1	Ozone enhancement due to photo-disassociation of
2	nitrous acid in eastern China
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4 Xuexi Tie ^{1,2} , Xin Long ^{1,5} , Guohui Li ¹ , Shuyu Zhao ¹ , Junji Cao ¹ , Jianmi	Xuexi Tie ^{1,2} , Xin Long ^{1,5} , Guohui Li ¹ , Shuyu Zhao ¹ , Junji Cao ¹ , Jianming Xu ^{3,4}
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8 9	¹ KLACP, SKLLQG, Institute of Earth Environment, Chinese Academy of Sciences, Xi'an 710061, China
10 11 12 13	² Center for Excellence in Urban Atmospheric Environment, Institute of Urban Environment, Chinese Academy of Sciences, Xiamen 361021, China
13 14 15	³ Shanghai Meteorological Service, Shanghai, 200030, China
16 17	⁴ Shanghai Key Laboratory of Meteorology and Health, Shanghai, 200030, China
18 19 20 21	⁵ School of Environment Science and Engineering, Southern University of Science and Technology, Shenzhen 518055, China
22	Correspondence to: XueXi Tie (tiexx@ieecas.cn) or
23	Jianming Xu (<u>metxujm@163.cn</u>)
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28 Abstract

29 PM_{2.5}, a particulate matter with a diameter of 2.5 micrometers or less, is one of the major components of the air pollution in eastern China. In the past few years, China's 30 government made strong efforts to reduce the PM_{2.5} pollutions. However, another 31 32 important pollutant (ozone) becomes an important problem in eastern China. Ozone (O_3) is produced by photochemistry, which requires solar radiation for the formation 33 of O₃. Under heavy PM_{2.5} pollution, the solar radiation is often depressed, and the 34 photochemical production of O₃ is prohibited. This study shows that during late 35 spring and early fall in eastern China, under heavy PM_{2.5} pollutions, there were often 36 strong O₃ photochemical productions, causing a co-occurrence of high PM_{2.5} and O₃ 37 concentrations. This co-occurrence of high PM2.5 and O3 is un-usual and is the main 38 39 focus of this study. Recent measurements show that there were often high HONO surface concentrations in major Chinese mega cities, especially during daytime, with 40 maximum concentrations ranging from 0.5 to 2 ppbv. It is also interesting to note that 41 the high HONO concentrations were occurred during high aerosol concentration 42 43 periods, suggesting that there were additional HONO surface sources in eastern China. Under the high daytime HONO concentrations, HONO can be photo-dissociated to be 44 45 OH radicals, which enhance the photochemical production of O₃. In order to study the above scientific issues, a radiative transfer model (TUV; Tropospheric 46 Ultraviolet-Visible) is used in this study, and a chemical steady state model is 47 48 established to calculate OH radical concentrations. The calculations show that by 49 including the OH production of the photo-dissociated of HONO, the calculated OH concentrations are significantly higher than the values without including this 50 production. For example, by including HONO production, the maximum of OH 51 concentration under the high aerosol condition (AOD=2.5) is similar to the value 52 53 under low aerosol condition (AOD=0.25) in the no-HONO case. This result suggests 54 that even under the high aerosol condition, the chemical oxidizing process for O₃ production can occurred, which explain the co-occurrence of high PM_{2.5} and high O₃ 55 in late spring and early fall seasons in eastern China. However, the O₃ concentrations 56 were not significantly affected by the appearance of HONO in winter. This study 57 58 shows that the seasonal variation of solar radiation plays important roles for controlling the OH production in winter. Because the solar radiation is in a very low 59 level in winter, adding the photolysis of HONO has smaller effect in winter than in 60

61	fall, and OH remains low values by including the HONO production term. This study
62	provides some important scientific highlights to better understand the O ₃ pollutions in
63	eastern China.
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65	Keywords; High PM _{2.5} and O ₃ , eastern China, HONO photolysis
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72 **1. Introduction**

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Currently, China is undergoing a rapid economic development, resulting in a higher 74 demand for energy and greater use of fossil fuels. As a result, the high emissions of 75 pollutants produce heavy pollutions in mega cities of eastern China, such as Beijing 76 and Shanghai. For example, in the city of Shanghai (a largest mega city in China), the 77 urban and economical developments of the city are very rapid. During 1990 to 2015, 78 79 the population increased from 13.3 to 24.1 million. The number of automobiles increased from 0.2 million (1993) to 2.0 million (2011). The rapid growing population 80 81 and energy usage caused a rapid increase in the emissions of pollutants, leading to severe air pollution problems in these mega cities (Zhang et al., 2006; Geng et al., 82 83 2007; Deng et al., 2008).

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85 Measurements, such as satellite observations have revealed much higher aerosol pollution in eastern China than in eastern US (Tie et al., 2006). The high aerosol 86 pollution causes a wide range of environmental consequences. According to a study 87 by Tie et al. (2009a), exposure to extremely high particle concentrations leads to a 88 89 great increase of lung cancer cases. High PM (particular matter) concentrations also significantly reduce the range of visibility in China's mega cities (Deng et al., 2008). 90 91 According to a recent study, the high aerosol pollution causes important effects on the 92 crop (rice and wheat) production in eastern China (Tie et al., 2016).

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In the troposphere, ozone formation is resulted from a complicated chemical process, 94 and requires ozone precursors, such as VOCs (volatile organic carbons) and $NO_X =$ 95 96 $NO + NO_2$ (nitrogen oxides) (Sillman, 1995). As the increase in industrial activity and number of automobiles, the precursors of ozone (O_3) and the global budget of 97 98 oxidization are also significantly increased (Huang et al., 2017; Huang et al., 2018). As a result, O₃ pollution becomes a serous pollution problem in Shanghai and other 99 Chinese mega cities (Geng et al., 2010; Tie 2009b; Tie et al., 2015). The effects on O₃ 100 production rate can be characterized as either NO_X-sensitive or VOC-sensitive 101 conditions. For the city areas, O₃ production is generally VOC-sensitive, while in the 102 103 remote area, O₃ production is generally NOx-sensitive in eastern China (Sillman, 104 1995; Zhang et al., 2003; Lei et al., 2004; Tie et al., 2013). Thus, better understanding

105 the trends of O_3 precursors (VOCs, NO_X) is important to determine the O_3 trends in 106 Shanghai (as well as many large cities in China).

107 In the past few years, China's government made strong efforts to reduce the PM_{2.5} pollutions. However, another important pollutant (O₃) becomes an important problem 108 in eastern China. Several studies regarding the O₃ formation are previously studied in 109 Shanghai. For example, Geng et al. (2007; 2008) study the relationship between O₃ 110 precursors (NOx and VOCs) for the ozone formation in Shanghai. Tie et al. (2009) 111 study the short-term variability of O₃ in Shanghai. Their study suggested that in 112 addition to the ozone precursors, meteorological conditions, such as regional transport, 113 have also strong impacts on the ozone concentrations. During September 2009, a 114 115 major field experiment (the MIRAGE-Shanghai) was conducted in Shanghai, and multiply chemical species were measured during the experiment. The summary of the 116 measurements by Tie et al (2013) suggests that the ozone formation in Shanghai is 117 under VOC-sensitive condition. However, if the emission ration of NOx/VOCs 118 119 reduces to a lower value (0.1-0.2), the ozone formation in Shanghai will switch from VOC-sensitive condition to NOx-sensitive condition. 120

Despite of some progresses have been made for the ozone formation in mega cities in 121 China, it is still lack of study of ozone development in large cities of China. For 122 example, this study shows that during late spring and early fall in eastern China, under 123 heavy PM_{2.5} pollutions, there were often strong O₃ chemical productions, causing the 124 co-occurrence of high PM_{2.5} and O₃ concentrations. Under heavy aerosol condition, 125 126 the solar radiation is depressed, significantly reducing the photochemical production of O₃. This co-occurrence of high PM_{2.5} and O₃ is an unusual and is the focus of this 127 128 study. He and Carmichael (1999) suggest that aerosol particles can enhance the scattering of solar radiation, enhancing the flux density inside the boundary layer. 129 130 Recent measurements also show that there were often high HONO concentrations in major Chinese mega cities, especially during daytime, with maximum concentrations 131 132 ranging from 0.5 to 2 ppbv (Huang et al., 2017). Zhang et al. (2016) suggest that there are several potential HONO sources, including surface emissions, conversion of NO₂ 133 134 at the ocean surface, etc., and adding these sources can improve the calculated HONO concentrations. It is also interesting to note that the high HONO surface 135 concentrations were occurred during high aerosol concentration periods, suggesting 136

that there are additional HONO surface sources in eastern China. Under the high daytime HONO concentrations, HONO can be photo-dissociated to be OH radicals, which enhance the photochemical production of O_3 .

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The paper is organized as follows: in Section 2, we describe the measurement of O_3 and $PM_{2.5}$. In Section 3, we describe the calculation of photo-dissociated rate of HONO and a steady state model for the calculation of OH, and the causes of high O_3 production under the heavy aerosol condition. Section 4 shows a brief conclusion of the results.

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2. Measurements of O₃ and PM_{2.5}

There are long-term measurements in Eastern China by Chinese Environment 149 Protection Agency (CEPA) for monitoring the air quality in China. In eastern China, 150 especially in the capital city of China (Beijing), there are often heavy air pollutions, 151 especially for fine particular matter ($PM_{2.5}$ – the radium of particle being less than 2.5 152 um). Figure 1 shows the measurement sites in Beijing, in which the measured 153 concentrations of PM_{2.5} and O₃ are used to the analysis. In the region, the air 154 pollutions were very heavy, especially in winter (Long et al., 2016; Tie et al., 2017). 155 The previous studies suggested that the both aerosol and O₃ pollutions became the 156 major pollutants in the region (Li et al., 2017). 157

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Figure 2 shows the daily averaged concentrations of PM_{2.5} and O₃ in the Beijing 159 160 region in 2015. The daily averaged concentrations show that there were strong daily and seasonal variations for both the concentrations of PM_{2.5} and O₃. Despite the daily 161 162 variation, the concentrations of PM_{2.5} existed a strong seasonal variation. For example, there were very high concentrations during winter, with maximum of $\sim 300 \ \mu g/m^3$. 163 While in summer, the maximum concentrations reduced to $\sim 150 \,\mu g/m^3$. The seasonal 164 variability of O₃ concentrations were opposite with the PM_{2.5} concentrations, with 165 lower concentrations in winter (< 50 μ g /m³) and higher concentrations in summer (> 166 150 μ g/m³). These seasonal variations of PM_{2.5} and O₃ have been studied by previous 167 168 studies (Tie and Cao, 2017; Li et al., 2017). Their results suggest that the winter high PM_{2.5} concentrations were resulted from the combination of both the high emissions 169

(heating season in the Beijing region), and poor meteorological ventilation conditions,
such as lower PBL (Planetary Boundary Layer) height (Quan et al., 2013; Tie et al.
2015). According to the photochemical theory of O₃ formation, the summer high and
winter low O₃ concentrations are mainly due to seasonal variation of the solar
radiation (Seinfeld, J. H. and Pandis, 2006).

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The heavy aerosol concentrations play important roles to reduce solar radiation, 176 causing the reduction of O₃ formation. (Bian et al., 2007). As we show in Fig. 3 177 (upper panel), during wintertime, the O₃ concentrations were strong anti-correlated 178 179 with the $PM_{2.5}$ concentrations, suggesting that the reduction of solar radiation by aerosol particles have important impact on the reduction of O₃ concentrations. Figure 180 3 (upper panel) also shows that the relationship between O_3 and $PM_{2.5}$ was not 181 linearly related. For example, when the concentrations of PM_{2.5} were less than 100 182 μ g/m³, O₃ concentrations rapidly decreased with the increase of PM_{2.5} concentrations. 183 In contrast, when the concentrations of PM_{2.5} were greater than 100 μ g/m³, O₃ 184 concentrations slowly decreased with the increase of PM2.5 concentrations. This is 185 consistent with the result of Bian et al (2007). 186

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188 It is interesting to note that from late spring to early fall periods, the correlation between PM_{2.5} and O₃ concentrations was positive relationship compared to the 189 negative relationship in winter (see Fig. 3 (lower panel)). This result suggests that O₃ 190 191 production was high during the heavy haze period, despite the solar radiation was greatly depressed. In order to clearly display this unusual event, we illustrate diurnal 192 variations of PM_{2.5} and O₃ and NO₂ during a fall period (from Oct.5 to Oc. 6, 2015). 193 Figure 4 shows that during this period (as a case study), the PM_{2.5} concentrations were 194 very high, ranging from 150 to 320 μ g/m³. Under such high aerosol condition, the 195 solar radiation should be significantly reduced, and O₃ photochemical production 196 would be reduced. However, the diurnal variation of O₃ was unexpectedly strong, 197 with high noontime concentration of >220 μ g/m³ and very low nighttime 198 concentration of ~25 μ g/m³. This strong diurnal variation was due to the 199 photochemical activity, which suggested that during relatively low solar conditions, 200 the photochemical activities of O₃ production was high. According to the theory of the 201 O₃ chemical production, the high O₃ production is related to high oxidant of OH 202 (Seinfeld and Pandis, 2006), which should not be occurred during lower solar 203

radiation. This result brings important issue for air pollution control strategy, because both $PM_{2.5}$ and O_3 are severe air pollutants in eastern China.

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To clearly understand the effect of the high aerosol concentrations on solar radiation, 207 we investigate the meteorological conditions, such as cloud covers, relation humidity 208 (RH), and solar radiation during the period of the case study (see Figs. 5 and 6). 209 Figure 5 shows that the cloud condition was close to the cloud free condition, but 210 there was a very heavy aerosol layer in the Beijing region, suggesting that cloud cover 211 212 played a minor role in the reduction of the solar radiation. The measured RH values (not shown) were generally higher than 60%, with a maximum of 95% during the 213 214 period. As a result, the high aerosol concentrations companied by high RH produced important effects on solar radiation. As shown in Fig. 6, the daytime averaged solar 215 radiation was significantly reduced (about 40% reduction in Oct. 5-6 period compared 216 217 with the value of Oct. 8).

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219 **3. Method**

[OH] = P[HOx]/L[HOx]*

In order to better understand the O_3 chemical production occurred in heavy aerosol condition in eastern China, the possible O_3 production in such condition is discussed. Ozone photochemical production (P[O₃]) is strongly related to the amount of OH radicals (Chameides et al., 1999). According to the traditional theory, the amount of surface OH radicals is proportional to the surface solar radiation, which is represented by

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Where [OH] represents the concentration of hydroxyl radicals ($\#/cm^3$); HOx represents the concentration of HO₂ + OH ($\#/cm^3$); P[HOx] represents the photochemical production of HOx ($\#/cm^3/s$); and L[HOx]* (1/s) represents the photochemical destruction of HOx, which is normalized by the concentrations of OH.

(R-1)

The major process for the photochemical production of P[HOx] is through the O_3 photolysis and follows by the reaction with atmospheric water vapor. It can be expressed as

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$$P[HOx] = J_1[O_3]/(k_1 \times am) \times 2.0 \times k_2[H_2O] = P_1[HOx]$$
 (R-2)

Where J₁ represents the photolysis of $O_3 + hv \rightarrow O^1D$; k₁ represents the reaction rate 240 of $O^1D + am \rightarrow O^3P$; and k₂ represents the reaction rate of $O^1D + H_2O \rightarrow 2OH$. As 241 we can see, this HOx production is proportional to the magnitude of solar radiation 242 (J_1) , and J_1 is the O₃ photolysis with the solar radiation. Figure 7 shows the 243 relationship between the values of J₁ and aerosol concentrations in October at 244 middle-latitude calculated by the TUV model (Madronich and Flocke, 1999). This 245 246 result suggests that under the high aerosol concentrations (AOD = 2.5), the J₁ value is strongly depressed, resulting in significant reduction of OH concentrations and O₃ 247 production. For example, the maximum J_1 value is about 2.7×10^{-5} (1/s) with lower 248 aerosol values (AOD = 0.25). According to the previous study, the surface $PM_{2.5}$ 249 concentrations were generally smaller than 50 μ g/m³ with this AOD value (Tie et al., 250 2017). However, when the AOD value increase to 2.5 (the PM_{2.5} concentrations are 251 generally >100 μ g/m³), the maximum J₁ value rapidly decreases to about 6x10⁻⁶ (1/s), 252 which is about 450% reduction compared to the value with AOD=0.25. This study 253 suggests that under high PM_{2.5} concentrations (>100 μ g/m³), the photochemical 254 255 production of OH (P[HOx]) is rapidly decreased, leading to low OH concentrations, which cannot initiate the high oxidation of O_3 production. As a result, the high O_3 256 production shown in Fig. 4 cannot be explained. Other sources for O₃ oxidation are 257 needed to explain this result. 258

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Recent studies show that the HONO concentrations are high in eastern China (Huang 260 et al., 2017). Because under high solar radiation, the photolysis rate of HONO is very 261 high, resulting in very low HONO concentrations in daytime (Seinfeld and Pandis, 262 2006). These measured high HONO concentrations are explained by their studies. 263 One of the explanations is that there are high surface HONO sources during daytime, 264 which produces high HONO concentrations (Huang et al., 2017). Zhang et al. (2016) 265 suggest that there are several potential HONO sources, including surface emissions, 266 conversion of NO₂ at the ocean surface, etc. Zhang et al. (2016) parameterized these 267 potential HONO sources in the WRF-Chem model, and the calculated HONO 268 concentrations are increased in the WRF-Chem model. 269

The version of the WRF-Chem model is based on the version developed by Grell et al. 271 (2015), and is improved mainly by Tie et al. (2017) and Li et al. (2011). The chemical 272 mechanism chosen in this version of WRF-Chem is the RADM2 (Regional Acid 273 Deposition Model, version 2) gas-phase chemical mechanism. For the calculation of 274 HONO, only the gas-phase chemistry of OH+NO is included to calculate HONO 275 276 concentrations. As shown in Fig. 8, the calculated HONO concentrations are significantly smaller than the measured HONO values in eastern China, suggesting 277 that in addition to the gas-reaction, there are missing HONO sources (surface sources 278 279 or others). Because these missing sources are not fully understood and large 280 uncertainty is remained, in the following calculation, we compare the OH concentrations due to both calculated HONO (without the missing sources) and the 281 282 measured HONO concentrations to illustrate the importance of these missing sources for the production of OH radicals and to suggest that further study to better 283 284 understand the missing sources is an urgent scientific issue.

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286 Figure 8 shows the measured HONO concentrations in three large cities in China (Shanghai, Xi'an, and Beijing) during fall and winter. It also shows the corresponding 287 288 PM_{2.5} and O₃ in the 3 cities (i.e., Fig. 8a for Beijing, Fig. 8b for Shanghai, and Fig. 8c 289 for Xian). It shows that the measured HONO concentrations were high, ranging from 290 sub-ppbv to a few ppbv, with higher values during morning, and lower values in daytime. The co-occurrences of high PM_{2.5} and O₃ happened in the 3 cities. As a 291 292 result, we think that the high HONO is a common event in large cities in eastern 293 China, especially in daytime. This high HONO is also measured by previous studies (Zhang et al. 2016; Huang et al. 2017). In this study, we make an assumption that the 294 co-occurrence between O₃ and PM_{2.5} occurred under high HONO concentrations. We 295 296 note that using this assumption may result in some uncertainties in estimating the effect of HONO on OH. For example, using the measured HONO in Xi'an and 297 298 Beijing could produce 1-2 times higher OH production by photolysis of HONO than the result by using the data from Shanghai. In this case, we use the measured HONO 299 300 from Shanghai to avoid the over estimate of the HONO effect, which can be considered as a low-limit estimation. 301

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303 It is also interesting to note that the high HONO concentrations were occurred during 304 high aerosol concentration periods. Figure 9 illustrates that when the $PM_{2.5}$ concentrations increased to 70-80 μ g/m³, and the HONO concentrations enhanced to 1.4-18 ppbv during September in Shanghai. This measured high HONO concentrations were significantly higher than the calculated concentrations (shown in Fig. 8), suggesting that some additional sources of HONO are needed. This result is consistent with the HONO measurements in other Chinese cities (Huang et al. 2017).

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The high HONO concentrations in daytime become a significant source of OH radicals. As a result, the OH production rate (P[HOx]) can be written to the following reactions.

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316 $P[HOx] = P_1[HOx] + P_2[HOx]$

 $P_2[HOx] = J_2 \times [HONO]$

317 $= J_1[O_3]/(k_1 \times am) \times 2.0 \times k_2[H_2O] + J_2 \times [HONO] \quad (R-4)$

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Because the chemical lifetime of OH is less than second, OH concentrations can be calculated according to equilibrium of chemical production and chemical loss. With the both OH chemical production processes, the OH concentrations can be calculated by the following equation (Seinfeld and Pandis, 2006).

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P1 + P2 = L1 + L2

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Where P1 and P2 are the major chemical productions, expressed in R-4, and L1 and L2 are the major chemical loss of OH, and represent by

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329 L1: $OH + NO_2 \rightarrow HNO_3$

 $0 L2: HO_2 + HO_2 \rightarrow H_2O_2 + O_2 (R-6)$

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332Under high NOx condition, such as in the large cities in eastern China, NOx333concentrations were often higher to 50 ppbv (as shown in Fig. 4). As a result, the L1334term is larger than L2. The OH concentrations can be approximately expressed as335 $[HO] = {J_1[O_3]/(k_1 \times am) \times 2.0 \times k_2[H_2O] + J_2 \times [HONO]}/{k_3[NO_2]}$ (R-5)

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(R-3)

(R-5)

339 Where k_3 is the reaction coefficient of OH + NO₂ \rightarrow HNO₃.

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341 **4. Result and analysis**

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4.1. OH productions in different HONO conditions

In order to quantify the individual effects of these two OH production terms (P1 and 345 P2) on the OH concentrations, the P1 and P2 are calculated under different daytime 346 HONO conditions (calculated low HONO and measured high HONO concentrations). 347 Figure 10 shows that under the low HONO condition, the P1 is significantly higher 348 than P2, and P2 has only minor contribution to the OH values. For example, the 349 maximum of P1 occurred at 13 pm, with a value of 65×10^6 #/cm³/s. In contrast, the 350 maximum of P2 occurred at 10 am, with a value of 15×10^6 #/cm³/s. However, under 351 high HONO condition, the P2 plays very important roles for the OH production. The 352 maximum of P2 occurred at 11 am, with a value of 350×10^6 #/cm³/s, which is about 353 500% higher than the P1 value. It is important to note that this calculation is based on 354 the high aerosol condition (AOD = 2.5) in September. This result can explain the high 355 356 O₃ chemical production in Fig. 4.

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4.2. OH in different aerosol conditions

360 In order to understand the effect of aerosol conditions, especially high aerosol conditions, on the OH concentrations. Figure 11 shows the OH concentrations with 361 and without HONO production of OH. With including the HONO production (i.e., 362 including P1 and P2), the calculated OH concentrations are significantly higher than 363 without including this production (i.e., only including P1). The both calculated OH 364 concentrations are rapidly changed with different levels of aerosol conditions. For 365 example, without HONO production, the maximum OH concentration is about 366 7.5×10^5 #/cm³ under low aerosol condition (AOD=0.25). In contrast, the maximum 367 OH concentration rapidly reduced to 1.5×10^5 #/cm³ under high aerosol condition 368 (AOD=2.5), and further decreased to 1.0×10^5 #/cm³ with the AOD value of 3.5. In 369 370 contrast, with including HONO production, the OH concentrations significantly increased. Under higher aerosol condition (AOD=2.5), the maximum of OH 371 concentration is about 7.5×10^5 #/cm³, which is the same value under low aerosol 372 condition in the no-HONO case. This result suggests that the measured high O₃ 373

production occurred in the high aerosol condition is likely due to the high HONOconcentrations in Shanghai.

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377 **4.3. Effects of clouds**

Cloud cover can have very important impacts on the photolysis of HONO, which can affect the effect of HONO on the OH radicals. The above calculations are based on the cloud-free condition, with heavy aerosol concentration in the Beijing region. As shown in Fig. 5, during the case study period (Oct 5 to 6, 2015) (see Fig. 4), the weather map shows that the cloud-free condition, with heavy aerosol condition.

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In order to understand the effects of cloud on the photolysis of HONO, we include 385 different cloud covers in the TUV model. The calculated results show in Fig. 12. 386 The results show that the thin cloud (with cloud cover in 2 km and cloud water of 10 387 g/m^3), could reduce the photolysis rate of HONO by about 40%, but the HONO could 388 still remain important effects. However, with dense cloud condition (with cloud 389 covers at 2 and 3 km and cloud water of 50 10 g/m^3), the photolysis rate of HONO 390 could reduce by 9-10 times by the cloud. In this case, adding photolysis rate of 391 HONO cannot produce important effect on OH radicals and the production of O₃. 392

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394 **<u>4.4. OH in winter</u>**

The measurement of O₃ also shows that the concentrations in winter were always low 396 (see Fig. 2), suggesting that the O_3 concentrations were not significantly affected by 397 the appearance of HONO. Figure 13 shows the OH concentrations in September and 398 399 December. It shows that under different aerosol conditions, OH concentrations in December were very low compared with the values in September. Both the calculated 400 OH concentrations include the HONO production term. For example, under the 401 condition of AOD=2.5, the maximum OH is about 7.5×10^5 #/cm³ in September, while 402 it rapidly reduces to 1.5×10^5 #/cm³ in December. Under the condition of AOD=3.5, 403 the maximum OH is still maintaining to a relative high level $(4.5 \times 10^5 \text{ } \#/\text{cm}^3)$ in 404 September. However, the maximum OH values are extremely low in December, with 405 maximum value of 0.5×10^5 #/cm³ in December. Because both the OH chemical 406 productions (P1 and P2) are strongly dependent upon solar radiation (see equation 407 408 R-4), the seasonal variation of solar radiation plays important roles for controlling the

OH production in winter (see Fig. 13). Because the solar radiation is in a very low
level in winter, adding the photolysis of HONO has smaller effect in winter than in
other seasons and OH remains low values by including the HONO production term.

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413 **Summary**

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Currently, China is undergoing a rapid economic development, resulting in a high 415 demand for energy, greater use of fossil fuels. As a result, the high emissions of 416 pollutants produce heavy aerosol pollutions (PM_{2.5}) in eastern China, such as in the 417 mega city of Beijing. The long-term measurements show that in addition to the heavy 418 419 aerosol pollution, the O₃ pollution becomes another major pollutants in the Beijing 420 region. The measured results show that there were very strong seasonal variation in 421 the concentrations of both PM_{2.5} and O₃ in the region. During winter, the seasonal variability of O₃ concentrations were anti-correlated with the PM_{2.5} concentrations. 422 423 However, from late spring to early fal, the correlation between PM_{2.5} and O₃ concentrations was positive compared to the negative in winter. This result suggests 424 425 that during heavy aerosol condition (the solar radiation was depressed), the O_3 chemical production was still high, appearing a co-occurrence of high PM_{2.5} and O₃ in 426 some cases from late spring to early fall. This co-occurrence of high PM2.5 and O3 is 427 the focus of this study. The results are highlighted as follows; 428

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(1) There are high daytime HONO concentrations in major Chinese mega cities, such
as in Beijing and Shanghai. It is also interesting to note that the high HONO
concentrations were occurred during high aerosol concentration periods. Under
the high daytime HONO concentrations, HONO can be photo-dissociated to be
OH radicals, and becomes an important process to produce OH.

(2) With including the OH production of measured HONO concentrations, the 435 calculated OH concentrations are significantly higher than without including this 436 production. For example, without HONO production, the maximum OH 437 concentration is about 7.5×10^5 #/cm³ under low aerosol condition (AOD=0.25), 438 and rapidly reduced to 1.5×10^5 #/cm³ under high aerosol condition (AOD=2.5) in 439 September. In contrast, by including HONO production, the OH concentrations 440 significantly increased. For example, under higher aerosol condition (AOD=2.5), 441 the maximum of OH concentration is about 7.5×10^5 #/cm³, which is similar to the 442

443 value under low aerosol condition in the no-HONO case. This result suggests that 444 even under the high aerosol conditions, the chemical oxidizing process for O_3 445 production can be active. This result is likely for explaining the co-occurrence of 446 high PM_{2.5} and high O_3 from late spring to early in eastern China.

(3) The measurement of O₃ also shows that the concentrations in winter were always
low, suggesting that the O₃ concentrations were not significantly affected by the
appearance of HONO. The calculated result shows that the seasonal variation of
solar radiation plays important roles for controlling the OH production in winter.
Because the solar radiation is in a very low level in winter, adding the photolysis
of HONO has smaller effect in winter than in other seasons, and OH remains low
values by including the HONO production term.

In recent years, the $PM_{2.5}$ pollutions are reduced due to the large control efforts by the Chinese government, the O₃ pollutions become another severe pollution problem in eastern China. This study is important, because it provides some important scientific highlights to better understand the O₃ pollutions in eastern China.

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459 Data availability. The data used in this paper can be provided upon request from
460 Xuexi Tie (tiexx@ieecas.cn).

461

462 **Author contributions**. XT came up with the original idea of investigating the 463 scientific issue. XT and JX designed the analysis method. XL, GL and SZ provided 464 the observational data and helped in discussion. XT prepared the manuscript with 465 contributions from all co-authors.

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- 581

- 583 **Figure Caption**
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Fig. 1. The geographic locations of the measurement sites in Beijing, in which the measured concentrations of $PM_{2.5}$ and O_3 are used to the analysis.

Fig. 2. The daily averaged concentrations of $PM_{2.5}$ and O_3 in the Beijing region in 2015. The concentrations are averaged over all sites shown in Fig. 1. The blue lines represent the $PM_{2.5}$ concentrations ($\mu g/m^3$), and the red bars represent the O_3 concentrations ($\mu g/m^3$). The rectangles show some typical events during winter (green), spring and fall (orange), and summer (red).

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Fig. 3. The correlation between O_3 and $PM_{2.5}$ concentrations during winter (upper panel) and from late spring to eraly fall (lower panel). During winter, O_3 concentrations were strong anti-correlated with the $PM_{2.5}$ concentrations. From late spring to early fall, O_3 concentrations were correlated with the $PM_{2.5}$ concentrations. 598

Fig. 4. The diurnal variations of $PM_{2.5}$ (blue line) and O_3 (red line), and NO_2 (green line) during a fall period (from Oct.5 to Oc. 6, 2015). It shows that with high $PM_{2.5}$ condition, there was a strong O_3 diurnal variation.

Fig. 5. The cloud condition during the period of the case study (between Oct 5 and 6, 2015) in the Beijing region. The bright white color shows the cloud covers, and the grey white shows the haze covers. The Beijing region was under the heavy haze conditions during the period.

Fig. 6. The measured solar radiation (W/m^2) from Oct. 3 to Oct. 9, 2015 in Beijing. The upper panel shows hourly values, and the lower panel shows the daytime averaged values.

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Fig. 7. The effect of aerosol levels with AOD = 0.25 (black line), AOD = 2.5 (red line), AOD = 3.5 (blue line), and AOD = 4.0 (green line) on the O_3 photolysis calculated by the TUV model in October at middle-latitude.

Fig. 8a. The measured HONO concentrations (ppbv) and the $PM_{2.5}$ and O_3 daily concentrations in Beijing. The upper panel shows the measured daily concentrations of $PM_{2.5}$ and O_3 as shown in Fig.2. The dark-red line was measured HONO in Beijing from 1 to 27 January, 2014.

- Fig. 8b. The measured HONO concentrations (ppbv) and the $PM_{2.5}$ and O_3 daily concentrations in Shanghai. The upper panel shows the measured daily concentrations of $PM_{2.5}$ and O_3 in 2015. The dark-red line was measured in Shanghai from 9 to 18 September, 2009. The green line was calculated by the WRF-Chem model.
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Fig. 8c. The measured HONO concentrations (ppbv) and the $PM_{2.5}$ and O_3 daily concentrations in Xi'an. The upper panel shows the measured daily concentrations of $PM_{2.5}$ and O_3 in 2015. The red line was measured HONO in Xi'An from 24 July to August 6, 2015.

630 631 **Fig. 9.** The measured HONO (upper left panel), $PM_{2.5}$ concentrations (lower left 632 panel), and O₃ concentrations (upper right panel) in fall in Shanghai. It illustrates that

- 633 the high HONO concentrations were corresponded with high PM_{2.5} concentrations.
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Fig. 10. The calculated OH production P(HOx) (#/cm³/s) by using the model calculated HONO (low concentrations) (in the upper panel) and by using the measured HONO (high concentrations) (in the lower panel). The red bars represent the calculation of the P1 term, and the red bars represent the calculation of the P2 term (OH production from HONO).

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Fig. 11. The calculated OH concentrations (#/cm³) with (upper panel) and without
(lower panel) HONO production of OH, under different aerosol levels. Dark red
(AOD=0.25), red (AOD=2.5)), red (AOD=3.5)), and red (AOD=4.0).

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Fig. 12. The effect of cloud cover on the photolysis rate of HONO (J[HONO]). The blue, red, and green lines represent the cloud water vapor of 0 (cloud-free), $10 (g/m^3$ thin cloud), and 50 (g/m³ – thick cloud), respectively. The left panel (A) represents the light aerosol condition, with AOD of 0.25, and the right panel (B) represents the heavy aerosol condition, with AOD of 2.5.

Fig. 13. The calculated OH concentrations in September (blue bars) and December(dark red bars), under different aerosol levels.

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Figures

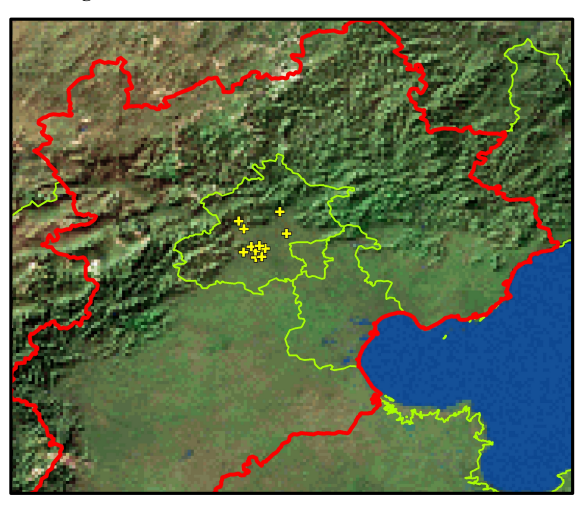


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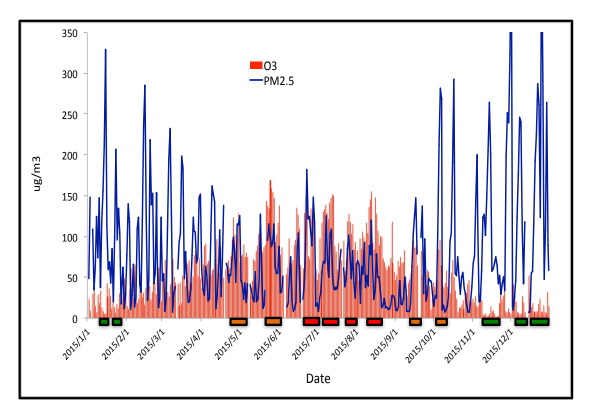


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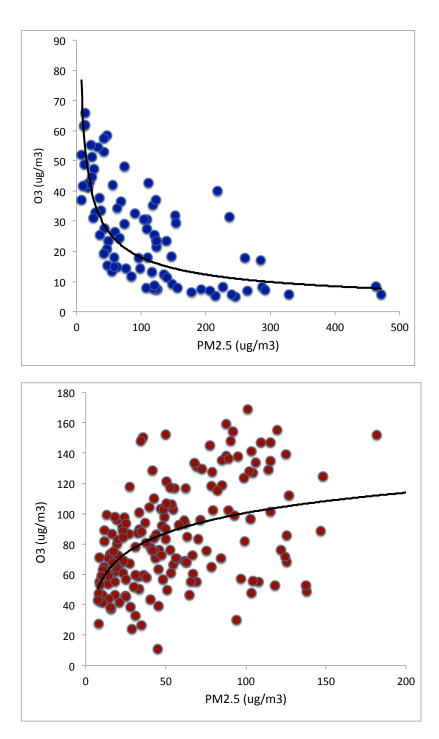




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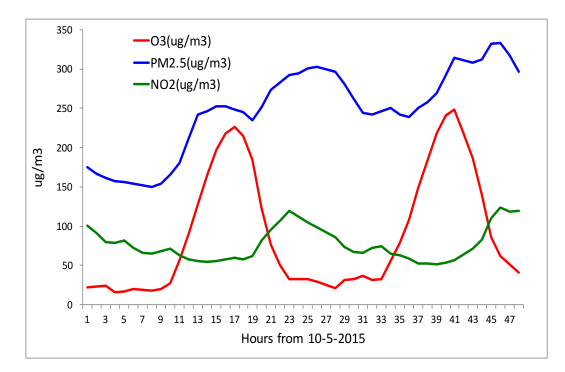
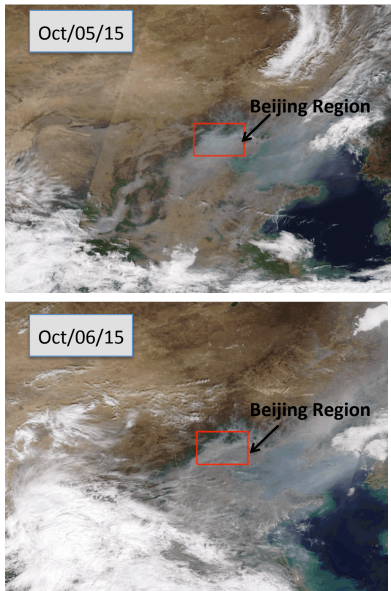


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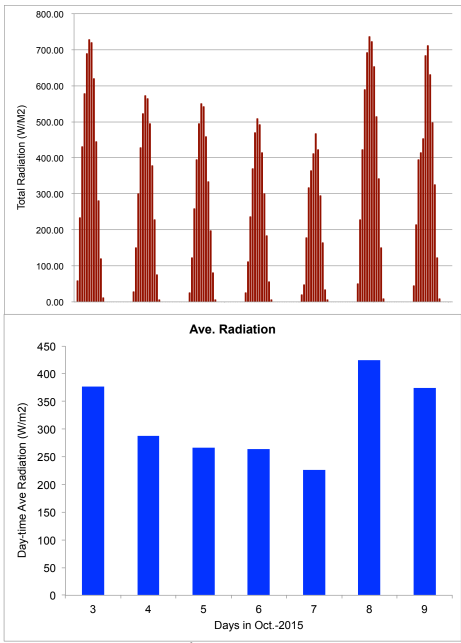


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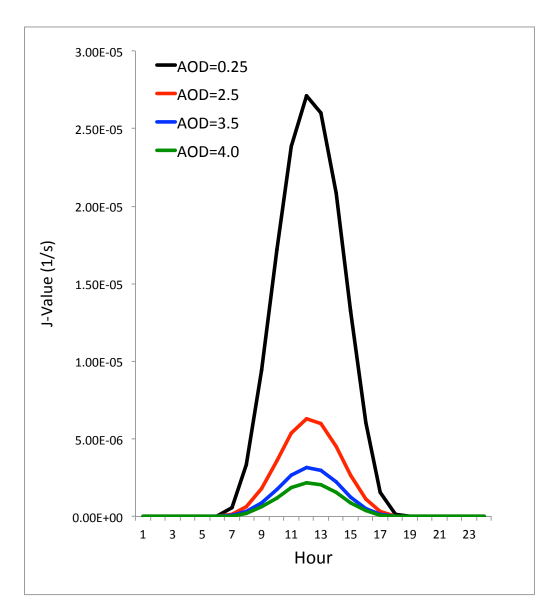
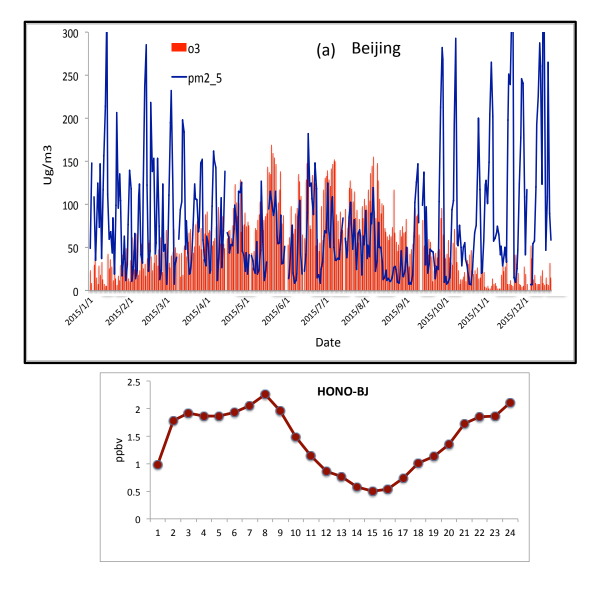


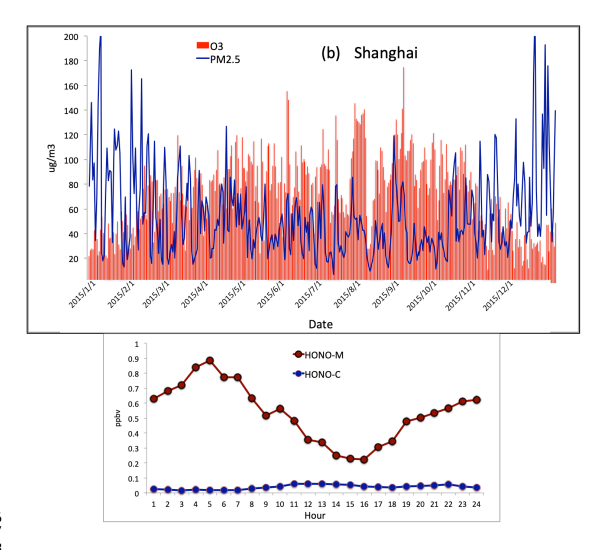


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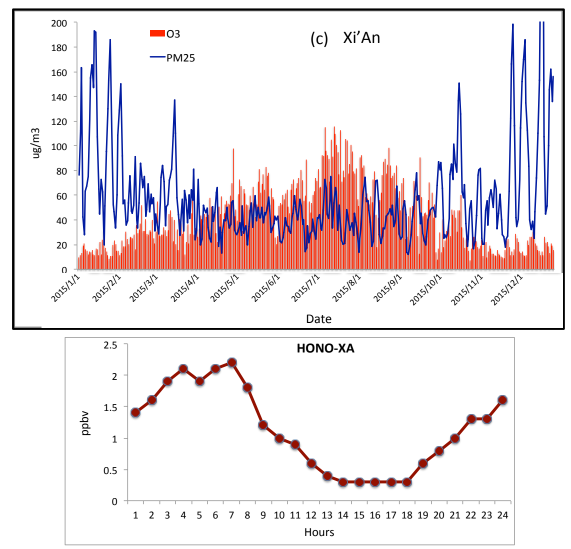


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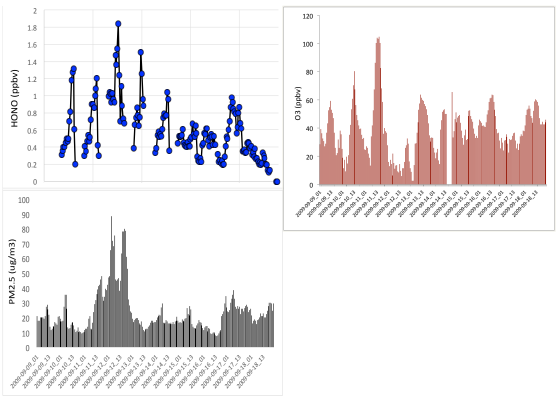


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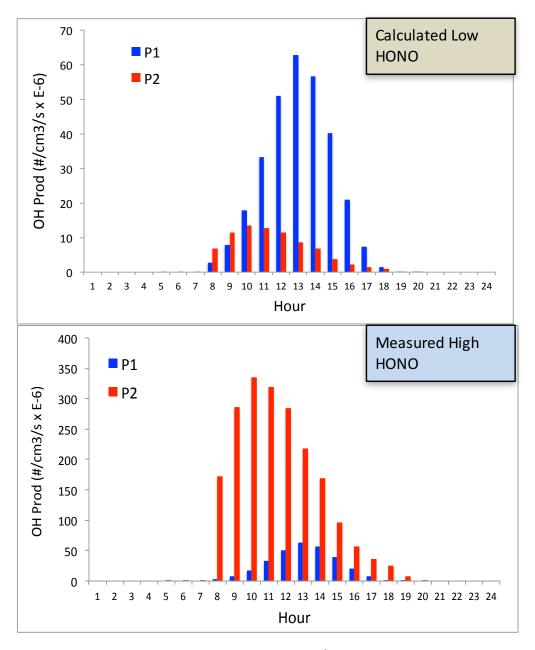


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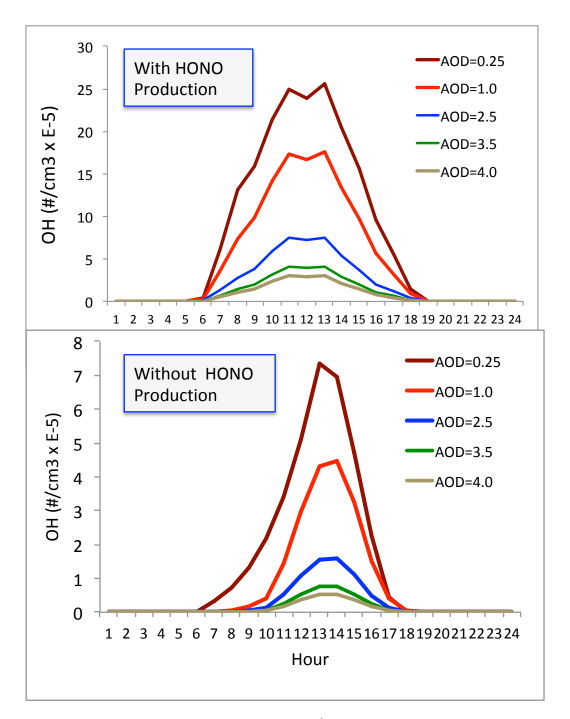


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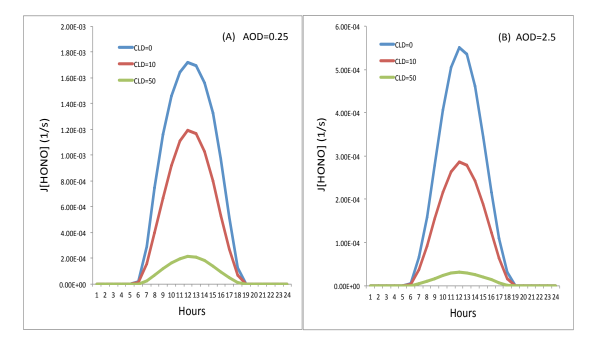




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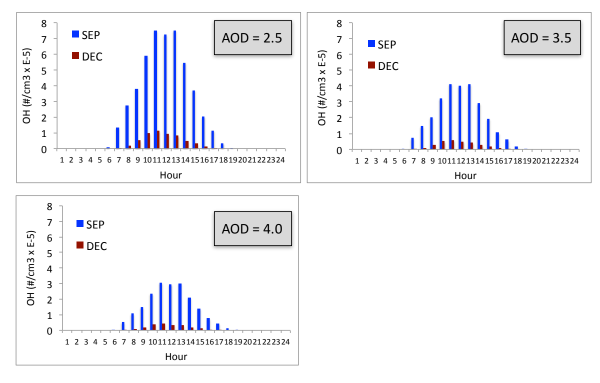




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