the autumn and winter. Pollution gases, dust, and suspended particle  
10 matters resulting from crop residue burning emissions result in poor air quality that threaten human  
11 health and terrestrial ecosystems. The Chinese presidential decree included the prohibition of crop  
12 residue burning into the Law of the People's Republic of China on the Prevention and Control of  
13 Atmospheric Pollution in August 2015 (<http://www.chinalaw.gov.cn>, last access on 17 July 2019),  
14 and since then the crop residue burning events were banned throughout China. Therefore, we obtain  
15 a decrease in fire numbers in China since 2015.

## 16 **6 Conclusion**

17 The first multiyear measurements of HCN in the polluted troposphere in densely populated  
18 eastern China have been presented. Tropospheric HCN columns were derived from solar spectra  
19 recorded with ground-based high spectral resolution Fourier transform infrared (FTIR) spectrometer  
20 at Hefei (117°10'E, 31°54'N) between 2015 and 2018. The seasonality and interannual variability  
21 of tropospheric HCN columns in eastern China have been investigated. The potential sources that  
22 drive the observed HCN seasonality and interannual variability were determined by using the  
23 GEOS-Chem tagged CO simulation, the global fire maps and the PSCFs (Potential Source  
24 Contribution Function) calculated using HYSPLIT back trajectories.

25 The tropospheric HCN columns over eastern China showed significant seasonal variations with  
26 three monthly mean peaks throughout the year. The magnitude of the tropospheric HCN peak in  
27 May > September > December. The tropospheric HCN column reached a maximum monthly mean  
28 of  $(9.8 \pm 0.78) \times 10^{15}$  molecules/cm<sup>2</sup> in May and a minimum monthly mean of  $(7.16 \pm 0.75) \times 10^{15}$   
29 molecules/cm<sup>2</sup> in November. In most cases, the tropospheric HCN columns at Hefei (32°N) are  
30 higher than the NDACC FTIR observations. Enhancements of the tropospheric HCN columns were  
31 observed between September 2015 and July 2016 compared to the same period of measurements in  
32 other years. The magnitude of the enhancement ranges from 5 to 46% with an average of 22%.  
33 Enhancement of tropospheric HCN ( $\Delta$ HCN) is correlated with the coincident enhancement of  
34 tropospheric CO ( $\Delta$ CO), indicating that enhancements of tropospheric CO and HCN were due to  
35 the same sources.

36 The GEOS-Chem tagged CO simulation, the global fire maps and the PSCFs analysis revealed  
37 that the seasonal maxima in May is largely due to the influence of biomass burning in South Eastern  
38 Asia (SEAS) ( $41 \pm 13.1\%$ ), Europe and Boreal Asia (EUBA) ( $21 \pm 9.3\%$ ) and Africa (AF) ( $22 \pm$   
39  $4.7\%$ ). The seasonal maxima in September is largely due to the influence of biomass burnings in  
40 EUBA ( $38 \pm 11.3\%$ ), AF ( $26 \pm 6.7\%$ ), SEAS ( $14 \pm 3.3\%$ ) and NA ( $13.8 \pm 8.4\%$ ). For the seasonal  
41 maxima in December, dominant contributions are from AF ( $36 \pm 7.1\%$ ), EUBA ( $21 \pm 5.2\%$ ), and  
42 NA ( $18.7 \pm 5.2\%$ ).

43 The enhancements of both tropospheric HCN and CO columns between September 2015 and  
44 July 2016 at Hefei (32°N) were attributed to an elevated influence of biomass burnings in SEAS,  
45 EUBA, and Oceania (OCE) in this period. In particular, an elevated fire numbers in OCE in the  
46 second half of 2015 dominated the tropospheric HCN enhancement in September – December 2015.  
47 An elevated fire numbers in SEAS in the first half of 2016 dominated the tropospheric HCN  
48 enhancement in January – July 2016.

49 Most high resolution FTIR instruments are located in Europe and Northern America, whereas  
50 the number of sites in Asia, Africa, and South America is very sparse. As one of few FTIR stations  
51 on Asian continent, the long-term observations of trace gases at Hefei are crucial to understand  
52 global warming, regional pollution, long term transport, and contribute to the evaluation of satellite  
53 data and model simulations.

54  
55 **Data availability.** The CO and HCN measurements at the selected NDACC sites can be found by  
56 the link <http://www.ndaccdemo.org>, and the CO and HCN measurements at Hefei are available on  
57 request.

1 **Author contributions.** YS conceived the concept and prepared the paper with inputs from all  
2 coauthors. CL, WW, CS, HY, XX, MZ, and JL carried out the experiments. The rest authors  
3 contributed to this work via provide refined data or constructive comments.

4 **Competing interests.** The authors declare that they have no conflict of interest.

5 **Acknowledgements.** This work is jointly supported by the National High Technology Research and  
6 Development Program of China (No.2019YFC0214802, No.2017YFC0210002, No.  
7 2016YFC0203302, 2018YFC0213201, 2019YFC0214702, 2016YFC0200404), the National  
8 Science Foundation of China (No. 41877309, No.41775025, No. 41575021, No. 51778596, No.  
9 91544212, No. 41722501, No. 51778596), the Sino-German Mobility programme (M-0036), and  
10 Outstanding Youth Science Foundation (No. 41722501). The processing and post processing  
11 environment for SFIT4 are provided by National Center for Atmospheric Research (NCAR),  
12 Boulder, Colorado, USA. The NDACC network is acknowledged for supplying the SFIT software  
13 and the HCN and CO data. The LINEFIT code is provided by Frank Hase, Karlsruhe Institute of  
14 Technology (KIT), Institute for Meteorology and Climate Research (IMK-ASF), Germany. The  
15 MeteoInfo software is provided by Prof. Yaqiang Wang, Chinese Academy of Meteorological  
16 Sciences. The authors acknowledge the NOAA Air Resources Laboratory (ARL) for making the  
17 HYSPLIT transport and dispersion model available on the Internet. The Mauna Loa (20°N) FTIR  
18 site is operated by National Center for Atmospheric Research (NCAR), U.S.A., and the Lauder  
19 (45°S) and Arrival Heights (78°S) sites are operated by National Institute of Water & Atmospheric  
20 Research, New Zealand. The multi-decadal monitoring program of ULiege at the Jungfrauoch  
21 station has been primarily supported by the F.R.S.-FNRS and BELSPO (both in Brussels, Belgium)  
22 and by the GAW-CH programme of MeteoSwiss. The International Foundation High Altitude  
23 Research Stations Jungfrauoch and Gornergrat (HFSJG, Bern) supported the facilities needed to  
24 perform the FTIR observations.

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