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Impacts of biogenic and anthropogenic emissions on summertime ozone formation in the Guanzhong Basin, China

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Abstract. This study is the first attempt to understand the synergistic impact of anthropogenic and biogenic emissions on summertime ozone (O₃) formation in the Guanzhong (GZ) Basin where Xi'an, the oldest and the most populous city (with a population of 9 million) in northwestern China, is located. Month-long (August 2011) WRF-Chem simulations with different sensitivity experiments were conducted and compared with near-surface measurements. Biogenic volatile organic compounds (VOCs) concentrations was characterized from six surface sites among the Oinling Mountains, and urban air composition was measured in Xi'an city at a tower 100 ma.s. The WRF-Chem control experiment reasonably reproduced the magnitudes and variations of observed O₃, VOCs, NO_x, PM_{2.5}, and meteorological parameters, with normalized mean biases for each parameter within ± 21 %. Subsequent analysis employed the factor separation approach (FSA) to quantitatively disentangle the pure and synergistic impacts of anthropogenic and/or biogenic sources on summertime O₃ formation. The impact of anthropogenic sources alone was found to be dominant for O₃ formation. Although anthropogenic particles reduced NO₂ photolysis by up to 60%, the anthropogenic sources contributed $19.1\,\mathrm{ppb}$ O_3 formation on average for urban Xi'an. The abundant biogenic VOCs from the nearby forests promoted O_3 formation in urban areas by interacting with the anthropogenic NO_x . The calculated synergistic contribution (from both biogenic and anthropogenic sources) was up to $14.4\,\mathrm{ppb}$ in urban Xi'an, peaking in the afternoon. Our study reveals that the synergistic impact of individual source contributions to O_3 formation should be considered in the formation of air pollution control strategies, especially for big cities in the vicinity of forests.

1 Introduction

Elevated ozone (O₃) levels in China have drawn increasing attention in recent years (e.g., Xue et al., 2014; Hu et al., 2016; Wang et al., 2017, 2016). O₃, a secondary pollutant, is mainly formed by complex photochemical reactions of nitrogen oxides (NO_x = NO + NO₂) and volatile organic compounds (VOCs). High O₃ concentration at ground level is

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harmful to human health and ecosystems (WHO, 2005; Feng et al., 2015; Brauer et al., 2016). O₃ in the troposphere is an important greenhouse gas that has the third-highest radiative forcing after CO₂ and CH₄ (Stevenson et al., 2013; IPCC, 2013). In addition, O₃ is the primary source of the hydroxyl radical (OH), which has a major influence on the oxidizing capacity of the atmosphere and thus impacts the oxidation chemistry of secondary pollutants (e.g., sulfate and secondary organic aerosol) (Ehhalt et al., 2000; Rohrer et al., 2006). In recent years, the surface O₃ level has been increasing in most Chinese cities. For instance, in the highly urbanized areas of China, maximum 8h O₃ concentration increased by 19% (16.9 ppb) from 2015 to 2017 in the Beijing-Tianjin-Hebei (BTH) region, the growth trend was 14 % (11.3 ppb) for the Pearl River delta (PRD) region and 12 % (10.5 ppb) for the Yangtze River delta (YRD) region (http://106.37.208.233:20035/, last access: 29 May 2018). In the future, the pollution trend is likely to worsen due to potential changes of climate and emissions (Wang et al., 2013; Liu et al., 2013; Zhu et al., 2016, 2017).

On the global scale, VOC emissions from natural vegetation is estimated to be one order greater than that from anthropogenic activities, in spite of large uncertainties in different studies (Guenther et al., 2006; Wu et al., 2007, 2008; Jiang et al., 2013; Zhu et al., 2017). In addition, biogenic VOCs (e.g., isoprene) are highly reactive, reacting more efficiently with OH than most anthropogenic VOC species (Carlo et al., 2004). Previous studies have demonstrated the significant impacts of biogenic VOCs on surface O3 formation under strong solar radiation, high temperature, and NO_x level (e.g., Fiore et al., 2005, 2011; Wang et al., 2008; Curci et al., 2009; Geng et al., 2011; Strong et al., 2013; Squire et al., 2014; Lee et al., 2014; Zhang et al., 2017). In China, biogenic VOC emissions are estimated to be 17–44 Tg C yr⁻¹ (Guenther et al., 1995; Lin et al., 2008; Fu et al., 2012, 2104; Li et al., 2014) and are concentrated in the warm summer season. Current studies report that biogenic VOCs contribute to surface O₃ concentrations in China (Geng et al., 2011; Qu et al., 2013, 2014) and play an important role in intercontinental transport of O_3 (Zhu et al., 2017).

The impacts of biogenic VOCs on O₃ formation may vary in different regions and different seasons (Im et al., 2011; Strong et al., 2013; Wagner et al., 2014; Lee et al., 2014). Qu et al. (2013) employed the RAQM model to examine the influence of biogenic emissions on daily maximum surface O₃ concentration in China. Their calculations showed that in general the impact from biogenic sources on O₃ were more obvious in South China than in North China, but the O₃ increments in different regions did not follow the same seasonality. Geng et al. (2011) used the WRF-Chem model to evaluate the effect of biogenic emissions on O₃ production in Shanghai in summer, and suggested that the carbonyls produced by the continuous oxidation of isoprene have important impacts on O₃ level in the city. In addition, some studies suggest that biogenic emissions may increase due to global warming and

land-use change, and the impact on O₃ formation could be more significant in the future (Lin et al., 2008; Fiore et al., 2011; Liu et al., 2013; Fu et al., 2015). Fiore et al. (2011) pointed out that the potential increases in biogenic isoprene in North America (NA) could offset the regional and intercontinental surface O₃ decreases produced by controls on NA anthropogenic emissions during warm seasons.

The Guanzhong (GZ) Basin is the most developed region in northwestern China. In the past few years, air pollution has grown up to be a severe issue in the GZ Basin (Wang et al., 2012; Feng et al., 2016; Xue et al., 2017), due to its specific basin topography and abundant anthropogenic emissions (Li et al., 2017). According to the data from national environmental monitoring stations in the GZ Basin, 43 % of days in 2013–2017 have AQI > 100 (e.g., unhealthy air quality category), and in summer O_3 was regarded as the primary pollutant in 70% of polluted cases. In this study, we employed a regional chemical model WRF-Chem to simulate O₃ concentration in the GZ Basin for summer 2011. Our aim is the source apportionment of urban O₃ formation in this city surrounded by forests, specifically quantifying the individual and synergistic contributions of anthropogenic and biogenic sources. The paper is organized as follows. We first describe the sampling campaign, the chemical model and the emission data used for driving the model (Sect. 2). We then evaluate the model performance by comparing the observed urban air quality and biogenic VOCs with the simulated results (Sect. 3). Finally, we analyze the sensitivity of summertime O₃ formation to biogenic and anthropogenic sources (Sect. 4).

2 Methodology

2.1 Sampling sites and descriptions

The study was conducted in Xi'an, one of the oldest cities in the world. Xi'an is the most populous city in the northwest of China, with a population greater than 9 million. It is located in the heartland of the GZ Basin between the Qinling Mountains and the Loess Plateau (Fig. 1). The topographical features result in air pollutants often being trapped in the valley with limited dispersion. The city borders the northern foot of the Qinling Mountains, around a 50 km distance from the city center to the foothills. The Qinling Mountains are an east-west mountain range of 1600 km in length and 300 km in width and are regarded as a natural boundary between northern and southern China. Climate and culture are significantly different from the north to the south. Xi'an city has high temperatures and strong solar intensity in summer, making it an ideal location to assess the importance of biogenic contributions to urban air quality.

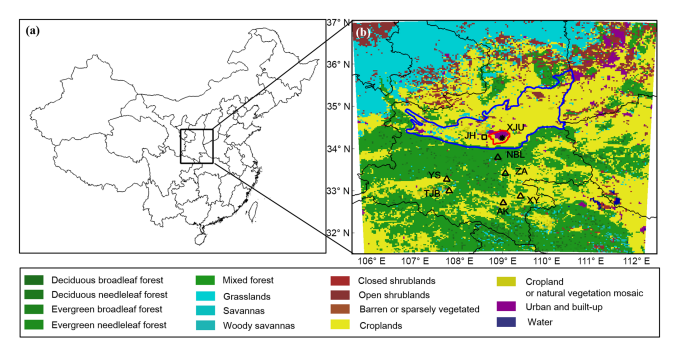


Figure 1. The simulation domain and the locations of six biogenic VOCs sites (triangles), one meteorological site (square) and one air quality site (snowflake). Colors indicate land types from MODIS. The area of red line indicates the urban area of Xi'an city. The area of blue line indicates the GZ Basin.

Table 1. Ambient biogenic VOCs observations in the Qinling Mountains during 6–7 August 2011.

Site ^a	Date	Start Time ^b	Location	Isoprene (ppb)		Monoterpenes (ppb)		Dominant monoterpenes ^c
				Observation	Simulation	Observation	Simulation	
NBL	6 Aug 2011	10:20	33.78° E, 108.88° N	3.8	4.5	0.42	0.29	α-pinene
ZA	6 Aug 2011	12:44	33.40° E, 109.05° N	0.1	0.9	0.16	0.22	α -pinene, limonene, menthone
XY	6 Aug 2011	16:14	32.87° E, 109.40° N	1.0	0.3	0.24	0.13	α -pinene
AK	7 Aug 2011	09:45	32.71° E, 109.01° N	0.8	0.4	0.10	0.08	α -pinene
TJB	7 Aug 2011	13:00	32.99° E, 107.78° N	0.5	1.1	0.27	0.26	α -pinene
YS	7 Aug 2011	14:50	33.27° E, 107.73° N	1.6	0.9	0.04	0.32	α-pinene
Average				1.3	1.4	0.21	0.22	

a Site names: NBL (Niubeiliang), ZA (Zhenan), XY (Xunyang), AK (Ankang), TJB (Tangjiaba), and YS (Youshui). The sampling duration is 30 min. Cher monoterpenes detected (0.1–9%) include tricyclene, α-thujene, camphene, sabinene, myrcene, α-phellandrene, Δ-carene, ο-cymene, β-ocimene, cineole, isopulegol, and isomenthone.

2.1.1 Biogenic VOC measurements in the Qinling Mountains

We selected six field sites in the Qinling Mountains (Fig. 1b, the triangles) and collected one ambient air sample at each site on 6–7 August 2011 under sunny weather conditions (details are presented in Table 1). Sampling was conducted between 9:30 to 16:30 local time (LT; each sampling lasted for 30 min) to target expected daily maximum isoprene concentrations. At each site, ambient air samples were pulled in

parallel onto three cartridges filled with Tenax GR and Carbograph 5TD solid adsorbents using a mass flow controlled pump for 30 min. Samples were shipped to the lab at NCAR (Boulder, CO, USA) for chemical analysis. Cartridges were desorbed using an UltraTM TD auto sampler with a Unity thermal desorption system (MARKES International Series 2, Llantrisant, UK) interfaced with a temperature programmed Agilent 7890A series Gas Chromatograph with a 5975C Electron Impact Mass Spectrometer and a Flame Ionization Detector (GC-MS/FID, Agilent Technologies, Santa Clara,

CA, USA). We used nitrogen as a carrier gas at the flow rate of 3 mL min⁻¹. Isoprene and monoterpene identifications were based on the comparison of retention time of authentic standards and mass spectra in the National Institute of Standards and Technologies (NIST) databases. Quantifications were calculated using FID calibrated with a NIST traceable standard.

2.1.2 Air quality monitoring in urban Xi'an

We set up an urban air quality monitoring site (Fig. 1b) at the roof (107 m above ground) of the main building (34.245° N, 108.984° E) on the campus of Xi'an Jiaotong University to minimize the ground level influences of local emissions. The campus is in the southeastern part of the downtown surrounded by residential areas. We obtained reliable observations of the concentrations of O₃, NO_x, and PM_{2.5} during 15-30 August 2011. Gases were measured by Ecotech analyzers (Ecotech Pty Ltd, Australia). O₃ were measured by an UV photometric analyzer EC9801. NO_x was measured by a gas-phase chemiluminescence detection analyzer EC9841, coupled with a hot molybdenum converter. We collected 24 h PM_{2.5} filter samples with a mini-volume sampler (Airmetrics, USA) at a flow rate of 5Lmin⁻¹ using both 47 mm quartz-fiber (Whatman, Middlesex, UK) and teflonmembrane (Gelman, Ann Arbor, MI) filters. We calculated the PM_{2.5} mass concentrations gravimetrically by weighing the teflon-membrane filters pre- and post-collection at least 4 times using an electronic microbalance (MC5, Sartorius, Göttingen, Germany) with $\pm 1 \,\mu g$ sensitivity under controlled conditions. EC and OC concentrations were analyzed based on a 0.5 cm² punch from the quartz-fiber filter following the IMPROVE A (Interagency Monitoring of Protected Visual Environments) thermal/optical reflectance (TOR) protocol (Chow et al., 2007) using a DRI model 2001 Carbon Analyzer. The concentrations of ions were quantified from a 10.8 cm² of the teflon-membrane filter by a Dionex DX-600 ion chromatography (Dionex Inc., Sunnyvale, CA, USA) (Zhang et al., 2014).

2.2 The WRF-Chem model

We employed the WRF-Chem model to study biogenic VOC emissions from the Qinling Mountains and their contributions to regional O₃ formation in urban Xi'an. WRF-Chem is a 3-D online-coupled meteorology and chemistry model consisting of the components of emission, transport, chemical transformation, photolysis and radiation (Tie et al., 2003; Li et al., 2011), dry and wet deposition (Wesely, 1989), and aerosol interactions (replaced with CMAQ aerosol module, Binkowski and Roselle, 2003; Li et al., 2010). WRF is a non-hydrostatic mesoscale dynamical system with various options for physical parameterizations (Skamarock et al., 2008). The chemical modules were implemented into the

WRF framework obeying the same schemes for the simultaneous simulations (Grell et al., 2005).

The simulated domain (Fig. 1) is $600 \, \mathrm{km} \times 600 \, \mathrm{km}$ centered on urban Xi'an with 3 km horizontal grid spacing. We set up 28 vertical layers from the surface up to $50 \, \mathrm{hPa}$ with 7 layers below 1 km to assure a high near-ground vertical resolution. The National Centers for Environmental Prediction (NCEP) FNL Operational Global Analysis data provided the initial and boundary fields of meteorology. Initial and boundary conditions of chemistry were derived by a global chemical transport model (Model for Ozone and Related chemical Tracers, MOZART) (Emmons et al., 2010). We considered the first 7 days as spin-up period, and the study focused on 6–7 and 15–30 August 2011 because of the available field observation datasets (as described in Sect. 2.1).

We adopted RADM2 (Regional Acid Deposition Model) as the gas phase chemical mechanism to predict O₃ formation. RADM2 is an aggregated species type using the reactivity based weighting scheme to adjust for lumping (Stockwell et al., 1990). The mechanism implemented in our WRF-Chem model covers 158 reactions among 36 species, containing the complete reaction paths for isoprene, monoterpenes and the relevant inorganic reactions. As an explicit species, isoprene chemistry is based on an updated CB4 gasphase mechanism (Carter and Atkinson., 1996). We noted some advances in isoprene chemistry recently. Some studies pointed out that isoprene reacts with OH radical to form hydroperoxy radicals (ISOPO2). Subsequently, in the presence of NO_x, ISOPO2 reacts with NO leading to the production of hydroxynitrates (ISOPN) by a minor branch, which sequesters NO_x and thus regulates O₃ formation. A number of laboratory filed observation and simulation studies (e.g., Paulot et al., 2009a, b, 2012; Horowitz et al., 2007; Hudman et al., 2009; Fisher et al., 2016; Travis et al., 2016) highlighted the importance of isoprene nitrate chemistry and all agreed there were still large uncertainties (for example, the estimates of ISOPN yield (4–15%)). Horowitz et al. (2007) found a 4 % ISOPN yield, best captured the alkyl and multifunctional nitrates measured by aircraft, and Hudman et al. (2009) pointed out that unreasonably high ISOPN yield (18%) would let ISOPN be a terminal sink for NO_x . In addition, the production of radicals from ISOPO2 is regarded to be important for HO_x and further for secondary pollutants (e.g., O₃ and secondary organic aerosol (SOA)) (e.g., Squire et al., 2015). Kanaya et al. (2012) quantified the formation of hydroperoxy-aldehydes (HPALDs) from ISOPO2 and subsequent rapid release of HO_x . They revised the model by adding detailed reactions of isomerization of ISOPO2 and photolysis of HPALDs, following Peeters and Müller (2010), and found the revision could increase OH and HO2 concentrations by 28-38 % for daytime. However, Kanaya et al. (2012) also pointed out that isomerization of ISOPO2 at the rates proposed by Peeters and Müller (2010) might be overestimated.

Back to our study, these advances in isoprene chemistry were not contained in the standard RADM2 mechanism. To assess the potential uncertainties, we modified the RADM2 mechanism by adding formation pathway of ISOPN from ISOPO2 + NO (the yield of 4%) following Horowitz et al. (2007) and ISOPO2 isomerization reaction following Li et al. (2018). Details of these updates can be found in Table S1 in the Supplement. We conducted four sensitivity simulations (15-17 August), namely ISO0 (standard RAMD2 mechanism), ISO1 (including the ISOPO2 isomerization reaction), ISO2 (including reactions of isoprene nitrate), and ISO3 (including both revisions in ISO1 and ISO2), to explore the effects on O₃ in the GZ Basin. The results show that near surface O₃ concentration was decreased by 2.9 ppb (7%) averaged for urban Xi'an in August after implementing these updates (ISO3-ISO0), and we thought this would not crucially impact conclusions of this study. Details of the sensitivity analysis results can be found in Table S2 and Fig. S1 in the Supplement.

2.3 Biogenic and anthropogenic emissions

Biogenic emissions were quantified by the widely used model MEGAN (Model of Emissions of Gas and Aerosols from Nature) (Guenther et al., 2006). MEGAN coupled into the WRF-Chem model, referred to here as WRF-MEGAN2, provides on-line estimates of the net landscape-averaged biogenic emissions from terrestrial ecosystems into the abovecanopy atmosphere. The on-line estimated emissions of isoprene, individual monoterpenes and other biogenic VOCs serve as the inputs for the further chemistry simulation. To drive MEGAN, we need the following inputs: emission factors (EFs), leaf area index (LAI), plant functional types (PFTs), as well as meteorology conditions. The meteorology was obtained from WRF simulations and the LAI and PFT data were extracted from MODIS (Tian et al., 2004). We adopted the canopy-scale emission factors of dominant species from Guenther et al. (2006).

Estimated across the year 2011 by WRF-MEGAN2, the isoprene and monoterpene emissions in the Qinling Mountains were mostly concentrated in summer (71 and 58%, respectively). During our simulation period, the isoprene emission from the domain is 157 Gg month $^{-1}$, accounting for $\sim 80\%$ of total biogenic VOC emissions (Table S3 in the Supplement). The rest are monoterpenes and other biogenic VOCs (e.g., acetone and MBO (2-methyl-3-buten-2-ol)). Figure 2a and b show the spatial distributions of biogenic isoprene and monoterpenes emission fluxes during the simulation period, indicating the high emission zone of isoprene in the Qinling Mountains lying to the south of Xi'an city.

The anthropogenic emissions were obtained from the Multi-resolution Emission Inventory for China (MEIC, Li et al., 2017) for the year of 2010, which was downscaled to a resolution of 3 km using locations of point sources

and various spatial proxies (Geng et al., 2017). The upgraded highly resolved emission data were based on a collection of statistics and newly developed emission factors. The emission inventory used in our model includes all major anthropogenic sources, but excluded open biomass burning which occupies a low proportion in the GZ Basin during our simulating period (estimated by Fire Inventory from NCAR, https://www2.acom.ucar.edu/modeling/ finn-fire-inventory-near, last access: 29 May 2018). The anthropogenic emission sources are composed of power, industry, residential, transportation, and agriculture. The emission estimates and uncertainties of VOCs, SO₂, NO_x, NH₃, and PM_{2.5} in the domain during the simulation period are summarized in Table S3, and the potential impacts of emission uncertainty on simulation will be discussed in Sect. 3.3. The estimated anthropogenic VOCs emissions are 72.2 Gg, contributing up to $\sim 30\%$ of total VOC emissions. Figure 2c and d present the spatial distributions of anthropogenic VOC and NO_x emissions in the simulation period. The highest emission intensity of anthropogenic VOCs and NO_x are in Xi'an city and the GZ Basin due to the frequent vehicle and industrial activities of this area.

2.4 Factor separation technique

 O_3 is formed by complicated nonlinear reactions of anthropogenic and biogenic precursors (NO $_x$ and VOCs) in the presence of sunlight. The approach referred to as the "brute-force" method (sensitivity analysis used to measure the model output response to emission changes) is traditionally used in air quality models to identify source contributions from specific non-reactive species in a linear process, but it cannot straightforwardly apply to secondary species due to the non-linearity in responses. In practice, the actual impact of one factor in a nonlinear process in the presence of others can be separated into (1) pure impact from the factor and (2) interactional impacts from all those factors. In this study, we adopted the factor separation approach (FSA) (Stein and Alpert, 1993) to decompose the pure contribution of a factor from its interaction with other factors.

We considered anthropogenic and biogenic sources as two interactional factors to influence the O_3 formation. $f_{\text{anth-bio}}$, f_{anth} , f_{bio} , and f_0 are the simulation results including both anthropogenic and biogenic sources, anthropogenic source only, biogenic source only, and neither, respectively. Pure contributions of anthropogenic and biogenic sources are expressed as Eqs. (1) and (2), respectively:

$$f_{\text{anth}}' = f_{\text{anth}} - f_0, \tag{1}$$

$$f'_{\text{bio}} = f_{\text{bio}} - f_0. \tag{2}$$

The calculated result including both anthropogenic and biogenic sources should include both pure contributions of the two factors, the synergistic impact, and the impact of back-

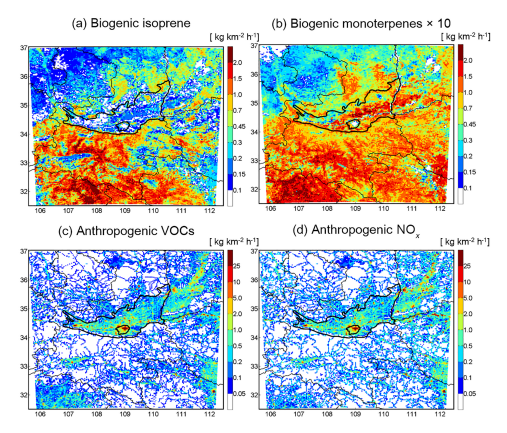


Figure 2. Monthly mean emissions of (a) biogenic isoprene, (b) biogenic monoterpenes, (c) anthropogenic VOCs, and (d) anthropogenic NO_{χ} in the GZ Basin and surrounding areas in August 2011.

ground transport (Eq. 3):

$$f_{\text{anth-bio}} = f'_{\text{anth}} + f'_{\text{bio}} + f'_{\text{anth-bio}} + f_0. \tag{3}$$

Thus, the synergistic effect between anthropogenic and biogenic sources is represented as follows:

$$f'_{\text{anth-bio}} = f_{\text{anth-bio}} - f'_{\text{anth}} - f'_{\text{bio}} - f_{0}$$

$$= f_{\text{anth-bio}} - (f_{\text{anth}} - f_{0}) - (f_{\text{bio}} - f_{0}) - f_{0}$$

$$= f_{\text{anth-bio}} - f_{\text{anth}} - f_{\text{bio}} + f_{0}. \tag{4}$$

Based on the FSA, we conducted four simulations, namely BASE, ANTH, BIO, and NEITHER, to explore the pure and synergistic impacts of anthropogenic and/or biogenic sources on O₃ production in the GZ Basin. Detailed simulation settings and the various contribution definitions are summarized in Table 2.

3 Observation data and model validation

3.1 Meteorology

The specific topographical features of Xi'an make the meteorological conditions crucial for the accumulation and dispersion of urban pollutants. To validate our model performance

in wind, temperature, and relative humidity, we compared the hourly meteorological data (http://www.meteomanz.com, last access: 29 May 2018) observed at the Jinghe site (34.26° N, 108.58° E west of Xi'an) with model simulations. Figure 3c illustrates the observed and simulated near-surface wind speed and directions during 15–30 August 2011. The WRF-Chem model successfully captured the prevailing wind direction from north and northeast, consistent with the in situ observations. It should be noted that in our simulation period the prevailing wind blew from south, which enhanced the transport of biogenic emissions from the Qinling Mountains to urban Xi'an. In addition, a continuous rainfall event during 18–22 August (green shadow in Fig. 3) was characterized by lower temperature and near-saturated humidity.

We conducted the statistical verification of meteorological variables in Table 3, including the r (correlation coefficient), NMB (normalized mean bias) and RMSE (root mean square errors). Modeled meteorological variables were in good agreement with observations (Fig. 3a–c) with the NMB less than $\pm 6\%$.

Table 2. Summary of different simulation settings and definitions of the various contributions from anthropogenic and/or biogenic sources.

Simulation	Simulation results	Anthropogenic emission	Biogenic emission			
BASE	$f_{ m anth-bio}$	✓	√			
ANTH	f_{anth}	\checkmark	X			
BIO	$f_{ m bio}$	X	\checkmark			
NEITHER	f_0	X	X			
Contribution						
$f_{\text{anth-bio}} - f_{\text{bio}}$		Actual contribution of anthropogenic e	emissions			
$f_{\text{anth-bio}} - f_{\text{anth}}$		Actual contribution of biogenic emissions				
		The contribution of background transport				
$f_0' = f_0$ $f_{\text{anth}}' = f_{\text{anth}} - f_0$		Pure contribution of anthropogenic emissions				
$f'_{\text{bio}}^{\text{intil}} = f_{\text{bio}} - f_0$ $f'_{\text{anth-bio}} = f_{\text{anth-bio}} - (f_{\text{anth}} + f_{\text{bio}}) + f_0$		Pure contribution of biogenic emissions				
$f'_{\text{anth-bio}} = f_{\text{anth-bio}} -$	$(f_{\text{anth}} + f_{\text{bio}}) + f_0$	Synergistic contribution of anthropogenic and biogenic emissions				

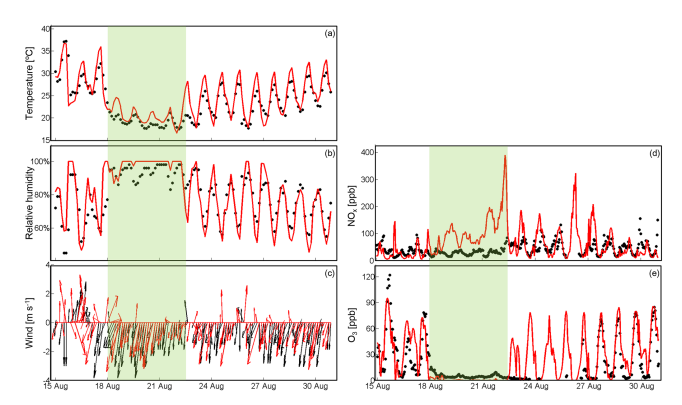


Figure 3. Observed (black) and simulated (red) temporal patterns of temperature (a), relative humidity (b), and wind (c) at the Jinghe site and NO_x (d) and O₃ (e) concentrations at Xi'an Jiaotong University, during the period from 15 to 30 August 2011. The green shadow (18-22 August) indicates rainy days.

3.2 **Biogenic VOC concentrations in the Qinling Mountains**

Samples from the Qinling Mountains show that the dominant VOC species was isoprene, and α -pinene was the main constituent of monoterpenes (Table 1). The ratio of isoprene to monoterpenes varies considerably. In general, different terpene emitters are not homogeneously distributed in a kilometer-scale grid and the point measurements are influenced by the microenvironment and meteorology (Zare et al., 2012; Kota et al., 2015). However, in this study, our goal is to estimate the biogenic effects on urban O₃ 50 km away from the foothills, which requests more concern on the regional scale VOC level, rather than the microenvironmentscale variability in either the observation or the simulation. Thus, we compared the average of VOC measurements with model simulations to validate whether the calculated results were reasonable. The isoprene mean concentration simulated in the six grids (corresponding to the time of observations)

	Me	r d	NMB ^d	RMSE d	
	Observation	Simulation			
Meteorology ^b					
Wind speed (m s ⁻¹)	2.6	2.5	_	-6%	1.8
Temperature (°C)	25.1	24.2	0.86	4 %	2.5
Relative humidity	73.6 %	74.2%	0.72	1 %	12 %
Air quality ^c					
NO_{χ} (ppb)	47.0	46.6	0.36	-1%	18.1
O ₃ (ppb)	31.5	38.7	0.72	21 %	8.1
$PM_{2.5} (\mu g m^{-3})$	107	94.6	_	-12%	49.3

Table 3. Statistics of meteorological and air quality variables over the GZ Basin in August 2011^a.

was 1.4 ppb, which is close to the observed average value of 1.3 ppb at the six sampling sites. Monoterpenes performed quite similarly, simulated 0.22 ppb comparing with observed 0.21 ppb. We also analyzed the temporal variation of simulated biogenic VOC during the whole simulation period and found the sub-month variability was relatively small (the SD < 25 %). The evaluation indicates that biogenic VOCs simulations reasonably agreed with the observations in the Qinling Mountains, on average, which provides a basis for us to further evaluate biogenic effects on O₃.

3.3 Gaseous and particulate pollutants in urban Xi'an

The sampling campaign was organized in summer. Based on the gaseous and particulate pollutant observations, the daily mean PM_{2.5} concentration was $90.0 \pm 53.5 \,\mu g \, m^{-3}$, with 57% of days exceeding the WHO Interim target-1 (IT-1) 75 $\,\mu g \, m^{-3}$. The daily mean NO_x concentrations were 25.8–63.2 ppb, with 40% of days exceeding the guideline 48.7 ppb ($\approx 100 \,\mu g \, m^{-3}$, GB 3095–2012). The maximum 8 h O₃ concentration was 3.5–95.6 ppb, with most of the values around the national first grade standard of 46.6 ppb ($\approx 100 \,\mu g \, m^{-3}$, GB 3095-2012). Summer in Xi'an is monitored as the least polluted season of the year, and the case we picked is regarded as a typical situation in summer Xi'an.

Figure 3 compares the simulated hourly O_3 and NO_x concentrations with in situ observations. During the rainy episode (the green shadow in Fig. 3), our model overestimated NO_x concentration and underestimated O_3 concentration. The deviation can be explained by the failure to simulate the precipitation in the WRF model resulting in underestimates in wet deposition. So, we focused our analysis on the period excluding the rainy period. During the non-raining days, our model well reproduced the diurnal variations and magnitudes of O_3 and NO_x concentrations. The calculated

 O_3 averaged for non-raining periods was 38.7 ppb, $\sim 20 \%$ higher than the observed value of 31.5 ppb. Our simulated O₃ also reproduced the temporal variation of the observed O_3 (r = 0.72). For NO_x , the simulated hourly NO_x averaged for the non-raining period was 46.6 ppb, close to the observed 47.0 ppb (NMB = -1%), which suggests no systematic bias in NO_x emissions. It is worth noting that the observed NO_x were detected by a chemiluminescence analyzer coupled with a hot molybdenum converter (MoO), and this method was recognized to cause higher NO2 concentration, due to the positive interference of other nitrogencontaining components (NO_z, e.g., PAN, HNO₃ and HONO). Xu et al. (2013) found that the uncertainty caused by the MoO converter was much smaller at urban sites (less than 10%) than that at suburban and background sites (more than 30 %). In the GZ Basin, to evaluate the uncertainty, we estimated the ratio of $NO_7/(NO_x + NO_7)$ from the model. The calculated results indicated that the NO_z accounted for 11 % of the total $NO_x + NO_z$ in urban Xi'an during the nonraining period. We noted the uncertainty in our NO_x measurements, but considered this would not crucially impact the model-measurement comparison.

Unfortunately, the anthropogenic VOC was not included in our samples as the observations were primarily targeted at biogenic VOC in the Qinling forest. Alternatively, three sensitivity simulations were conducted (for 15–17 August), namely VOC0 (using standard MEIC emission estimates), VOC1 (with an increase of anthropogenic VOC emission by 50%) and VOC2 (with a decrease of anthropogenic VOC emission by 33%) to explore the sensitivity of simulated VOC and O₃ concentrations to anthropogenic VOC emissions. We found that 50% increases of anthropogenic VOC emission could lead to a 22% increase of urban VOC concentration, while the 33% emission decrease resulted in a 24% decrease of concentration (Table S4 in the Supple-

^a Averaged for the period from 15 to 30 August 2011, excluding the rainy days. ^b Meteorological data were obtained from the hourly surface measurements at Jinghe station (34.26° N, 108.58° E). ^c Air quality data were measured at the roof (107 m above ground) of the main building (34.25° N, 108.98° E) on the campus of Xi'an Jiaotong University. ^d r: correlation coefficient; NMB: normalized mean bias; RMSE: root mean square errors.

ment). It is worth noting that the concentration of O_3 stayed almost the same (because the O_3 production regime is NO_x -limit). We addressed that the uncertainties of VOC emission obviously affected the VOC concentrations; however, MEIC inventory is the most updated available emission for China so far, and quantifying its uncertainties can be done in future studies (possibly with satellite-based measurement of HCHO, Miller et al., 2008).

We analyzed PM_{2.5} concentration and composition (sulfate, nitrate, ammonium, EC, organic matter) with the filter-base measurements. The model predicted PM_{2.5} concentration to be $94.6 \pm 28.2 \,\mu \text{g m}^{-3}$, slightly lower (NMB = -12 %) than measured $107 \mu g m^{-3}$ averaged for the non-raining period, but did not perform well in capturing temporal correlation (r = 0.17). The simulated PM_{2.5} showed the similar compositions to the observation (Fig. S2b and c). Sulfate is the dominant constituent of both simulated (32%) and observed (37%) PM_{2.5}. High sulfate concentration was mainly attributed to the high SO₂ emission in the GZ Basin as well as the humid weather conditions (Wang et al., 2014). The secondary constituent of observed PM_{2.5} is organic matter, which accounted for 16% of the total observed PM_{2.5}, close to the simulated result (14%). Secondary organic matter contributed half to total simulated organic matter, mainly due to the abundant precursor (i.e., VOCs) emissions and the high atmospheric oxidation capacity in summer.

4 Impacts of biogenic and anthropogenic sources on O₃ formation

In this section, we analyze the results from the four simulations (BASE, ANTH, BIO and NEITHER; see Table 2) to characterize the fate of O_3 and its precursors in the GZ Basin and to quantify the pure and synergistic impacts of anthropogenic and/or biogenic sources on summertime O_3 formation.

4.1 Base simulation of O₃

Firstly, we discuss the spatial and temporal characteristics of the simulated O_3 and the precursors (VOCs and NO_x) in the GZ Basin in the BASE simulation.

Figure 4a shows spatial distribution of the simulated VOCs during the non-raining period, overlaid with the simulated wind vectors. The highest concentration (more than 50 ppb, with ethane being the dominant species) was in urban Xi'an and its downwind region (the southwest of urban Xi'an), due to anthropogenic activities. In addition, another high-value area (~30 ppb, with isoprene being the dominant species) was found in the Qinling Mountains, which was probably due to biogenic sources. To better understand the composition of VOCs, we analyzed some typical individual VOC species. Figure S3a in the Supplement shows the spatial

distribution of xylenes, representing anthropogenic VOCs, and Fig. S4a and S5a in the Supplement show isoprene and monoterpenes, representing biogenic VOCs. The anthropogenic xylenes were mainly distributed in the GZ Basin, while the high biogenic isoprene and monoterpene concentrations were found over the Qinling Mountains. These results explain the spatial feature of total VOCs and the dominant sources. Detailed discussion of source apportionment is given in Sect. 4.2

The spatial distribution of NO_x was slightly different (Fig. 5a). The highest concentrations of NO_x were in the GZ Basin (average of 11.1 ppb), especially in urban Xi'an (averaged of 30.1 ppb), while among the Qinling Mountains, NO_x was low and dominated by biogenic sources.

PM, even though not directly involved in the formation pathways of O₃, influences the chemical equilibrium indirectly. In the daytime, NO_2 photolysis frequency $(J(NO_2))$ is determined by the solar radiation influenced by PM via scattering and absorption. Figure 6 shows the changes of J (NO₂) (calculated by model track output photor_no2) with the participation of PM (concentration, 102 µg m⁻³; aerosol optical depth (AOD) at 550 nm, 1.92; single scattering albedo (SSA) at 550 nm, 0.92) averaged for urban Xi'an. J (NO₂) was reduced by 40-60 %, most significantly in morning and evening rush hours. In the night time, PM_{2.5} can remove N_2O_5 from the NO_x cycle via heterogeneous reactions, as one of the major NO_x sinks in the atmosphere (Xue et al., 2014). Figure S6a in the Supplement shows the spatial feature of PM_{2.5}. The densest area was urban Xi'an (averaged for $102 \,\mu\mathrm{g}\,\mathrm{m}^{-3}$) followed by the western part of the GZ Basin. The spatial distribution of high-values of PM_{2.5} was similar to that of NOx, but covered a wider area mostly in the downwind region of urban Xi'an, which is expected due to longer lifetime of aerosols compared with NO_x and the time required for secondary aerosol formation, thus further dispersion.

The typical diurnal variation of O₃ (Fig. 7b) demonstrates there are higher concentrations in the afternoon and lower at night. For better understanding of O₃ concentration characteristics and source or sink mechanisms, we discussed two different time scales: (1) O₃ peak time (14:00–18:00 LT) (Fig. 8) and (2) O₃ 24 h average (Fig. 9). During the peak time, simulated near-surface O₃ was high in the GZ Basin, with averaged concentration of 75 ppb. In the downwind region of high NO_x and VOCs in the west of urban Xi'an, the concentration reached up to 110 ppb. We employed the ratio of H₂O₂/HNO₃ to investigate the chemistry regime of O₃ formation (Sillman, 1995; Wang et al., 2017). If the ratio is greater than 0.5, the O₃ production regime is considered NO_x-controlled, otherwise VOC-controlled if the ratio less than 0.3. The range between 0.3 and 0.5 is defined as the transition regime from NO_x- to VOC-controlled, indicating the competition of both NO_x and VOCs in O_3 production. Figure 10a shows the spatial distribution of the simulated H₂O₂/HNO₃ ratio during the O₃ peak time. The west

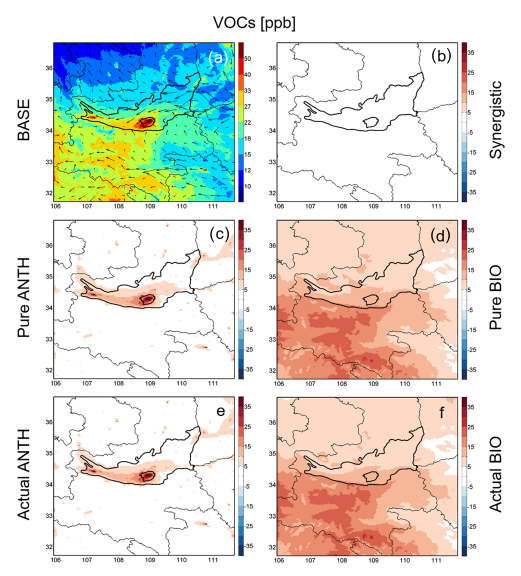


Figure 4. Spatial distributions of monthly mean concentrations of VOCs in August 2011. Panel (a) is the result from the BASE simulation, overlaid with simulated wind vectors. Panels (b–f) are simulated VOCs concentrations contributed from synergistic anthropogenic and biogenic, pure anthropogenic, pure biogenic, actual anthropogenic, and actual biogenic sources, respectively.

and southeast of the GZ Basin were right in the transition regime with a complicated O_3 production mechanism sensitive to both NO_x and VOCs. Most of the rest of the simulation region was VOC-controlled, excluding the Yuncheng and Hejing cities in the neighboring Shanxi provinces.

On the 24 h average scale, the spatial distribution of O_3 presented a different picture (Fig. 9). The original high-value area during the peak time in the GZ city cluster shifted to low-value region due to the consumption of O_3 by abundant NO_x emissions. At night time, the titration effect of freshly emitted NO dominates, and the O_3 concentration tends to drop to a lower level. The high value of 24 h averaged O_3 converged in the south and northwest outside of the GZ Basin. Those areas have elevated O_3 due to high daytime pro-

duction, similar to the nearby zone of peak O_3 , but also have lower emissions of NO resulting in lower loss of O_3 .

4.2 Pure impact of biogenic or anthropogenic sources

Using the FSA method, we evaluated the pure contribution of anthropogenic or biogenic sources to the summertime O_3 formation in the GZ Basin. In the scenario of pure contribution of anthropogenic emissions, the VOC concentrations were mostly distributed over the GZ city cluster (8.0 ppb), especially in urban Xi'an (26.4 ppb) (Fig. 4c). In the scenario of pure contribution of biogenic emissions, the VOCs were widely dispersed over the Qinling Mountains (Fig. 4d), with a calculated 9.9 ppb for the GZ Basin and 12.4 ppb for urban Xi'an. NO_x concentration has the similar pattern as VOCs

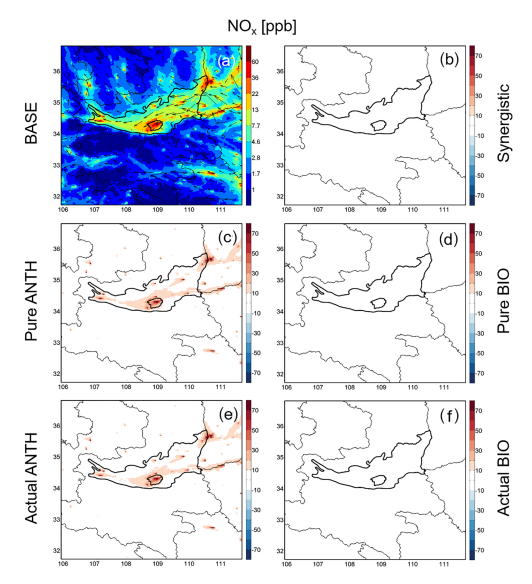


Figure 5. Spatial distributions of monthly mean concentrations of NO_x in August 2011. Panel (a) is the result from the BASE simulation, overlaid with simulated wind vectors. Panels (**b-f**) are simulated NO_x concentrations contributed from synergistic anthropogenic and biogenic, pure anthropogenic, pure biogenic, actual anthropogenic and actual biogenic sources, respectively.

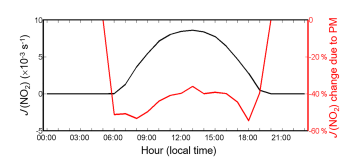


Figure 6. Diurnal variations of J (NO₂) (black) and the changes in J (NO₂) (red) averaged in urban Xi'an due to PM effects in August 2011.

in the scenario of pure contribution of anthropogenic emissions, with averaged concentrations of 11.0 ppb for the GZ Basin and 30.3 ppb for urban Xi'an (Fig. 5c). However, in the GZ Basin and urban Xi'an, biogenic sources contributed less than 0.2 ppb to NO_x concentration (Fig. 5d). In the scenario of pure contribution of anthropogenic emissions, $PM_{2.5}$ spread over a wider area (Fig. S6c), due to the time required for secondary aerosol formation. In the scenario of pure contribution of biogenic emissions, $PM_{2.5}$ was mostly distributed among the Qinling Mountains (Fig. S6d), but the concentration was lower by one order of magnitude (Table 4, Fig. S7 in the Supplement).

In the scenario of pure contribution of anthropogenic emissions, daily peak O_3 accumulated in the downwind region in

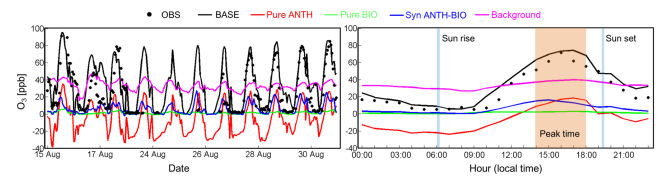


Figure 7. Temporal patterns of the simulated O_3 concentrations and the tested contributing components during the period from 15 to 30 August 2011, excluding the rainy days (18–22 August). The orange shadow (14:00–18:00 LT) indicates daily O_3 peak time.

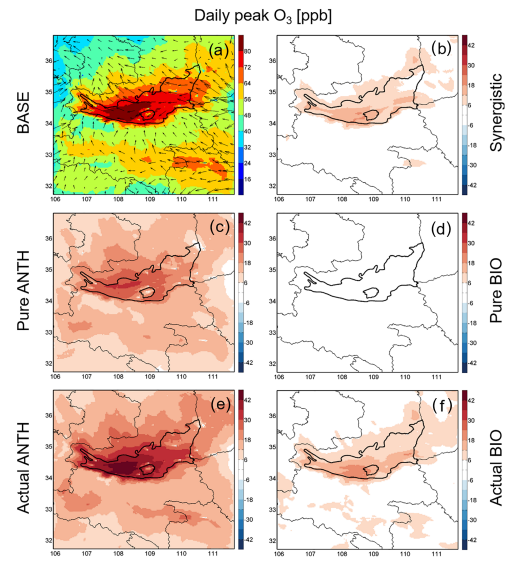


Figure 8. Spatial distributions of monthly mean concentrations of daily peak O_3 in August 2011. Panel (a) is the result from the BASE simulation, overlaid with simulated wind vectors. Panels (**b**-**f**) are simulated daily peak O_3 concentrations contributed from synergistic anthropogenic and biogenic, pure anthropogenic, pure biogenic, actual anthropogenic, and actual biogenic sources, respectively.

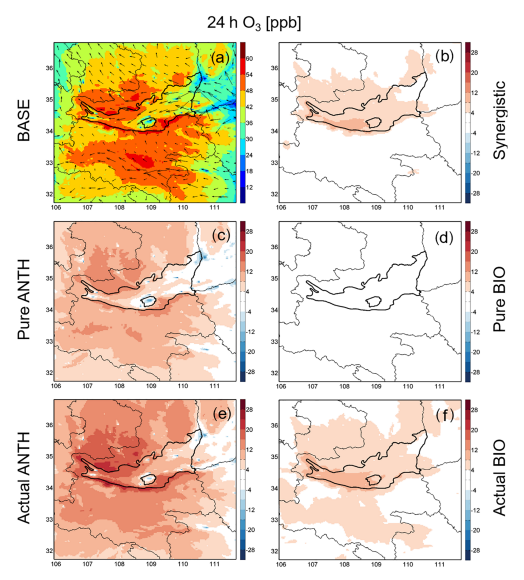


Figure 9. Spatial distributions of monthly mean concentrations of 24 h averaged O₃ in August 2011. Panel (a) is the result from the BASE simulation, overlaid with simulated wind vectors. Panels (b–f) are simulated 24 h averaged O₃ concentrations contributed from synergistic anthropogenic and biogenic, pure anthropogenic, pure biogenic, actual anthropogenic, and actual biogenic sources, respectively.

the central and western GZ due to high VOCs and NO_x concentrations. Daily peak O_3 concentrations reached 22.6 ppb for GZ and 19.1 ppb for urban Xi'an (Fig. 8c). Lower daily peak O_3 concentration was found outside of GZ Basin, where less anthropogenic VOCs and NO_x were emitted. In contrast, daily peak O_3 was negligible (less than 3 ppb) in the scenario of pure contribution of biogenic emissions (Fig. 8d) due to the low NO_x emissions. However, the distribution of 24 h averaged O_3 was different from daily peak O_3 . 24 h averaged O_3 concentration in the scenario of pure contribution of anthropogenic emissions was more diluted in the GZ city cluster than for surrounding areas (Fig. 9c). Due to the abundant NO emission and its titration effect on O_3 , the pure ef-

fect of anthropogenic sources was negative, calculated to be -2.2 ppb in urban Xi'an.

4.3 Synergistic impact of the interaction between biogenic and anthropogenic sources

The synergistic impact on O_3 formation includes the interactions between anthropogenic and biogenic sources. In other words, it reflects the potential production trend of either " O_3 -promoted" or " O_3 -suppressed" under the natural coexistence of all emission sources. In the cases of NO_x , VOCs and $PM_{2.5}$, the synergistic impacts contributed less than ± 3 % of total concentrations (Figs. 4b, 5b, and S5b and S6 in the Supplement, as well as Table 4). However, the synergistic impact on O_3 played a remarkable role showing positive impacts for

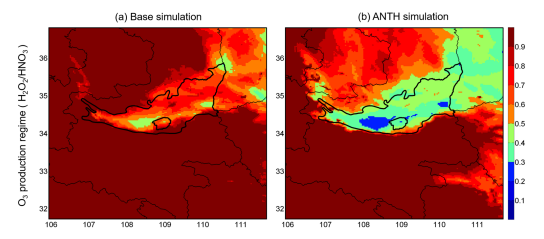


Figure 10. The monthly mean ratio of H_2O_2/HNO_3 during the daily O_3 peak time (14:00–18:00 LT) in August 2011 in the (a) base simulation and (b) the simulation without biogenic sources.

Table 4. The various contribution components of the simulated O_3 (and the precursors) and $PM_{2.5}$ in August 2011.

	NO_X	VOCs	O ₃ [ppb	PM _{2.5}	
	[ppb]	[ppb]	daily peak*	24 h	$[\mu g m^{-3}]$
The GZ Basin					
Base	11.1	24.5	74.1	44.4	65.1
Pure ANTH	11.0	8.0	22.6	5.0	52.0
Pure BIO	0.1	9.9	2.0	1.1	3.3
Actual ANTH	10.6	7.7	33.0	10.8	53.5
Actual BIO	-0.3	9.6	12.5	7.0	4.9
Syn ANTH-BIO	-0.4	-0.3	10.5	5.8	1.5
Urban Xi'an					
Base	30.1	44.8	74.7	38.7	102
Pure ANTH	30.3	26.4	19.1	-2.2	88.7
Pure BIO	0.15	12.4	2.6	1.4	3.4
Actual ANTH	29.6	26.2	33.4	4.6	91.1
Actual BIO	-0.6	12.2	16.8	8.2	5.8
Syn ANTH-BIO	-0.7	-0.2	14.3	6.8	2.4

^{*} Daily O_3 peak time is from 14:00-18:00 LT.

both daily peak (Fig. 8b) and 24 h averaged O_3 (Fig. 9b). It means that the mixed state of anthropogenic and biogenic sources potentially enhanced the O_3 production more than each single source. To make it more specific, we started the discussion from the result of ANTH simulation without the biogenic sources (Table 2). Figure 10b shows the O_3 production regime in the ANTH simulation. VOC-controlled O_3 production regime covered the west of urban Xi'an and the southeast of the GZ Basin. In the rest of the GZ Basin and the neighboring Shanxi provinces, the O_3 production was in the transition regime, controlled by both NO_x and VOC. NO_x -controlled O_3 production regime dominated the rest of the region. After we included biogenic VOC emissions in the simulation, the O_3 concentration was significantly enhanced

in the VOC-controlled regions, and partly enhanced in the mix-controlled region. However, in the VOC-controlled region, the synergistic impact contributed little.

The synergistic impact is of great importance, approximately the same magnitude as the impact from pure contributions of anthropogenic sources. The synergistic impact contributed daily peak O₃ concentrations of 10.5 ppb for the GZ Basin and 14.3 ppb for urban Xi'an, while the pure anthropogenic impact contributed 22.6 ppb for GZ Basin and 19.1 ppb for urban Xi'an. However, the extent was $\sim 50 \,\%$ smaller on the 24 h averaged scale, but still increased O₃ concentration by 5.8 ppb for the GZ Basin and 6.8 ppb for urban Xi'an. Figure 7 shows the diurnal variation of the observed and simulated O₃ concentration at Xi'an Jiaotong University, as well as the tested contributing components. Transport dominated O₃, constantly contributing 30–40 ppb as background. The impact of pure anthropogenic sources was positive on O₃ production during 13:00–19:00 LT but negative during the rest of the time, and the impact of pure biogenic sources was negligible. Synergistic impact of both anthropogenic and biogenic sources resulted in a positive contribution during 10:00-21:00 LT, comparable to the impact of pure anthropogenic sources.

It is worth noting that the biogenic contribution to $PM_{2.5}$ is not obvious (less than 3 %) in GZ Basin, which might be different from some other regions (e.g., Fu et al., 2012; Li et al., 2013). The main reasons are that (1) organic matter, the most important biogenic $PM_{2.5}$ component, only accounted for 14–16 % of $PM_{2.5}$ in GZ Basin in August, (2) undeniably, uncertainties still exist in organic matter simulations in the model.

5 Conclusions

The GZ Basin is a representative region in the northwest of China, suffering serious air pollution in recent years. Geo-

graphically, the GZ Basin borders the northern foot of the Qinling Mountains. For this reason, in addition to the anthropogenic emissions from metropolitan areas, biogenic emissions are of great importance in the region, especially in warm season with active photochemistry. In this study, we used the WRF-Chem model to simulate O₃ in the GZ Basin and compared the results to near-surface measurements, with the aim of quantifying the pure and synergistic impacts of anthropogenic and/or biogenic sources on summertime O₃ formation. The simulation was driven by the best currently available inventory of anthropogenic emissions and online calculated biogenic emissions. Near-surface measurements were captured from six surface sites among the Qinling Mountains for biogenic VOCs and one 100 m-high site in Xi'an city for air quality (NO_x, VOCs, O₃ and PM_{2.5}).

Our model successfully reproduced the observed air quality and meteorological parameters. The biogenic VOCs simulation showed a reasonable agreement. Our model also well-reproduced the magnitudes and variations of O_3 , NO_x , and $PM_{2.5}$ concentrations excluding rainy days, with normalized mean bias less than $\pm 21\,\%$.

We further conducted three scenario simulations to explore the pure and synergistic impacts of anthropogenic and/or biogenic sources on O₃ and the precursors, by using the factor separation approach (FSA). The results concluded that, for the precursors, pure impact of anthropogenic source contributed 99 % of NO_x, 80 % of PM_{2.5}, and 33 % of VOCs in the GZ Basin, and pure impact of biogenic source contributed 40 % of VOCs but only 1–5 % of PM_{2.5} and NO_x. Meanwhile, synergistic impacts from the combination of anthropogenic and biogenic sources did not bring significant changes on NO_x , VOCs and $PM_{2.5}$ (less than $\pm 4\%$). In the case of daily peak O₃, the pure impact of anthropogenic source remained the dominant contributor (19.1 ppb for urban Xi'an), even after anthropogenic particles reduced the NO₂ photolysis by up to 60 %. The abundant biogenic VOCs from the nearby forests promoted the O₃ formation by interaction with anthropogenic NO_x , contributing 14.4 ppb to O₃ in urban Xi'an. This synergistic impact presented a positive contribution to O₃ production throughout the day and the positive effect was much more prominent during 12:00-19:00 LT.

O₃ pollution in China has been raising increasing concern in recent years. Some scientists hold the view that excessive concentration of PM_{2.5} suppressed the formation of O₃ in the past, hiding the problem temporally. However, with the effective control of PM_{2.5}, O₃ pollution is manifested. The phenomenon can also be demonstrated by the government control action during G20 summit (The Group of Twenty Finance Ministers and Central Bank Governors) in Hangzhou in 2016. The concentration of PM_{2.5} was depressed sharply under the strict emission control, but O₃ concentration was even higher than usual. Better understanding of O₃ pollution sources and sinks and formation mechanisms in high PM_{2.5} exposed area in China will benefit and guide the implemen-

tation of PM_{2.5}/O₃ cooperative control. Our results suggest that, in big cities geographically close to forest, O₃ pollution can be enhanced by the synergistic impact from the combination of biogenic and anthropogenic sources. The synergistic contribution of each single source to O₃ formation cannot be neglected when making pollution control strategies.

Data availability. Observational datasets and modeling results are available upon request to the corresponding author (linan@nuist.edu.cn).

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/acp-18-7489-2018-supplement.

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

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