

Reduction in C₂H₆ from 2015 to 2020 over Hefei, eastern China points to air quality improvement in China

Youwen Sun^{1,2}, Hao Yin^{1,2,*}, Cheng Liu^{1,2,3,*}, Emmanuel Mahieu⁴, Justus Notholt⁵, Yao Té⁶, Xiao Lu⁷, Mathias Palm⁵, Wei Wang¹, Changgong Shan¹, Qihou Hu¹, Min Qin¹, Yuan Tian⁸, and Bo Zheng⁹

¹Key Laboratory of Environmental Optics and Technology, Anhui Institute of Optics and Fine Mechanics, HFIPS, Chinese Academy of Sciences, Hefei 230031, China

²Key Laboratory of Precision Scientific Instrumentation of Anhui Higher Education Institutes, University of Science and Technology of China, Hefei, 230026, China

³Anhui Province Key Laboratory of Polar Environment and Global Change, USTC, Hefei, 230026, China

⁴Institute of Astrophysics and Geophysics, University of Liège, Belgium

⁵University of Bremen, Institute of Environmental Physics, P. O. Box 330440, 28334 Bremen, Germany

⁶Laboratoire d'Etudes du Rayonnement et de la Matière en Astrophysique et Atmosphères (LERMA-IPSL), Sorbonne Université, CNRS, Observatoire de Paris, PSL Université, 75005 Paris, France

⁷School of Atmospheric Sciences, Sun Yat-sen University, Zhuhai, 519082, China

⁸Institutes of Physical Science and Information Technology, Anhui University, Hefei 230601, China

⁹Institute of Environment and Ecology, Tsinghua Shenzhen International Graduate School, Tsinghua University, Shenzhen 518055, China

*Corresponding authors.

E-mail addresses: chliu81@ustc.edu.cn (C. Liu) and yhyh95@mail.ustc.edu.cn (H. Yin)

Abstract

Ethane (C₂H₆) is an important greenhouse (GHG) gas and plays a significant role in tropospheric chemistry and climate change. This study first presents and quantifies the variability, source, and transport of C₂H₆ over densely populated and highly industrialized eastern China by using ground-based high-resolution Fourier transform infrared (FTIR) remote sensing together with atmospheric modelling techniques. We obtained a retrieval error of 6.21 ± 1.2 (1 σ)% and degrees of freedom (DOFS) of 1.47 ± 0.2 (1 σ) in retrieval of C₂H₆ tropospheric column-averaged dry-air mole fraction (troDMF) over Hefei, eastern China (117°E, 32°N, 30 m a.s.l.). The observed C₂H₆ troDMF reached a minimum monthly mean value of 0.36 ± 0.26 ppbv in July and a maximum monthly mean value of 1.76 ± 0.35 ppbv in December, and showed a negative change rate of -2.60 ± 1.34 %/yr from 2015 to 2020. The dependencies of C₂H₆ troDMF on meteorological and emission factors were analyzed by using generalized additive models (GAMs). Generally, both meteorological and emission factors have positive influences on C₂H₆ troDMF in cold season (DJF/MAM) and negative influences on C₂H₆ troDMF in warm season (JJA/SON). GEOS-Chem chemical model simulation captured the observed C₂H₆ troDMF variability and was thus used for source attribution. GEOS-Chem model sensitivity simulations concluded that the anthropogenic emissions (fossil fuel plus biofuel emissions) and the natural emissions (biomass burning plus biogenic emissions) accounted

1 for 48.1% and 39.7% of C₂H₆ troDMF variability over Hefei, respectively. The observed C₂H₆
2 troDMF variability mainly results from the emissions within China (74.1%), where central, eastern,
3 and northern China dominated the contribution (57.6%). Seasonal variability in C₂H₆ transport
4 inflow and outflow over the observation site is largely related to the mid-latitude westerlies and
5 Asian monsoon system. Reduction in C₂H₆ abundance from 2015 to 2020 mainly results from the
6 decrease in local and transported C₂H₆ emissions, which points to air quality improvement in China
7 in recent years.

8 Keywords: Remote sensing; FTIR; Ethane; Climate change; GAMs

9 1. Introduction

10 Ethane (C₂H₆) is an important greenhouse (GHG) gas and one of the most abundant Volatile
11 Organic Compounds (VOCs) in the atmosphere (Abad et al. 2011; Singh et al. 2001; Steinfeld 1998).
12 Although C₂H₆ is much less abundant than methane (CH₄) and also less efficient relative to mass,
13 it plays a significant role in tropospheric chemistry and climate change (Tzompa-Sosa et al. 2017).
14 In the presence of nitrogen oxides (NO_x = NO + NO₂), the C₂H₆ oxidation can enhance tropospheric
15 ozone (O₃) generation, which shows a positive radiative influence on climate (Sun et al. 2018a) and
16 threatens crop yields (Sun et al. 2018a; Van Dingenen et al. 2009) and human health (Sun et al.
17 2018a; Tzompa-Sosa et al. 2017). In addition, as a major source of acetaldehyde (CH₃CHO), C₂H₆
18 has a great impact on the production of peroxyacetyl nitrate (PAN) which is a key reservoir species
19 of NO_x (Fischer et al. 2014). The main sink of tropospheric C₂H₆ is predominantly destruction via
20 reaction with the hydroxyl radical (OH)(Xiao et al. 2008), which determines the residence time of
21 most tropospheric species (Steinfeld 1998). As a result, tropospheric C₂H₆ can decrease the
22 atmospheric oxidative capacity and indirectly impact the climate by extending the CH₄ lifetime
23 (Monks et al. 2018; Taylor et al. 2020). Atmospheric C₂H₆ has a relatively long residence time of a
24 few months (Franco et al. 2016), allowing it to undergo intercontinental transport. As a result,
25 observations of C₂H₆ can be assimilated into a chemical transport model to estimate nonlocal
26 emissions and air quality, and provide valuable insights into model biases of C₂H₆ simulations
27 (Tzompa-Sosa et al. 2017).

28 On a global scale, the main sources of C₂H₆ are leakage from production, processing, and
29 transport of natural gas (62%), biofuel combustion (20%) and biomass burning emission (18%),
30 largely occurred in the Northern Hemisphere (NH) (Franco et al. 2016; Xiao et al. 2008). Additional
31 minor sources of C₂H₆ are from biogenic and oceanic sources. However, on a regional scale, the
32 proportion of each C₂H₆ source may show large difference. Natural gas leakage contribution can
33 reach 80% of C₂H₆ emissions in regions with active oil and natural gas production (Gilman et al.
34 2013), where C₂H₆ emissions are highly correlated with CH₄ emissions. In such regions, C₂H₆ can
35 be applied as a tracer for separation of fossil fuel CH₄ emissions from multiple methane (CH₄)
36 sources (e.g., oil and gas, cows, wetlands, and rice yield) (McKain et al. 2015; Roscioli et al. 2015).
37 The C₂H₆ abundance in the Southern Hemisphere (SH) is much lower than those in the NH since
38 the anthropogenic C₂H₆ sources are low in the SH, and the residence time of C₂H₆ is shorter than
39 the interhemispheric exchange rate. Many studies concluded that C₂H₆ in the SH is primarily emitted
40 from biomass burning, and is closely correlated with CO and HCN emissions (Notholt et al. 2000;
41 Rinsland et al. 2002; Vigouroux et al. 2012; Zeng et al. 2012).

42 C₂H₆ is one of the target gases of a global ground-based Fourier transform infrared

1 spectroscopy (FTIR) network, namely the infrared working group (IRWG) of the Network for
2 Detection of Atmospheric Composition Change (NDACC) (De Mazière et al. 2018). FTIR time
3 series of C₂H₆ with different time periods have been reported in many stations for validation of
4 satellite data or chemical model simulation (Abad et al. 2011; Franco et al. 2015; Franco et al. 2016;
5 Glatthor et al. 2009), or evaluation of local air quality and air pollutants transport caused by
6 anthropogenic emission and biomass burning (Angelbratt et al. 2011; Lutsch et al. 2016; Lutsch et
7 al. 2019; Nagahama and Suzuki 2007; Rinsland et al. 2002; Simpson et al. 2012; Viatte et al. 2015;
8 Viatte et al. 2014; Vigouroux et al. 2012; Zeng et al. 2012; Zhao et al. 2002). Several FTIR sites
9 have observed the decrease in C₂H₆ over 1990 – 2010, and characterized consistent interannual
10 trends in the –1 to –2.7% yr⁻¹ range (Franco et al. 2015; Franco et al. 2016; Simpson et al. 2012;
11 Zeng et al. 2012). This declining trend has been largely attributed to the reduction of global fugitive
12 emissions (Franco et al. 2015; Simpson et al. 2012). Recently, several studies concluded that the
13 long-term decline in C₂H₆ in the NH reversed from 2009 onwards (Franco et al. 2015; Franco et al.
14 2016). Using ground-based FTIR C₂H₆ total columns derived at five selected NDACC sites, Franco
15 et al. (2016) characterized the C₂H₆ evolution from 2009 – 2015 and determined growth rates of
16 ~3% yr⁻¹ at remote sites and of ~5% yr⁻¹ at mid-latitudes. This change is mainly attributed to the
17 exploitation of shale gas and tight oil reservoirs in North America (Franco et al. 2016; Helmig et al.
18 2016).

19 The NDACC network has been operating for almost three decades around the globe (De
20 Mazière et al. 2018; Sun et al. 2018a). However, most instruments are located in Europe and
21 Northern America, but the number of observation sites in the rest parts of world remains sparse, and
22 there is only one qualified observations site in China, i.e., the Hefei site (117°E, 32°N, 30 m a.s.l.)
23 located in a densely populated and highly industrialized area in eastern China (Sun et al. 2018a).
24 The Hefei site is not yet affiliated to the NDACC network but its observation routine is following
25 the NDACC standard convention since 2015 (Sun et al. 2018a). As the consequence of a series of
26 actions for emission control, air pollution over China in recent years has been significantly
27 decreased (Zhang et al. 2019; Zheng et al. 2018). However, the atmospheric pollution over densely
28 populated and highly industrialized eastern China is still in severe situation (Zhang et al. 2019;
29 Zheng et al. 2018). The complexity, extension, and severity of the atmospheric pollution in eastern
30 China are still unrivaled compared to the rest of world (Lu et al. 2018; Zheng et al. 2018). FTIR
31 observations at Hefei have been used extensively for evaluation of satellite data (Tian et al. 2018;
32 Wang et al. 2017), chemical model simulation (Tian et al. 2018; Yin et al. 2020; Yin et al. 2019),
33 local air quality (Shan et al. 2019; Sun et al. 2018a) and the transport of air pollutants caused by
34 anthropogenic and biomass burning emissions (Sun et al. 2018a; Sun et al. 2021; Sun et al. 2020).

35 In this study, we first present and quantify the variability, source, and transport of C₂H₆ over
36 densely populated and highly industrialized eastern China by using FTIR observation, GEOS-Chem
37 model simulation, and the analysis of the meteorological fields. The seasonality and interannual
38 variability of C₂H₆ over Hefei, eastern China from 2015 – 2020 are investigated. The dependencies
39 of C₂H₆ on meteorological and co-emitted gases (hereafter emission factors) are analyzed by using
40 generalized additive models (GAMs)(Wood and Simon 2004). The ground-based FTIR C₂H₆ time
41 series are for the first time applied to evaluate the GEOS-Chem model for the specifics of C₂H₆
42 simulation over eastern China. We further run a series of GEOS-Chem sensitivity simulations to
43 quantify relative contributions of various source categories and regions to the observed C₂H₆
44 variability. The three-dimensional (3D) transport inflow and outflow pathways of C₂H₆ over the

1 observation site are finally determined by the GEOS-Chem sensitivity simulations and the analysis
2 of the meteorological fields. This study can not only enhance the understanding of regional emission,
3 transport, and air clean actions over eastern China, but also contribute to form new reliable remote
4 sensing data in this sparsely-monitored regions for climate change research.

5 The next section describes the retrieval for FTIR tropospheric column-averaged dry-air mole
6 fraction (troDMF) of C_2H_6 , the configuration of GEOS-Chem model simulation, and the GAMs
7 regression approach. Section 3 reports the variability of C_2H_6 troDMF and comparison with the
8 GEOS-Chem simulation. Section 4 reports the GAMs regression results and the interpretation.
9 Section 5 reports the results for source attribution using GEOS-Chem sensitivity simulation and the
10 analysis of the meteorological fields. We conclude the study in Section 6.

11 2. Methods and data

12 2.1 C_2H_6 troDMF retrieval

13 The C_2H_6 troDMF time series were calculated by using direct solar absorption spectra saved
14 with a FTIR spectrometer in operation at Hefei, eastern China (Sun et al. 2018a; Tian et al. 2018).
15 Site description and instrumentation can be found in (Sun et al. 2018a). Briefly, the FTIR
16 observatory includes a high resolution FTIR spectrometer (IFS125HR, Bruker) and a solar tracker
17 (Solar Tracker-A 547, Bruker). The instrument has been operating continuously since its installation;
18 however, short data gaps of up to eight months have occurred due to a scanner problem between
19 November 2016 and July 2017. This FTIR observatory alternately saved near infrared (NIR) and
20 middle infrared (MIR) solar spectra in routine observations, with spectral ranges of 4,000 to 11,000
21 cm^{-1} and 500 to 8,500 cm^{-1} , respectively (Tian et al. 2018). The NIR and MIR spectra are saved
22 with different spectral resolutions but both of them can be used to retrieve total columns and volume
23 mixing ratio (VMR) profiles of a variety of trace gases in the atmosphere. The MIR spectra used in
24 present work are saved with a spectral resolution of 0.005 cm^{-1} following the requirements of
25 NDACC standard convention (<http://www.ndacc.org/>, last accessed on 27 December 2020). The
26 FTIR instrument is equipped with a KBr beam splitter, a filter centered at 2800 cm^{-1} , and an InSb
27 detector for C_2H_6 measurements. The number of C_2H_6 measurements on each measurement day
28 varied from 1 to 17 with an average of 6. In total, there were 743 days of qualified measurements
29 between 2015 and 2020.

30 In this study, the VMR profile of C_2H_6 was first retrieved by using the SFIT4 algorithm updated
31 from SFIT2 (Pougetchev et al. 1995) and implementing the optimal estimation method (Rodgers
32 2000). The C_2H_6 troDMF was then calculated by taking a weighting average of the C_2H_6 VMR
33 profile and the air mass using a fixed tropospheric altitude. The C_2H_6 VMR profile was retrieved in
34 a broad window of 2976 – 2978 cm^{-1} . The VMR profiles of CH_4 and H_2O and column of O_3 were
35 also retrieved together with the C_2H_6 VMR profile for minimizing the atmospheric absorption
36 interference. Spectroscopic absorption parameters of all gases are based on the atm16 line list from
37 the compilation of Geoffrey Toon (Sun et al. 2021). The *a priori* vertical profiles of temperature
38 H_2O , and pressure were interpolated from the National Centers for Environmental Protection (NCEP)
39 reanalysis data and the *a priori* vertical profiles of other gases were from the statistical averages of
40 the Whole-Atmosphere Community Climate Model version 6 (WACCM) simulations from 1980 to
41 2020. The diagonal elements of the *a priori* covariance matrices S_a and the measurement noise
42 covariance matrices S_e were set to standard deviation (SD) of the WACCM simulations and the

1 inverse square of the signal-to-noise ratio (SNR) of each spectrum, respectively (Franco et al. 2015).
2 The non-diagonal elements of both \mathbf{S}_a and \mathbf{S}_e were set to zero. The instrument line shape (ILS) of
3 the FTIR instrument deduced from optical path alignment diagnosis with a low-pressure HBr cell
4 was adopted in the retrieval (Hase 2012; Sun et al. 2018b).

5 For each retrieval, the averaging kernels reflect the sensitivity of the retrieved profile to the
6 real profile. The area of the averaging kernels at a specific height is calculated as the sum of the
7 elements of the corresponding averaging kernels (Pougetchev et al. 1995). It represents the fraction
8 of the retrieval at that height which comes from the measurement rather than from the *a priori*
9 information (Rodgers, 2000). A value close to unity at a specific height indicates that the retrieved
10 profile at that height is nearly independent of the *a priori* profile and is thus from the measurement.
11 The trace of the averaging kernel matrix is defined as degrees of freedom for signal (DOFS) and it
12 quantifies the number of independent information in the retrieved profile. Fig. 1 shows the averaging
13 kernels as well as their area, cumulative sum of DOFS, and VMR profile of randomly selected C_2H_6
14 retrieval at Hefei. Ground-based FTIR C_2H_6 observations at Hefei have a sensitivity of larger than
15 0.7 from ground to about 10 km altitude, indicating that the retrievals are mainly sensitive to the
16 troposphere. This also means that the retrieved profile information below 10 km comes for more
17 than 70% from the measurement, or in other words, that the *a priori* signal impacts the retrieval by
18 less than 30% [Fig 1(a)]. The typical DOFS obtained at Hefei over the total atmosphere for C_2H_6 is
19 1.69 ± 0.29 (1σ), meaning that we can roughly provide two pieces of information on the vertical
20 profile [Fig 1(b)]. The shape of the retrieved profile is heavily weighted toward the lower
21 troposphere. As shown in Fig.1(c), the C_2H_6 concentration decreased by 72.7% with an increase in
22 the height from surface to 2 km and kept decreasing slowly in the rest part of the atmosphere till
23 approaching around zero in the stratosphere and above. The C_2H_6 partial column below 10 km
24 accounted for 88.6% of C_2H_6 total column. This percentage is expected to show less seasonal
25 variation since the shape of the retrieved profile is similar to the shape of the *a priori* profile due to
26 the low DOFS [Fig. 1 (c)]. As a result, in subsequent analysis, the C_2H_6 VMRs averaged between
27 the surface and 10 km are selected as representatives of C_2H_6 troDMF. The selected tropospheric
28 layer (from surface up to 10 km) corresponds to 1.47 ± 0.2 (1σ) of DOFS and can be used with
29 confidence. This selected layer is totally within the tropopause height (~ 16 km) at Hefei over four
30 seasons (Sun et al. 2020). The Hefei site is located in the northeastern margin of a GEOS-Chem grid
31 cell [Fig. 2]. This selected layer also ensures the line of sights of all observations are totally within
32 the same grid cell.

33 We calculated the error budget for C_2H_6 retrieval at Hefei following the formalism of Rodgers
34 (2000), and separated all error components into systematic or random errors according to whether
35 they vary steadily or randomly over consecutive measurements. The random, systematic, and the
36 combined error budgets for the selected tropospheric layer (from surface up to 10 km) are
37 summarized in Table 1. The input covariance matrix of temperature is based on the differences
38 between Sonde and an ensemble of NCEP temperature profiles near Hefei, leading to 2 to 5 K in
39 the troposphere and 3 to 7 K in the stratosphere. For each interfering gases, the corresponding
40 covariance matrix is obtained with the WACCM v6 climatology. The input covariance matrix of
41 measurement error is based on the inverse square of the SNR of each spectrum. We regularly use a
42 low-pressure HBr cell to diagnose the misalignment of the FTIR spectrometer and to realign the
43 instrument when indicated. The FTIR spectrometer at Hefei is assumed to be not far from the ideal
44 condition, and the input uncertainties for zero level, background curvature, field of view, optical

1 path difference, solar zenith angle, interferogram phase, and ILS are estimated to be 1%. For the
2 C₂H₆ spectroscopic absorption coefficients, the line list in atm16 follows HITRAN 2012 (Rothman
3 et al., 2013), and we use 5% for line intensity, pressure-, and temperature-broadening coefficients.
4 For the retrieval parameter and smoothing error, the input covariance matrix are prescribed from the
5 optimal estimation retrieval outputs. To estimate the retrieval error of C₂H₆ troDMF at Hefei, the
6 elements of all gain matrices were set to zero for the altitudes outside the selected layer. The
7 contributions of all error components to C₂H₆ troDMF retrieval at Hefei are summarized in Table 1.
8 The dominant random errors are from temperature uncertainty (1.69%) and the zero level
9 uncertainty (1.45%), and the dominant systematic error is from the line intensity uncertainty
10 (5.12%). Total random and systematic errors are estimated to be 2.32% and 5.48%, respectively.
11 Total retrieval error calculated as square root sum of the squares of total random and systematic
12 errors is estimated to be 6.21%.

13 In order to exclude the measurements that seriously affected by instable weather conditions or
14 by the *a priori* profile due to low measurement information content in less favourable observational
15 conditions, e.g., around noontime when the probed atmosphere is thinner, or in summer when C₂H₆
16 is less abundant. The FTIR measurements saved with a solar intensity variation (SIV) of larger than
17 10% or retrievals with total DOFS of less than 0.7 or the root-mean-square (RMS) of fitting residuals
18 of larger than 2%, which accounted for 11.2% of total measurements, were excluded in this study.

19 **2.2 GEOS-Chem sensitivity simulation**

20 Relative contribution of various source categories and regions to the observed C₂H₆ variability
21 were quantified by a series of sensitivity simulations using the GEOS-Chem chemical model version
22 12.2.1 (Bey et al. 2001) (<http://geos-chem.org>, last access on 24 August 2020). All simulations
23 implemented a universal tropospheric-stratospheric Chemistry (UCX) mechanism (Eastham et al.
24 2014; Fisher et al. 2017) and were driven by the Goddard Earth Observing System-Forward
25 Processing (GEOS-FP) meteorological fields at a degraded horizontal resolution of 2°×2.5°. The
26 temporal resolutions are 1 hour (hr) for surface meteorological variables and planetary boundary
27 layer height (PBLH), and 3 hr for other meteorological variables. The time step used in the model
28 are 10 minutes for transport and 20 minutes for chemistry and emissions, as recommend for the
29 GEOS-Chem full-chemistry simulation at 2°×2.5° (Philip et al., 2016). The non-local scheme for
30 the boundary layer mixing process are described in Lin and McElroy (2010). Dry deposition was
31 calculated by the resistance-in-series algorithm (Wesely 1989; Zhang et al. 2001) and wet deposition
32 followed that of Liu et al. (2001). The photolysis rates were available from the FAST-JX v7.0
33 photolysis scheme (Bian and Prather 2002). All simulations were spun up for one year (July 2014
34 to July 2015) and output hourly mean C₂H₆ VMR profiles globally ranging from the surface to 0.01
35 hPa at a horizontal resolution of 2°×2.5°. This study only considered the C₂H₆ simulations from
36 2015 to 2020 in the grid box containing Hefei (31.52°–32.11°N by 116.53°–118.02°E).

37 In the recent past, the inventories led to significant underestimation of the C₂H₆ simulation
38 (e.g., HTAP2 in Franco et al. 2016). Since then, some efforts have improved the situation (e.g.,
39 Tzompa-Sosa et al. 2017). In this study, we refer to Sun et al. (2020a) for more details on
40 implementation of emission inventories. Briefly, global anthropogenic emissions were from the
41 Community Emissions Data System (CEDS) inventory which overwrites Asia by the latest Multi-
42 resolution Emission Inventory for China (MEIC) (Hoesly et al. 2018; Li et al. 2017; Lu et al. 2019;

1 Zheng et al. 2018). Global biomass burning and biogenic emissions were from the Global Fire
2 Emissions Database (GFED v4) inventory (Giglio et al. 2013) and the Model of Emissions of Gases
3 and Aerosols from Nature (MEGAN version 2.1) inventory (Guenther et al. 2012), respectively.
4 The CH₄ emission fields are prescribed based on NOAA measurements for 1983–2016 and are
5 extended to 2020 using the linear extrapolation of local 2011–2016 change rates (Murray, 2016; Lu
6 et al., 2019).

7 Particular improvements have been done for the latest bottom-up MEIC emission inventory in
8 the accuracy of vehicle emission modelling (Zheng et al. 2014), power plant emission calculation
9 (Liu et al. 2015), and the non-methane VOCs (NMVOCs) speciation method (Li et al. 2014). Many
10 studies have verified that the MEIC emission inventory can reasonably represents the anthropogenic
11 emissions over Asia (Hoesly et al. 2018; Li et al. 2017; Lu et al. 2019; Sun et al. 2021; Tian et al.
12 2018; Yin et al. 2020; Yin et al. 2019; Zheng et al. 2018). Anthropogenic C₂H₆ emissions in China
13 by region and category for the 2015 and 2019 MEIC emission inventories are summarized in Table
14 2. All subdivided geographical regions are shown in Fig. 2 and the resulting delimitations are
15 summarised in Table 3. The delimitations of these geographical regions are based on the levels of
16 urbanization and industrialization in China. Region ① in Fig. 2 only covers a few sparsely city
17 clusters representing the region with least population and industrialization in China (Lu et al. 2019).
18 Regions ②, ④, and ⑤ cover the North China Plain (NCP), Yangtze River Delta (YRD), and
19 Pearl River Delta (PRD) city clusters, respectively, which are the three most developed city clusters
20 facing severe air pollution in China. Region ③ covers the Sichuan Basin (SCB) and central
21 Yangtze River (CYR) city clusters with newly emerging severe air pollution in China. Total annual
22 Chinese anthropogenic emissions of C₂H₆ in 2015 and 2019 are 0.883 Tg and 0.859 Tg, respectively.
23 In both years, anthropogenic C₂H₆ emissions in China are dominated by industry and residential
24 emissions. The highest anthropogenic C₂H₆ emission rates (calculated as the ratio of total C₂H₆
25 emission to the coverage) are in densely populated and industrialized region clusters in eastern part
26 of China (including northern China (NR), eastern China (ER), central China (CR), southern China
27 (SR), and adjacent regions) with large seasonal variation [Fig.S1]. The highest anthropogenic C₂H₆
28 emissions typically occur in winter [Fig.S1]. The anthropogenic emissions in WR region are
29 typically lower than those in other parts of China because of lower population and industries in the
30 region (Lu et al. 2019; Zheng et al. 2018).

31 In order to quantify the contributions of different source categories and regions to the observed
32 C₂H₆, we first conducted a reference full chemistry simulation (BASE) with implementation of all
33 emission inventories as described above. Then, we conducted a series of sensitivity simulations to
34 assess the change of each sensitivity simulation relative to the BASE simulation. Model
35 configurations in this study were similar to those in Sun et al. (2021) but with a different emission
36 perturbation method. When an emission inventory was shut off in Sun et al. (2021), the emissions
37 of all atmospheric compounds in that inventory were suppressed. In contrast, this work only
38 suppressed C₂H₆ in each case except for biogenic and biomass burning emission perturbations,
39 which suppressed all atmospheric compounds since we cannot separate C₂H₆ emission from current
40 biogenic and biomass burning emission inventories. Model configurations in this study are
41 summarised in Table 3 and were described as follows.

42 (i) To analyse the contributions of different emission categories, we shut off C₂H₆ in each
43 individual emission inventory to evaluate the change of the simulation in the presence of C₂H₆ in
44 other emission inventories. As a result, the relative contribution of each emission category was

1 estimated as the relative difference between the GEOS-Chem simulation in the presence and
2 absence of C_2H_6 in that emission inventory. We have conducted four such sensitivity simulations by
3 shutting off C_2H_6 emissions in (1) fossil fuel emission inventory (including emissions from
4 agriculture, industry, power plant, residential, and transport), (2) biogenic emission inventory, (3)
5 biomass burning emission inventory, and (4) biofuel emission inventory (Table 3). The sum of fossil
6 fuel and biofuel C_2H_6 emissions is defined as anthropogenic C_2H_6 source and the sum of biogenic
7 and biomass burning C_2H_6 emissions is referred to as natural C_2H_6 source.

8 (ii) To analyse the contributions of different geographical regions, we shut off all categories of
9 C_2H_6 emissions (i.e., the aforementioned anthropogenic plus natural C_2H_6 sources) within each
10 geographical region to assess the change of the simulation in the presence of C_2H_6 emissions outside
11 that geographical region. Thus, the relative contribution of each geographical region was estimated
12 as the relative difference between the GEOS-Chem simulation in the presence and absence of C_2H_6
13 emissions within that geographical region. We have conducted five such sensitivity simulations by
14 shutting off C_2H_6 emissions within five geographical regions shown in Fig. 2.

15 **2.3 Generalized additive models (GAMs) regression**

16 In this study, we investigate the dependencies of C_2H_6 on meteorological and emission factors
17 by using the GAMs regression (Wood and Simon 2004; Wood 2004). Regression analysis is
18 proceeded using the thin plate smoothing spline function (Pearce et al. 2011). Smoothing parameters
19 and confidence intervals are calculated according to the Restricted Maximum Likelihood standard
20 (REML) and the unconditional Bayesian method, respectively (Pearce et al. 2011). The GAMs
21 regression is better than the traditional statistical models in dealing with nonlinear fittings (Veaux
22 and Richard 2012). For climate change applications, where there are many non-linear relationships
23 between variables, the GAMs regression is particularly attractive (Zhang et al. 2019).

24 We introduced a variety of potential meteorological and emission factors into the GAMs
25 regression one at a time and performed significance tests based on the Akaike's Information Criteria
26 (AIC) values (Wood and Simon 2004). The explanatory variables which passed the significance
27 tests with the smallest AIC values were included into the final GAMs model. Furthermore,
28 explanatory variables in GAMs regression may interact with each other and result in unstable
29 fittings due to the internal multicollinearity. For the explanatory variables that show a strong
30 collinearity with each other, we only included one of them in the final GAMs model. The degree of
31 multicollinearity can be quantified by the variance inflation factor (VIF) (Ma et al. 2020). Generally,
32 a stronger collinearity between the explanatory variables results in a larger VIF, and the VIF of an
33 explanatory variable could be 1.0 if it is not correlated with other explanatory variables (Ma et al.
34 2020). In this study, we included all the meteorological and emission factors in the GAMs and
35 calculated the VIF for all the influencing factors. The multicollinearity diagnosis concluded that the
36 main causes of multicollinearity are between the HCN and CO, and between the tropopause height
37 and PBLH. Including either of these two data pairs in the GAMs regression showed significant
38 collinearities with the VIF values of greater than an empirical threshold of 4.0, indicating an unstable
39 regression (Lin et al. 2018). After omitting HCN and PBLH in the final GAMs model, the adjusted
40 VIF values of all the variables were less than 4.0 and the variables uniformly passed the significance
41 tests. As a result, the final GAMs model in the context of the C_2H_6 troDMF time series y can be
42 described as (Pearce et al. 2011):

$$\log(y) = \beta + S(ua) + S(va) + S(omega) + S(qv) + S(troph) + S(pres) + S(temp) + S(ch_4) + S(co) + \varepsilon \quad (1)$$

where β and ε are the mean response constant and the fitting residual, respectively. $S(ua)$, $S(va)$, $S(omega)$, $S(qv)$, $S(troph)$, $S(pres)$, $S(temp)$, $S(ch_4)$, and $S(co)$ are the smoothing functions of daily average zonal wind (with a unit of m s^{-1}), meridional wind (m s^{-1}), vertical wind (Pa s^{-1}), water vapor concentration (%), tropopause height (km), pressure (hPa), temperature ($^{\circ}\text{C}$), CH_4 troDMF (ppbv), and CO troDMF (ppbv). Positive values of ua , va , and $omega$ represent northward, eastward, and upward winds, respectively. The sum of $S(ch_4)$ and $S(co)$ is referred to as the emission influences, and the sum of remaining smoothing functions is referred to as the meteorological influences.

For driving the GAMs regression, we first derived CH_4 and CO VMR profiles from direct solar absorption spectra similar to that of C_2H_6 , see section 2.1. The spectra for CH_4 retrievals are exactly the same as those of C_2H_6 but the spectra for CO are saved at a different filter channel. The respective VMR profiles were then converted to troDMF values following the method of C_2H_6 . The retrieval configurations, waveband selections and the interfering gases considerations for CH_4 and CO can be found in Sun et al. (2018b). The DOFS of the retrievals between surface and 10 km altitude for both CH_4 and CO are larger than 1.5 and the corresponding retrieval errors are less than 8% (Sun et al., 2018b). All meteorological factors are from the GEOS-FP meteorological fields at their native resolution of $0.25^{\circ} \times 0.3125^{\circ}$ ranging from the surface to 0.01 hPa at a temporal resolution of 1 hr. Since the meteorological fields and C_2H_6 concentration are not uniformly distributed along the altitude, the summing mean values of the meteorological fields cannot properly characterize the meteorological influences. In this study, we use a method similar to that of Shaiganfar et al. (2017) to increase the influence weighting toward lower troposphere. As a result, all meteorological parameters except tropopause height ($troph$) are converted into the C_2H_6 profile weighting averaged value ω_{avg} through Eq. (2):

$$\omega_{avg} = \frac{\sum_i \omega(z_i) \cdot c(z_i) \cdot \mathbf{Airmass}(z_i)}{\sum_i c(z_i) \cdot \mathbf{Airmass}(z_i)} \quad (2)$$

where $\omega(z_i)$, $c(z_i)$, and $\mathbf{Airmass}(z_i)$ represent the value of the meteorological factor, C_2H_6 concentration, and the air mass at the altitude z_i .

3. Variability and comparison with GEOS-Chem model data

We have compared the observed daily mean time series, seasonal cycle and interannual variability of C_2H_6 troDMF to the GEOS-Chem BASE simulations for investigating the chemical model performance for the specifics of polluted regions over eastern China. As the vertical resolution of GEOS-Chem is different from the FTIR observation, smoothing correction has been done for the GEOS-Chem profiles (Rodgers and Connor 2003). First, the GEOS-Chem daily mean profiles of C_2H_6 have been interpolated to the FTIR altitude grid for ensuring a common altitude grid. In order to match the FTIR observations which only operates during daytime, the average for GEOS-Chem simulations was only performed during daytime from 9:00 to 17:00 local time (LT). The interpolated profiles were then smoothed by the concurrent seasonal mean values of the FTIR averaging kernels and *a priori* profiles (Rodgers 2000; Rodgers and Connor 2003). The smoothed GEOS-Chem profiles were subsequently converted to troDMF values by using the corresponding regridded air density profiles from the model with the method described in section 2.1. Finally, the GEOS-Chem C_2H_6 troDMF time series only for the days with available FTIR observations were

1 averaged by month and compared with the FTIR monthly mean data.

2 Fig. 3 (a) shows the comparison of daily mean time series of C₂H₆ troDMF between the FTIR
3 observation and the smoothed GEOS-Chem model simulation from 2015 to 2020. Fig. 3 (b)
4 compares the seasonal cycles derived from Fig. 3 (a) for the days with available FTIR observations
5 only. Generally, the measured features in terms of seasonality and interannual variability can be
6 reproduced by the GEOS-Chem simulations with a correlation coefficient (*r*) of 0.88 [Fig.S2]. Large
7 GEOS-Chem vs. FTIR differences tended to occur in the trough and peak of the observations. For
8 instance, the observed monthly mean value of C₂H₆ troDMF was overestimated by 35.6%
9 (calculated as (GEOS-Chem – FTIR)/FTIR) in July and underestimated by 14.6 % in December by
10 the GEOS-Chem. These discrepancies may be mainly attributed to uncertainties in the horizontal
11 transport and vertical mixing schemes simulated by the GEOS-Chem model at a relatively coarse
12 spatial resolution, which are difficult to match column observation over a single point. In addition,
13 the number of C₂H₆ measurements at Hefei on each measurement day varied a wide range from 1
14 to 17 depending on the weather condition, but GEOS-Chem simulations are available consecutively
15 by hour for the same day. This difference in the temporal resolution of GEOS-Chem and FTIR could
16 also cause inconsistencies due to the high variability of atmospheric C₂H₆. However, considering
17 the concurrent data pairs only (± 30 min), the averaged difference between GEOS-Chem and FTIR
18 data (GEOS-Chem minus FTIR) is (-0.02 ± 0.05) ppbv (-1.6 ± 4.2) %, which is within the FTIR
19 uncertainty budget. As a result, GEOS-Chem can simulate C₂H₆ concentration and variability for
20 the specifics of polluted regions over eastern China. With improved emission inventories, previous
21 studies have also found that global chemistry transport models were able to reproduce the absolute
22 values as well as seasonal cycles of the ground-based FTIR C₂H₆ observations in the other parts of
23 the world (Franco et al., 2015; Franco et al., 2016; Tzompa-Sosa et al. 2017).

24 As typically observed, the peak-to-peak amplitude of the modulation with respect to the
25 monthly mean of C₂H₆ troDMF spanned a large range of -16.0% to 72.8% depending on season
26 [Fig. 3 (b)]. The observed C₂H₆ troDMF roughly decreases over time for the first half of the year
27 and increases over time for the second half of the year [Fig. 3 (b)]. High levels of C₂H₆ troDMF
28 occur in the late autumn to early spring and low levels of C₂H₆ troDMF occur in late spring to early
29 autumn. The observed C₂H₆ troDMF reached a minimum monthly mean value of (0.36 ± 0.26) ppbv
30 in July and a maximum monthly mean value of (1.76 ± 0.35) ppbv in December. C₂H₆ troDMFs in
31 December were on average 4.9 times higher than those in July. Since the tropospheric OH oxidation
32 capability in summer is higher than that in winter, the C₂H₆ seasonality characterized by a winter
33 maximum and a summer minimum was also observed in other FTIR stations (Angelbratt et al. 2011;
34 Franco et al. 2015; Franco et al. 2016; Lutsch et al. 2019; Nagahama and Suzuki 2007; Rinsland et
35 al. 2002; Simpson et al. 2012; Viatte et al. 2015; Viatte et al. 2014; Vigouroux et al. 2012; Zeng et
36 al. 2012; Zhao et al. 2002). We have used the bootstrap resampling method of Gardiner et al. (2008)
37 to evaluate the seasonality and interannual variability of C₂H₆ troDMF, where a 3rd Fourier series
38 plus a linear function was used to fit daily mean time series of C₂H₆ troDMF by both FTIR
39 observations and GEOS-Chem model simulations. We incorporated the errors arising from the
40 autocorrelation in the residuals into the uncertainties in the change rates following the procedure of
41 Santer et al. (2008). The observed C₂H₆ troDMFs from 2015 to 2020 showed a negative change rate
42 of $(-2.60 \pm 1.34)\%$ yr⁻¹, which is in reasonable agreement with the modelled change rate of $(-2.1 \pm$
43 $0.7)\%$ yr⁻¹ [Fig. 3 (a)].

4. GAMs regression results and interpretation

C₂H₆ troDMF time series from 2015 to 2020 over Hefei by the FTIR and the GAMs regression model are shown in Fig. 4. The observed C₂H₆ variability can be reproduced by the GAMs regression model with a good agreement, as confirmed by a correlation coefficient (r) of 0.90 [Fig. 4(a); Fig. S3]. Meanwhile, the observed C₂H₆ troDMF time series also show high correlations with both the accumulated meteorological factor ($r=0.88$) [Fig. 4 (b)] and the accumulated emission factor ($r = 0.70$) [Fig. 4(c)], indicating both meteorological and emission influences are important factors for driving the C₂H₆ troDMF variability. Generally, both meteorological and emission factors show positive influences on C₂H₆ troDMFs in cold season (DJF/MAM) and negative influences on C₂H₆ troDMFs in warm season (JJA/SON). However, the seasonal dependency of meteorological influence is stronger than that of emission influence. During the studied years, the year to year differences in meteorological influence are small, while the positive emission influence showed an overall decreasing change rate since 2016, which probably drives a decreasing change rate in C₂H₆ troDMF in recent years.

The influence of each explanatory variable x_i in GAMs regression calculated as $100\% \cdot [e^{s(x_i)} - 1]$ are shown in Fig. 5, which reflects the influence of each individual variable on the relative change of C₂H₆ troDMF. The corresponding DOFS of each smoothing function are also shown in Fig.5. If an explanatory variable is linearly correlated with the C₂H₆ troDMF, the DOFS of the resulting smoothed variable could be equal to 1, and the larger the slope the higher the linear response. Otherwise, the larger the deviation of DOFS relative to 1, the more significant the nonlinear relationship is (Veaux and Richard 2012). During the studied years, the DOFS of zonal wind (ua), convection wind ($omega$), pressure ($pres$), tropopause height ($troph$), temperature ($temp$), and CO troDMF (co) were close to 1, reflecting a roughly linear relationship of these explanatory variables with the C₂H₆ troDMF. In contrast, the DOFS of meridional wind (va), H₂O troDMF (qv) and CH₄ troDMF (ch_4) were much greater than 1, reflecting a significant nonlinear relationship with the C₂H₆ troDMF.

The observed C₂H₆ troDMF was influenced by many factors. The zonal wind (ua), meridional wind (va), CH₄ troDMF (ch_4), and CO troDMF (co) showed positive influences and the other explanatory variables showed negative influences on the observed C₂H₆ troDMF variability. The results show that the northward, eastward, and downward convection winds facilitate the accumulation of C₂H₆ over the observation site and result in higher C₂H₆ troDMF. Since most anthropogenic and biomass burning sources of CO, and fossil fuel source of CH₄ are common sources of C₂H₆, C₂H₆ troDMF gradually went up as CH₄ and CO troDMFs increased. In contrast, meteorological conditions as high temperature, high humidity, and low pressure are more favorable to C₂H₆ oxidation, resulting in lower C₂H₆ troDMF. Meanwhile, deep upward convection wind and high tropopause height facilitate removal of C₂H₆ over the observation site and result in lower C₂H₆ troDMF.

5. Source attribution

5.1 Contributions of different source categories and regions

The absolute and relative seasonal contributions of fossil fuel, biogenic, biomass burning, and biofuel emissions to C₂H₆ variability from 2015 to 2020 over Hefei are shown in Fig. 6. The GEOS-Chem annual mean C₂H₆ troDMF simulations were decreased by 0.51, 0.27, 0.32, and 0.20 ppbv in

1 the absence of fossil fuel, biogenic, biomass burning, and biofuel C₂H₆ emission inventories, which
2 correspond to 34.6, 18.4, 21.3, and 13.5% of relative contribution to the modelled C₂H₆ variability,
3 respectively. The anthropogenic emissions account for 48.1% and the natural emissions account for
4 39.7% of the C₂H₆ variability. The remaining contribution calculated as the difference between the
5 BASE simulation and the sum of anthropogenic and natural contributions is 0.17 ppbv (12.2%).
6 This missing rest can be largely attributed to nonlinear interactional effects among different sources
7 which were not captured by the sensitivity simulations. Indeed, shutting off an emission inventory
8 may induce significantly lower concentrations in atmospheric compounds (i.e., C₂H₆ for noFF and
9 noBIOF or all suppressed compounds for noBVOC and noBB simulations) globally. On the one
10 hand, some of them may react with OH, which would lead to higher OH concentrations available
11 for the oxidation of C₂H₆, and eventually enhances the C₂H₆ destruction from other emission
12 categories. On the other hand, some of them may form OH by a series of oxidation reaction, which
13 would lead to lower OH concentrations available for the oxidation of C₂H₆, and eventually mitigates
14 the C₂H₆ destruction from other emission categories. However, it is difficult to quantify the
15 nonlinear impact of each individual emission category, since the concentration and spatial
16 distribution of C₂H₆ in each emission category are different. Especially when biogenic and biomass
17 burning emissions are suppressed, the impacts become hard to assess, since all NMVOCs
18 compounds play a key role in both OH formation and destruction. Investigating the nonlinear impact
19 of each individual emission category would require additional work that is beyond the scope of the
20 present work.

21 The contributions of all emission sources are seasonal dependent, and the fossil fuel
22 contribution shows the largest seasonal difference, which consolidates the GAMs regression results
23 that the emission influences are seasonal dependent. The fossil fuel contribution in winter and spring
24 (DJF/MAM) are larger than those in summer and autumn (JJA/SON), with a maximum of 52.0% in
25 DJF and a minimum of 13.0% in JJA. The JJA/SON meteorological conditions which show stronger
26 solar radiation, higher temperature, wetter atmospheric condition, and lower pressure than those in
27 DJF/MAM are more favorable for increasing VOCs emissions from biogenic sources (BVOCs),
28 which consolidates the fact that C₂H₆ abundance from biogenic source in JJA/SON are larger than
29 those in DJF/MAM. The missing rest contributes to a maximum of 32% in JJA when the C₂H₆
30 oxidation reaches the seasonal maximum and is thus more sensitive to the on-off state of different
31 sources.

32 Fig. 7 explores the absolute and relative seasonal contributions of ER, CR, NR, WR, and SR
33 regions to the C₂H₆ variability from 2015 to 2020 over Hefei. The GEOS-Chem annual mean C₂H₆
34 troDMF simulations were decreased by 0.28, 0.22, 0.29, 0.07, and 0.12 in the absence of the C₂H₆
35 emissions in ER, CR, NR, WR, and SR regions, which correspond to 21.5%, 15.8%, 20.3%, 5.7%,
36 and 8.9%, of relative contribution to the modelled C₂H₆ variability, respectively. The contributions
37 of all geographical regions are also seasonal dependent. The results show that the observed C₂H₆
38 variability was largely attributed to emissions within China (74.1%), which show a maximum in
39 JJA/SON and a minimum in DJF/MAM. As vicinities of the observation site, the ER, CR, and NR
40 regions dominated the contribution within China (57.6%). The remaining contribution calculated as
41 the difference between the BASE simulation and the total contributions of above individual source
42 regions is 0.42 ppbv (25.9%). This contribution is the sum of C₂H₆ emissions outside China (ROW)
43 and the nonlinear interactional effects among the geographical sensitivity simulations. This rest
44 contribution in DJF/MAM are ~ 4.0 times larger than those in JJA/SON. Considering the nonlinear

1 interactional effects mainly occur in JJA/SON but this rest contribution in the meantime shows the
2 seasonal minimum value, this remaining contribution can be largely attributed to ROW
3 contributions.

4 As a relatively long lifetime species (a few months), C₂H₆ emissions originating from either
5 nearby or in distant areas can be transported to Hefei site under favourable weather conditions, and
6 thus contribute to the observed C₂H₆ variability. In addition, atmospheric compounds originating
7 either nearby or in distant areas, which affect the chemistry of C₂H₆ oxidation, could also affect the
8 observed C₂H₆ variability. For contributions within China, the lowest contribution of the WR region
9 to the observed C₂H₆ variability is largely attributed to the lowest C₂H₆ emission rates in this region
10 (Table 2). A smaller contribution of the SR region to the observed C₂H₆ variability in DJF/MAM in
11 comparison with the ER, CR, and NR regions is mainly attributed to less air masses that originated
12 in south China under the dry winter monsoon conditions, see section 5.2.

13 **5.2 Transport inflow and outflow pathways**

14 The direct GEOS-Chem sensitivity simulations can clearly characterize the 3D transport inflow
15 and outflow pathways of C₂H₆ over the observation site. Fig.8 shows the spatial distribution of
16 GEOS-Chem C₂H₆ BASE simulations around China along with horizontal wind vectors at 900 hPa
17 in different seasons. General atmospheric circulation patterns over eastern China are typically
18 affected by mid-latitude westerlies and Asian monsoon, including the East Asian summer monsoon
19 and South Asian summer monsoon (Chen et al. 2009; Liang et al. 2005; Liu et al. 2003). Fig. 9
20 illustrates the latitude – height distributions of C₂H₆ VMR over China from six source regions along
21 with the 3D atmospheric circulation patterns in different seasons. The 3D transport inflow and
22 outflow pathways of C₂H₆ over the observation site are thus deduced as follows.

23 As indicated by the arrows in Figs. 8 and 9, the high pressure system over the Eurasian
24 continent in DJF triggers the descending of strongly cold air over eastern China and results in air
25 masses converging toward the observation site from western and northern areas, while the high
26 pressure system over the Indian ocean and Pacific in JJA triggers the ascending of strongly heated
27 air over eastern China and results in air masses converging toward the observation site from South
28 Asia and East Asia (SEAS) (Liang et al. 2004; Liu et al. 2003). In DJF, the mid-latitude westerlies
29 extend to tropics (about 15 °N) over middle and upper troposphere, and subtropics (about 30°N)
30 near the surface, while the easterlies mainly prevail in tropics and are weak over eastern China [Figs.
31 8 and 9]. In the summer monsoon season, the atmospheric circulation patterns over eastern China
32 change dramatically and is dominated by surface wind regime originating from Pacific, South China
33 Sea, or Arabian Sea [Figs.8 and 9]. Meanwhile, the mid-latitude westerlies recede to the North
34 Temperate Zone (north of 30°N) and the westerly jet center shifts to north of 50°N in JJA (from ~
35 30°N in DJF) [Fig. 8]. In JJA, the tropical region is characterized by the strong easterlies in the
36 upper troposphere and by the southwesterly air flow in the lower troposphere [Fig. 9]. The prevailing
37 winds in the transition seasons in MAM and SON are still westerlies with frequent cold fronts [Figs.
38 8 and 9]. These above seasonal circulation patterns determine the transport inflow and outflow of
39 C₂H₆ around the observation site. However, the transported scales are also influenced by source
40 location and strength, travel trajectory and travel time (Liu et al. 2003).

41 Generally, C₂H₆ emissions from CR, NR, and WR regions can be transported to the observation
42 site by the strong westerlies throughout the year [Fig. 9]. C₂H₆ from the SR region can be transported

1 to the observation site by deep convection followed by northward transport into the mid-latitude
2 westerlies in MAM or driven by the South Asian summer monsoon or westerlies in other seasons
3 (Liu et al. 2003)[Fig. 8]. The observed C₂H₆ transport inflow originating from the local ER region
4 is mainly driven by the local circulation cell or Asian monsoon. The driver for C₂H₆ transport inflow
5 originating from the northern ROW (> 32°N) is the same as those from the CR, NR, and WR regions.
6 The driver for C₂H₆ transport inflow originating from the southern ROW (< 32°N) is the same as
7 that from the SR region. However, C₂H₆ originating from ROW relative to those from China reaches
8 a higher altitude due to longer transport distances. The ROW contributions in JJA/SON are lower
9 than those in DJF/MAM can be partly attributed to a stronger removal along the long-range transport
10 pathway by the abundant wet precipitation and oxidation during the summer monsoon and post
11 monsoon season.

12 Seasonal variability in C₂H₆ transport outflow over the observation site is mainly associated
13 with the monsoon system. In DJF, C₂H₆ over the observation site is capped in the lower troposphere
14 by the subsidence over eastern China and is swept by northeasterly southwestward into southwestern
15 areas, where they are lifted up into the free troposphere by convection and then flow away
16 northeastward [Fig. 9]. In JJA, the observed C₂H₆ is transported northeastward by the Asian
17 monsoon and is undergone frequent lifting into the upper troposphere by deep convection [Figs. 8
18 and 9]. Frequent cold fronts are common phenomena during meteorologically transitional periods
19 in MAM and SON. In SON, the winter monsoon builds continental high system, and sweeps the
20 observed C₂H₆ in the lower troposphere northward to relatively high latitudes where they can be
21 lifted up into the free troposphere by deep convection or cold fronts. In MAM, convection over
22 lower latitudes at Asian continent starts to rise, which lifts up the observed C₂H₆ into the free
23 troposphere and then flow away southward.

24 **5.3 Potential factors driving interannual variability of C₂H₆**

25 China has implemented a series of active clean air policies since 2013 to mitigate severe air
26 pollution problems (Sun et al. 2020; Zhang et al. 2019; Zheng et al. 2018). Since then the emissions
27 of major air pollutants have decreased, and the overall air quality has greatly improved (Sun et al.
28 2020; Zhang et al. 2019; Zheng et al. 2018). Many air pollutants, such as NO₂, sulphur dioxide
29 (SO₂), particulate matter 2.5 (PM_{2.5}), PM₁₀, CO, black carbon (BC), and organic carbon (OC),
30 showed negative trends in recent years (Lu et al., 2019; Zhang et al. 2019; Zheng et al. 2018). We
31 follow the method of Zheng et al. (2018) to estimate the relative change rate of anthropogenic C₂H₆
32 emissions in China during 2015 – 2019 using the MEIC emission inventory. As tabulated in Table
33 2, anthropogenic C₂H₆ emissions in all geographical regions showed a decreasing change rate during
34 2015 – 2019 except those in WR region where industrialization, urbanization, land use, and
35 infrastructure construction have expanded rapidly in recent years, resulting in an increasing change
36 rate of anthropogenic C₂H₆ emissions in the region (Ran et al. 2014). The relative change rates of
37 anthropogenic C₂H₆ emissions in China during 2015 – 2019 are estimated as: 12.12% for WR, –
38 5.32% for NR, –1.03% for CR, –7.66% for ER, –5.01% for SR, and –2.74% in total. The major
39 driving forces for the decline in C₂H₆ emissions over China are attributed to the reductions from
40 residential and transport sectors, with relative change rates of –14.53% and –3.93%, respectively
41 (Table 2).

42 The interannual change rate of anthropogenic C₂H₆ emissions in China in recent years is

1 estimated to be $-0.69\% \text{ yr}^{-1}$ (calculated as $-2.74\%/(2019 - 2015)$), which is much lower than the
2 observed decreasing change rate in C_2H_6 tropDMF ($-2.60 \pm 1.34\% \text{ yr}^{-1}$), indicating additional
3 driving forces could exist, e.g., reductions in natural C_2H_6 emissions in China or reductions in long
4 range transport of C_2H_6 emissions originating either anthropogenic or natural sources outside China.
5 On the one hand, the Law of the People's Republic of China on the Prevention and Control of
6 Atmospheric Pollution included the prohibition of crop residue burning over China in 2015 because
7 crop residue burning emissions can result in poor air quality (<http://www.chinalaw.gov.cn>, last
8 access on 19 June 2020). Since then the crop residue burning events over China decreased
9 dramatically (Sun et al. 2020). Meanwhile, biomass burning events in Africa, SEAS, and Oceania
10 regions in 2015 were higher than those in onward years due to the El Niño Southern Oscillation
11 (ENSO) (Sun et al. 2020). The decreased global biomass burning emissions could probably also
12 contribute to the observed decreasing change rate in C_2H_6 tropDMF over Hefei since 2015. On the
13 other hand, large oil price fluctuations in recent years probably tightened oil and gas development
14 around the globe which can cause a reduction in C_2H_6 leakage around the globe. However, C_2H_6
15 observations around the globe and more statistical data are needed to support this deduction, which
16 is beyond the scope of this paper and requires further study.

17 **6. Conclusions**

18 Ethane (C_2H_6) is an important greenhouse (GHG) gas and plays a significant role in
19 tropospheric chemistry and climate change. As a relatively long residence time species (a few
20 months), observations of C_2H_6 can reflect regional and hemispheric changes in emissions and
21 climate, and can be assimilated into a chemical transport model to assess nonlocal emissions and
22 provide valuable insights into model biases of C_2H_6 simulations.

23 This study for the first time presents and quantifies the variability, source, and transport of
24 C_2H_6 over densely populated and highly industrialized eastern China by using ground-based high-
25 resolution Fourier transform infrared (FTIR) observations. Seasonal and interannual variabilities of
26 C_2H_6 over Hefei, eastern China from 2015 – 2020 have been investigated. The dependencies of
27 C_2H_6 on meteorological and emission factors were analyzed by using generalized additive models
28 (GAMs). The FTIR C_2H_6 time series are for the first time applied to evaluate the standard GEOS-
29 Chem full-chemistry model for the specifics of C_2H_6 simulation over eastern China. GEOS-Chem
30 model simulation with the state-of-the art inventory is in good agreement with the FTIR observation.
31 The GEOS-Chem model was further run in a sensitivity mode to quantify relative contribution of
32 various source categories and regions to the observed C_2H_6 variability. The three-dimensional (3D)
33 transport inflow and outflow pathways of C_2H_6 over the observation site were finally determined
34 by the GEOS-Chem sensitivity simulation and the analysis of the meteorological fields.

35 We obtained a retrieval error of 6.21 ± 1.2 (1σ)% and degrees of freedom (DOFS) of $1.47 \pm$
36 0.2 (1σ) in retrieval of C_2H_6 tropospheric column-averaged dry-air mole fraction (troDMF). The
37 observed C_2H_6 troDMF reached a minimum monthly mean value of (0.36 ± 0.26) ppbv in July and
38 a maximum monthly mean value of (1.76 ± 0.35) ppbv in December, and showed a negative change
39 rate of $(-2.60 \pm 1.34) \%/yr$ from 2015 to 2020. Generally, both meteorological and emission factors
40 have positive influences on C_2H_6 troDMF in cold season (DJF/MAM) and negative influences in
41 warm season (JJA/SON). GEOS-Chem model sensitivity simulations concluded that the
42 anthropogenic emissions (fossil fuel plus biofuel) accounted for 48.1% and the natural emissions

1 (biomass burning plus biogenic) accounted for 39.7% of C₂H₆ variability over Hefei. The observed
2 C₂H₆ variability over Hefei was mainly attributed to the emissions within China (74.1%), where
3 central, eastern, and northern China dominated the contribution (57.6%). Seasonal variability in
4 C₂H₆ transport inflow and outflow over the observation site is largely related to the mid-latitude
5 westerlies and Asian monsoon system. Reduction in C₂H₆ from 2015 to 2020 mainly results from
6 the decrease in local and transported C₂H₆ emissions, which points to air quality improvement in
7 China in recent years.

8 This study can not only enhance the insights of regional emission, transport, and air clean
9 actions over China, but also contribute to form new reliable remote sensing dataset in this sparsely-
10 monitored regions for climate change research.

11 **Code and data availability.** The new ground-based Fourier transform infrared (FTIR) spectroscopic
12 remote sensing dataset for atmospheric C₂H₆ over Hefei, eastern China in this study can be accessed
13 from <https://doi.org/10.6084/m9.figshare.13020545>. The MEIC emission inventories used in this
14 study are available from <http://meicmodel.org/>.

15 **Author contributions.** YS designed and wrote the paper with inputs from all coauthors. HY carried
16 out the GEOS-Chem simulations and GAMs regression. BZ provided the latest MEIC emission
17 inventory. The rest authors contributed to this work by providing constructive comments.

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

19 **Acknowledgements.** The processing and post processing environment for SFIT4 are provided by
20 National Center for Atmospheric Research (NCAR), Boulder, Colorado, USA. The NDACC
21 network is acknowledged for supplying the SFIT software. The LINEFIT code is provided by Frank
22 Hase, Karlsruhe Institute of Technology (KIT), Institute for Meteorology and Climate Research
23 (IMK-ASF), Germany. We thank the senate of Bremen, Germany for support. We thank the FTIR
24 group at university of Wollongong, Australia for help in setting up and operating the FTIR
25 spectrometer at Hefei. We thank the GEOS-Chem team and Tsinghua University, China for
26 providing the latest MEIC inventory.

27 **Financial support.** This work is jointly supported by the National Key Research and Development
28 Program of China (No.2019YFC0214802, No.2017YFC0210002, No. 2016YFC0203302,
29 2018YFC0213201, 2019YFC0214702, 2016YFC0200404), the National Science Foundation of
30 China (No.41775025, No. 41575021, No. 51778596, No. 91544212, No. 41722501, No. 51778596),
31 and the Sino-German Mobility programme (M-0036). Emmanuel Mahieu is a Senior Research
32 Associate with the Fonds de la Recherche Scientifique -FNRS. His contribution has been primarily
33 supported by the F.R.S.-FNRS under Grant no J.0147.18.

34 **References**

35 Abad, G. G., Allen, N. D. C., Bernath, P. F., Boone, C. D., McLeod, S. D., Manney, G. L., Toon, G. C.,

1 Carouge, C., Wang, Y., Wu, S., Barkley, M. P., Palmer, P. I., Xiao, Y., and Fu, T. M.: Ethane, ethyne
2 and carbon monoxide concentrations in the upper troposphere and lower stratosphere from ACE and
3 GEOS-Chem: a comparison study, *Atmos. Chem. Phys.*, 11, 9927-9941, 2011.

4 Angelbratt, J., Mellqvist, J., Simpson, D., Jonson, J. E., Blumenstock, T., Borsdorff, T., Duchatelet, P.,
5 Forster, F., Hase, F., Mahieu, E., De Maziere, M., Notholt, J., Petersen, A. K., Raffalski, U., Servais,
6 C., Sussmann, R., Warneke, T., and Vigouroux, C.: Carbon monoxide (CO) and ethane (C₂H₆) trends
7 from ground-based solar FTIR measurements at six European stations, comparison and sensitivity
8 analysis with the EMEP model, *Atmos. Chem. Phys.*, 11, 9253-9269, 2011.

9 Bey, I., Jacob, D. J., Yantosca, R. M., Logan, J. A., Field, B. D., Fiore, A. M., Li, Q. B., Liu, H. G. Y.,
10 Mickley, L. J., and Schultz, M. G.: Global modeling of tropospheric chemistry with assimilated
11 meteorology: Model description and evaluation, *J. Geophys. Res.-Atmos.*, 106, 23073-23095, 2001.

12 Bian, H. S., and Prather, M. J.: Fast-J2: Accurate simulation of stratospheric photolysis in global chemical
13 models, *J. Atmos. Chem.*, 41, 281-296, 2002.

14 Chen, D., Wang, Y., McElroy, M. B., He, K., Yantosca, R. M., and Le Sager, P.: Regional CO pollution
15 and export in China simulated by the high-resolution nested-grid GEOS-Chem model, *Atmos. Chem.*
16 *Phys.*, 9, 3825-3839, 2009.

17 De Mazière, M., Thompson, A. M., Kurylo, M. J., Wild, J. D., Bernhard, G., Blumenstock, T., Braathen,
18 G. O., Hannigan, J. W., Lambert, J. C., Leblanc, T., Mcgee, T. J., Nedoluha, G., Petropavlovskikh,
19 I., Seckmeyer, G., Simon, P. C., Steinbrecht, W., and Strahan, S. E.: The Network for the Detection
20 of Atmospheric Composition Change (NDACC): history, status and perspectives, *Atmos. Chem.*
21 *Phys.*, 18, 4935-4964, 2018.

22 Eastham, S. D., Weisenstein, D. K., and Barrett, S. R. H.: Development and evaluation of the unified
23 tropospheric-stratospheric chemistry extension (UCX) for the global chemistry-transport model
24 GEOS-Chem, *Atmos. Environ.*, 89, 52-63, 2014.

25 Fischer, E. V., Jacob, D. J., Yantosca, R. M., Sulprizio, M. P., Millet, D. B., Mao, J., Paulot, F., Singh, H.
26 B., Roiger, A., Ries, L., Talbot, R. W., Dzepina, K., and Deolal, S. P.: Atmospheric peroxyacetyl
27 nitrate (PAN): a global budget and source attribution, *Atmos. Chem. Phys.*, 14, 2679-2698, 2014.

28 Fisher, J. A., Murray, L. T., Jones, D. B. A., and Deutscher, N. M.: Improved method for linear carbon
29 monoxide simulation and source attribution in atmospheric chemistry models illustrated using
30 GEOS-Chem v9, *Geosci. Model. Dev.*, 10, 4129-4144, 2017.

31 Franco, B., Bader, W., Toon, G. C., Bray, C., Perrin, A., Fischer, E. V., Sudo, K., Boone, C. D., Bovy, B.,
32 Lejeune, B., Servais, C., and Mahieu, E.: Retrieval of ethane from ground-based FTIR solar spectra
33 using improved spectroscopy: Recent burden increase above Jungfraujoch, *J. Quant. Spectrosc. Ra.*,
34 160, 36-49, 2015.

35 Franco, B., Mahieu, E., Emmons, L. K., Tzompa-Sosa, Z. A., Fischer, E. V., Sudo, K., Bovy, B., Conway,
36 S., Griffin, D., Hannigan, J. W., Strong, K., and Walker, K. A.: Evaluating ethane and methane
37 emissions associated with the development of oil and natural gas extraction in North America,
38 *Environ. Res. Lett.*, 11, 2016.

39 Gardiner, T., Forbes, A., de Maziere, M., Vigouroux, C., Mahieu, E., Demoulin, P., Velazco, V., Notholt,
40 J., Blumenstock, T., Hase, F., Kramer, I., Sussmann, R., Stremme, W., Mellqvist, J., Strandberg, A.,
41 Ellingsen, K., and Gauss, M.: Trend analysis of greenhouse gases over Europe measured by a
42 network of ground-based remote FTIR instruments, *Atmos. Chem. Phys.*, 8, 6719-6727, 2008.

43 Giglio, L., Randerson, J. T., and van der Werf, G. R.: Analysis of daily, monthly, and annual burned area
44 using the fourth-generation global fire emissions database (GFED4), *J. Geophys. Res.-Biogeo.*, 118,

1 317-328, 10.1002/jgrg.20042, 2013.

2 Gilman, J. B., Lerner, B. M., Kuster, W. C., and de Gouw, J. A.: Source Signature of Volatile Organic
3 Compounds from Oil and Natural Gas Operations in Northeastern Colorado (vol 47, pg 1297, 2013),
4 Environ. Sci. & Tech., 47, 10094-10094, 2013.

5 Glatthor, N., von Clarmann, T., Stiller, G. P., Funke, B., Koukouli, M. E., Fischer, H., Grabowski, U.,
6 Hopfner, M., Kellmann, S., and Linden, A.: Large-scale upper tropospheric pollution observed by
7 MIPAS HCN and C₂H₆ global distributions, Atmos. Chem. Phys., 9, 9619-9634, 2009.

8 Guenther, A. B., Jiang, X., Heald, C. L., Sakulyanontvittaya, T., Duhl, T., Emmons, L. K., and Wang, X.:
9 The Model of Emissions of Gases and Aerosols from Nature version 2.1 (MEGAN2.1): an extended
10 and updated framework for modeling biogenic emissions, Geosci. Model. Dev., 5, 1471-1492, 2012.

11 Hase, F.: Improved instrumental line shape monitoring for the ground-based, high-resolution FTIR
12 spectrometers of the Network for the Detection of Atmospheric Composition Change, Atmos. Meas.
13 Tech., 5, 603-610, 2012.

14 Helmig, D., Rossabi, S., Hueber, J., Tans, P., Montzka, S. A., Masarie, K., Thoning, K., Plass-Duelmer,
15 C., Claude, A., Carpenter, L. J., Lewis, A. C., Punjabi, S., Reimann, S., Vollmer, M. K., Steinbrecher,
16 R., Hannigan, J., Emmons, L. K., Mahieu, E., Franco, B., Smale, D., and Pozzer, A.: Reversal of
17 global atmospheric ethane and propane trends largely due to US oil and natural gas production, Nat.
18 Geosci., 9, 490-495, 2016.

19 Hoesly, R. M., Smith, S. J., Feng, L. Y., Klimont, Z., Janssens-Maenhout, G., Pitkanen, T., Seibert, J. J.,
20 Vu, L., Andres, R. J., Bolt, R. M., Bond, T. C., Dawidowski, L., Kholod, N., Kurokawa, J., Li, M.,
21 Liu, L., Lu, Z. F., Moura, M. C. P., O'Rourke, P. R., and Zhang, Q.: Historical (1750-2014)
22 anthropogenic emissions of reactive gases and aerosols from the Community Emissions Data
23 System (CEDS), Geosci. Model. Dev., 11, 369-408, 2018.

24 Lin J. T., Mcelroy M. B.: Impacts of boundary layer mixing on pollutant vertical profiles in the lower
25 troposphere: Implications to satellite remote sensing, Atmos. Environ., 44(14):1726-1739, 2010.

26 Li, M., Zhang, Q., Streets, D. G., He, K. B., Cheng, Y. F., Emmons, L. K., Huo, H., Kang, S. C., Lu, Z.,
27 Shao, M., Su, H., Yu, X., and Zhang, Y.: Mapping Asian anthropogenic emissions of non-methane
28 volatile organic compounds to multiple chemical mechanisms, Atmos. Chem. Phys., 14, 5617-5638,
29 2014.

30 Li, M., Zhang, Q., Kurokawa, J., Woo, J. H., He, K. B., Lu, Z. F., Ohara, T., Song, Y., Streets, D. G.,
31 Carmichael, G. R., Cheng, Y. F., Hong, C. P., Huo, H., Jiang, X. J., Kang, S. C., Liu, F., Su, H., and
32 Zheng, B.: MIX: a mosaic Asian anthropogenic emission inventory under the international
33 collaboration framework of the MICS-Asia and HTAP, Atmos. Chem. Phys., 17, 935-963, 2017.

34 Liang, Q., Jaegle, L., Jaffe, D. A., Weiss-Penzias, P., Heckman, A., and Snow, J. A.: Long-range transport
35 of Asian pollution to the northeast Pacific: Seasonal variations and transport pathways of carbon
36 monoxide, J. Geophys. Res.-Atmos., 109, 2004.

37 Liang, Q., Jaegle, L., and Wallace, J. M.: Meteorological indices for Asian outflow and transpacific
38 transport on daily to interannual timescales, J. Geophys. Res.-Atmos., 110, 2005.

39 Lin, X., Liao, Y., and Hao, Y.: The burden associated with ambient PM_{2.5} and meteorological factors in
40 Guangzhou, China, 2012–2016: A generalized additive modeling of temporal years of life lost,
41 Chemosphere, 212, 705-714, 2018.

42 Liu, F., Zhang, Q., Tong, D., Zheng, B., Li, M., Huo, H., and He, K. B.: High-resolution inventory of
43 technologies, activities, and emissions of coal-fired power plants in China from 1990 to 2010,
44 Atmos. Chem. Phys., 15, 13299-13317, 2015.

- 1 Liu, H. Y., Jacob, D. J., Bey, I., and Yantosca, R. M.: Constraints from Pb-210 and Be-7 on wet deposition
2 and transport in a global three-dimensional chemical tracer model driven by assimilated
3 meteorological fields, *J. Geophys. Res.-Atmos.*, 106, 12109-12128, 2001.
- 4 Liu, H. Y., Jacob, D. J., Bey, I., Yantosca, R. M., Duncan, B. N., and Sachse, G. W.: Transport pathways
5 for Asian pollution outflow over the Pacific: Interannual and seasonal variations, *J. Geophys. Res.-*
6 *Atmos.*, 108, 2003.
- 7 Lu, X., Hong, J. Y., Zhang, L., Cooper, O. R., Schultz, M. G., Xu, X. B., Wang, T., Gao, M., Zhao, Y. H.,
8 and Zhang, Y. H.: Severe Surface Ozone Pollution in China: A Global Perspective, *Environ. Sci.*
9 *Tech. Let.*, 5, 487-494, 2018.
- 10 Lu, X., Zhang, L., Chen, Y. F., Zhou, M., Zheng, B., Li, K., Liu, Y. M., Lin, J. T., Fu, T. M., and Zhang,
11 Q.: Exploring 2016-2017 surface ozone pollution over China: source contributions and
12 meteorological influences, *Atmos. Chem. Phys.*, 19, 8339-8361, 2019.
- 13 Lutsch, E., Damers, E., Conway, S., and Strong, K.: Long-range transport of NH₃, CO, HCN, and C₂H₆
14 from the 2014 Canadian Wildfires, *Geophys. Res. Lett.*, 43, 8286-8297, 2016.
- 15 Lutsch, E., Strong, K., Jones, D. B. A., Blumenstock, T., Conway, S., Fisher, J. A., Hannigan, J. W., Hase,
16 F., Kasai, Y., Mahieu, E., Makarova, M., Morino, I., Nagahama, T., Notholt, J., Ortega, I., Palm, M.,
17 Poberovskii, A. V., Sussmann, R., and Warneke, T.: Detection and Attribution of Wildfire Pollution
18 in the Arctic and Northern Mid-latitudes using a Network of FTIR Spectrometers and GEOS-Chem,
19 *Atmos. Chem. Phys. Discuss.*, 2019, 1-57, 10.5194/acp-2019-881, 2019.
- 20 Ma, Y. X., Ma, B. J., Jiao, H. R., Zhang, Y. F., Xin, J. Y., and Yu, Z.: An analysis of the effects of weather
21 and air pollution on tropospheric ozone using a generalized additive model in Western China:
22 Lanzhou, Gansu, *Atmos. Environ.*, 224, 2020.
- 23 McKain, K., Down, A., Raciti, S. M., Budney, J., Hutyra, L. R., Floerchinger, C., Herndon, S. C.,
24 Nehrkorn, T., Zahniser, M. S., Jackson, R. B., Phillips, N., and Wofsy, S. C.: Methane emissions
25 from natural gas infrastructure and use in the urban region of Boston, Massachusetts, *P. Natl. Acad.*
26 *Sci. USA*, 112, 1941-1946, 2015.
- 27 Monks, S. A., Wilson, C., Emmons, L. K., Hannigan, J. W., Helmig, D., Blake, N. J., and Blake, D. R.:
28 Using an Inverse Model to Reconcile Differences in Simulated and Observed Global Ethane
29 Concentrations and Trends Between 2008 and 2014, *J. Geophys. Res.-Atmos.*, 123, 11262-11282,
30 2018.
- 31 Nagahama, Y., and Suzuki, K.: The influence of forest fires on CO, HCN, C₂H₆, and C₂H₂ over northern
32 Japan measured by infrared solar spectroscopy, *Atmos. Environ.*, 41, 9570-9579, 2007.
- 33 Notholt, J., Toon, G. C., Rinsland, C. P., Pougatchev, N. S., Jones, N. B., Connor, B. J., Weller, R.,
34 Gautrois, M., and Schrems, O.: Latitudinal variations of trace gas concentrations in the free
35 troposphere measured by solar absorption spectroscopy during a ship cruise, *J. Geophys. Res.-*
36 *Atmos.*, 105, 1337-1349, 2000.
- 37 Philip, S., Martin, R. V., and Keller, C. A.: Sensitivity of chemistry-transport model simulations to the
38 duration of chemical and transport operators: a case study with GEOS-Chem v10-01, *Geosci. Model*
39 *Dev.*, 9(5), 1683-1695. <https://doi.org/10.5194/gmd-9-1683-2016>, 2016.
- 40 Pearce, J. L., Beringer, J., Nicholls, N., Hyndman, R. J., and Tapper, N. J.: Quantifying the influence of
41 local meteorology on air quality using generalized additive models, *Atmos. Environ.*, 45, 1328-
42 1336, 2011.
- 43 Pougatchev, N. S., Connor, B. J., and Rinsland, C. P.: Infrared Measurements of the Ozone Vertical-
44 Distribution above Kitt Peak, *J. Geophys. Res.-Atmos.*, 100, 16689-16697, 1995.

1 Ran, L., Lin, W. L., Deji, Y. Z., La, B., Tsering, P. M., Xu, X. B., and Wang, W.: Surface gas pollutants
2 in Lhasa, a highland city of Tibet – current levels and pollution implications, *Atmos. Chem.*
3 *Phys.*, 14, 10721-10730, 10.5194/acp-14-10721-2014, 2014.

4 Rinsland, C. P., Jones, N. B., Connor, B. J., Wood, S. W., Goldman, A., Stephen, T. M., Murcray, F. J.,
5 Chiou, L. S., Zander, R., and Mahieu, E.: Multiyear infrared solar spectroscopic measurements of
6 HCN, CO, C₂H₆, and C₂H₂ tropospheric columns above Lauder, New Zealand (45 degrees S latitude),
7 *J. Geophys. Res.-Atmos.*, 107, 2002.

8 Rodgers, C.: *Inverse Methods for Atmospheric Sounding - Theory and Practice*, in, 2000.

9 Rodgers, C. D., and Connor, B. J.: Intercomparison of remote sounding instruments, *J. Geophys. Res.-*
10 *Atmos.*, 108, Artn 411610.1029/2002jd002299, 2003.

11 Rothman, L. S., Gordon, I. E., Babikov, Y., Barbe, A., Benner, D. C., Bernath, P. F., Birk, M.,
12 Bizzocchi, L., Boudon, V., Brown, L. R., Campargue, A., Chance, K., Cohen, E. A., Coudert, L.
13 H., Devi, V. M., Drouin, B. J., Fayt, A., Flaud, J.-M., Gamache, R. R., Harrison, J. J., Hartmann,
14 J.-M., Hill, C., Hodges, J. T., Jacquemart, D., Jolly, A., Lamouroux, J., Roy, R. J. L., Li, G.,
15 Long, D. A., Lyulin, O. M., Mackie, C. J., Massie, S. T., Mikhailenk, S., Müller, H. S. P.,
16 Naumenko, O. V., and Nikitin, A. V.: The HITRAN2012 molecular spectroscopic database, *J.*
17 *Quant. Spectrosc. Ra.*, 130, 4–50, <https://doi.org/10.1016/j.jqsrt.2013.07.002>, 2013.

18 Roscioli, J. R., Yacovitch, T. I., Floerchinger, C., Mitchell, A. L., Tkacik, D. S., Subramanian, R.,
19 Martinez, D. M., Vaughn, T. L., Williams, L., Zimmerle, D., Robinson, A. L., Herndon, S. C., and
20 Marchese, A. J.: Measurements of methane emissions from natural gas gathering facilities and
21 processing plants: measurement methods, *Atmos. Meas. Tech.*, 8, 2017-2035, 2015.

22 Santer, B. D., Thorne, P. W., Haimberger, L., Taylor, K. E., Wigley, T. M. L., Lanzante, J. R., Solomon,
23 S., Free, M., Gleckler, P. J., Jones, P. D., Karl, T. R., Klein, S. A., Mears, C., Nychka, D., Schmidt,
24 G. A., Sherwood, S. C., and Wentz, F. J.: Consistency of modelled and observed temperature trends
25 in the tropical troposphere, *Int. J. Climatol.*, 28, 1703-1722, 2008.

26 Shaiganfar, R., Beirle, S., van der Gon, H. D., Jonkers, S., Kuenen, J., Petetin, H., Zhang, Q., Beekmann,
27 M., and Wagner, T.: Estimation of the Paris NO_x emissions from mobile MAX-DOAS observations
28 and CHIMERE model simulations during the MEGAPOLI campaign using the closed integral
29 method, *Atmos. Chem. Phys.*, 17, 7853-7890, 2017.

30 Shan, C. G., Wang, W., Liu, C., Sun, Y. W., Hu, Q. H., Xu, X. W., Tian, Y., Zhang, H. F., Morino, I.,
31 Griffit, D. W. T., and Velazco, V. A.: Regional CO emission estimated from ground-based remote
32 sensing at Hefei site, China, *Atmos. Res.*, 222, 25-35, 2019.

33 Simpson, I. J., Andersen, M. P. S., Meinardi, S., Bruhwiler, L., Blake, N. J., Helmig, D., Rowland, F. S.,
34 and Blake, D. R.: Long-term decline of global atmospheric ethane concentrations and implications
35 for methane, *Nature*, 488, 490-494, 2012.

36 Singh, H., Chen, Y., Staudt, A., Jacob, D., Blake, D., Heikes, B., and Snow, J.: Evidence from the Pacific
37 troposphere for large global sources of oxygenated organic compounds, *Nature*, 410, 1078-1081,
38 2001.

39 Steinfeld, J. I.: *Atmospheric Chemistry and Physics: From Air Pollution to Climate Change*,
40 *Environment: Science and Policy for Sustainable Development*, 40, 26-26,
41 10.1080/00139157.1999.10544295, 1998.

42 Sun, Y., Yin, H., Liu, C., Zhang, L., Cheng, Y., Palm, M., Notholt, J., Lu, X., Vigouroux, C., Zheng, B.,
43 Wang, W., Jones, N., Shan, C., Qin, M., Tian, Y., Hu, Q., Meng, F., and Liu, J.: Mapping the drivers
44 of formaldehyde (HCHO) variability from 2015 to 2019 over eastern China: insights from Fourier

1 transform infrared observation and GEOS-Chem model simulation, *Atmos. Chem. Phys.*, 21, 6365–
2 6387, <https://doi.org/10.5194/acp-21-6365-2021>, 2021.

3 Sun, Y. W., Liu, C., Palm, M., Vigouroux, C., Notholt, J., Hui, Q. H., Jones, N., Wang, W., Su, W. J.,
4 Zhang, W. Q., Shan, C. G., Tian, Y., Xu, X. W., De Maziere, M., Zhou, M. Q., and Liu, J. G.: Ozone
5 seasonal evolution and photochemical production regime in the polluted troposphere in eastern
6 China derived from high-resolution Fourier transform spectrometry (FTS) observations, *Atmos.*
7 *Chem. Phys.*, 18, 14569-14583, 2018a.

8 Sun, Y. W., Palm, M., Liu, C., Hase, F., Griffith, D., Weinzierl, C., Petri, C., Wang, W., and Notholt, J.:
9 The influence of instrumental line shape degradation on NDACC gas retrievals: total column and
10 profile, *Atmos. Meas. Tech.*, 11, 2879-2896, 2018b.

11 Sun, Y. W., Liu, C., Zhang, L., Palm, M., Notholt, J., Hao, Y., Vigouroux, C., Lutsch, E., Wang, W., Shan,
12 C. G., Blumenstock, T., Nagahama, T., Morino, I., Mahieu, E., Strong, K., Langerock, B., De
13 Maziere, M., Hu, Q. H., Zhang, H. F., Petri, C., and Liu, J. G.: Fourier transform infrared time series
14 of tropospheric HCN in eastern China: seasonality, interannual variability, and source attribution,
15 *Atmos. Chem. Phys.*, 20, 5437-5456, 2020.

16 Taylor, T. E., Eldering, A., Merrelli, A., Kiel, M., Somkuti, P., Cheng, C., Rosenberg, R., Fisher, B., Crisp,
17 D., Basilio, R., Bennett, M., Cervantes, D., Chang, A., Dang, L., Frankenberg, C., Haemmerle, V.
18 R., Keller, G. R., Kurosu, T., Laughner, J. L., Lee, R., Marchetti, Y., Nelson, R. R., O'Dell, C. W.,
19 Osterman, G., Pavlick, R., Roehl, C., Schneider, R., Spiers, G., To, C., Wells, C., Wennberg, P. O.,
20 Yelamanchili, A., and Yu, S.: OCO-3 early mission operations and initial (vEarly) XCO₂ and SIF
21 retrievals, *Remote Sens. Environ.*, 251, 112032, <https://doi.org/10.1016/j.rse.2020.112032>, 2020.

22 Tian, Y., Sun, Y. W., Liu, C., Wang, W., Shan, C. G., Xu, X. W., and Hu, Q. H.: Characterisation of
23 methane variability and trends from near-infrared solar spectra over Hefei, China, *Atmos. Environ.*,
24 173, 198-209, 2018.

25 Tzompa-Sosa, Z. A., Mahieu, E., Franco, B., Keller, C. A., Turner, A. J., Helmig, D., Fried, A., Richter,
26 D., Weibring, P., Walega, J., Yacovitch, T. I., Herndon, S. C., Blake, D. R., Hase, F., Hannigan, J.
27 W., Conway, S., Strong, K., Schneider, M., and Fischer, E. V.: Revisiting global fossil fuel and
28 biofuel emissions of ethane, *J. Geophys. Res.-Atmos.*, 122, 2493-2512, 2017.

29 Van Dingenen, R., Dentener, F. J., Raes, F., Krol, M. C., Emberson, L., and Cofala, J.: The global impact
30 of ozone on agricultural crop yields under current and future air quality legislation, *Atmos. Environ.*,
31 43, 604-618, 2009.

32 Veaux, D., and Richard, D.: Generalized Additive Models, *Technometrics*, 34, 225-226, 2012.

33 Viatte, C., Strong, K., Walker, K. A., and Drummond, J. R.: Five years of CO, HCN, C₂H₆, C₂H₂, CH₃OH,
34 HCOOH and H₂CO total columns measured in the Canadian high Arctic, *Atmos. Meas. Tech.*, 7,
35 1547-1570, 2014.

36 Viatte, C., Strong, K., Hannigan, J., Nussbaumer, E., Emmons, L. K., Conway, S., Paton-Walsh, C.,
37 Hartley, J., Benmergui, J., and Lin, J.: Identifying fire plumes in the Arctic with tropospheric FTIR
38 measurements and transport models, *Atmos. Chem. Phys.*, 15, 2227-2246, 2015.

39 Vigouroux, C., Stavrakou, T., Whaley, C., Dils, B., Duflot, V., Hermans, C., Kumps, N., Metzger, J. M.,
40 Scolas, F., Vanhaelewyn, G., Muller, J. F., Jones, D. B. A., Li, Q., and De Maziere, M.: FTIR time-
41 series of biomass burning products (HCN, C₂H₆, C₂H₂, CH₃OH, and HCOOH) at Reunion Island
42 (21 degrees S, 55 degrees E) and comparisons with model data, *Atmos. Chem. Phys.*, 12, 10367-
43 10385, 2012.

44 Wang, W., Tian, Y., Liu, C., Sun, Y. W., Liu, W. Q., Xie, P. H., Liu, J. G., Xu, J., Morino, I., Velasco, V.

1 A., Griffith, D. T., Notholt, J., and Warneke, T.: Investigating the performance of a greenhouse gas
2 observatory in Hefei, China, *Atmos. Meas. Tech.*, 10, 2627-2643, 2017.

3 Wesely, M. L.: Parameterization of Surface Resistances to Gaseous Dry Deposition in Regional-Scale
4 Numerical-Models, *Atmos. Environ.*, 23, 1293-1304, 1989.

5 Wood, and Simon, N.: Stable and Efficient Multiple Smoothing Parameter Estimation for Generalized
6 Additive Models, *Journal of the American Statistical Association*, 99, 673-686, 2004.

7 Wood, S. N.: Stable and efficient multiple smoothing parameter estimation for generalized additive
8 models, *Journal of the American Statistical Association*, 99, 673-686, 2004.

9 Xiao, Y. P., Logan, J. A., Jacob, D. J., Hudman, R. C., Yantosca, R., and Blake, D. R.: Global budget of
10 ethane and regional constraints on US sources, *J. Geophys. Res.-Atmos.*, 113, 2008.

11 Yin, H., Sun, Y. W., Liu, C., Zhang, L., Lu, X., Wang, W., Shan, C. G., Hu, Q. H., Tian, Y., Zhang, C. X.,
12 Su, W. J., Zhang, H. F., Palm, M. A., Notholt, J., and Liu, J. G.: FTIR time series of stratospheric
13 NO₂ over Hefei, China, and comparisons with OMI and GEOS-Chem model data, *Opt. Express*,
14 27, A1225-A1240, 2019.

15 Yin, H., Sun, Y. W., Liu, C., Lu, X., Smale, D., Blumenstock, T., Nagahama, T., Wang, W., Tian, Y., Hu,
16 Q. H., Shan, C. G., Zhang, H. F., and Liu, J. G.: Ground-based FTIR observation of hydrogen
17 chloride (HCl) over Hefei, China, and comparisons with GEOS-Chem model data and other ground-
18 based FTIR stations data, *Opt. Express*, 28, 8041-8055, 2020.

19 Zeng, G., Wood, S. W., Morgenstern, O., Jones, N. B., Robinson, J., and Smale, D.: Trends and variations
20 in CO, C₂H₆, and HCN in the Southern Hemisphere point to the declining anthropogenic emissions
21 of CO and C₂H₆, *Atmos. Chem. Phys.*, 12, 7543-7555, 2012.

22 Zhang, C. X., Liu, C., Hu, Q. H., Cai, Z. N., Su, W. J., Xia, C. Z., Zhu, Y. Z., Wang, S. W., and Liu, J. G.:
23 Satellite UV-Vis spectroscopy: implications for air quality trends and their driving forces in China
24 during 2005-2017, *Light-Sci. Appl.*, 8, 2019.

25 Zhang, L. M., Gong, S. L., Padro, J., and Barrie, L.: A size-segregated particle dry deposition scheme for
26 an atmospheric aerosol module, *Atmos. Environ.*, 35, 549-560, 2001.

27 Zhao, Y., Strong, K., Kondo, Y., Koike, M., Matsumi, Y., Irie, H., Rinsland, C. P., Jones, N. B., Suzuki,
28 K., Nakajima, H., Nakane, H., and Murata, I.: Spectroscopic measurements of tropospheric CO,
29 C₂H₆, C₂H₂, and HCN in northern Japan, *J. Geophys. Res.-Atmos.*, 107, 2002.

30 Zheng, B., Huo, H., Zhang, Q., Yao, Z. L., Wang, X. T., Yang, X. F., Liu, H., and He, K. B.: High-
31 resolution mapping of vehicle emissions in China in 2008, *Atmos. Chem. Phys.*, 14, 9787-9805,
32 2014.

33 Zheng, B., Tong, D., Li, M., Liu, F., Hong, C. P., Geng, G. N., Li, H. Y., Li, X., Peng, L. Q., Qi, J., Yan,
34 L., Zhang, Y. X., Zhao, H. Y., Zheng, Y. X., He, K. B., and Zhang, Q.: Trends in China's
35 anthropogenic emissions since 2010 as the consequence of clean air actions, *Atmos. Chem. Phys.*,
36 18, 14095-14111, 10.5194/acp-18-14095-2018, 2018.

37

38

1 **Tables**

2 **Table 1.** Error budget and degrees of freedom (DOFS) for signal of randomly selected C₂H₆ troDMF retrieval at
3 Hefei, eastern China

Error source	Input value	Error budget
Temperature uncertainty	1σ of NCEP	1.69%
Zero level uncertainty	1%	1.45%
Retrieval parameters uncertainty	*	< 0.1%
Measurement error	1/SNR ²	0.53%
Interfering species uncertainty	SD of WACCM	0.11%
Smoothing uncertainty	*	0.37%
Total random error	/	2.32%
Background curvature uncertainty	1%	0.14%
Field of view uncertainty	1%	< 0.1%
Optical path difference uncertainty	1%	< 0.1%
Solar zenith angle uncertainty	1%	< 0.1%
Phase uncertainty	1%	< 0.1%
ILS uncertainty	1%	< 0.1%
Line temperature broadening uncertainty	5%	0.4%
Line intensity uncertainty	5%	5.12%
Line pressure broadening uncertainty	5%	0.93%
Total systematic error	/	5.48%
Total errors	/	6.21%
DOFS (-)	/	1.47

4 * These input values for error budget estimation are based on the retrieval output

5 **Table 2.** Anthropogenic C₂H₆ emissions in China by region and category for the 2015 and 2019 MEIC emission
6 inventories

Region		Industry (Tg yr ⁻¹)	Power plant (Tg yr ⁻¹)	Residential (Tg yr ⁻¹)	Transport (Tg yr ⁻¹)	Sum (Tg yr ⁻¹)
WR	2015	0.084	<0.01	0.011	<0.01	0.1
	2019	0.097	<0.01	0.011	<0.01	0.112
	change	15.36%	82.54%	- 6.61%	- 3.41%	12.12%
NR	2015	0.241	<0.01	0.125	0.026	0.394
	2019	0.241	<0.01	0.105	0.025	0.373
	change	0.04%	2.51%	- 15.96%	- 4.38%	- 5.32%
CR	2015	0.144	<0.01	0.041	<0.01	0.189
	2019	0.15	<0.01	0.033	<0.01	0.187
	change	4.68%	- 7.00%	-20.75%	- 1.13%	- 1.03%
ER	2015	0.07	<0.01	0.026	0.01	0.11
	2019	0.067	<0.01	0.022	0.01	0.097
	change	-4.83%	5.40%	-16.79%	- 3.70%	-7.66%
SR	2015	0.06	<0.01	0.026	0.01	0.09
	2019	0.056	<0.01	0.027	0.01	0.09
	change	- 7.94%	- 9.26%	1.52%	- 4.24%	- 5.01%
China	2015	0.60	<0.01	0.231	0.05	0.883
	2019	0.612	<0.01	0.197	0.048	0.859
	change	1.91%	4.04%	-14.53%	-3.93%	-2.74%

7
8

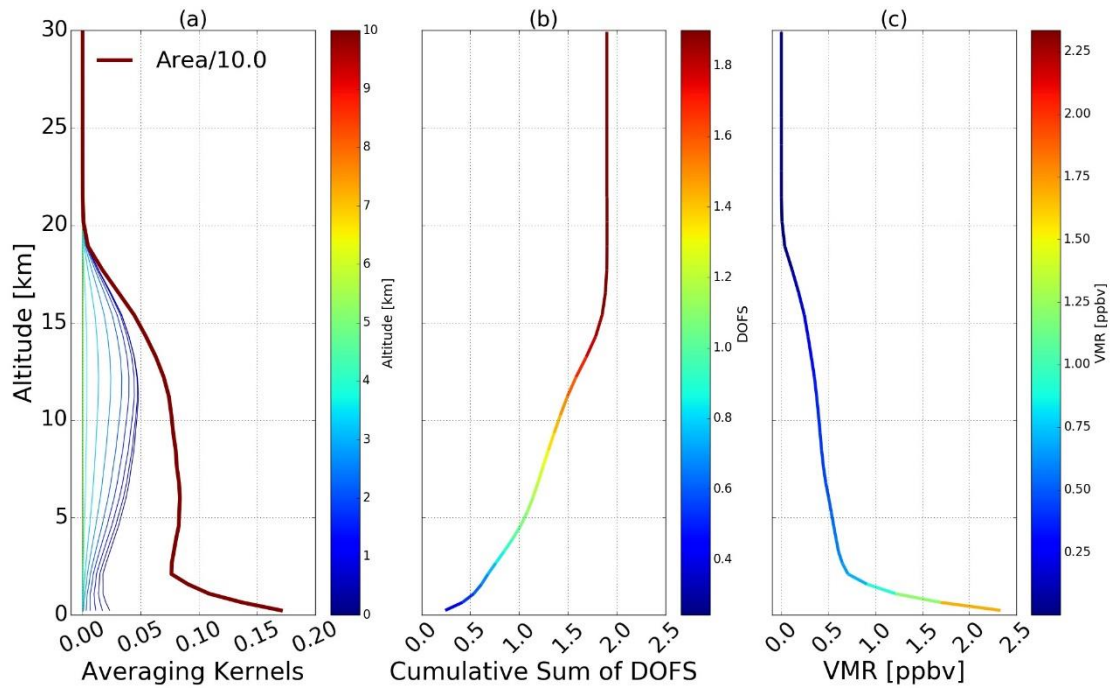
1
2

Table 3. GEOS-Chem model configurations and delimitations of all geographical regions used in sensitivity simulations.

Simulation	Region	Description
BASE	Global	Standard simulation with all anthropogenic and natural C ₂ H ₆ emissions. The BASE simulation is taken as the reference and used for model evaluation
noFF	Global	Turn off global fossil fuel C ₂ H ₆ emissions in BASE simulation
noBVOC	Global	Turn off global biogenic C ₂ H ₆ emissions in BASE simulation
noBB	Global	Turn off global biomass burning C ₂ H ₆ emissions in BASE simulation
noBIOF	Global	Turn off global biofuel C ₂ H ₆ emissions in BASE simulation
Rest	Global	Difference between BASE and the sum of FF, BVOC, BB, and BIOF contributions
noWR	78.6° E – 103.4° E; 27.6°N - 48.8°N	Turn off fossil fuel, biogenic, biomass burning, and biofuel C ₂ H ₆ emissions within western China (WR), i.e., region ① in Fig. 2, in BASE simulation
noNR	103.4°E – 129.8°E; 34.6°N – 53.5°N	Turn off fossil fuel, biogenic, biomass burning, and biofuel C ₂ H ₆ emissions within northern China (NR), i.e., region ② in Fig. 2, in BASE simulation
noCR	103.4°E – 115.6°E; 27.6°N – 34.6°N	Turn off fossil fuel, biogenic, biomass burning, and biofuel C ₂ H ₆ emissions within central China (CR), i.e., region ③ in Fig. 2, in BASE simulation
noER	115.6°E – 123.6°E; 21.0°N – 34.6°N	Turn off fossil fuel, biogenic, biomass burning, and biofuel C ₂ H ₆ emissions within eastern China (ER), i.e., region ④ in Fig. 2, in BASE simulation
noSR	98.1°E – 115.6°E; 21.0°N – 27.6°N	Turn off fossil fuel, biogenic, biomass burning, and biofuel C ₂ H ₆ emissions within southern China (SR), i.e., region ⑤ in Fig. 2, in BASE simulation
ROW	Rest of world	Difference between BASE and the sum of WR, NR, CR, ER, and SR contributions

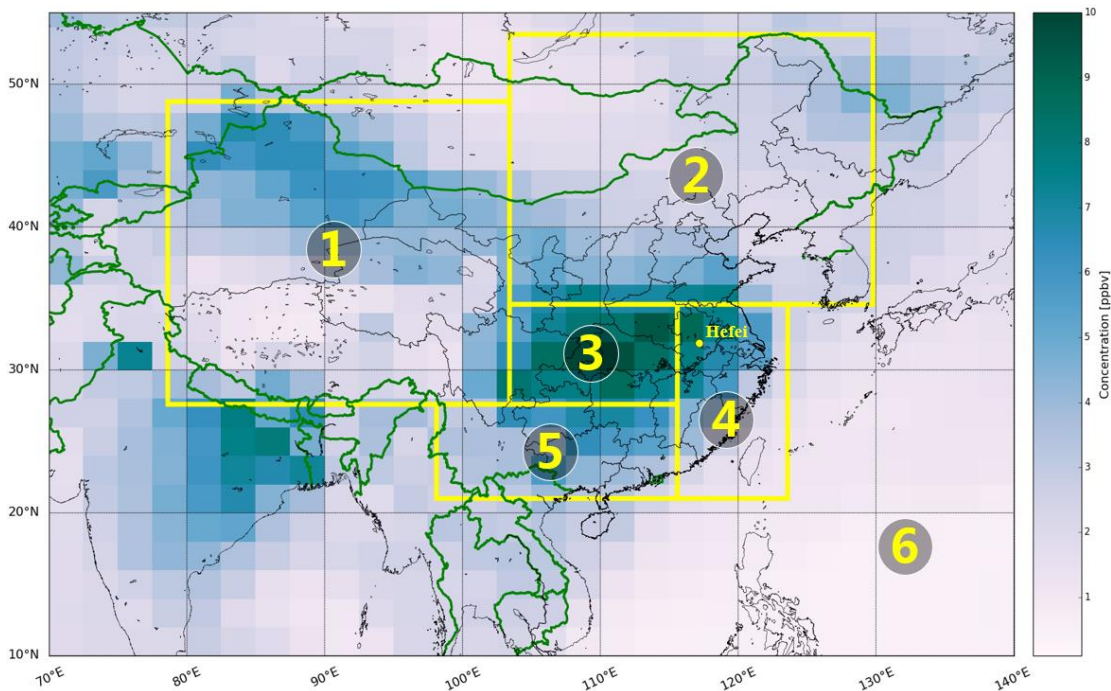
3
4

1 **Figures**



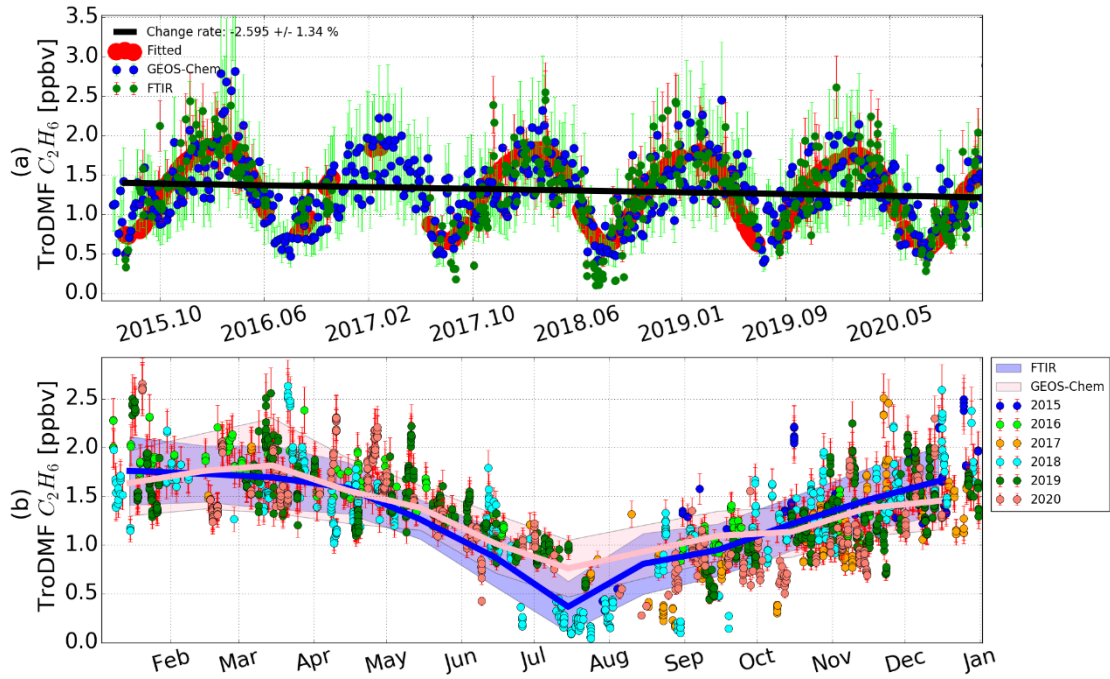
2

3 **Fig. 1.** (a) Averaging kernels and their area scaled by a factor of 0.1, (b) cumulative sum of degrees of freedom for
 4 signal (DOFS), and (c) volume mixing ratio (VMR) profile of randomly selected C_2H_6 retrieval over Hefei, eastern
 5 China.



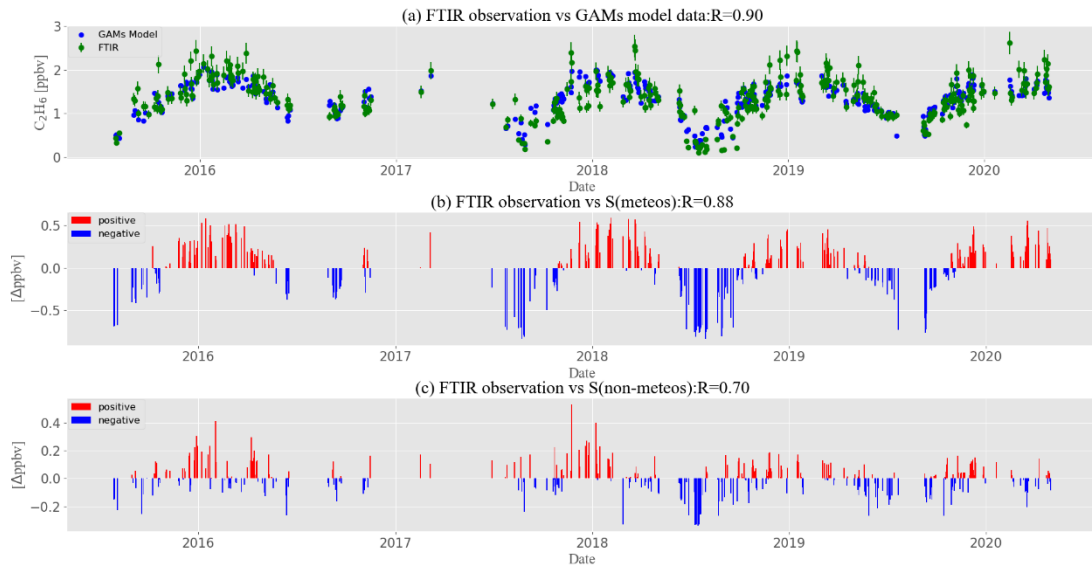
6

7 **Fig. 2.** Geographical regions used for GEOS-Chem sensitivity simulations. The numbers ①–⑥ represent western,
 8 northern, central, eastern, and southern China, and the rest of world, respectively. See Table 3 for latitude and
 9 longitude delimitations. Daily mean values of C_2H_6 troDMF on 1 January 2017 provided by GEOS-Chem BASE
 10 simulation was selected as a representative of wintertime enhancement in eastern China. The base map of this figure
 11 is created by the Basemap package of Python.



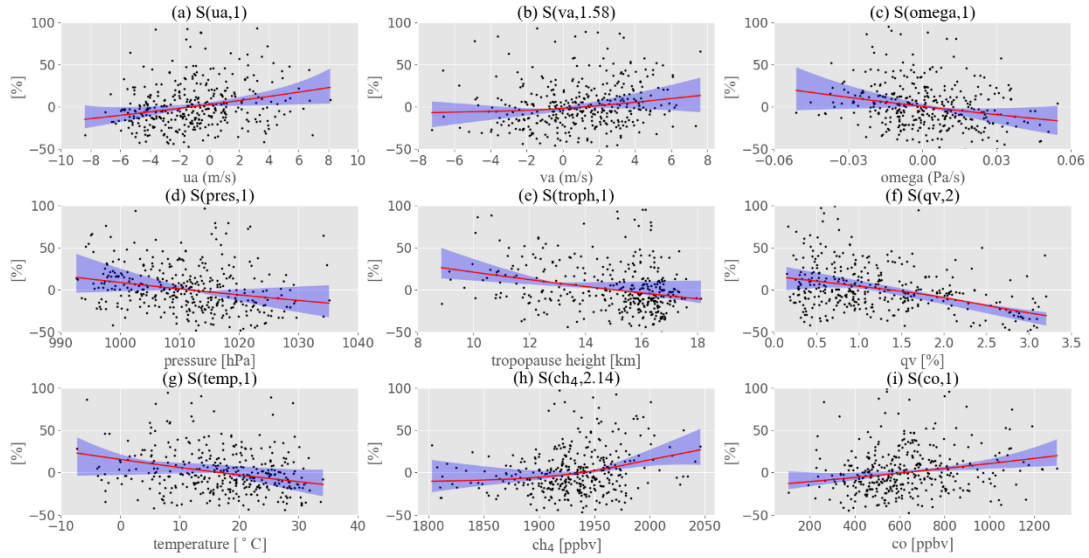
1
2
3
4
5
6
7
8
9

Fig. 3. (a) C_2H_6 troDMF time series comparison between FTIR observation and GEOS-Chem model BASE simulation from 2015 to 2020 over Hefei, eastern China. The seasonality and interannual variability are represented by red dots and black line, respectively, which are fitted by using a bootstrap resampling model with a 3rd Fourier series plus a linear function. (b) Seasonal variations of C_2H_6 troDMF by FTIR and GEOS-Chem simulation. Bold curves and the shadows are monthly mean values and the 1- σ standard variations, respectively. Vertical error bars for FTIR and GEOS-Chem represent retrieval uncertainties and diurnal variabilities, respectively. A retrieval error of 6.21% in Table 1 was used to estimate the absolute uncertainties of all observations. As a result, the absolute uncertainties in winter are larger than those in summer due to a higher C_2H_6 level in the season.



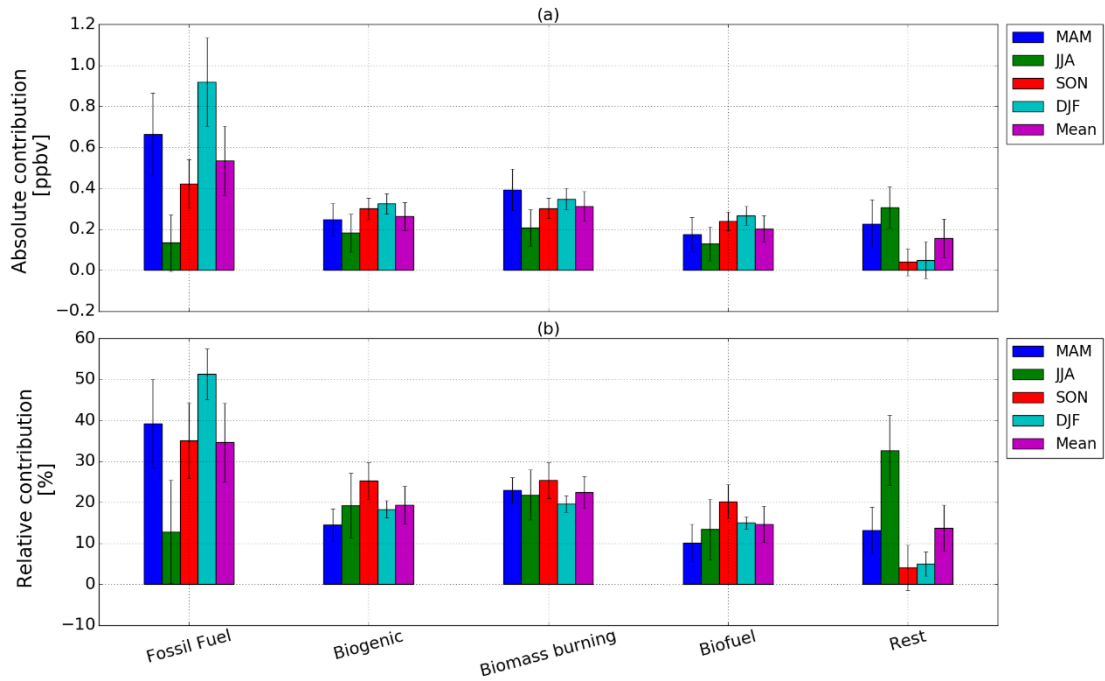
10
11
12
13
14
15

Fig. 4. (a) C_2H_6 troDMF time series from 2015 to 2020 over Hefei, eastern China by FTIR and GAMs regression model. (b) Time series of accumulated meteorological smooth functions ($S(meteos)$), and (c) time series of accumulated emission smooth functions ($S(non-meteos)$). Positive and negative influences are indicated with red and blue bars, respectively. Correlation coefficients for the total, meteorological, and emission influences are also shown.



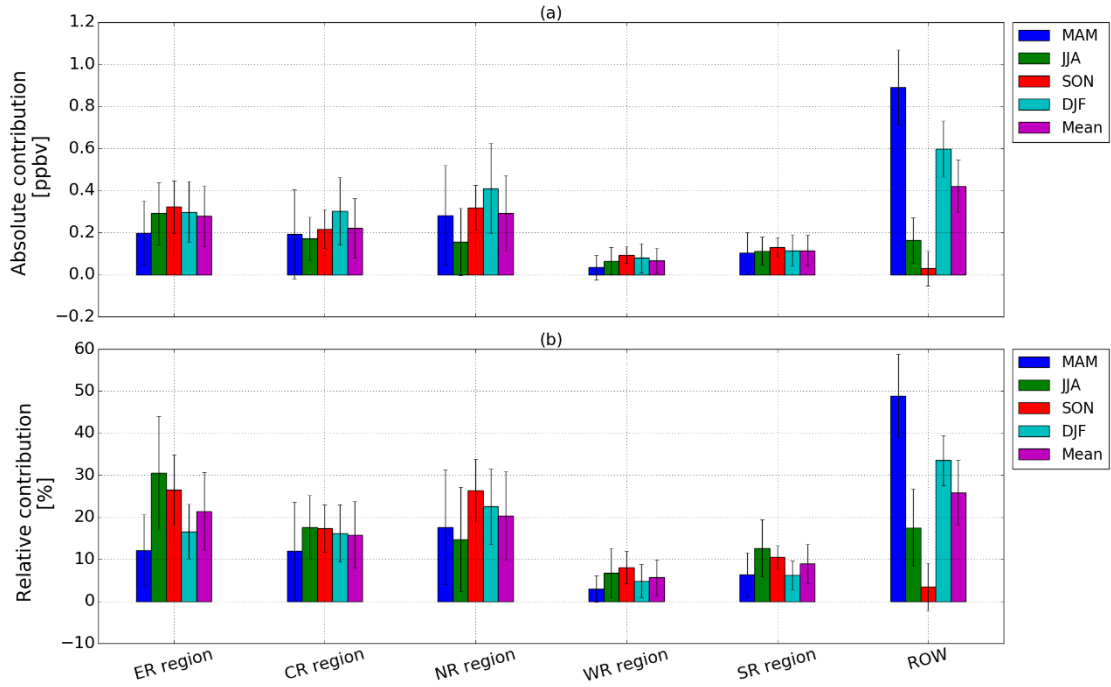
1
2
3
4
5
6
7

Fig. 5. Influence of each individual variable in the GAMs model on C_2H_6 troDMF from 2015 to 2020 over Hefei, eastern China. (a) to (i) are for zonal wind (ua), meridional wind (va), convection wind ($omega$), pressure ($pres$), tropopause height ($troph$), H_2O troDMF (qv), temperature ($temp$), CH_4 troDMF (ch_4), and CO troDMF (co), respectively. The DOFS of each smoothing function is included in the bracket in each panel. The x-axis represents variation range of each variable and the y-axis represents relative percentage change of C_2H_6 troDMF relative to its annual mean value.

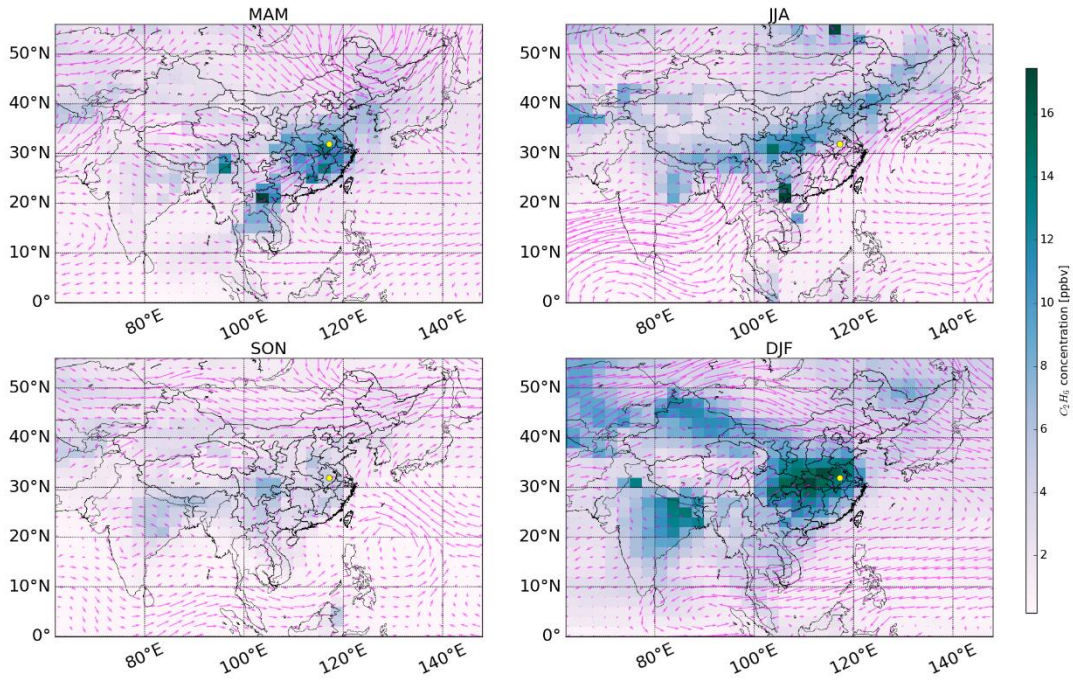


8
9
10
11
12

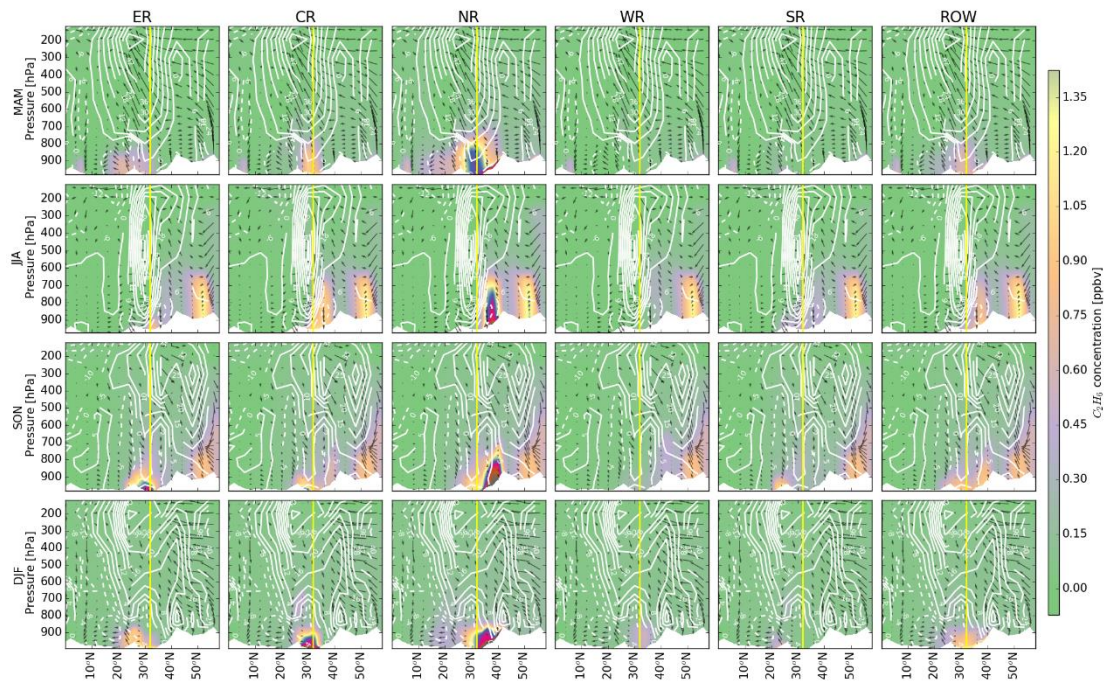
Fig. 6. Absolute (a) and relative (b) contributions of fossil fuel, biogenic, biomass burning, and biofuel emission sources to the observed C_2H_6 variability from 2015 to 2020 over Hefei, eastern China. The remaining contribution calculated as the difference between the BASE simulation and the sum of anthropogenic and natural contributions is also shown. All contributions are grouped by season. Vertical error bars represent 1- σ standard variation.



1
2 **Fig. 7.** Absolute (a) and relative (b) contributions of ER, CR, NR, WR, SR, and ROW regions to the observed C_2H_6
3 variability from 2015 to 2020 over Hefei, eastern China. All contributions are grouped by season. Geographical
4 definition of each region is summarised in Table 3. Vertical error bars represent 1- σ standard variation.



5
6 **Fig. 8.** Spatial distribution of C_2H_6 troDMF in the GEOS-Chem BASE simulations in different seasons. The arrows
7 indicate horizontal wind vectors at 900 hPa; the observation site is marked with a yellow dot. Meteorological fields
8 are from the GEOS-FP $0.25^\circ \times 0.3125^\circ$ dataset.



1

2

3

4

5

6

7

8

Fig. 9. The first row shows the latitude–height distributions of C_2H_6 VMR averaged over $113\text{--}121^\circ$ E in spring (MAM), originating in different source regions (corresponding to different columns). See Table 3 for latitude and longitude definitions. The white area indicates topography, and the white contours at intervals of 6 m s^{-1} are the easterly (dashed) and westerly (solid) mean meridional winds; the wind vectors (consisting of zonal wind in m s^{-1} and vertical velocity in units of Pa s^{-1}) are represented by arrows; the observation site is marked with a yellow line. The second to fourth rows are the same as the first row but for summer (JJA), autumn (SON), and winter (DJF), respectively. Meteorological fields are from the GEOS-FP $0.25^\circ \times 0.3125^\circ$ dataset.