
1 **Measurement and model analyses of the ozone variation during 2006 to 2015 and its response**
2 **to emission change in megacity Shanghai, China**

3
4 Jianming Xu^{1,2}, Xuexi Tie^{3,4}, Wei Gao^{1,2}, Yanfen Lin⁵, and Qingyan Fu⁵

5
6 ¹ Yangtze River Delta Center for Environmental Meteorology Prediction and Warning, Shanghai
7 Meteorological Service, Shanghai, 200135, China

8 ² Shanghai Key Laboratory of Health and Meteorology, Shanghai Meteorological Service, Shanghai,
9 200135, China

10 ³ Key Laboratory of Aerosol Chemistry & Physics, SKLLQG, Institute of Earth Environment, Chinese
11 Academy of Science, Xi'an, 710061, China

12 ⁴ Center for Excellence in Urban Atmospheric Environment, Institute of Urban Environment,
13 Chinese Academy of Science, Xiamen, 361021, China

14 ⁵ Shanghai Environmental Monitoring Center, Shanghai, 200135, China

15
16
17
18
19 Correspondence: Xuexi Tie (tiexx@ieecas.cn)

20

21 **Abstract.** The fine particles ($PM_{2.5}$) in China decrease significantly in recent years as a result
22 of the implement of Chinese Clean Air Action Plan since 2013, while the O_3 pollution is getting
23 worse, especially in megacities such as Beijing and Shanghai. Better understanding the elevated
24 O_3 pollution in Chinese megacities and its response to emission change is important for
25 developing an effective emission control strategy in future. In this study, we analyze the
26 significant increasing trend of daily maximum O_3 concentration from 2006 to 2015 in the
27 megacity Shanghai with the variability of 0.8-1.3 ppbv yr^{-1} . It is likely attributed to the notable
28 reduction of NO_x concentration with the decreasing rate of 1.86-2.15 ppbv yr^{-1} accompanied with
29 the little change of VOCs during the same period by excluding the weak trends of meteorological
30 impacts on local dispersion (wind speed), regional transport (wind direction) and O_3 photolysis
31 (solar radiation). It is further illustrated by using a state of the art regional chemical/dynamical
32 model (WRF-Chem) to explore the O_3 variation response to the reduction of NO_x emission in
33 Shanghai. The control experiment conducted for September of 2009 shows very excellent
34 performance for O_3 and NO_x simulations including both the spatial distribution pattern, and the
35 day by day variation through comparing with 6 in-situ measurements from MIRAGE-shanghai
36 field campaign. Sensitive experiments with 30% reduction of NO_x emission from 2009 to 2015 in
37 Shanghai estimated by Shanghai Environmental Monitoring Center shows that the calculated O_3
38 concentrations exhibit obvious enhancement by 4-7 ppbv in urban zones with the increasing
39 variability of 0.96-1.06 ppbv yr^{-1} , which is well consistent with the observed O_3 trend as a result
40 of the strong VOC-limited condition for O_3 production. The large reduction of NO_x combined with
41 less change of VOCs during the past ten years promotes the O_3 production in Shanghai to move
42 towards NO_x -limited regime. Further analysis of WRF-Chem experiments and O_3 isopleths
43 diagram suggests that the O_3 production in downtown is still under VOC-limited regime after
44 2015 despite of the remarkable NO_x reduction, while moves to the transition regime between
45 NO_x -limited and VOC-limited in sub-urban zones. Supposing the insignificant VOCs variation
46 persists, the O_3 concentration in downtown would keep increasing till 2020 with the further 20%
47 reduction of NO_x emission after 2015 estimated by Shanghai Clean Air Action Plan. The O_3
48 production in Shanghai will switch from VOC-limited to NO_x -limited regime after 2020 except
49 downtown area which is likely close to the transition regime. As a result the O_3 concentration will
50 decrease by 2-3 ppbv in sub-urban zones, and more than 4 ppbv in suburb response to 20%
51 reduction of NO_x emission after 2020, whereas is not sensitive to both NO_x and VOCs changes in
52 downtown. This result reveals that the control strategy of O_3 pollution is a very complex process,
53 and needs to be carefully studied.

54
55 **Key Words:** O_3 pollution in Shanghai, Long-term O_3 trend, WRF-Chem

56

57 1 Introduction

58 Ozone (O₃) in the troposphere plays the important role in the oxidation of chemically and
59 climatically relevant trace gases, hence regulating their lifetime in the atmosphere (Monks et al.,
60 2015). In the lower troposphere, O₃ is produced from photochemical reactions involving volatile
61 organic compounds (VOCs, broadly including CO) and nitrogen oxides (NO_x = NO + NO₂) in the
62 presence of sunlight (Brasseur et al., 1999). As a strong oxidant, O₃ at ground level is detrimental
63 to human health and vegetation (Tai et al., 2014), and has been received continuous attention
64 from both the scientific and regulatory communities in the past three decades.

65 Shanghai has emerged as one of the largest megacities in the world over the last two
66 decades. The city has a fleet of over 3.6 million vehicles and the population of over 2400 million
67 permanent residents, which results in high emissions of NO_x, VOCs, and primary particulate
68 matter (PM) to the atmosphere from industrial and commercial activities, leading to the
69 photochemical smog formation. Persistent high level of surface O₃ and PM were observed in
70 Shanghai during the past ten years (Geng et al., 2007; Ran et al., 2009; Tie et al., 2009a; Xu et al.,
71 2015). In order to mitigate the adverse impacts from severe air pollution, the Clean Air Action
72 Plan was issued in the end of 2013 to implement the emission reduction program in Shanghai
73 and its neighboring area. As a result, the annual mean PM_{2.5} (particles with diameter $\leq 2.5 \mu\text{m}$)
74 mass concentration has decreased from 50 $\mu\text{g m}^{-3}$ in 2013 to 39 $\mu\text{g m}^{-3}$ in 2017. However O₃
75 pollution has been continuously worsen, with the non-attainment days (daily maximum O₃
76 concentration exceeding 200 $\mu\text{g m}^{-3}$, or daily maximum 8h-O₃ concentration exceeding 100 $\mu\text{g m}^{-3}$)
77 increased from 99 d in 2014 to 129 d in 2016. As a result, O₃ becomes the primary air pollutant
78 affecting the ambient air quality instead of PM_{2.5} in Shanghai. Similar issue has also been
79 occurred in other cities in the eastern China (Lu et al., 2018). For example, the mean PM_{2.5} mass
80 concentration over the 74 major cites decreased by 40% from 2013 to 2017, whereas the
81 maximum daily 8-h average O₃ concentration in summer exceeds the Chinese National Ambient
82 Air Quality Stand (GB3095-2012) over most of eastern China (Li et al., 2019). Thus better
83 understanding the causes of elevated O₃ in China is important for developing effective O₃ control
84 strategies, especially in megacities such as Shanghai.

85 A prerequisite to an effective emission-based O₃ control strategy is to understand the
86 temporal and spatial relationship between O₃ and its precursors, and the response of O₃
87 concentrations to the changes in emissions of O₃-precursors (such as NO_x and VOCs, Lin et al.,
88 1988). The relationship of O₃ and O₃-precursors can be clarified as NO_x-limited or VOC-limited
89 chemistry of O₃ formation, which is usually defined based on the relative impact of a given
90 percent reduction in NO_x relative to VOCs in the context of urban chemistry (Sillman, 1999).

91 Some observational and modeling works on O₃ chemical formation and transformation have
92 been carried out in Shanghai since 2007. The O₃ production in Shanghai city is clearly under
93 VOC-limited regime (Geng et al., 2007), in which the aromatics and alkenes play the dominant
94 roles (Geng et al., 2008a). The aircraft measurements in Yangtze River Delta (YRD) region show
95 the strong anti-correlation between NO_x and O₃ during noontime, indicating the similar
96 VOC-limited regime for O₃ production in the area neighboring Shanghai (Geng et al., 2008b). Thus
97 either NO_x reduction or VOCs growth is favorable for O₃ enhancement in Shanghai. Gao et al.
98 (2017) reported that O₃ concentration in Shanghai downtown increased 67% from 2006 to 2015,
99 whereas NO_x concentration decreased about 38%. This is also consistent with the results of Lin et
100 al. (2017) that, the median of the maximum daily 8-h average O₃ concentration in Shanghai

101 increased notably from 2006 to 2016, with the rate of 1.4 ppbv yr^{-1} , while the NO_2 decreased
102 from 66.7 to $42.1 \mu\text{g m}^{-3}$ with about 20% reduction. These previous studies provide the useful
103 information regarding the O_3 chemical formation and transformation in Shanghai. However, such
104 O_3 variation responding to emission change has not been clearly investigated. Considering that O_3
105 formation is a complicated process including chemistry, transport, emission, deposition and their
106 interactions, the chemical transport model is the powerful tool to gain an understanding of these
107 interacting processes. For example, Lei et al. (2007), Ying et al. (2009) and Song et al. (2010)
108 investigated the O_3 production rate and its sensitivity to emission changes of O_3 precursors by
109 CAMx model in Mexico City Metropolitan Area (MCMA). Tie et al. (2013) analyzed the
110 comprehensive data of the MIRAGE-Shanghai field campaign by WRF-Chem model, and
111 quantified the threshold value by the emission ratio of NO_x/VOCs for switching from VOC-limited
112 to NO_x -limited in Shanghai. Recently Li et al. (2019) suggested an important cause of the
113 increasing O_3 in North China Plain (NCP) during 2013 to 2017 as the significant decrease in $\text{PM}_{2.5}$
114 slowing down the sink of hydroperoxy radicals and thus speeding up the O_3 production by
115 GOES-CHEM model. However such implication for O_3 trend is not pervasive in YRD and other
116 regions. Moreover, the 5-year O_3 records seem rather short to examine the inter-annual
117 variability of O_3 concentration. The GOES-CHEM experiment with 50 km resolution maybe is
118 suitable for the O_3 simulation at regional scale but is too coarse to resolve the local O_3 budget at
119 urban scale, such as Beijing or Shanghai. To our knowledge, there are no peer-reviewed modeling
120 studies focus on the past long term O_3 variations response to emission changes conducted in
121 Shanghai. Thus this paper extends the study of Tie et al. (2013) and Gao et al. (2017) to not only
122 further examine the inter-annual O_3 variations from a larger scale with more comprehensive
123 measurements, but also explore the O_3 enhancement response to NO_x reduction in Shanghai and
124 predict the future O_3 variations by models. The effects of emission changes on long term O_3
125 variability are evaluated by WRF-Chem model with high resolution and compared with
126 measurements. The shift of O_3 photochemical regime relative to the variations of NO_x and VOCs
127 concentrations in the past ten years is discussed by O_3 isopleths diagram combined with
128 WRF-Chem model to provide more insights into the O_3 control strategy. Moreover, the future O_3
129 levels and its possible chemical regime in Shanghai are also discussed according to the Shanghai
130 Clean Air Action Plan.

131 The paper is constructed as follows. The measurements and models used for this study are
132 described in Sect. 2. The analysis on the long-term in-situ measurements of O_3 and its precursors,
133 as well as the model sensitive experiments are presented and discussed in Sect. 3-6. The
134 conclusion is summarized in Sect. 7.

135 136 **2 Measurements and models**

137 **2.1 Measurements**

138 The measurements of O_3 and NO_x are collected from 6 sites (XJH, PD, JS, BS, SS, DT) over
139 Shanghai (Fig. 1 a) under different influence of air pollutant emissions. The XJH site is located at
140 the downtown of Shanghai, which is strongly influenced by emission of transportation. The PD
141 site is located at the sub-urban area near a big park, which is influenced by the mixed emissions
142 of transportation and residential. The JS site is located in the south of Shanghai with several large
143 chemical industries. The BS site is located in the north of Shanghai with some big steel and power

144 plants. The SS site is located at the top of the sole hill (100 m a.g.l) in Shanghai, which has minor
145 effect from local emissions, and is influenced by regional transport. The DT site is located at a
146 remote island without anthropogenic activities. These O₃ and NO_x measurements are used for
147 the evaluation on WRF-Chem performance. In addition, the VOCs are sampled at the downtown
148 site XJH and the sub-urban site PD, and are analyzed at a chemistry laboratory. The study on the
149 O₃ chemical production in this paper is limited at XJH and PD by the intensive measurements of
150 O₃ and its precursors (VOCs and NO_x) from 2006 to 2015. The meteorological measurements
151 including wind speed and direction, solar radiation and temperature are collected at BS site,
152 which is the only climatology observatory in Shanghai. The meteorological measurements in BS
153 are used for international exchange of meteorological data representing Shanghai, sponsored by
154 the World Meteorological Organization (WMO).

155 **2.2 Instruments**

156 O₃ is measured using an EC 9810 Ozone Analyzer, together with a UV photometer, which
157 accurately and reliably measures O₃ concentration in ambient air. The oxides of nitrogen analyzer
158 (EC9841B/ECOTECH) have a heated molybdenum NO₂ to NO converter. The resulting NO
159 concentration is quantified using the chemiluminescence technique. This instrument has
160 automated to set to be zero, and include an optional external valve manifold and external
161 calibration sources. Quality control checks are performed every 3 days, including inspection of
162 the shelter and instruments as well as zero, precision and span checks. Filter is replaced once
163 every two weeks and calibration is made every month. The O₃ concentrations are recorded every
164 1 min.

165 VOCs concentrations are sampled for 24 h every day with a 6 L silonite canister with a
166 silonite coated valve (model 29-10622). The internal silonite coating improves long-term VOC
167 storage. The instrument has a large volume to detect volatile chemicals down to low pptv range.
168 Absorption is eliminated by using nupropackless valves and by eliminating teflon tape in the valve
169 stem. These canisters are recognized to meet or exceed the technical specifications required for
170 EP methods TO14-A and TO15. Gases samples are pre-processed using Model 7100 VOC
171 preconcentrator. Samples are analyzed for VOCs using a gas chromatography system (Agilent
172 GC6890) coupled with mass-selective detection (Agilent MSD5975 N) with length of 60 m,
173 diameter of 0.32 mm, and film thickness of 1.0 um. This measurement system can detect VOCs
174 concentrations down to low pptv range.

175 These instruments to measure O₃, NO_x and VOCs concentrations are calibrated carefully.
176 Detail information for the instruments and the procedures to perform data quality control are
177 described by Geng et al. (2007), Ran et al. (2009), Tie et al. (2013) and Gao et al. (2017). These
178 data have been widely used to investigate the diurnal, seasonal and inter-annual variations of O₃
179 in Shanghai (Geng et al., 2007; 2015; Tang et al., 2008; Ran et al., 2009; Gao et al., 2017) and its
180 chemical mechanism (Geng et al., 2008a; 2008b; Tie et al., 2009a; 2013).

181 **2.3 WRF-Chem model**

182 The regional chemical/transport model (Weather Research and Forecasting Chemical model-
183 WRF-Chem) (Grell et al., 2005) is used to investigate the O₃ variations response to emission
184 changes in Shanghai. The version of the model is improved mainly by Tie et al. (2007) and Li et al.
185 (2010; 2011). The chemical mechanism chosen in WRF-Chem is the RADM2 (Regional Acid

186 Deposition Model, version 2) gas-phase chemical mechanism (Stockwell et al., 1990), which
187 includes 158 reactions among 36 species. The fast radiation transfer module (FTUV) is developed
188 and used to calculate photolysis rates (Tie et al., 2003), considering the impacts of aerosols and
189 clouds on the photochemistry (Li et al., 2011). The aerosol modules are developed by EPA CMAQ
190 (version 4.6) (Binkowski and Roselle, 2003). The wet deposition of chemical species is calculated
191 using the method in the CMAQ module and the dry deposition parameterization follows Wesely
192 et al. (1989). The ISORROPIA version 1.7 is used to calculate the inorganic aerosols (Nenes et al.,
193 1998). The secondary organic aerosol (SOA) is predicted using a non-traditional SOA module,
194 including the volatility basisset (VBS) modeling approach in which primary organic components
195 are assumed to be semi-volatile and photochemically reactive and are distributed in
196 logarithmically spaced volatility bins. The partitioning of semi-volatile organic species is
197 calculated supposing the bulk gas and particle phases are in equilibrium and all condensable
198 organics form a pseudoideal solution. Nine surrogate species with saturation concentrations from
199 10^{-2} to $10^6 \mu\text{g m}^{-3}$ at room temperature are used for the primary organic aerosol (POA)
200 components. The SOA contributions from glyoxal and methylglyoxal is also included. The major
201 physical processes employed in WRF are summarized as the Lin microphysics scheme (Lin et al.,
202 1983), the Yonsei University (YSU) PBL scheme (Hong et al., 2006), the Noah Land surface model
203 (Chen and Dudhia, 2001), and the long wave radiation parameterization (Dudhia, 1989).

204 The domain is set up to covered a region (centered at 32.5°N, 118°E) of 356×345 grids with
205 a horizontal resolution of 6 km (Zhou et al., 2017). The initial and lateral boundary conditions of
206 the meteorology are extracted from the NCEP FNL reanalysis data. The lateral meteorological
207 boundary is updated every 6 h. The chemical lateral boundary conditions are constrained by the
208 global chemical transport model (MOART–Model for Ozone and Related chemical Tracers) with
209 aerosol formation modules (Tie et al., 2001; Emmonset al., 2010). Both the chemical and
210 dynamical integration step is set as 60 s. The Multi-resolution Emission Inventory for China (MEIC)
211 developed by Zhang et al. (2009) is used in WRF-Chem for the domain except Shanghai with 0.25°
212 resolution. The anthropogenic emissions (including CO, NO_x, SO₂ and VOCs) for Shanghai are
213 developed by Tie et al. (2013) with 0.16° resolution based on the MIRAGE-shanghai field
214 campaign. NO_x and SO₂ emissions in YRD region are adjusted by Zhou et al. (2017) according to
215 the performance evaluation of WRF-Chem prediction for about 195 cities during 2014-2015. The
216 distribution of NO_x emission in 2009 in Shanghai is depicted in Fig. 1b. The biogenic emissions are
217 calculated online using the MEGAN (Model of Emissions of Gases and Aerosol from Nature)
218 model developed by Guenther et al. (2006).

219

220 **Figure 1.** (a) The distribution of land-use category in Shanghai. The blue dots denote the locations
221 of 6 sties (XJH, BS, PD, SS, JS, DT). (b) The NO_x emission of 2009 scenario in Shanghai.

222 **2.4 OZIPR model**

223 The ozone isopleths diagram for Shanghai is plot by OZIPR (Ozone Isopleths Plotting Package
224 Research) model (Gery and Crouse, 2002). The OZIPR model employs a trajectory-based air
225 quality simulation model in conjunction with the Empirical Kinetics Modeling Approach (EKMA)
226 to relate O₃ concentrations levels of organic and nitrogen oxide emissions. OZIPR simulates
227 complex chemical and physical processes of the lower atmosphere through a trajectory model.
228 The physical representation is a well-mixed column of air extending from the ground to the top of

229 the mixed layer. Emissions from the surface are included as the air column passes over different
230 emission sources, and air from above the column is mixed in as the inversion rises during the day.
231 O₃ precursor concentrations and ambient information such as temperature, relative humidity and
232 boundary layer height from measurements in Shanghai were specified for each single run.
233 Therefore a series of simulations were performed to calculate peak O₃ concentration as a
234 function of initial precursor concentrations (Tang et al., 2008; Geng et al., 2008b).

235

236 **3 Variability of O₃ and its precursors measured in Shanghai**

237 **3.1 Variation of O₃ concentration**

238 Fig. 2a and b show the annual variation of daily maximum O₃ concentration at downtown site XJH
239 and sub-urban site PD respectively from 2006 to 2015. The daily maximum O₃ concentrations
240 increase notably during the past ten years with the increasing rate of 0.808 ppbv yr⁻¹ at XJH and
241 1.374 ppbv yr⁻¹ at PD respectively. In similar the daily maximum 8h-O₃ concentration also
242 increased at the rate of 1.06 and 1.4 ppbv yr⁻¹ respectively. It is consistent with the reported O₃
243 increasing trend ranging from 1-2 ppbv yr⁻¹ at background and urban sites in eastern China during
244 2001 to 2015 (Tang et al., 2009; Ma et al., 2016; Sun et al., 2016). In 2006, the mean daily
245 maximum O₃ concentrations at XJH and PD are 25.2 ppbv and 32.7 ppbv respectively. While in
246 2017, the mean daily maximum O₃ concentrations at the two sites increase to 41.3 ppbv and 51.8
247 ppbv respectively, with 64% and 58% enhancement compared with that in 2006. The mean daily
248 maximum O₃ concentration at downtown site XJH during 2006 to 2015 is 39.2 ppbv, which is
249 significantly lower than that at sub-urban site PD of 50.7 ppbv, suggesting the O₃ is depressed in
250 downtown area. Geng et al. (2007) suggested that the O₃ production in the city of Shanghai was
251 under VOC-limited regime, thus higher NO_x in downtown resulted in lower O₃ concentration.
252 Considering the inhomogeneous spatial distribution of the precursors of O₃ in Shanghai (Geng et
253 al. 2008a), we extend the analysis on O₃ variations to a broader scope by using the O₃
254 measurements from 31 sites provided by Shanghai Environmental Monitoring Center, covering
255 the entire Shanghai area. It is shown in Fig. 2c that the median of the O₃-8h concentration also
256 increases significantly from 2006 to 2015, with the increasing rate of 1.571 ppbv yr⁻¹, indicating
257 that the significant increasing trend of O₃ concentration not only occurs in the city of Shanghai,
258 but also expanded to a larger area nearby Shanghai. Li et al. (2019) also reported a regional O₃
259 increasing phenomena in summer during 2013 to 2017 from Shanghai to Beijing in eastern China.

260 In order to analyze the individual contribution to the long-term O₃ trend, the variations of O₃
261 precursors, and meteorological parameters are measured and showed in the following sections.

262

263 **Figure 2.** The annual variation of daily maximum O₃ concentration (ppbv) from 2006 to 2015 at (a)
264 downtown site XJH and (b) sub-urban site PD, both presenting the significant increasing trends
265 with 0.808 ppbv yr⁻¹ at XJH and 1.374 ppbv yr⁻¹ at PD. The variation of the median 8-h O₃
266 concentration (ppbv) from 2006 to 2015 averaged for 31 sites over Shanghai (c), also shows the
267 increasing variability of 1.571 ppbv yr⁻¹.

268 **3.2 Variations of the precursors (NO_x and VOCs)**

269 It is well known that the tropospheric O₃ formation is throughout a complicated photochemical
270 process, and is strongly related to the precursors of O₃ (VOCs and NO_x). According to the previous

271 studies (Geng et al., 2007; Ran et al., 2009), the chemical formation of O₃ in Shanghai is revealed
272 to be under VOC-limited. Thus either enhancement of VOCs or reduction in NO_x would both
273 result in the growth of O₃ concentration. In order to better understanding the factors possibly
274 driving the O₃ increasing trend depicted in Fig. 2, the variations of NO_x and VOCs concentrations
275 at XJH and PD in the same period are presented in Fig. 3. The NO_x concentrations present
276 significant decreasing trend from 2006 to 2015 at both XJH and PD sites, which is opposite to the
277 increasing trend of O₃ variations in Fig. 2. At XJH, the decreasing rate of NO_x is 2.15 ppbv yr⁻¹,
278 which is more remarkable than that at PD site of 1.86 ppbv yr⁻¹. According to the studies by Lin et
279 al (2017), the reduction of NO_x concentration in Shanghai was likely attributed to the
280 implementation of stringent emission control strategy for transportation, including improvement
281 of gas quality, popular usage of electricity cars, and limitation of heavy cars into the urban zones.
282 These regulations significantly decrease the emissions of NO_x into the atmosphere, resulting in
283 lower NO_x concentrations. Zheng et al. (2018) also reported the 30% reduction of NO_x emission in
284 the past 5 years in YRD region. In comparison, the VOCs concentrations at XJH and PD decrease
285 very slightly during 2006 to 2015. At XJH, the mean VOCs concentration during 2013 to 2015 is
286 about 20 ppbv, which is some lower than that during 2009 to 2012 of 23 ppbv. At PD, the VOCs
287 concentration shows strong inter-annual variations, ranging from 16 to 22 ppbv. Generally the
288 VOCs concentration at the downtown site XJH is higher than that at the sub-urban site PD by 14%.
289 It is consistent with the studies of Cai et al. (2010), suggesting that about 25% of VOCs is
290 attributed to the vehicles in shanghai urban zones.

291

292 **Figure 3.** The mean annual concentrations (ppbv) of NO_x (dots) and VOCs (bars) from 2006 to
293 2015 at (a) downtown site XJH and (b) sub-urban site PD respectively. The NO_x concentrations at
294 XJH and PD both present obvious decreasing trends with -2.1 ppbv yr⁻¹ and -1.87 ppbv yr⁻¹. While
295 the VOCs concentrations at both sites present no clear inter-annual trends.

296

297 3.3 Meteorological impacts on O₃ photolysis, dispersion and transport

298 In addition to the precursors, meteorology such as solar radiation and wind speed and directions
299 also plays the important roles in O₃ concentration through the photochemical and physical
300 processes. Fig. 4 shows the annual variation of wind speed and total solar radiation from 2006 to
301 2015. The solar radiation presents weak annual variations ranging from 140 to 150 Wm⁻²,
302 exhibiting a large variability but without a significant trend. As a result, the variation of solar
303 radiation cannot explain the significant change of O₃ concentration on the view of photolysis. The
304 wind speed is usually regarded as the indicator for the dispersion capacity for air pollutants.
305 Several studies reported that the wind speed in winter in eastern China presented decreasing
306 variability during the past 40 years due to the decadal variation of winter monsoon affecting the
307 haze occurrence (Wang et al., 2016; Zhao et al., 2016; Xu et al., 2017). While high O₃ events
308 usually occur in summer season for middle-latitude cities such as Shanghai (Wang et al., 2017).
309 The mean summer wind speed in Fig. 4a fluctuates between 3.3 ms⁻¹ to 3.9 ms⁻¹ during 2006 to
310 2015 except the minimum value in 2014 (2.9 ms⁻¹) due to fewer typhoon in the period. Without
311 2014, the variability of summer wind speed is insignificant, with a trend of -0.02 m s⁻¹ yr⁻¹, which
312 could not be regarded as the dominant factor to interpret the increasing O₃ trend. Local O₃
313 concentration would be affected by transport of upstream plumes usually determined by wind

314 direction. Geng et al. (2011) suggested that O₃ concentration was higher in west wind compared
315 with other wind sectors in Shanghai indicating the possible O₃ transport from western area out of
316 Shanghai. Fig. 5 presents the annual wind rose at Baoshan site from 2006 to 2015, presenting the
317 very similar pattern of wind direction in each year. The mean wind direction concentrates in the
318 sector between 60-80 degree, suggesting the dominant wind in Shanghai is easterly accounting
319 for 50%. The east wind in Shanghai usually carries with the clean air mass from the sea to
320 improve the local air quality (Xu et al., 2015). The frequency of west wind changes little during
321 2006 and 2015 ranging from 10-15%, suggesting that the regional transport is not a major factor
322 driving the O₃ increasing. Based on the above analysis, it is speculated that the rapid O₃
323 increasing during 2006–2015 in shanghai is likely attributed to the reduction of NO_x
324 concentration as a result of the VOC-limited condition for O₃ production.

325

326 **Figure 4.** The annual variation of (a) summer wind speed (m s^{-1}) and (b) total solar radiation (W
327 m^{-2}) from 2006 to 2015 in Shanghai. Both wind speed and the solar radiation present weak
328 inter-annual variations but without significant trends.

329

330 **Figure 5.** The wind rose in each year from 2006 to 2015 in Shanghai. The red line means the
331 resultant vector suggesting the dominant wind direction.

332

333 **3.4 Different O₃ variability in nighttime and daytime**

334 The mean diurnal variations of O₃ concentrations between 2006 and 2015 are compared in Fig.
335 6a at XJH and PD sites respectively. The maximum and minimum O₃ concentrations occur in the
336 afternoon (14-15 pm) and early morning (6-7 am) respectively at both sites. In addition, the
337 diurnal O₃ concentrations at XJH and PD sites all increase significantly from 2006 to 2015. For
338 example, the peak O₃ concentration at XJH increases from 21 ppbv to 37 ppbv, meanwhile the
339 minimum O₃ concentration rises from 5 ppbv to 14 ppbv, exhibiting higher increasing rate. Similar
340 diurnal O₃ enhancement is also observed at PD site during the same period. The O₃ chemical
341 mechanism in daytime includes both production and loss processes. In contrast, in nighttime, the
342 photochemical production ceases, and there mainly exists loss process for O₃. In addition both
343 dry deposition and nighttime turbulence also have the influence in the nighttime O₃
344 concentration according to the work by Hu et al. (2013). Fig. 6b shows the annual change rate of
345 the diurnal O₃ concentration from 2006 to 2015 at XJH and PD sites respectively. The O₃
346 concentrations present increasing trends both in daytime (8:00-18:00, LST) and nighttime
347 (19:00-07:00, LST) at XJH and PD sites, which is consistent with the results in Fig. 2. The nighttime
348 O₃ concentrations increase more significantly than daytime O₃ at XJH, with the increasing rate of
349 1.239 and 0.956 ppbv yr⁻¹ respectively. While at PD site the O₃ concentrations increase by 1.338
350 ppbv yr⁻¹ in daytime which is higher than that in nighttime by 1.028 ppbv yr⁻¹. In comparison, the
351 nighttime O₃ concentrations exhibit higher increasing rate at downtown site XJH than that at
352 sub-urban site PD due to more NO emissions or more intensified urbanization (Hu et al., 2013) at
353 urban center. These results suggest that the reduction of NO_x concentration from 2006 to 2015
354 has different effects on daytime and nighttime O₃ variations. The O₃ concentration in nighttime is
355 more sensitive to NO_x reduction at downtown site, resulting in less O₃ lost compared with that in
356 daytime. The results in Fig. 6b also show that the increasing rate of nighttime O₃ in downtown

357 site XJH is higher than that at sub-urban site PD due to the more reduction of NO_x concentration
358 in downtown area. Furthermore, the seasonal variability of daytime and nighttime O₃
359 concentrations at XJH site are illustrated in Fig. 7. Both daytime and night O₃ concentrations
360 present increasing trends in all seasons. In comparison, the larger increasing rates of nighttime O₃
361 concentration are observed in spring, summer and autumn than that in daytime. For example,
362 the nighttime O₃ concentrations increase at 1.341, 1.159 and 1.525 ppbv yr⁻¹ in spring, summer
363 and autumn respectively, which are more significant than that of 1.008, 0.378 and 1.370 ppbv yr⁻¹
364 in daytime. The variability of winter O₃ concentrations in daytime and nighttime are generally
365 close perhaps due to the lower O₃ photochemical productions. Hu et al. (2016) suggested that
366 the nighttime boundary layer tended to be less stable resulted from the enhanced sensible heat
367 flux in urban area, thus leading to more active nighttime turbulence. The sounding
368 measurements at 20:00 (LST) in Shanghai are used to calculate the vertical temperature gradient
369 between 1000 hPa and 925 hPa to indicate the intensity of nighttime turbulence, while
370 presenting no significant trend from 2010 to 2015. Furthermore the PBL height retrieved from
371 Lidar measurements at 20:00 (LST) presents the similar results as soundings. Based on the above
372 measurements, the variation of turbulence at night may have only minor contribution to the
373 nighttime O₃ increasing in Shanghai. However the effect of dry deposition could not be excluded
374 by lacking of measurements, which need further investigation.

375

376 **Figure 6.** (a) The mean diurnal variation of O₃ concentration (ppbv) compared between 2006 and
377 2015 in XJH (red dots) and PD (blue dots). (b) The annual change rate of diurnal O₃ concentration
378 (ppbv.yr⁻¹) from 2006 to 2015 at downtown site XJH (red bars) and sub-urban site PD (blue bars).

379

380 **Figure 7.** The daytime (8:00-18:00, LST) and nighttime (19:00-07:00, LST) O₃ variability from 2006
381 to 2015 at downtown site XJH in (a) spring, (b) summer, (c) autumn and (d) winter.

382

383 **4 WRF-Chem study on the O₃ variation response to emission change**

384 **4.1 Design of the model experiments scheme**

385 To better understand the role of NO_x emission reduction in O₃ variation, the WRF-Chem model is
386 utilized to calculate the changes of O₃ concentrations. Lin et al. (2017) suggested that the NO_x
387 emission was reduced in Shanghai in recent years resulted from the implementation of the
388 Shanghai Clean Air Action Plan. The NO_x emission in 2015 is estimated at 33.4×10^4 ton in
389 Shanghai, reduced significantly by 30% compared with that in 2009 of 44.9×10^4 ton. Thus it
390 provided the good opportunity to examine the O₃ variation response to the reduction of NO_x
391 emission in Shanghai. The NO_x emissions in 2009 and 2015 are put into WRF-Chem model
392 respectively to calculate the O₃ concentration. The other emissions (including gas and particulate
393 matter) and meteorology used in WRF-Chem are set to be same. As a result, the difference of O₃
394 concentrations calculated by WRF-Chem is solely attributed to the change of NO_x emission
395 between 2009 and 2015, which is furthermore compared with the measurements.

396 The MIRAGE-shanghai field campaign was conducted in September of 2009 to explore the
397 O₃ chemical formation and transformation in Shanghai (Tie et al., 2013). The mean temperature,
398 mean wind speed and total precipitation in this month is 25 °C, 2.85 m s⁻¹ and 89.5 mm
399 respectively, which is very close to the climatological conditions during the past ten years from

400 2006 to 2015, with 24.7 °C for mean temperature, 2.81 m s⁻¹ for mean wind speed, and 126 mm
401 for total precipitation respectively. In addition, Shanghai is located at the typical sub-tropical area.
402 The meteorology in September is characterized as the low cloud cover and rain occurrence, the
403 slight wind speed and humidity, as well as the moderate solar radiation intensity. As suggested by
404 Tie et al. (2013), the chemical age of O₃ plume in Shanghai urban area in September of 2009 was
405 very young, indicating that the O₃ production was more dependent on the local emissions under
406 such kind of meteorology, hence providing more insights into the O₃ chemical mechanism
407 response to the local emission changes. We chose the meteorology in September of 2009 as the
408 atmospheric background for all the sensitive experiments by WRF-Chem.

409 Tie et al. (2009a; 2013) highlighted that the WRF-Chem model was capable of studying the
410 chemical and physical processed of O₃ in September of 2009 during the MIRAGE-Shanghai
411 campaign. The calculated O₃, NO_x, VOCs and aerosols by WRF-Chem in clean and polluted
412 episodes are fairly in agreement with the measurements except HONO, suggesting that the
413 emission inventory in 2009 used in the model is reasonable for the Shanghai region. Moreover
414 the VOCs emission in Shanghai is greatly improved according to the measurements from the
415 MIRAGE-shanghai field campaign by Tie et al. (2013). Such emission from Tie et al. (2013)
416 representing 2009 scenario is used in this study to conduct the control experiment (T1) as the
417 baseline to simulate the O₃ and NO_x concentrations in September of 2009. The T1 experiment is
418 composed of 30 model runs for each day in September of 2009. Each model run is initiated at the
419 20:00 (LST) and performed for 52 h integrations. The first 28 h integration is regarded as model
420 spin-up periods, the results from the later 24 h integration is captured hourly and averaged for
421 mean daily concentration of O₃ and NO_x. The aim of the T1 experiment is to further evaluate the
422 reliability of the emission inventory in 2009 used in WRF-Chem by fully comparing the calculated
423 O₃ and NO_x concentrations with in-situ measurements of 6 sites over Shanghai.

424 **4.2 The NO_x emission in 2009 used for base experiment**

425 The distribution of NO_x emission of 2009 scenario (Tie et al., 2013) in Shanghai used in
426 WRF-Chem model has been showed in Fig. 1b. The NO_x emission is mostly distributed in the
427 urban zones, suggesting that transportation is the important source. The NO_x is largely exported
428 in downtown and two neighboring sub-urban zones in the east and north respectively. The
429 maximum NO_x emission is estimated at 16 kg hr⁻¹ km⁻² at downtown, compared with 2-6 kg hr⁻¹
430 km⁻² in the sub-urban area. In addition, there is a small town located in the south of Shanghai
431 with the similar intensity of NO_x emission as the sub-urban zones. The total NO_x emission of 2009
432 scenario in Shanghai (Fig. 1b) is estimated at 41.4 × 10⁴ ton in the model, which is close to the
433 47.8 × 10⁴ ton suggested by Lin et al. (2017) according to the Shanghai Environmental Year Book.

434 **4.3 Performance evaluation on the base experiment**

435 The mean daytime and nighttime O₃ concentrations in September 2009 are calculated by
436 WRF-Chem and compared with measurements over 6 sites in Shanghai, which are presented in
437 Fig. 8a and b respectively. Both modeled and measured O₃ concentrations in daytime are higher
438 than that in nighttime. The calculated daytime O₃ concentration is about 10-18 ppbv higher than
439 that in nighttime in urban region, which is consistent with the measured difference of 12-14 ppbv
440 at XJH and PD sites. The observed daytime and nighttime O₃ concentrations at remote site DT
441 show the minimum difference of 5 ppbv which is also captured by WRF-Chem model due to the

442 less impact of anthropogenic emissions. In Fig. 8a, there exists a large O₃ plume with high
443 concentration of 40-48 ppbv in daytime in the west of Shanghai and its neighboring area from
444 WRF-Chem simulations. It is also illustrated by the daytime O₃ measurements at SS site with 40
445 ppbv. However such daytime O₃ plume dissipates at night (Fig. 8b) leading to the significant
446 difference of O₃ concentration between day and night. Tie et al. (2013) suggested the
447 enhancement of O₃ concentration in the downwind of Shanghai due to the considerable O₃
448 formation in the aged city plume transported westerly in September resulted from the dominant
449 west winds. According to the study of Tie et al. (2013), the O₃ concentrations had a minimum
450 within 20 km of the city, and enhanced at the west of 100–150 km away from the city in daytime,
451 which was consistent with the results of daytime O₃ distribution simulated by WRF-Chem in Fig.
452 8a. In addition, both model simulations and in-situ measurements in daytime and nighttime
453 highlight the lower O₃ concentration in urban zones than that in suburb. The simulated daytime
454 and nighttime O₃ concentration in downtown is 28-32 ppbv and 12-14 ppbv respectively,
455 significantly lower than that at sub-urban (36-38 ppbv and 26-28 ppbv respectively) and rural
456 area (40-42 ppbv and 36-38 ppbv respectively). Similarly, the measured daytime O₃ concentration
457 at downtown site XJH is 28 ppbv, lower than that at sub-urban site PD and remote site DT by 12
458 ppbv and 21 ppbv respectively. Geng et al. (2007) suggested that under VOC-limited regime, the
459 lower O₃ concentration in downtown was resulted from the higher NO_x emission, which
460 depressed the O₃ production process. Under high NO_x conditions, the OH radicals are lost by the
461 reaction of $\text{NO}_2 + \text{OH} \rightarrow \text{HNO}_3$ (Sillman, 1995). As a result, higher NO_x concentration in urban
462 area leads to lower OH concentration, which results in smaller O₃ production. Tang et al. (2008)
463 also suggested that the O₃ concentration in Shanghai downtown was higher on weekends than
464 that on weekdays due to the reduced NO_x concentration. However the discrepancy is also evident
465 between model results and measurements. For example, the modeled nighttime O₃
466 concentrations at XJH and PD are about 2-6 ppbv lower than the measurements, perhaps due to
467 the uncertainty of NO_x and VOCs emission in urban area suggested by Tie et al. (2009a). In
468 addition, the calculated daytime O₃ concentrations in the remote site DT and chemical site JS are
469 lower than measurements by 10 ppbv and 6 ppbv respectively. The former is resulted from the
470 overestimation of the wind speed by WRF-Chem model leading to excessive O₃ transport for
471 underestimation (Zhou et al., 2017). While the latter is mainly due to the prominent
472 underestimation of the VOCs emission in the chemical zones suggested by Tie et al. (2009a).

473

474 **Figure 8.** The calculated distribution of (a) daytime and (b) nighttime O₃ concentration by
475 WRF-Chem (shade) in September of 2009 compared with measurements (circles) of 6 sites over
476 Shanghai. The minimum O₃ concentrations in daytime and nighttime both occur in urban center.

477

478 Fig. 9a and b show the daily variations of O₃ and NO_x concentrations compared between
479 WRF-Chem simulations and the in-situ measurements over 5 sites. The statistical analysis of
480 model performance for O₃ and NO_x is listed in Table 1 and Table 2 respectively. The calculated
481 magnitude and daily variation of O₃ concentrations agree well with the measurements,
482 suggesting that both meteorology and photochemistry are well reproduced by WRF-Chem model.
483 For example, the Root Mean Square Error (RMSE) calculated between modeled and measured O₃
484 concentration are 7.4, 10.5, 12, 8.6, 9.2 ppbv for XJH, JS, DT, PD and BS respectively, and the
485 difference between the simulation results and in-situ measurement is below 10%, which are very

486 satisfactory compare with the similar works by Geng et al (2007) and Tie et al. (2013). The
487 correlated coefficients (R) for the mean daily O₃ concentration range from 0.6 to 0.8 above 99%
488 confidence over 5 sites, indicating good consistency of day by day variations between the model
489 results and measurements. Comparably the O₃ concentration is best simulated by WRF-Chem at
490 the downtown site XJH and sub-urban site PD with the lower RMSE and better R. However the
491 discrepancy of daily O₃ concentration between the model and measurements is also evident. For
492 example, a rapid change of O₃ concentration from 16 to 19 in September was observed over all
493 sites, indicating it's a regional event instead of a local phenomenon. The O₃ concentration firstly
494 increases significantly during 16-19 (episode-1), then sharply decreased during the later 4 days
495 (episode-2). The similar rapid O₃ change in Shanghai was also reported by Tie et al. (2009a), and
496 their explanation is that this episode was mainly related to the intensity of the sub-tropical
497 high-pressure system on Pacific Ocean in summer. The model captures the O₃ variations and
498 magnitudes during the both risen and fallen episodes very well at downtown site XJH, but
499 substantially underestimates the increasing variability of O₃ concentration during episode-1 at
500 sub-urban and rural sites by 10-15 ppbv. Geng et al. (2008a) suggested the "chemical transport of
501 O₃" from Shanghai downtown area to the distance of 18-36 km far away, which increased the O₃
502 concentration at sub-urban or rural sites. This "chemical transport of O₃" is difficult to be
503 reflected by WRF-Chem model due to the current inventory is too coarse to accurately reflect the
504 detailed distribution and variation of NO_x emission, e.g. the NO_x emission from mobile source in
505 the city. In addition, the underestimation of the O₃ concentration at suburb of Shanghai in
506 summer is possibly attributed to the model bias of sea breeze simulations. Under the condition of
507 weak sub-tropical pressure, the sea breeze develops at noontime to yield a cycling wind pattern
508 in Shanghai, leading to the rapid accumulation of high O₃ concentration. The WRF-Chem usually
509 underestimates the sea surface temperature, which tends to accelerate the sea breeze
510 development and weak the O₃ trapping in the city (Tie et al., 2009a). The calculated daily NO_x
511 concentration by WRF-Chem compared with measurements are shown in Fig. 9b. Both the
512 modeled and measured NO_x concentrations at the remote site DT are very low, with the average
513 of 1.4 and 2.9 ppbv respectively due to seldom anthropogenic emissions there. The calculated
514 NO_x concentration at XJH and PD are generally well consistent with the measurements with the
515 excellent R of 0.8 and 0.82 and small RMSE of 6.9 and 7.5 ppbv respectively. However the NO_x
516 concentration is underestimated by WRF-Chem at sub-urban site BS in the steel zone. The
517 calculated NO_x concentration at BS is 16.1 ppbv, which is lower than the measurements by 5 ppbv.
518 The difference of NO_x concentrations between the model and observations is generally above
519 10%, suggesting the performance of NO_x simulation is somewhat lower than that of O₃. It was
520 also reported by Tie et al. (2007; 2009b; 2013), during the evaluation of the NO_x calculations by
521 WRF-Chem in MIRAGE-Shanghai and MIRAGE-mex campaign studies. The lifetime of NO_x at the
522 surface is about 1-2 days, shorter than O₃. Thus the NO_x concentration is determined by the
523 detailed emissions and dynamical factors, which need to develop the advanced inventory with
524 higher resolution to reproduce both the spatial distributions and temporal variations of NO_x
525 emission.

526

527 **Figure 9.** The calculated mean daily concentrations (ppbv) of (a) O₃ and (b) NO_x at 5 sites in
528 September of 2009 by WRF-Chem (red circles) and compared with measurements (blue circles).

529

530 4.4 Sensitive study on the O₃ variability response to the emission change

531 The T1 experiment shows the excellent performance for O₃ and NO_x simulations, including the
532 spatial distribution pattern, and the day by day variation and magnitude. It is indicated that the
533 emission in 2009 scenario used in WRF-Chem is reasonable, and the model is efficient for
534 conducting the sensitive studies on O₃ variation response to the emission change. In order to
535 better understand the measured long-term trend of O₃ concentration during the past ten years in
536 Shanghai and its relationship to the emission reduction, several sensitive studies are conducted in
537 this study (Table 3). The control study of T1 is conducted based on the NO_x emission in 2009
538 scenario in Shanghai. According to the study of Lin et al. (2017), the NO_x emission in 2015 in
539 Shanghai is reduced by 30% compared with that in 2009. Thus we conduct the sensitive
540 experiment T2 by WRF-Chem, cutting the NO_x emission by 30% compared with T1, whereas
541 keeping the other emissions and meteorology same as T1. As a result, the calculated O₃
542 difference between T1 and T2 is likely attributed to the NO_x emission reduction between 2015
543 and 2009.

544 Fig. 10a shows the distribution of the difference of O₃ concentration simulated by T1 and T2
545 (T2-T1). The reduction of NO_x emission has the obvious effect on the magnitude and distribution
546 of O₃ concentration. The O₃ concentration increases notably in urban area corresponding to the
547 higher NO_x emissions in Fig. 1, ranging from 2-7 ppbv. The enhancement of O₃ concentration is
548 most significant in downtown and neighboring sub-urban zones, as well as the southern town,
549 generally more than 4 ppbv. For example, the maximum increase in O₃ concentration is 6.4 ppbv
550 occurred at downtown site XJH, followed by 4-5 ppbv at sub-urban site PD. The increasing rates
551 of O₃ trend at XJH and PD are estimated at 1.06 ppbv yr⁻¹ and 0.96 ppbv yr⁻¹ from 2009 to 2015
552 by WRF-Chem, which is consistent to the observed O₃ growth variability of 1-1.3 ppbv yr⁻¹. The
553 response of O₃ concentration to the NO_x reduction is not evident in the rural area including the
554 eastern part of Shanghai and the island with low NO_x emissions. The comparison of T1 and T2
555 further illustrates the speculation that the significant increasing trend of O₃ concentration during
556 the past ten years in Shanghai is mostly attributed to the reduction of NO_x emission as a result of
557 the implementation of Shanghai Clean Air Action Plan.

558 The O₃ chemical formation is strongly related to NO_x and VOCs concentrations. As discussed
559 by Geng et al. (2008a) the O₃ chemical formation is clearly under VOC-limited regime in Shanghai
560 and its neighboring area. Under the high NO_x condition, NO tends to react with O₃ instead of NO₂,
561 flowing by $\text{NO}_2 + \text{OH} \rightarrow \text{HNO}_3$, causing the decrease of the reactivity and ensuing O₃
562 concentrations. Thus reduced NO_x emission would result in increase in O₃ concentration, which
563 has been discussed in Fig. 10a.

564 Despite of minor change of VOCs in the last ten years, it is worth to investigate the effect of
565 the VOCs changes on O₃ concentrations in Shanghai. For this purpose, we conduct a sensitive
566 study (T3), with 50% increase of VOCs emission compared with T1, but keeping NO_x and other
567 emissions as well as the meteorology same as T1. For RADM2 gas mechanism used in WRF-Chem,
568 the VOCs are surrogated into 14 species, such as alkane, alkene, aromatic, formaldehyde, etc. All
569 the species of VOCs are increased by 50% at every model grid over Shanghai and at every hour.
570 The difference of O₃ concentration between T3 and T1 (T3-T1) is shown in Fig. 10b. As we
571 expected, the O₃ concentration in Shanghai is sensitive to the enhancement of VOCs emission,
572 increased by 3-4 ppbv in urban area due to more NO is converted to NO₂ by reaction with RO₂
573 and HO₂. Furthermore, the abundant O₃ plumes produced in the urban zones significantly

574 transport to the downwind areas about 100-200 km away, resulting in elevated O₃ concentration
575 in the western Shanghai by about 2 ppbv. According to Tie et al. (2013), the O₃ plume released in
576 Shanghai urban area can be transported to downwind of the city by about 100-150 km away in
577 the MIRAGE-shanghai field campaign. The model studies of T1, T2 and T3 highlight that under the
578 emission of 2009 scenario, the O₃ chemical production is clearly under VOC-limit regime, either
579 decreasing NO_x concentration or increasing VOCs concentration would result in the O₃
580 enhancement. The analysis on in-situ measurements and model experiments jointly suggests that
581 the significant O₃ increasing trend during the past ten years in Shanghai is mainly attributed to
582 the large reduction of NO_x emission.

583

584 **Figure 10.** The difference of O₃ concentration (ppbv) between (a) T2 and T1 (T2-T1), (b) T3 and
585 T1 (T3-T1) respectively conducted by WRF-Chem model. The difference between T2 and T1 lies in
586 the NO_x emissions set in T2 (2015 scenario) is 30% lower than that in T1 (2009 scenario), which is
587 estimated by Lin et al. (2017) according to the Shanghai Environment Yearbook. The difference
588 between T3 and T1 is dependent on that the VOCs emission in T3 is 50% higher than that in T1.

589

590 **4.5 The variation of O₃ production regime response to emission change**

591 The O₃ chemical mechanism in Shanghai was explored by several studies based on the in-situ
592 measurements around 2008 and 2009. Geng et al. (2008a; 2008b), Ran et al. (2009) and Tie et al.
593 (2009a) all revealed that the O₃ production around 2008 and 2009 in Shanghai was clearly under
594 VOC-limit regime which was further illustrated by the above model studies. As indicated in Fig. 3,
595 the significant decrease of NO_x concentration is observed from 2009 to 2015 in Shanghai, while
596 the VOCs concentration changed little during the same period. As we know, the O₃ chemical
597 formation is strongly related to NO_x and VOCs concentrations with nonlinearity. Thus the
598 different variability of NO_x and VOCs concentration from 2009 to 2015 inevitably has the large
599 effect on the O₃ production regime, which need to be investigated deeply.

600 The complex relationship among NO_x, VOCs and O₃ concentrations is usually depicted by O₃
601 isopleths diagram. The O₃ isopleths plot (Fig. 11) in Shanghai used in this study is constructed by
602 the OZIPR model based on the in-situ measurements of O₃, NO_x, VOCs and meteorology. Under
603 high VOCs and low NO_x condition (low NO_x/VOCs ratio), the O₃ production is not sensitive to
604 VOCs, while positively correlated to NO_x concentration, which is viewed as NO_x-limited regime. By
605 contrast, under low VOCs and high NO_x condition (high NO_x/VOCs ratio), the O₃ production tends
606 to increase with the VOCs growth or NO_x reduction, which is regarded as VOC-limited regime. The
607 NO_x-limited and VOC-limited regime is divided by a ridge line (the dot-dash line in Fig. 11) in the
608 O₃ isopleths plot. The O₃ production is not sensitive to neither NO_x concentration nor VOCs
609 concentration when near the ridge line, which is regarded as the transition regime.

610 The O₃ chemical production regime at XJH and PD in 2009 and 2015 is positioned
611 respectively in Fig. 11. In 2009 the O₃ production at both XJH and PD sites (marked as red and
612 blue hollow circle respectively) are clearly under VOC-limited regime. Thus decrease in NO_x
613 concentration leads to the O₃ enhancement, which is highlighted by the previous in-situ
614 measurements and model experiments. Since then the O₃ production regime tends to move
615 toward the dot-dash line due to the significant reduction of NO_x concentration accompanied with
616 the relative less change of VOCs at the two sites. In 2015 the O₃ production at XJH (marked as red

617 solid circle) is still under VOC-limited regime, but for PD (marked as blue solid circle), it is close to
618 the dot-dash line, approaching the transition regime between VOC-limited to NO_x-limited. This
619 result suggests that if the NO_x emission keeps reduction after 2015 assuming the VOCs
620 concentration keeps constant, the O₃ concentration will continue to increase at XJH, while at PD
621 the O₃ concentration is supposed to be insensitive to the NO_x change. According to the O₃
622 chemical regime depicted in Fig. 11, if the NO_x concentration decreases by 5 ppbv after 2015, the
623 peak O₃ concentration at XJH will further increase by 3 ppbv, whereas at PD it seems to change
624 very slightly. To better understand this further change, more sensitive studies of WRF-Chem are
625 conducted in the following sections.

626

627 **Figure 11.** The O₃ chemical production at downtown site XJH and sub-urban site PD in 2009 and
628 2015 depicted by O₃ isopleths diagram. The hollow and solid red circles denote O₃ production
629 regime at XJH in 2005 and 2019 respectively. The hollow and solid blue circles denote O₃
630 production regime at PD in 2005 and 2019 respectively

631

632 **5 The future O₃ evaluation**

633 **5.1 The O₃ level in 2020**

634 According to the Shanghai Clean Air Action Plan, the NO_x emission in Shanghai will be further
635 reduced by 20% in 2020 compared with that in 2015. According to the above analysis based on
636 the O₃ isopleths plot (Fig. 11), the O₃ concentrations in downtown and sub-urban seem to have
637 distinct different responses to further NO_x reduction after 2015. In order to better understand
638 the future O₃ variation, the sensitive experiment T4 is conducted by WRF-Chem with 20%
639 reduction of NO_x emission compared with T2. T2 and T4 represent the NO_x emission in 2015 and
640 2020 respectively. The other emissions and meteorology are set to be same as T1. The difference
641 of O₃ concentration between T2 and T4 (T4-T2) is presented in Fig. 12a. The O₃ concentration
642 keeps increasing in downtown area such as XJH site, ranging from 2-4 ppbv. However, for the
643 sub-urban zones such as the PD site, the O₃ concentration changes very little response to the
644 further NO_x reduction, ranging from 0-1 ppbv. As discussed in Fig. 11, in 2015 the O₃ production
645 at PD is possibly under the transition regime from VOC-limited to NO_x-limited near the ridge line.
646 As a result, the O₃ concentration is not sensitive to the variation of NO_x concentration. However
647 the O₃ concentration in the suburb zones generally decreases by 1ppbv, indicating that with the
648 further NO_x reduction after 2015 the O₃ chemical production transfers from VOCs-limited to
649 NO_x-limited regime in the rural of Shanghai.

650 It is suggested in Fig.11 that the O₃ production at downtown site XJH in 2015 is still under
651 VOC-limited regime despite of the significant NO_x reduction. The O₃ concentration would be also
652 sensitive to the variation of VOCs concentration. Thus the sensitive experiment T5 is conducted
653 by WRF-Chem model with 50% enhancement of VOCs emission compared with T2 (representing
654 the emission in 2015 scenario). It is presented in Fig. 12b that the O₃ concentration increases by
655 2-3 ppbv in downtown area due to the enhancement of VOCs, suggesting that the O₃ production
656 at downtown in 2015 is still under VOC-limited regime, which is consistent with the results in Fig.
657 11. Moreover the O₃ plumes produced in urban area transport to the downwind area to
658 accumulate the high O₃ concentration in the western area to Shanghai by 2 ppbv. While at
659 sub-urban site PD, the O₃ concentration changes less than 1 ppbv response to the increase in

660 VOCs emission, which is similar as the very weak O₃ variations relative to the NO_x reduction in Fig.
661 12a. Overall, the models studies of T4 and T5 jointly suggest that the O₃ concentration at
662 sub-urban site PD in 2015 is not sensitive to either NO_x or VOCs variations due to the O₃
663 production is under the transition regime depicted in the O₃ isopleths plot.

664

665 **Figure 12.** The difference of O₃ concentration (ppbv) between (a) T4 and T2 (T4-T2), (b) T5 and
666 T2 (T5-T2) respectively conducted by WRF-Chem model. The difference between T4 and T2 is
667 that the NO_x emissions set in T4 (2020 scenario) is 20% lower than that in T2 (2015 scenario),
668 which is estimated according to the Shanghai Clean Air Action Plan. The difference between T5
669 and T2 lies in that the VOCs emission in T5 is 50% higher than that in T2.

670

671 5.2 The O₃ chemical production after 2020

672 The above study shows that the O₃ production at sub-urban site PD in 2020 will likely transfer
673 from VOCs-limited regime to NO_x-limited regime without consideration of possible VOCs changes.
674 For the purpose of the O₃ pollution control strategy, it is worth to estimate the O₃ level response
675 to emission change after 2020 in Shanghai. It is also essential to access how many NO_x emission
676 need to be cut after 2020 will cease the O₃ enhancement in downtown area. Thus the sensitive
677 experiment T6 is conducted by further 20% reduction of NO_x emission from 2020 scenario (T4).
678 The difference of O₃ concentration between T6 and T4 (T6-T4) is shown in Fig. 13a. It is clear that
679 the O₃ concentration at downtown keeps nearly constant regardless of the further reduction of
680 NO_x emission after 2020. That is to say the increasing trend of O₃ in downtown with the NO_x
681 reduction ceases after 2020, indicating that the O₃ production likely approaches the transition
682 regime. In addition, the O₃ concentration decreases significantly out of the downtown area,
683 ranging from 2-3 ppbv in sub-urban zones, and more than 4 ppbv in suburb, indicating that the
684 O₃ production in Shanghai transfers to NO_x-limited regime after 2020 except the downtown area
685 where the O₃ production is likely near the transition zone. On the other hand, if the NO_x emission
686 is kept constant after 2020 as T4, while the VOCs emission is increased by 50% conducted in T7
687 experiment, the O₃ concentration (Fig. 13b) changes little in both urban and suburb area in
688 Shanghai which is different from the previous model study of T5 the T3 when O₃ production is
689 under VOC-limited condition. It is suggested that the O₃ concentration after 2020 is not sensitive
690 to the variation of VOCs concentration because the continuous reduction of NO_x emission keeps
691 in promoting the O₃ production to transfer into NO_x-limited regime. Thus further reduction of
692 NO_x tends to decrease the O₃ concentration in Shanghai.

693

694 **Figure 13.** The difference of O₃ concentration (ppbv) between (a) T6 and T4 (T6-T4), (b) T7 and
695 T4 (T7-T4) respectively conducted by WRF-Chem model. The NO_x emissions set in T6 is 20% lower
696 than that in T4 (2020 scenario). The VOCs emission in T7 is 50% higher than that in T4.

697

698 Conclusions

699 O₃ pollution is a serious issue in China. Better understanding the elevated O₃ and its response to
700 emission change is important for Chinese megacities. In this study, we analyze the increasing
701 trend of O₃ concentration by long-term measurements of O₃ and its precursors as well as
702 meteorology in Shanghai combined with the WRF-Chem model. The O₃ production regime

703 response to the emission change in Shanghai during the past ten years is also explored by O₃
704 isopleths plot. In addition, the future O₃ variation and its chemical production in Shanghai are
705 evaluated by WRF-Chem model. The main conclusions are summarized as follows:

706 (1) The daily maximum O₃ concentration measured in Shanghai increased significantly from
707 2006 to 2015 with the rate of 0.808 ppbv yr⁻¹ at downtown site XJH and 1.374 ppbv yr⁻¹ at
708 sub-urban site PD respectively. The observed increasing trend of O₃ is not limited in the urban
709 zones but expanded to the larger scale covering the total Shanghai city. The NO_x and VOCs
710 concentrations presented different variability from O₃ during the same period, in which NO_x
711 concentration decreases significantly at both XJH and PD sites, whereas the VOCs changes very
712 little without evident trend.

713 (2) Because there are minor trends of measured O₃ photolysis, local dispersion and regional
714 transport resulted from meteorology, it is speculated that the significant O₃ increasing trend
715 during 2006 to 2015 in Shanghai is likely attributed to the reduction of NO_x concentration as a
716 result of the strong VOCs-limited regime for O₃ production. The nighttime O₃ is more sensitive to
717 NO_x reduction than that in daytime, because of more O₃ is depressed by NO_x in nighttime. As a
718 result, the observed nighttime O₃ concentration at XJH increases more rapidly than that in
719 daytime response to the NO_x reduction.

720 (3) The WRF-Chem model is utilized to calculate the long term O₃ variations response to
721 emission change. The sensitive experiments illustrate that either reduction of NO_x emission or
722 growth of VOCs emission conducted by WRF-Chem lead to the significant enhancement in O₃
723 concentration in urban zones in 2009 as the baseline, indicating the O₃ production is clearly
724 under VOC-limited regime. The calculated O₃ concentration increases by 1-7 ppbv in urban zones
725 from 2009 to 2015 resulted from 30% reduction of NO_x emission estimated by Shanghai
726 Environmental Monitoring Center. The enhancement of O₃ concentration is significant in urban
727 zones generally more than 4 ppbv, with the maximum elevation of 6-7 ppbv occurred at
728 downtown area, which is consistent with the measurements. The increasing rates of O₃ trend at
729 downtown site XJH and sub-urban site PD are estimated at 1.06 ppbv yr⁻¹ and 0.96 ppbv yr⁻¹ from
730 2009 to 2015 by WRF-Chem, which is close to the observed O₃ growth variability of 1-1.3 ppbv
731 yr⁻¹. This result suggests that the observed increasing trend of O₃ concentration during the past
732 ten years in Shanghai is likely attributed to the reduction of NO_x emission under the VOC-limited
733 condition for O₃ production.

734 (4) The model sensitive study suggests that significant decrease in NO_x concentration
735 combined with the obscure VOCs variation from 2006 to 2015 gradually promotes the O₃
736 chemical production in Shanghai from VOC-limited to NO_x-limited, which is consistent with the O₃
737 isopleths diagram. The O₃ isopleths plot shows that O₃ production is in VOC-limited regime in
738 both downtown site XJH and sub-urban site PD in 2009. With the 30% reduction of NO_x emission
739 from 2009 to 2015 estimated by Shanghai Environmental Monitoring Center, the O₃ production in
740 XJH is still under VOC-limited regime, while the O₃ production moves to the transition regime in
741 PD, suggesting that the O₃ concentration in sub-urban zones is not sensitive to the variation of
742 either NO_x or VOCs concentration.

743 (5) In order to better understand the O₃ control strategy in Shanghai, the future O₃
744 production is estimated by WRF-Chem. The O₃ concentration in Shanghai downtown would keep
745 increasing till 2020 with the 20% reduction of NO_x emission after 2015 estimated by Shanghai
746 Clean Air Action Plan. If the NO_x emission is further decreased by 20% after 2020, The O₃

747 concentration will decrease by 2-3 ppbv in sub-urban zones, and more than 4 ppbv in suburb.
748 While the O₃ concentration in downtown is not sensitive to either NO_x reduction or VOCs
749 enhancement after 2020, indicating the O₃ production in shanghai will transfer to NO_x-limited
750 regimes except downtown where the O₃ production is likely close to the transition regime.
751 Further reduction of NO_x emission after 2020 tend to mitigate the O₃ pollution in Shanghai.

752 (6) There are some uncertainties and limitations existed in the study. First, the
753 inhomogeneity of the NO_x reduction is not considered in the sensitive experiments by lacking of
754 the high resolution emission inventory (e.g. 1 km resolution). Second, the variation of VOCs
755 emission is not taken into account in the model experiments due to the more uncertainties
756 existed in the current VOCs emission inventory. While O₃ production in Shanghai is very sensitive
757 to some VOC species, especially aromatics. Thus the accurate emission inventory of VOCs need to
758 be developed and included in the future study. Third, the same meteorology is used for all
759 WRF-Chem simulations. However the O₃ photolysis, advection, and vertical diffusion are all
760 strongly affected by meteorology. The change of meteorology would be considered and
761 evaluated in the future studies for more deep investigation.

762

763 **Data availability.** The data used in this paper can be provided upon request from Jianming Xu
764 (metxujm@163.com).

765

766 **Author contributions.** XT came up with the original idea of investigating the impact of emission
767 change on long term O₃ variations by. XT and JX designed the analysis method. JX conducted the
768 analysis. WG, YL and QF provided the observational data and helped in discussion.

769

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

771

772 **Acknowledgements.** This study was funded by the National Key R&D Program of China (grant
773 2018YFC0213800), the National Natural Science Foundation of China (91644223, 41430424,
774 41730108 and 41801367).

775

776 **References**

- 777 Binkowski, F. S. and Roselle, S. J.: Models-3 community multi scale air quality (CMAQ) model
778 aerosol component – 1. Model description, *Journal of Geophysical Research*, 108 (D6),
779 4183, doi:10.1029/2001jd001409, 2003.
- 780 Brasseur, G. P., Orlando, J. J., and Tyndall, G. S.: *Atmospheric chemistry and global change*, Oxford
781 University Press, Cambridge, USA, 654 pp., 1999.
- 782 Cai, C. J., Geng, F. H., Tie, X. X., Yu, Q., and An J. L.: Characteristics and source apportionment of
783 VOCs measured in Shanghai, China, *Atmospheric Environment*, 44, 5005-5014, 2010.
- 784 Chen, F. and Dudhia, J.: Coupling an advanced land surface hydrology model with the Penn
785 State-NCAR MM5 modeling system, Part I: Model implementation and sensitivity, *Monthly*
786 *Weather Review*, 129, 569–585, 2001.
- 787 Dudhia, J.: Numerical study of convection observed during the winter monsoon experiment using
788 a mesoscale two-dimensional model, *Journal of the Atmospheric Sciences*, 46, 3077–3107,
789 1989.
- 790 Emmons, L. K., Walters, S., Hess, P. G., Lamarque, J. F., Pfister, G. G., Fillmore, D., Granier, C.,
791 Aguenther, A., Kinnison, D., Laepple, T., Orlando, J., Tie, X., Tyndall, G., Wiedinmyer, C.,
792 Baughcum, S. L., and Kloster, S.: Description and evaluation of the model for ozone and
793 related chemical tracers, version4 (MOZART-4), *Geoscientific Model Development*, 3, 43–67,
794 2010.
- 795 Gao, W., Tie, X. X., Xu, J. M., Huang, R. J., Mao, X. Q., Zhou, G. Q., and Chang, L. Y.: Long-term
796 trend of O₃ in a mega City (Shanghai), China: Characteristics, causes, and interactions with
797 precursors, *Science of the Total Environment*, 603-604, 425-433, 2017.
- 798 Geng, F. H., Zhao, C. S., Tang, X., Lu, G. L., and Tie, X. X.: Analysis of ozone and VOCs measured in
799 Shanghai: a case study, *Atmospheric Environment*, 41, 989–1001, 2007.
- 800 Geng, F. H., Tie, X., Xu, J., Zhou, G., Peng, L., Gao, W., Tang, X., Zhao, C.: Characterizations of ozone,
801 NO_x, and VOCs measured in Shanghai, China, *Atmospheric Environment*, 42, 6873–6883,
802 2008a.
- 803 Geng, F. H., Zhang, Q., Tie, X., Huang, M., Ma, X., Deng, Z., Quan, J., and Zhao, C.: Aircraft
804 measurements of O₃, NO_x, CO, VOCs, and SO₂ in the Yangtze River Delta region, *Atmospheric*
805 *Environment*, 43, 584–593, 2008b.
- 806 Geng, F. H., Tie, X., Guenther, A., Li, G., Cao, J., and Harley, P.: Effect of isoprene emissions from
807 major forests on ozone formation in the city of Shanghai, China, *Atmospheric Chemistry and*
808 *Physics*, 11, 10449–10459, 2011.
- 809 Geng, F. H., Mao, X. Q., Zhou, M. Y., Zhong, S. Y., and Lenschow, D.: Multi-year ozone
810 concentration and its spectra in Shanghai, China, *Science of the Total Environment*, 521-522,
811 135-143, 2015.
- 812 Gery, M. W., and Crouse, R. R.: *User’s Guide for Executing OZIPR*, Atmospheric Research and
813 Exposure Assessment Lab., Office of Research and Development, U.S. EPA, Research Triangle
814 Park, N. C., <http://www.epa.gov/scram001/models/other/oziprdme.txt>, 2002.
- 815 Grell, G. A., Peckham, S. E., Schmitz, R., McKeen, S. A., Frost, G., Skamarock, W. C., and Eder, B.:
816 Fully coupled “online” chemistry within the WRF model, *Atmospheric Environment*, 39,
817 6957–6975, 2005.
- 818 Guenther, A., Karl, T., Harley, P., Wiedinmyer, C., Palmer, P. I., and Geron, C.: Estimates of global
819 terrestrial isoprene emissions using MEGAN (Model of Emissions of Gases and Aerosols from

820 Nature), *Atmospheric Chemistry and Physics*, 6, 3181–3210, 2006.

821 Hong, S. Y. and Lim, J. O. J.: The WRF Single-Moment 6-Class Microphysics Scheme (WSM6),
822 *Journal of the Korean Meteorological Society*, 42, 129–151, 2006.

823 Hu, X. M., Klein, P. M., and Xue, M.: Evaluation of the updated YSU planetary boundary layer
824 scheme within WRF for wind resource and air quality assessments, *Journal of Geophysical*
825 *Research-Atmospheres*, 118, 10490-10505, 2013.

826 Hu, X. M., Xue, M., Klein, P. M., Illston, B. G., and Chen, S.: Analysis of Urban Effects in Oklahoma
827 City using a Dense Surface Observing Network, *Journal of Applied Meteorology and*
828 *Climatology*, 55, 723-741, 2016.

829 Lei, W., Foy, B. de, Zavala, M., Volkamer, R., and Molina, L. T.: Characterizing ozone production in
830 the Mexico City Metropolitan Area: a case study using a chemical transport model,
831 *Atmospheric Chemistry and Physics*, 7, 1347-1366, 2007.

832 Li, G., Lei, W., Zavala, M., Volkamer, R., Dusanter, S., Stevens, P., and Molina, L. T.: Impacts of
833 HONO sources on the photochemistry in Mexico City during the MCMA-2006/MILAGO
834 Campaign, *Atmospheric Chemistry and Physics*, 10, 6551–6567, 2010.

835 Li, G., Bei, N., Tie, X., and Molina, L. T.: Aerosol effects on the photochemistry in Mexico City
836 during MCMA-2006/MILAGRO campaign, *Atmospheric Chemistry and Physics*, 11,
837 5169–5182, 2011.

838 Li, K., Jacob, D. J., Liao, H., Shen, L., Zhang, Q., and Bates, K. H.: Anthropogenic drivers of
839 2013-2017 trends in summer surface ozone in China, *PANS*, 116, 2, 422-427, 2019.

840 Lin, X., Trainer, M., and Liu, S. C.: On the nonlinearity of the tropospheric ozone production,
841 *Journal of Geophysical Research Atmospheres*, 93 (D12), 15879–15888, 1988.

842 Lin, Y. F., Wang, Q., Fu, Q. Y., Duan, Y. S., Xu, J. M., Liu, Q. Z., Li, F., and Huang, K.: Temporal-spatial
843 characteristics and impact factors of ozone pollution in Shanghai, *Environmental Monitoring*
844 *in China (in Chinese)*, 33, 4, 60-67, 2017.

845 Lin, Y. L., Farley, R. D., and Orville, H. D.: Bulk parameterization of the snowfield in a cloud model,
846 *Journal of Climate and Applied Meteorology*, 22, 1065–1092, 1983.

847 Lu, X., Hong, J., Zhang, L., Cooper, O., Schultz, M., Xu, X., Wang, T., Gao, M., Zhao, Y., and Zhang, Y.:
848 Severe surface ozone pollution in China: A global perspective, *Environmental Science*
849 *&Technology Letters*, 5, 487–494, 2018.

850 Monks, P. S., Archibald, A. T., Colette, A., Cooper, O., Coyle, M., Derwent, R., Fowler, D., Granier, C.,
851 Law, K. S., Mills, G. E., Stevenson, D. S., Tarasova, O., Thouret, V., von Schneidmesser, E.,
852 Sommariva, R., Wild, O., and Williams, M. L.: Tropospheric ozone and its precursors from the
853 urban to the global scale from air quality to short-lived climate forcer, *Atmospheric*
854 *Chemistry and Physics*, 15, 8889–8973, 2015.

855 Ma, Z., Xu, J., Quan, W., Zhang, Z., Lin, W., and Xu, X.: Significant increase of surface ozone at a
856 rural site, north of eastern China, *Atmospheric Chemistry and Physics*, 16, 3969–3977, 2016.

857 Nenes, A., Pandis, S. N., and Pilinis, C.: ISORROPIA: A new thermodynamic equilibrium model for
858 multiphase multicomponent inorganic aerosols, *Aquatic Geochemistry*, 4, 123–152, 1998.

859 Ran, L., Zhao, C., Geng, F., Tie, X., Tang, X., Peng, L., Zhou, G., Yu, Q., Xu, J., and Guenther, A.:
860 Ozone photochemical production in urban Shanghai, China: analysis based on ground level
861 observations, *Journal of Geophysical Research Atmospheres*, 114, D15301, 2009.

862 Sillman, S.: The use of NO_y, H₂O₂, and HNO₃ as indicators for ozone-NO_x-hydrocarbon sensitivity
863 in urban locations, *Journal of Geophysical Research Atmospheres*, 100, 14175–14188, 1995.

864 Sillman, S.: The relation between ozone, NO_x and hydrocarbons in urban and polluted rural
865 environments, *Atmospheric Environment*, 33, 1821–1845, 1999.

866 Song, J., Lei, W., Bai, N., Zavala, M., de Foy, B., Volkamer, R., Cardenas, B., Zheng, J., Zhang, R., and
867 Molina L. T.: Ozone response to emission changes: a modeling study during the
868 MCMA-2006/MILAGRO Campaign, *Atmospheric Chemistry and Physics*, 10, 3827-3846, 2010

869 Stockwell, W. R., Middleton, P., Chang, J. S., and Tang, X.: The second generation regional acid
870 deposition model chemical mechanism for regional air quality modeling, *Journal of*
871 *Geophysical Research Atmospheres*, 95, 16343–16367, 1990.

872 Sun, L., Xue, L., Wang, T., Gao, J., Ding, A., Copper, O., Lin, M., Xu, P., Wang, Z., Wang, X., Wen, L.,
873 Zhu, Y., Chen, T., Yang, L., Wang, Y., Chen, J., and Wang, W.: Significant increase of
874 summertime ozone at Mount Tai in central eastern China, *Atmospheric Chemistry and*
875 *Physics*, 16, 10637–10650, 2016.

876 Tai, A. P. K., Martin, M. V., and Heald, C. L.: Threat to future global food security from climate
877 change and ozone air pollution, *Nature Climate Chang*, 4, 817–821, 2014.

878 Tang, G., Li, X., Wang, Y., Xin, J., and Ren, X.: Surface ozone trend details and interpretations in
879 Beijing, 2001–2006, *Atmospheric Chemistry and Physics*, 9, 8813–8823, 2009.

880 Tang, W. Y., Zhao, C. S., Geng, F. H., Peng, L., Zhou, G. Q., Gao, W., Xu, J. M., and Tie, X. X.: Study of
881 ozone "weekend effect" in Shanghai, *Science in China Series D: Earth Sciences*, 51, 9,
882 1354-1360, 2008.

883 Tie, X., Brasseur, G., Emmons, L., Horowitz, I., and Kinnison, D.: Effects of aerosols on
884 tropospheric oxidants: a global model study, *Journal of Geophysical Research Atmospheres*,
885 106, 22931–22964, 2001.

886 Tie, X., Madronich, S., Walters, S., Zhang, R. Y., Rasch, P., and Collins, W.: Effect of clouds on
887 photolysis and oxidants in the troposphere, *Journal of Geophysical Research Atmospheres*,
888 108, 4642, doi:10.1029/2003jd003659, 2003.

889 Tie, X., Madronich, S., Li, G., Ying, Z., Zhang, R., Garcia, A., Taylor, J., and Liu, Y.: Characterizations
890 of chemical oxidants in Mexico City: A regional chemical dynamical model (WRF-Chem)
891 study, *Atmospheric Environment*, 41, 1989-2008, 2007.

892 Tie, X., Geng, F. H., Peng, L., Gao, W., Zhao, C. S.: Measurement and modeling of O₃ variability in
893 Shanghai, China: application of the WRF-Chem model, *Atmospheric Environment*, 43,
894 4289–4302, 2009a.

895 Tie, X., Madronich, S., Li, G., Ying, Z., Weinheimer, A., Apel, E., and Campos, T.: Simulation of
896 Mexico City plumes during the MIRAGE-Mex field campaign using the WRF-Chem model,
897 *Atmospheric Chemistry and Physics*, 9, 4621-4638, 2009b.

898 Tie, X., Geng, F., Guenther, A., Cao, J., Greenberg, J., Zhang, R., Apel, E., Li, G., Weinheimer, A.,
899 Chen, J., and Cai, C.: Megacity impacts on regional ozone formation: observations and
900 WRF-Chem modeling for the MIRAGE-Shanghai field campaign, *Atmospheric Chemistry and*
901 *Physics*, 13, 5655–5669, doi:10.5194/acp-13-5655-2013, 2013.

902 Wang, H. J., and Chen, H. P.: Understanding the recent trend of haze pollution in eastern China:
903 roles of climate change, *Atmospheric Chemistry and Physics*, 16, 4205–4211, 2016

904 Wang, T., Xue, L., Brimblecombe, P., Lam Y. F., Li, L., and Zhang, L.: Ozone pollution in China: A
905 review of concentrations, meteorological influences, chemical precursors, and effects,
906 *Science of Total Environment*, 575, 1582-1596, 2017.

907 Wesely, M. L.: Parameterization of surface resistances to gaseous dry deposition in regional-scale

908 numerical models, *Atmospheric Environment*, 23, 1293–1304, 1989.

909 Xu, J. M., Yan, F. X., Xie, Y., Wang, F. Y., Wu, J. B., and Fu, Q. Y.: Impact of meteorological conditions
910 on a nine-day particulate matter pollution event observed in December 2013, Shanghai,
911 China, *Particuology*, 20, 69–79, 2015.

912 Xu, J. M., Chang, L. Y., Yan, F. X., and He, J. H.: Role of climate anomalies on decadal variation in
913 the occurrence of wintertime haze in the Yangtze River Delta, China, *Science of the Total
914 Environment*, 599-600, 918-925, 2017.

915 Ying, Z. M., Tie, X., and Li, G. H.: Sensitivity of ozone concentrations to diurnal variations of
916 surface emissions in Mexico City: A WRF/Chem modeling study, *Atmospheric Environment*,
917 43, 851–859, 2009.

918 Zhang, Q., Streets, D. G., Carmichael, G. R., He, K. B., Huo, H., Kannari, A., Klimont, Z., Park, I. S.,
919 Reddy, S., Fu, J. S., Chen, D., Duan, L., Lei, Y., Wang, L. T., Yao, Z. L.: Asian emissions in 2006
920 for the NASA INTEX-B mission, *Atmospheric Chemistry and Physics*, 9, 5131-5153, 2009.

921 Zhao, S., Li, J. P., Sun, C.: Decadal variability in the occurrence of wintertime haze in central
922 eastern China tied to the Pacific decadal oscillation, *Scientific Reports*, 6, 27424, 2016.

923 Zheng, B., Tong, D., Li, M., Liu, F., Hong, C., Geng, G., Li, H., Li, X., Peng, L., Qi, J., Yan, L., Zhang, Y.,
924 Zhao, H., Zheng, Y., He, K., and Zhang, Q.: Trends in China's anthropogenic emissions since
925 2010 as the consequence of clean air actions, *Atmospheric Chemistry and Physics*, 18,
926 14095-14111, <https://doi.org/10.5194/acp-18-14095-2018>, 2018.

927 Zhou, G. Q., Xu, J. M., Xie, Y., Chang, L. Y., and Gao, W.: Numerical air quality forecasting over
928 eastern China: An operational application of WRF-Chem, *Atmospheric Environment*, 153,
929 94-108, 2017.

930

931 **Table 1.** Statistical analysis on O₃ simulation in September of 2009 by WRF-Chem model
 932 compared with measurements of 5 sites (XJH, JS, DT, PD, BS) over Shanghai. MO and MM
 933 represent the mean value (unit: ppbv) of observed and modeled O₃ concentration respectively.
 934 RMSE and R are the Root Mean Square Error and correlated coefficient respectively calculated
 935 between modeled and measured O₃ concentration.

	MO	MM	RMSE	R (99% confidence)
		ppbv		\
XJH	21.6	23.0	7.2	0.78
JS	34.6	30.0	10.3	0.64
DT	47.3	40.3	12.0	0.61
PD	33.5	34.9	8.6	0.74
BS	31.7	31.2	9.3	0.67

937

938

939 **Table 2.** Statistical analysis on NO_x simulation in September of 2009 by WRF-Chem model
 940 compared with measurements of 5 sites (XJH, JS, DT, PD, BS) over Shanghai. MO and MM
 941 represent the mean value (unit: ppbv) of observed and modeled NO_x concentration respectively.
 942 RMSE and R are the Root Mean Square Error and correlated coefficient respectively calculated
 943 between modeled and measured NO_x concentration.

944

	MO	MM	RMSE	R (99% confidence)
		ppbv		\
XJH	32.1	33.7	7.0	0.74
JS	14.9	14.7	7.6	0.61
DT	3.0	1.5	2.3	0.6
PD	20.3	16.8	7.5	0.82
BS	21.6	16.1	9.8	0.8

945

946

947 **Table 3.** Scheme of WRF-Chem sensitivity simulations.

948

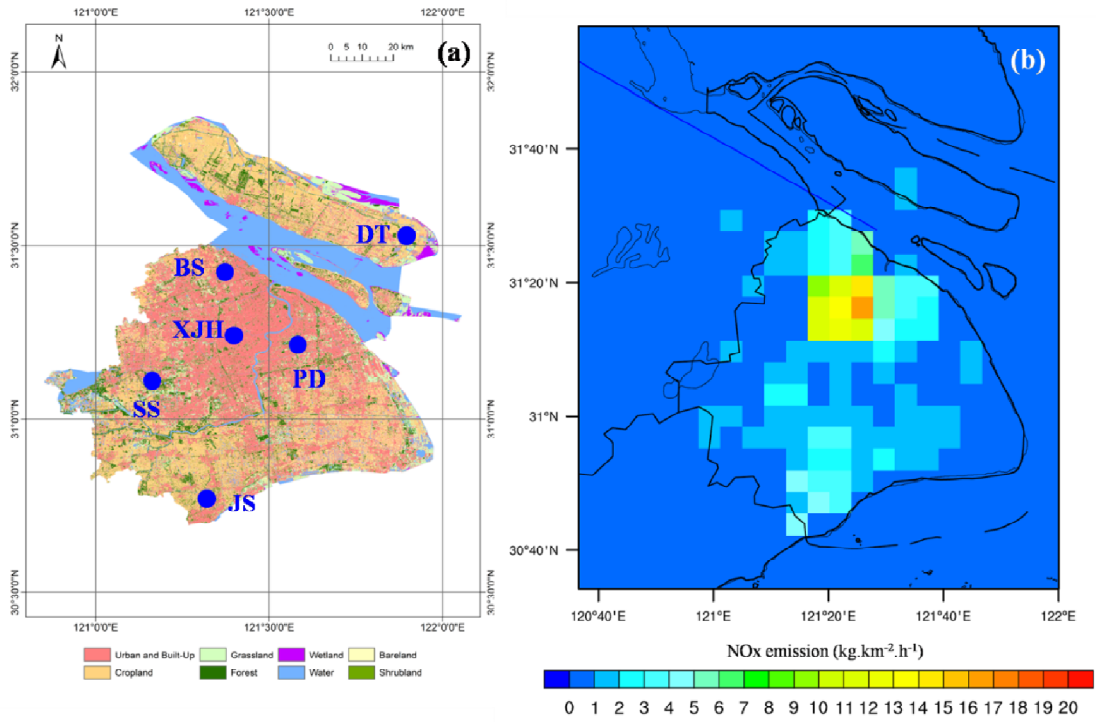
Simulation	NO _x EI	VOCs EI	Meteorology
T1 (Control Run)	2009	2009	September of 2009
T2	2015 (30% reduction)	2009	September of 2009
T3	2009	50% increasing	September of 2009
T4	2020 (50% reduction)	2009	September of 2009
T5	2015	50% increasing	September of 2009
T6	70% reduction	2009	September of 2009
T7	2020 (50% reduction)	50% increasing	September of 2009

949

950

951

952



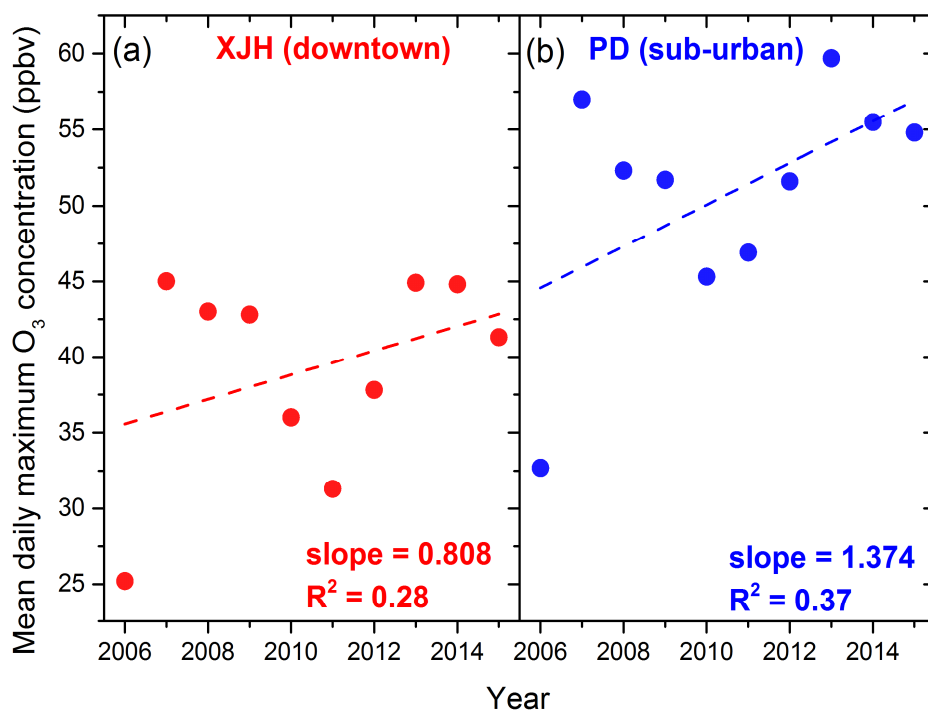
953

954 **Figure1** (a) The distribution of land-use category in Shanghai. The blue dots denote the locations

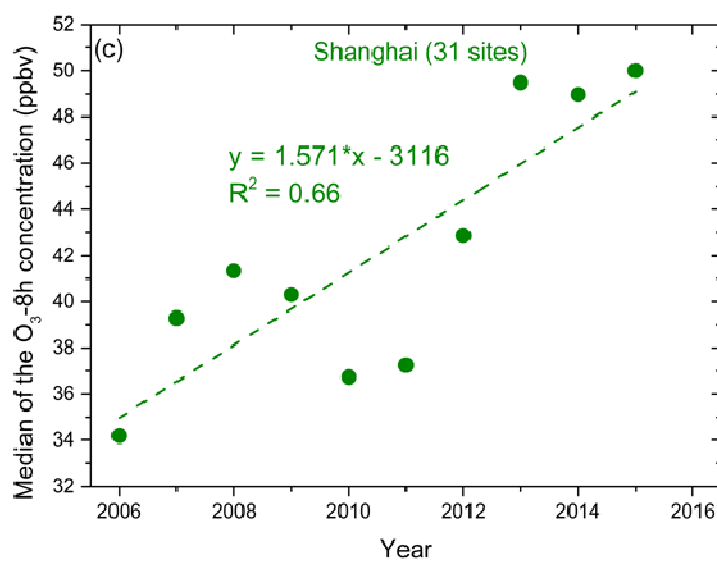
955

of 6 sites (XJH, BS, PD, SS, JS, DT). (b) The NO_x emission of 2009 scenario in Shanghai.

956

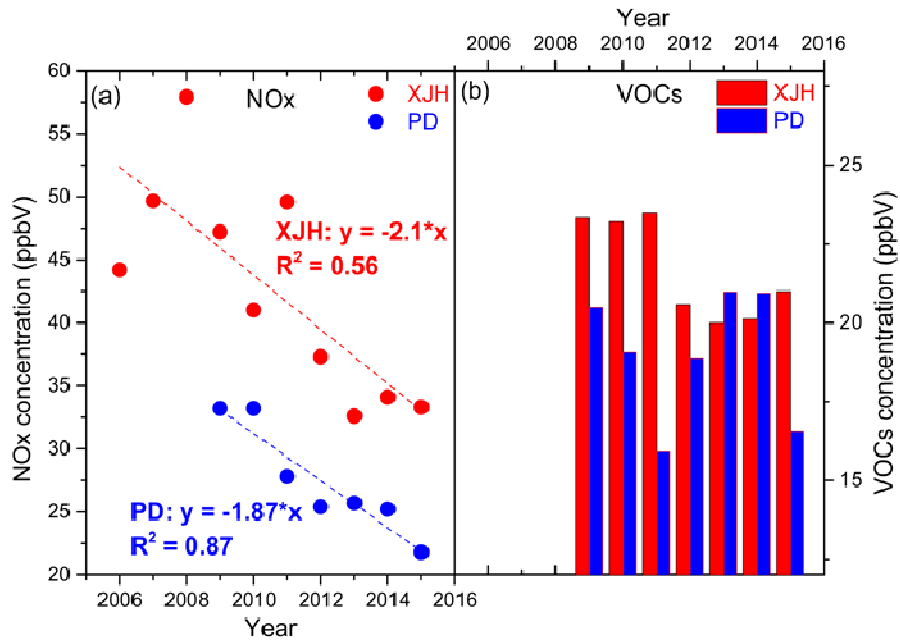


957



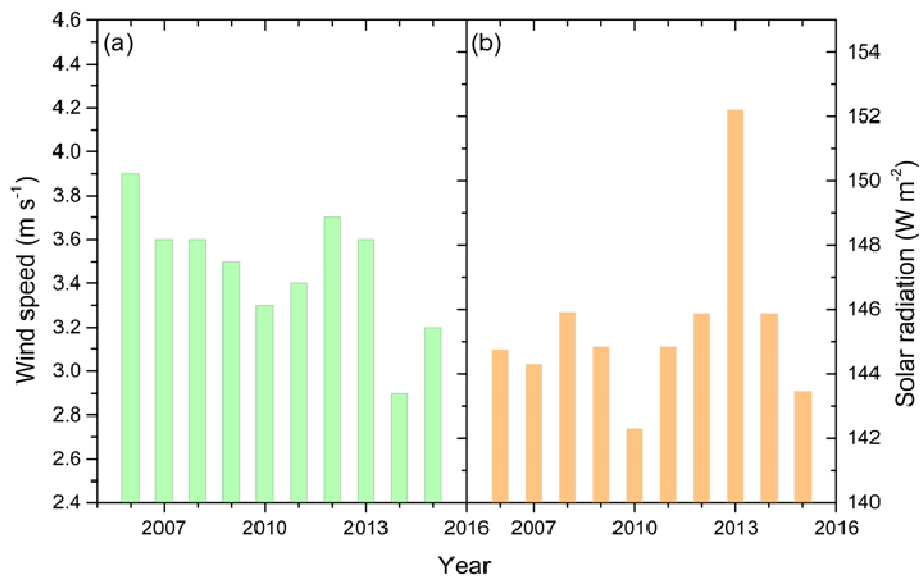
958

959 **Figure 2.** The annual variation of daily maximum O₃ concentration (ppbv) from 2006 to 2015 at (a)
 960 downtown site XJH and (b) sub-urban site PD, both presenting the significant increasing trends
 961 with 0.808 ppbv yr⁻¹ at XJH and 1.374 ppbv yr⁻¹ at PD. The variation of the median 8-h O₃
 962 concentration (ppbv) from 2006 to 2015 averaged for 31 sites over Shanghai (c), also shows the
 963 increasing variability of 1.571 ppbv yr⁻¹.



964

965 **Figure 3.** The mean annual concentrations (ppbv) of (a) NO_x (dots) and (b) VOCs (bars) from 2006
 966 to 2015 at downtown site XJH and sub-urban site PD respectively. The NO_x concentrations at XJH
 967 and PD both present obvious decreasing trends with 2.1 ppbv yr⁻¹ and 1.87 ppbv yr⁻¹. While the
 968 VOCs concentrations at both sites present no clear inter-annual trends.
 969



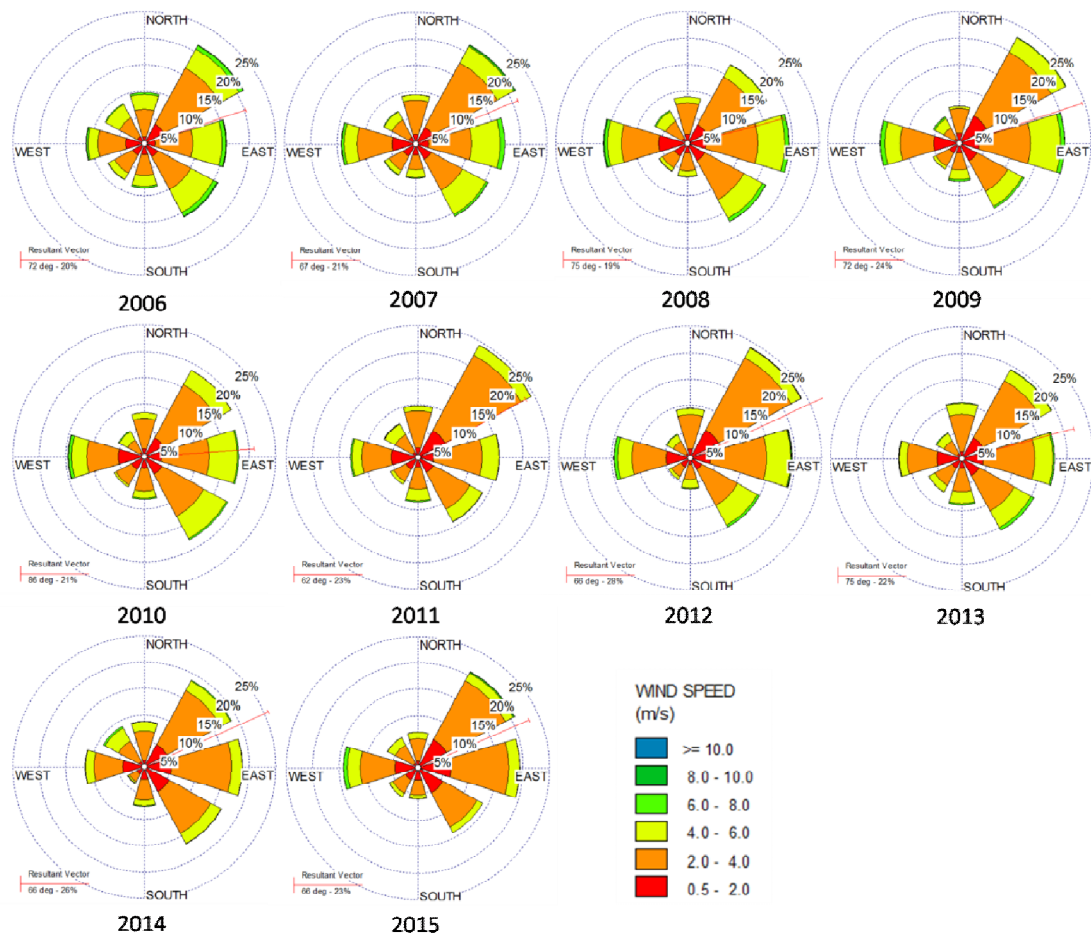
970

971 **Figure 4.** The annual variation of (a) summer wind speed (m s⁻¹) and (b) total solar radiation (W
 972 m⁻²) from 2006 to 2015 in Shanghai. Both wind speed and the solar radiation present weak
 973 inter-annual variations but without significant trends.

974

975

976



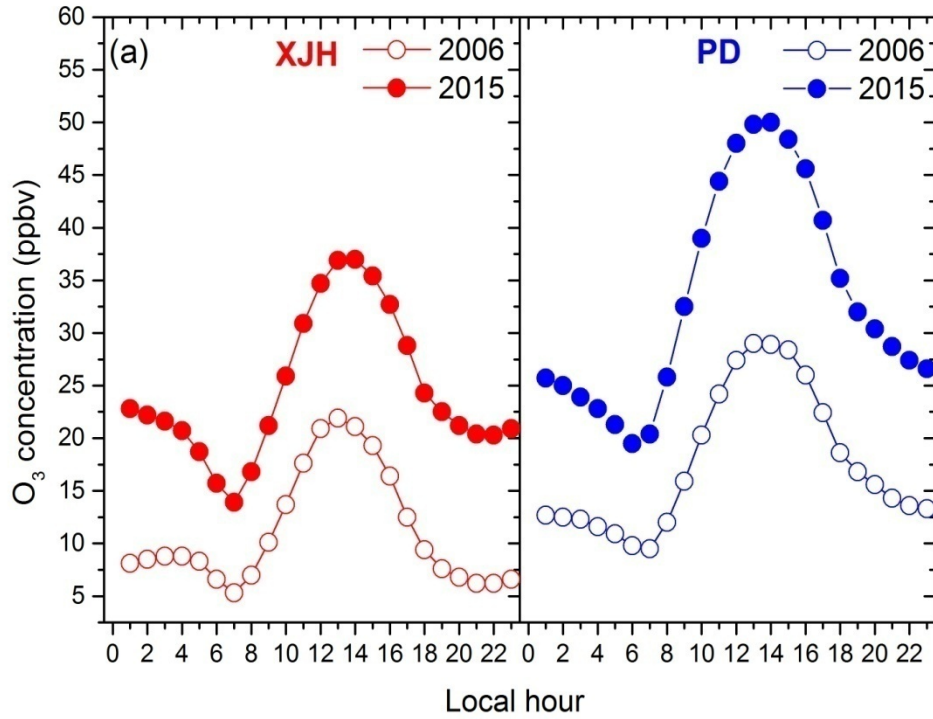
977

978 **Figure 5.** The wind rose of each year from 2006 to 2015 in Shanghai. The red line means the
 979 resultant vector suggesting the dominant wind direction.

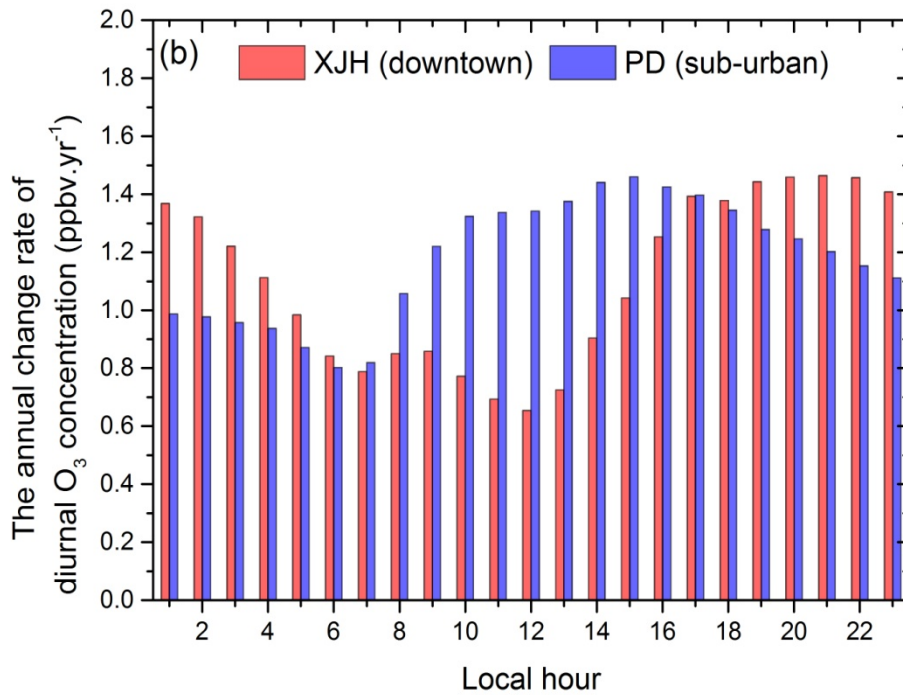
980

981

982

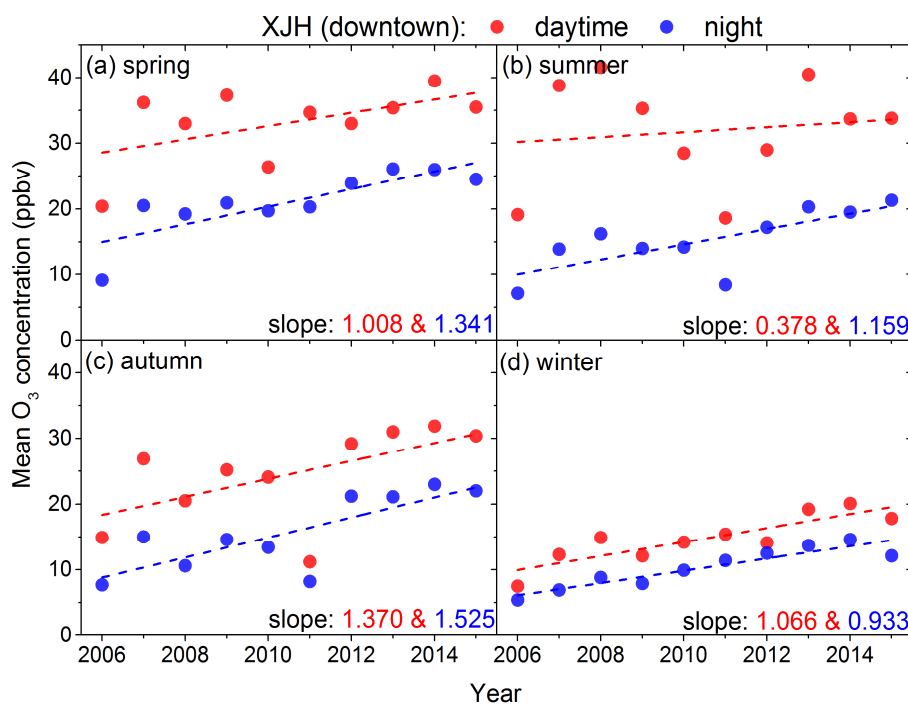


983



984

985 **Figure 6.** (a) The mean diurnal variation of O₃ concentration (ppbv) compared between 2006 and
 986 2015 in XJH (red dots) and PD (blue dots). (b) The annual change rate of diurnal O₃ concentration
 987 (ppbv.yr⁻¹) from 2006 to 2015 at downtown site XJH (red bars) and sub-urban site PD (blue bars).
 988

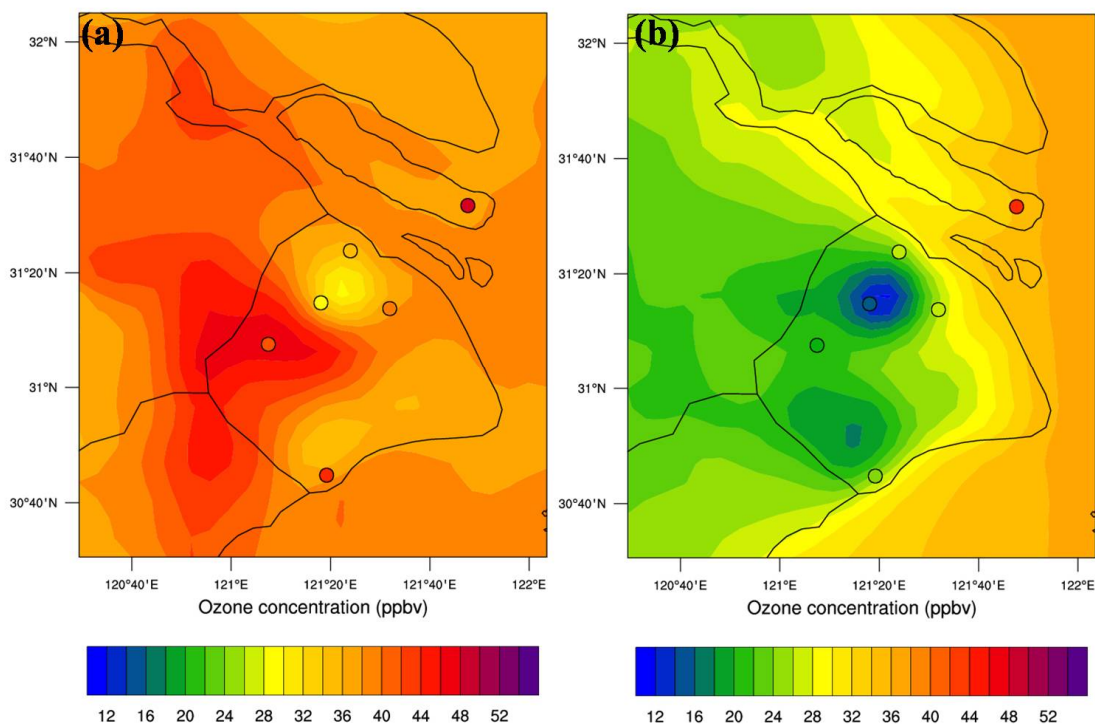


989

990 **Figure 7.** The daytime (8:00-18:00, BJT) and nighttime (19:00-07:00, BJT) O₃ variability from 2006
 991 to 2015 at downtown site XJH in (a) spring, (b) summer, (c) autumn and (d) winter.

992

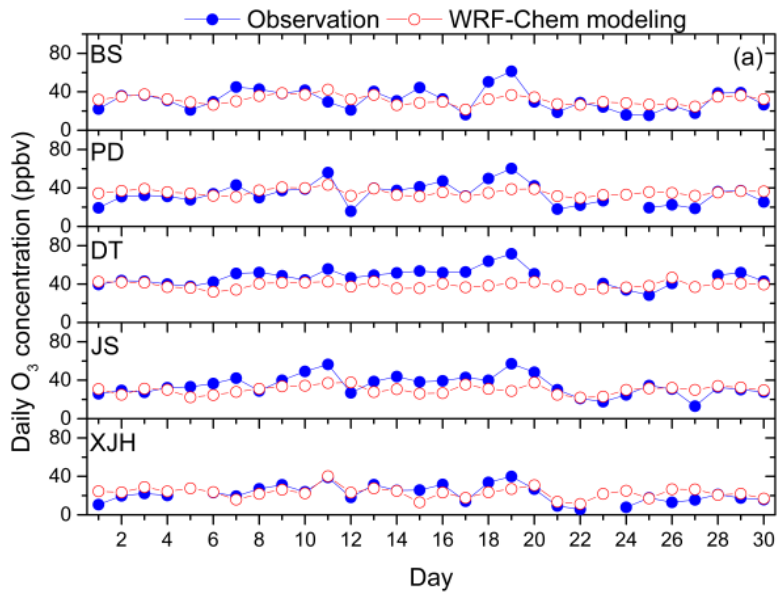
993



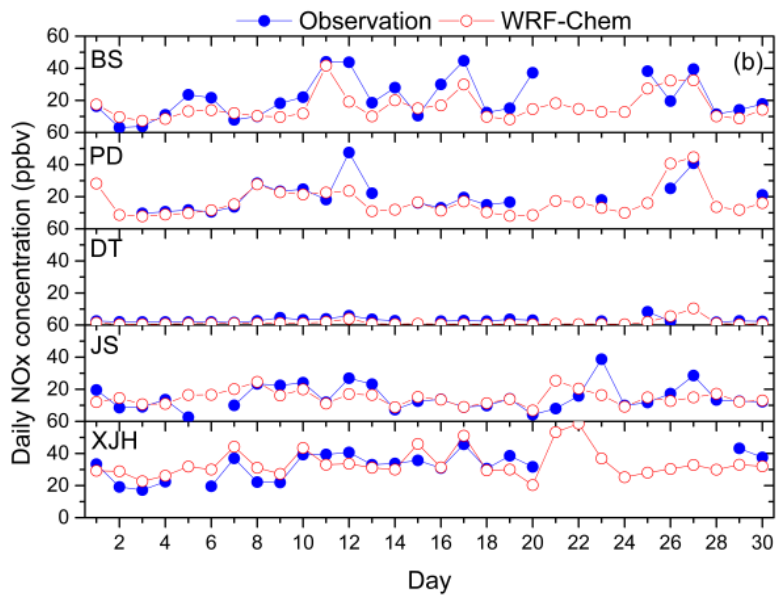
994

995 **Figure 8.** The calculated distribution of (a) daytime and (b) nighttime O₃ concentration by
 996 WRF-Chem (shade) in September of 2009 compared with measurements (circles) of 6 sites over
 997 Shanghai. The minimum O₃ concentrations in daytime and nighttime both occur in urban center.

998

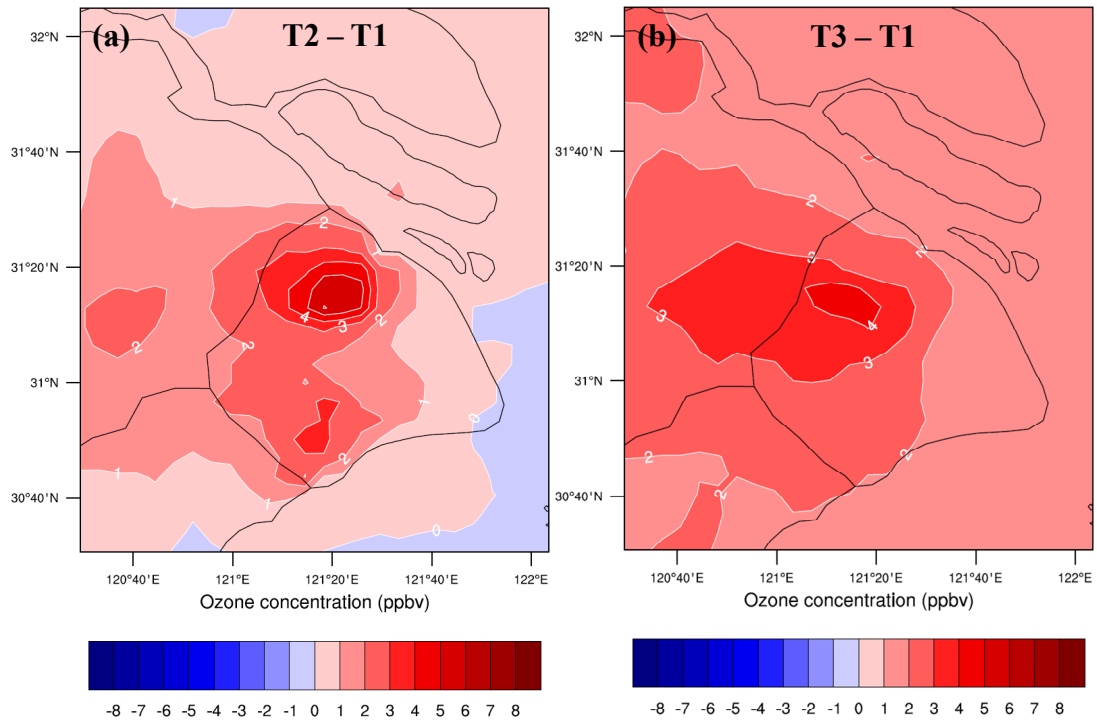


999



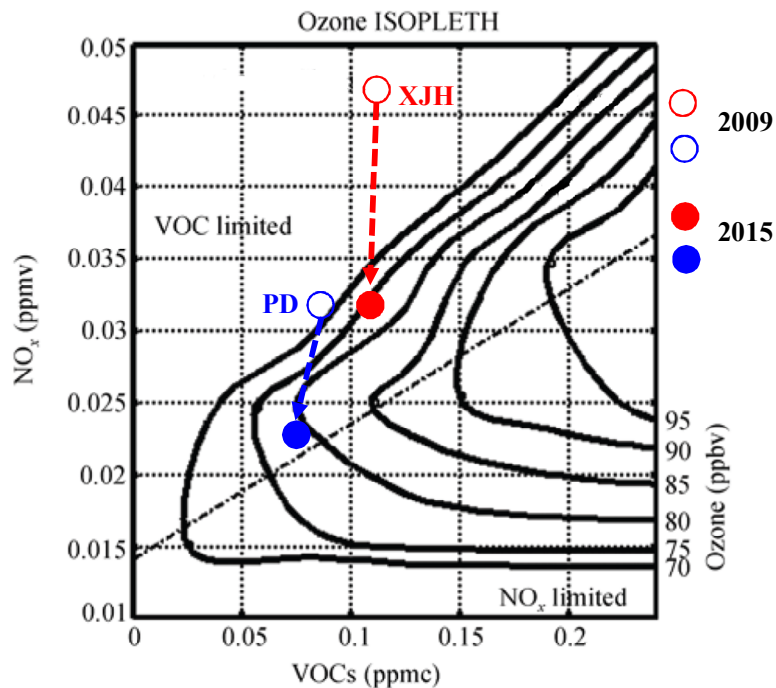
1000

1001 **Figure 9.** The calculated mean daily concentrations (ppbv) of (a) O_3 and (b) NO_x at 5 sites in
 1002 September of 2009 by WRF-Chem (red circles) and compared with measurements (blue circles).
 1003



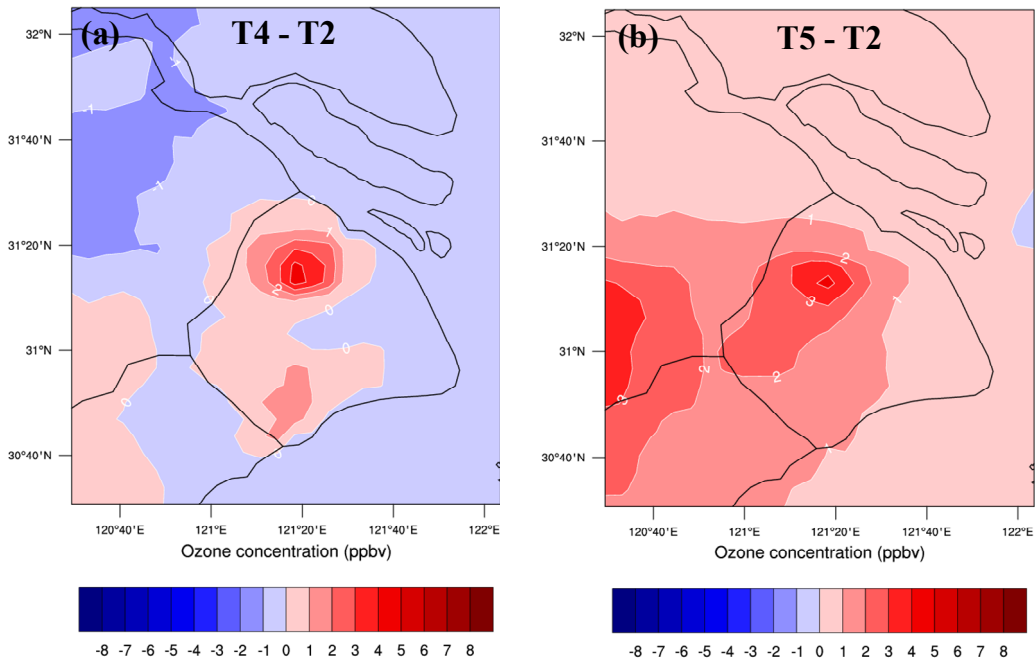
1004

1005 **Figure 10.** The difference of O₃ concentration (ppbv) between (a) T2 and T1 (T2-T1), (b) T3 and
 1006 T1 (T3-T1) respectively conducted by WRF-Chem model. The difference between T2 and T1 lies in the NO_x emissions set in T2 (2015 scenario) is 30% lower than that in T1 (2009 scenario), which is
 1007 estimated by Lin et al. (2017) according to the Shanghai Environment Yearbook. The difference
 1008 between T3 and T1 is dependent on that the VOCs emission in T3 is 50% higher than that in T1.
 1009



1010

1011 **Figure 11.** The O₃ chemical production at downtown site XJH and sub-urban site PD in 2009 and
 1012 2015 depicted by O₃ isopleths diagram. The hollow and solid red circles denote O₃ production
 1013 regime at XJH in 2005 and 2019 respectively. The hollow and solid blue circles denote O₃
 1014 production regime at PD in 2005 and 2019 respectively



1015

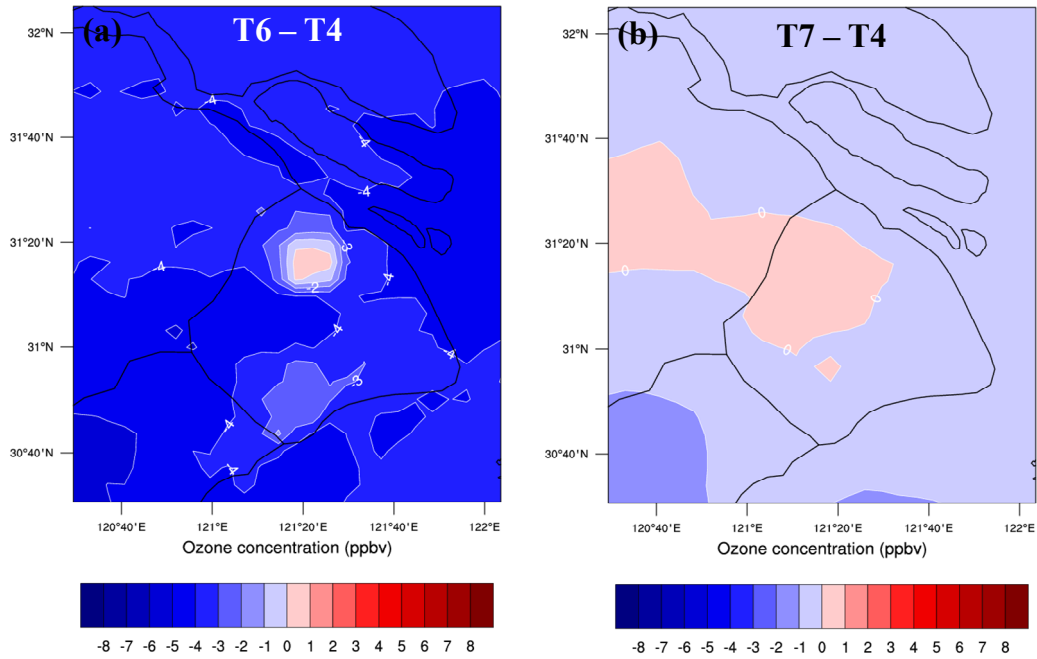
1016 **Figure 12.** The difference of O₃ concentration (ppbv) between (a) T4 and T2 (T4-T2), (b) T5 and
 1017 T2 (T5-T2) respectively conducted by WRF-Chem model. The difference between T4 and T2 is
 1018 that the NO_x emissions set in T4 (2020 scenario) is 20% lower than that in T2 (2015 scenario),
 1019 which is estimated according to the Shanghai Clean Air Action Plan. The difference between T5
 1020 and T2 lies in that the VOCs emission in T5 is 50% higher than that in T2.

1021

1022

1023

1024



1025

1026

1027

1028

Figure 13. The difference of O₃ concentration (ppbv) between (a) T6 and T4 (T6-T4), (b) T7 and T4 (T7-T4) respectively conducted by WRF-Chem model. The NO_x emissions set in T6 is 20% lower than that in T4 (2020 scenario). The VOCs emission in T7 is 50% higher than that in T4.