

# A Better Understanding of an Extremely High Ozone Episode with Ensemble Simulation

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**Abstract.** Severe ozone pollutions may occur in the Great Bay Area (GBA) when typhoons approach South China. However, numerical models often fail to capture the high ozone concentrations during the episodes, leading to uncertainties in understanding their formation mechanisms. This study conducted an ensemble simulation with 30 members (EMs) using the WRF-Chem model, coupled with a self-developed ozone source apportionment method, to analyze an extremely high ozone episode associated with Typhoon NIDA in the summer of 2016. The newly proposed index effectively distinguished between well-performing (good) and poorly performing (bad) EMs. Compared to the bad EMs, the good EMs accurately reproduced surface ozone variations, particularly capturing the extremely high concentrations observed in the afternoon of July 31. The formation of such high ozone levels was attributed to the retention of ozone in the residual layer at night and the enhanced photochemistry during daytime. As Typhoon NIDA approached, weak winds confined large amounts of ozone in the residual layer at night. The development of planetary boundary layer (PBL) facilitated the downward transport of ozone aloft, contributing to the rapid increase in surface ozone in the following morning. The enhanced photochemistry was primarily driven by increased ozone precursors resulting from favorable accumulation conditions and enhanced biogenic emissions. During the period of high ozone concentrations, contributions from local and surrounding regions increased. Additionally, ozone from southeastern Asia could transport to the GBA at high altitudes and then contribute to surface ozone when the PBL developed.

## 1 Introduction

Tropospheric ozone, especially within the planetary boundary layer (PBL), has attracted much public attention because of its detrimental effects on human health and vegetation (Monks et al., 2015; Feng et al., 2019; Lu et al., 2020). As a typical

secondary air pollutant, tropospheric ozone is primarily formed via photochemical reactions with the participation of nitrogen oxides (NO<sub>x</sub>) and volatile organic compounds (VOCs) (Sillman, 1995; Xie et al., 2014; Li et al., 2019; Wang et al., 2019). Furthermore, the meteorological conditions play important roles in the chemical production and accumulation of ozone in the atmosphere, leading to severe air pollution (Ding et al., 2013a; Wang et al., 2017; Wang et al., 2022).

In the Great Bay Area (GBA; including the Pearl River Delta, Hong Kong, and Macao), China, ozone pollution is closely linked to the western Pacific subtropical high (WPSH) and northwest Pacific typhoon in summer and autumn (Gong and Liao, 2019; Shao et al., 2022; Liu et al., 2023). In particular, when a typhoon approaches this region, severe ozone pollution easily occurs (Deng et al., 2019; Qu et al., 2021). Many previous studies have suggested that when the GBA is at the periphery of a typhoon (Hu et al., 2010; Jiang et al., 2015), high solar irradiance and high temperature promote the ozone photochemical production (So and Wang, 2003; Ding et al., 2004), while unfavorable diffusion conditions increase the accumulation of ozone (Jiang et al., 2008; Li et al., 2022). Thus, high-ozone episodes may occur and persist in the period leading up to a typhoon landfall. Numerical simulations can help us quantitatively study the evolution of this type of ozone pollution (Li et al., 2022; Ouyang et al., 2022; Wang et al., 2022). Furthermore, numerical simulations can also help us separate the impacts of typhoons by comparing base simulation with sensitive simulation that remove the typhoon system (Hu et al., 2019). However, two key issues must be considered, as they may directly affect the accuracy and credibility of the simulation results. The first problem (Prob. 1) is related to whether the model can capture the variations in this type of ozone pollution, especially the extremely high ozone concentration. The second problem (Prob. 2) occurs because this type of ozone pollution is controlled by the competitive effects of the WPSH and typhoon; hence, erasing the typhoon system may increase the impact of the WPSH, which may lead to a high bias in the simulation results or even affect the final conclusions (Gilliam et al., 2015; Chatani and Sharma, 2018).

Ensemble simulation by perturbing meteorological factors in the initial and boundary conditions can offer a solution to the above problems. This method has a greater probability to capture the spatial and temporal variations of air pollutants in extreme synoptic systems (Delle Monache et al., 2006; Zhang et al., 2007; Bei et al., 2014; Zhu et al., 2016), thereby providing more accurate data for relevant studies and yielding more reliable conclusions. More importantly, by comparing well-performing ensemble members (good EMs) with poorly performing EMs (bad EMs), we can better understand the impact of meteorology on ozone, which may help mitigate the uncertainties associated with Prob.2.

In this study, we applied the Weather Research and Forecasting model (WRF; Skamarock et al., 2008) with Chemistry (WRF-Chem; Grell et al., 2005) coupled with an ozone source apportionment method (WRF-Chem-O3tag; Gao et al., 2016; 2017). By perturbing meteorological factors in the initial and boundary condition files, we constructed an ensemble simulation (including 30 EMs) to simulate the extremely high-ozone episode that occurred in the GBA associated with the approach of Typhoon NIDA. By comparing the results of the EMs, we aimed to address the following issues: (1) more accurately capture the variation pattern and the extremely high concentration of surface ozone in the GBA; (2) elucidate the physical and chemical formation mechanisms of this ozone episode; and (3) quantify the changes in the ozone contributions from various geographical source regions during this pollution episode. The structure of this paper is as follows: Section 2

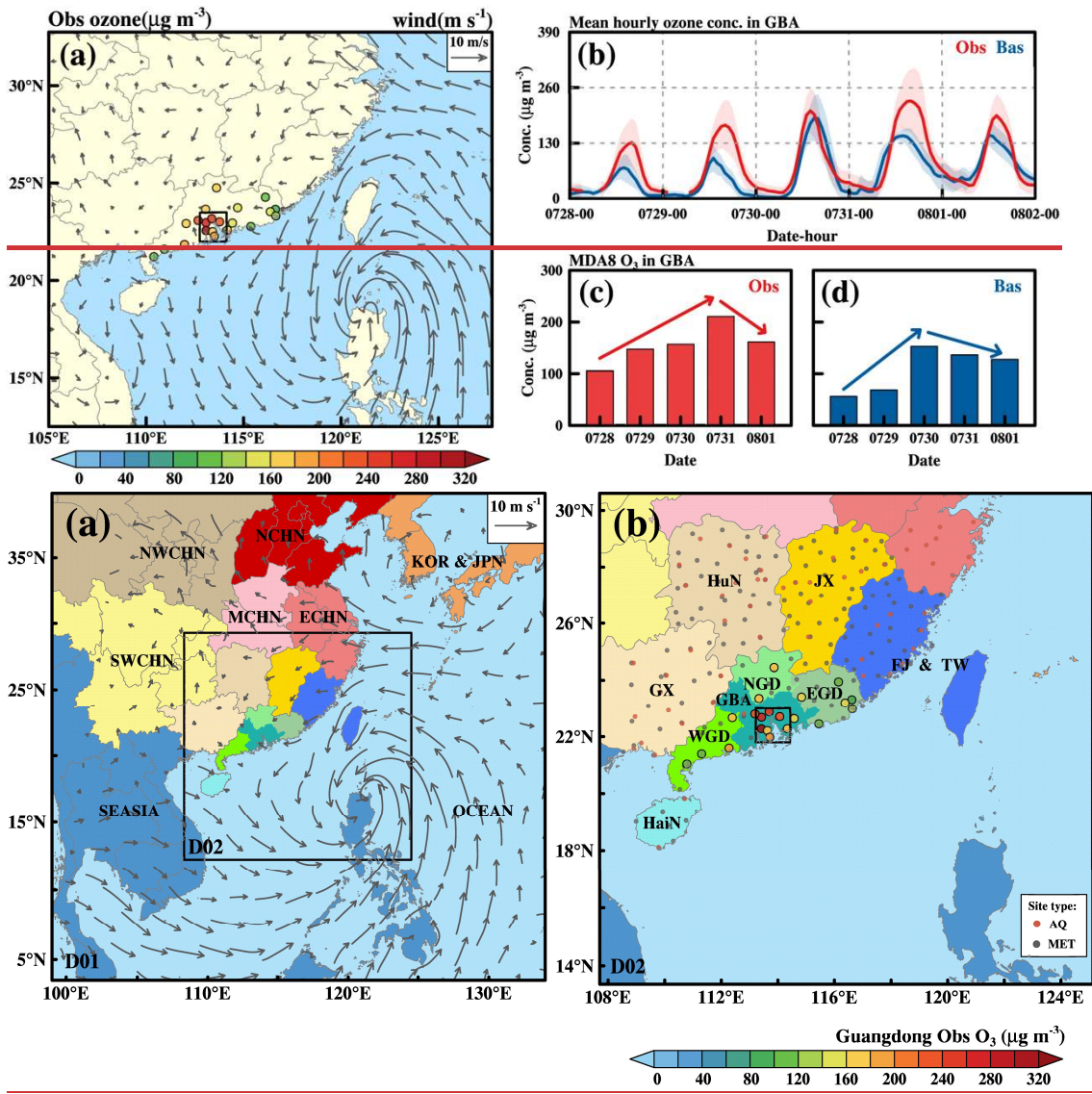
provides a description of the ozone episode, model system and ensemble simulation. The results and discussion are presented in section 3. The conclusions are summarized in section 4.

## **2 The high ozone episode and the methods applied in this study**

### **2.1 The extremely high-ozone episode and the basic simulation**

#### **70 2.1.1 Severe ozone pollution when Typhoon NIDA approached**

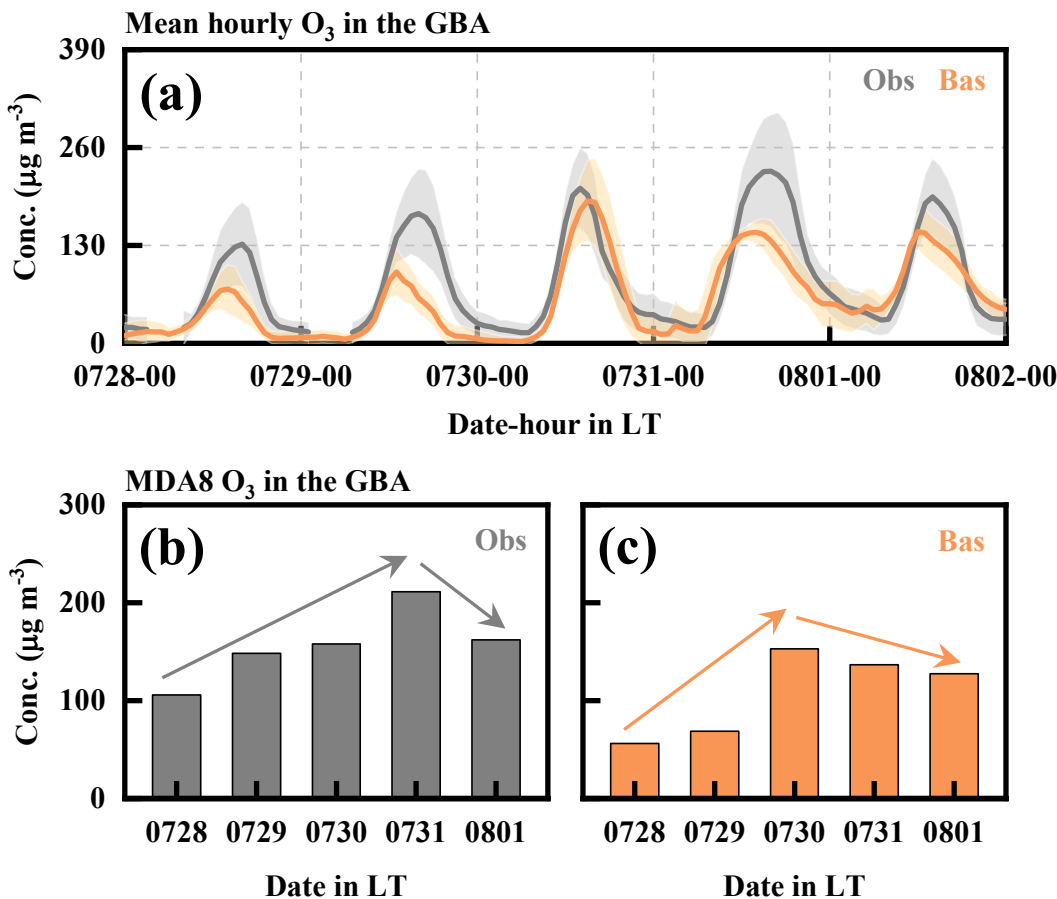
Between July 28 and August 3, 2016, Super Typhoon NIDA formed in the western Pacific and eventually made landfall along the coast of the GBA, resulting in severe disasters in South China, particularly in Guangdong and Guangxi Provinces. When Typhoon NIDA approached the GBA, a high-ozone episode occurred in this region. During the afternoon of July 31, the mean concentrations of surface ozone exceeded  $200 \mu\text{g m}^{-3}$  (grade II of the national standard for the hourly ozone concentration) in most areas of the GBA (black square [in Fig. 1b](#)). Notably, the maximum concentration reached  $366 \mu\text{g m}^{-3}$ , recorded in Jiangmen city [at 14:00 local time \(LT\) on July 31](#).





2.1.2 Basic simulation by using WRF-Chem model

In this study, we conducted a basic simulation using the WRF-Chem model. This model is a fully coupled 3D Eulerian model system that incorporates two-way online feedback between the meteorological model and the chemical transport model. It has been widely employed in air quality research and forecasting studies (Li et al., 2018; Gao et al., 2021; Yang et al., 2022; Huang et al., 2020; 2023). Regarding the model configuration, we established nested model domains (Fig. 2a1a). To simulate the lifetimes of both the ozone episode and Typhoon NIDA, the parent domain (D01) covered most parts of East Asia and Southeast Asia. The simulation period spanned from 00:00 on July 25 to 00:00 on August 4, 2016 (UTC). The first two days were designed as the spin-up period. The complete introduction of Other-detailedthe model configurations and descriptions-of-the-used-relevant datasets can be found in the sSupplementary Informationinformation.

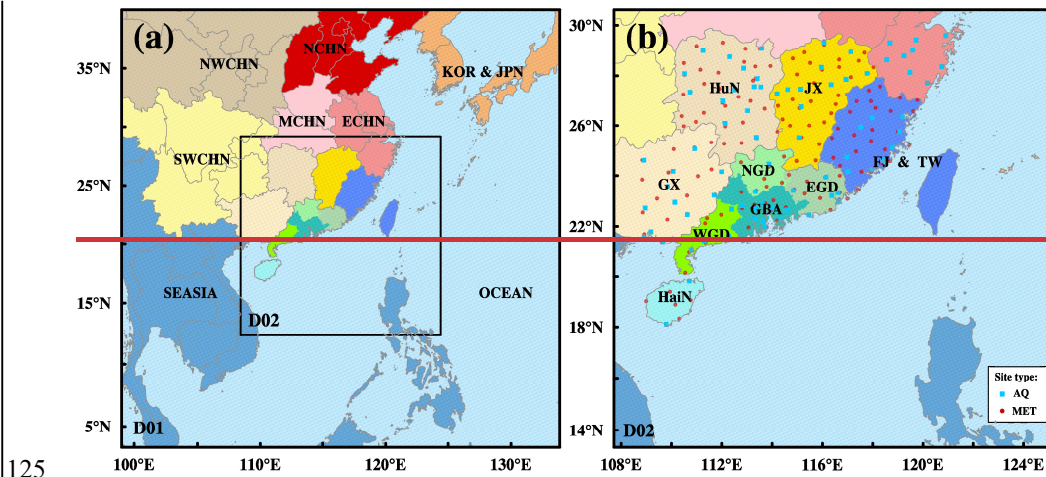


**Figure 2: (a) The comparisons of mean hourly ozone in the GBA between the observations (Obs) and the basic simulation (Bas); The mean maximum daily average ozone (MDA8 O<sub>3</sub>) of Obs (b) and Bas (c) in the GBA.**

As shown in Fig. 4b2a, the comparison of the hourly ozone concentrations revealed that the basic simulation results (Bas) did not show good agreement with the observations (Obs) in the GBA. Although Bas could capture the diurnal variation well, it failed to simulate the ozone peaks during this episode. In particular, Bas significantly underestimated the maximum concentration that occurred in the afternoon of July 31. Regarding the mean maximum daily 8-hour average ozone (MDA8 O<sub>3</sub>), the maximum value was reached on July 31 in Obs (Fig. 4e2b), whereas in Bas, the maximum value was reached on 30 July (Fig. 4d2c). The results suggested that Bas could not reproduce this ozone episode well, although the model settings have been widely employed and shown to be effective in many other air quality modeling studies. Hence, by using the same model settings, we conducted an ensemble simulation in this study. We believe that this approach can enhance simulation accuracy and facilitate a better understanding of this high ozone episode.

## 2.2 Ensemble simulation and ozone source apportionment

For the ensemble simulation, all EMs were generated with the “cv3” background error covariance option in the WRF-3DVAR package via perturbation of the meteorological factors in the initial and boundary condition files. The horizontal wind components, potential temperature, and water vapor mixing ratio were perturbed with standard deviations of 2 m s<sup>-1</sup>, 1 K, and 0.5 g kg<sup>-1</sup>, respectively (Zhu et al., 2016; Xiao et al., 2023). This ensemble initialization method has been widely utilized in data assimilation and weather analysis (Meng and Zhang, 2008a, b; He et al., 2019; Chan et al., 2022). In this study, we used this method to create 30 EMs and run them separately. By comparing the simulated ozone concentrations with the observations, “good” and “bad” EMs could be distinguished on the basis of their performance. The formation of the ozone episode and its source contributions could then be quantitatively examined by contrasting the results of the good and bad EMs. In addition, the ensemble initialization method suggested that the purpose of this work is to study the impacts of meteorological fields on the formation of the high ozone episode. Since we did not change anthropogenic emissions in the ensemble simulation, the impacts of anthropogenic emissions on ozone concentrations were not discussed.



~~Figure 2: Model domain and geographical source region setting. The locations of the air quality (AQ; square) and meteorology (MET; dot) observation stations are shown in (b).~~

2.3 Online ozone source apportionment method coupled with the WRF-Chem model

130 In this study, we also employed our ozone source apportionment method in the ensemble simulation to quantify the ozone contributions during this episode. Similar to the Ozone Source Apportionment Technology (OSAT; Yarwood et al., 1996), our method is a mass balance technique that aims to identify the ozone contributions from each geographical source region preset in the model domain. Due to its secondary pollutant properties, the photochemical production of ozone is attributed to the contributions of ozone precursors (NO<sub>x</sub> and VOCs) emitted from the geographical source regions. In this method, the  
135 model domain is divided into several source regions, where ozone and its precursors are treated as independent variables. Because of the nonlinear relationship of ozone photochemistry, the ozone formation sensitivity (NO<sub>x</sub>-limited or VOCs-limited) will be determined for each grid at each time step. Then, the chemical production of ozone will be allocated based on the concentration ratio of the key ozone precursor (NO<sub>x</sub> for NO<sub>x</sub>-limited and VOCs for VOCs-limited) from each source region. These variables will also undergo all the physical processes as the original variables experience during the simulation,  
140 but they will not interfere with the original simulation. In addition, the initial and boundary chemical conditions of the relevant species also need to be defined as sources to maintain the mass balance. More information on this method can be found in Gao et al. (2016, 2017).

Before initiating the simulation, we need to establish geographical source regions for the ozone apportionment method. There were 17 geographical source regions in the model domain (Fig. 21a). The GBA, where we studied in this paper, was  
145 designated as one region. The other cities in Guangdong Province were categorized into three regions based on their relative locations to the GBA (Fig. 1b): east Guangdong (EGD), north Guangdong (NGD) and west Guangdong (WGD). For other areas in China, the regional settings were defined according to administrative divisions. Source regions outside of China were defined based on national divisions. The marine areas in the model domain were established as one region. The detailed compositions of the various geographical source regions are listed in Table 1. In addition, the boundary conditions were  
150 defined as one region, which we referred to as O<sub>3</sub> inflow, as it can flow into the model domain and have a significant impact on ozone concentrations, which is typically treated as background ozone (Gao et al., 2017; 2020). The initial conditions of D01 and D02 were also defined as independent ozone contributions (INIT1 and INIT2, respectively).

Table 1: List of the geographical source regions in the model domain.

Source regions	Details
GBA	Pearl River Delta, Hong Kong and Macao, China
EGD	Cities in the east of Guangdong Province, China
NGD	Cities in the north of Guangdong Province, China

WGD	Cities in the west of Guangdong Province, China
FJ & TW	Fujian and Taiwan provinces, China
JX	Jiangxi Province, China
HuN	Hunan Province, China
GX	Guangxi Province, China
HaiN	Hainan Province, China
ECHN	East China, including Jiangsu, Zhejiang, Anhui Province and Shanghai city, China
MCHN	Middle China, including Henan and Hubei Province, China
SWCHN	Southwest China, including Yunnan, Sichuan, Guizhou, Xizang Province and Chongqing, China
NCHN	North China, including Hebei, Shanxi, Shandong, Liaoning, Jilin Province and Beijing, Tianjin cities
NWCHN	Northwest China, including Shaanxi, Inner Mongolia, Ningxia, Gansu, and Qinghai provinces
KOR & JPN	North Korea, South Korea, and Japan
SEASIA	Southeast Asia, including Brunei, Cambodia, India, Indonesia, Laos, Malaysia, Myanmar, Philippines, Thailand, and Vietnam
OCEAN	Ocean area, including the Bohai sea, Huanghai sea, Donghai sea, Nanhai, and parts of the western Pacific

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### 3. Results and Discussion

#### 3.1 EM selection and model validation

By comparing the observed ozone concentration with those simulated by each EM, we identified the EMs that could well reproduce the high ozone episode in the GBA by considering the following two rules: (1) whether the EM could reproduce the variation pattern of the mean surface ozone in the GBA and (2) whether the EM could capture the maximum concentration that occurred in the afternoon of July 31. Thus, two statistical metrics were determined to select good and bad EMs: the correlation coefficient (R) of the mean surface ozone in the GBA between the simulation and observation and the mean normalized bias (MNB) calculated from the simulated and observed ozone concentrations during the afternoon of July 31 (12:00~16:00 LT), when the maximum concentration occurred. The R and MNB of each EM were listed in Table 2. Among all the EMs, the values of R ranged from 0.79 to 0.92, indicating acceptable agreement with the variation pattern of observed ozone. The MNBs of all the EMs ranged from -46.99% to -10.92%. According to EPA (2005, 2007), the

recommended threshold values of MNB on ozone are  $\pm 15\%$ . Only EM17 could meet the criterion, while the other fulfill the requirement, other EMs didn't meet the benchmark not.

170 Table 2. The R and MNB of surface ozone in the GBA between each EM and the observation

<u>EMs</u>	<u>R (<math>\times 100</math>)</u>	<u>MNB (%)</u>
<u>EM01</u>	<u>86.23</u>	<u>-30.97</u>
<u>EM02</u>	<u>88.44</u>	<u>-27.40</u>
<u>EM03</u>	<u>90.67</u>	<u>-32.58</u>
<u>EM04</u>	<u>83.25</u>	<u>-24.80</u>
<u>EM05</u>	<u>90.79</u>	<u>-21.42</u>
<u>EM06</u>	<u>89.75</u>	<u>-35.62</u>
<u>EM07</u>	<u>86.60</u>	<u>-34.00</u>
<u>EM08</u>	<u>82.77</u>	<u>-33.74</u>
<u>EM09</u>	<u>88.52</u>	<u>-31.62</u>
<u>EM10</u>	<u>85.64</u>	<u>-29.89</u>
<u>EM11</u>	<u>88.48</u>	<u>-32.46</u>
<u>EM12</u>	<u>82.06</u>	<u>-36.74</u>
<u>EM13</u>	<u>92.30</u>	<u>-23.47</u>
<u>EM14</u>	<u>90.01</u>	<u>-35.37</u>
<u>EM15</u>	<u>85.06</u>	<u>-28.46</u>
<u>EM16</u>	<u>91.03</u>	<u>-18.35</u>
<u>EM17</u>	<u>92.08</u>	<u>-10.92</u>
<u>EM18</u>	<u>87.63</u>	<u>-30.95</u>
<u>EM19</u>	<u>83.59</u>	<u>-31.34</u>
<u>EM20</u>	<u>81.18</u>	<u>-37.64</u>
<u>EM21</u>	<u>84.54</u>	<u>-30.11</u>
<u>EM22</u>	<u>89.51</u>	<u>-29.16</u>
<u>EM23</u>	<u>85.15</u>	<u>-46.99</u>
<u>EM24</u>	<u>86.04</u>	<u>-26.11</u>
<u>EM25</u>	<u>88.55</u>	<u>-32.95</u>

<u>EM26</u>	<u>87.68</u>	<u>-29.10</u>
<u>EM27</u>	<u>87.89</u>	<u>-29.39</u>
<u>EM28</u>	<u>81.99</u>	<u>-40.32</u>
<u>EM29</u>	<u>79.68</u>	<u>-31.42</u>
<u>EM30</u>	<u>89.28</u>	<u>-28.72</u>

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It should be noted that both R and MNB were important to evaluate the model performance on ozone in this high ozone episode, however, their validation results didn't address a consistent conclusion. In order to drawing a common conclusion, by considering the effects of both R and MNB, In-addition, using R and MNB intervals of [-1, 1] and [-100, 100],

175 ~~respectively,~~ we introduced an index ( $I_{dis}$ ) to quantitatively assess the performance of each EM. For each EM  $k$ , the index  $I_{dis}^k$  can be expressed as:

$$I_{dis}^k = \text{sqrt}[(R^k \times 100 - R_{best} \times 100)^2 + (|MNB^k| \cancel{MNB^k} - MNB_{best})^2], \quad \text{—————}$$

(1)

where  $R^k$  and  $MNB^k$  are the correlation coefficient and MNB between the observed ozone concentration and the simulated  
180 ozone concentration obtained from EM  $k$ , respectively. The values of  $R_{best}$  and  $MNB_{best}$  are 1 and 0, respectively. Thus, as shown in Fig. 3a,  $I_{dis}^k$ , calculated using equation (1), represents the “distance” of the model performance of EM  $k$  from the best performance (red star). A smaller  $I_{dis}$  value indicates better performance of the EM. For clarity,  $I_{dis}^k$  is also represented by colors in Fig. 3a, where red indicates that the index is close to the “best”, while blue indicates that the index is far from the “best”.

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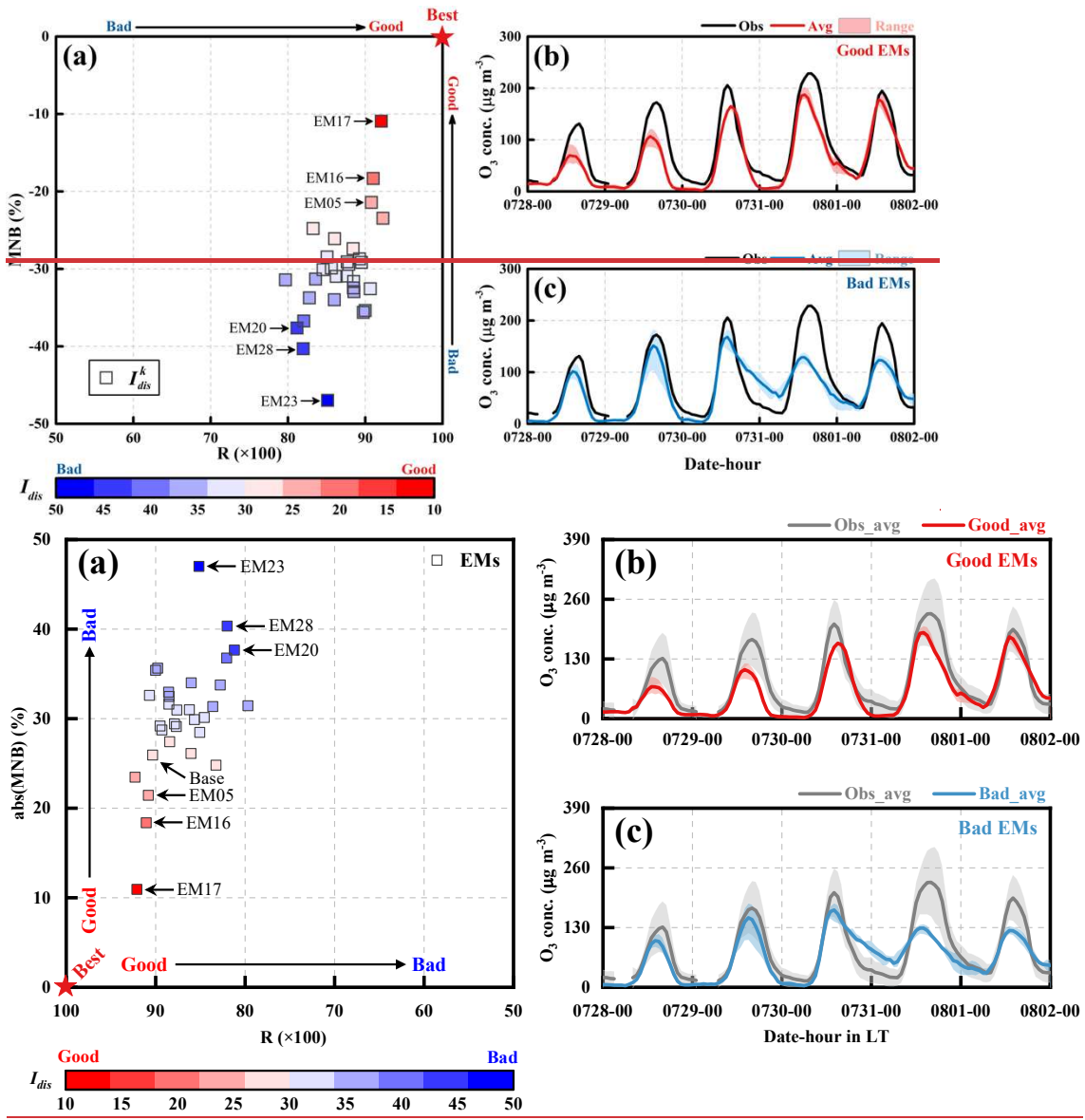


Figure 3: (a) Model performance of each EM on surface ozone in the GBA; comparisons between the observed and simulated surface ozone concentrations in the GBA for (b) good EMs and (c) bad EMs.

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In Fig. 3a,  $I_{dis}$  (square) indicated that EM05, EM16, and EM17 were the top three members with performances closest to the best, suggesting that these three EMs more effectively reproduce the variations in ozone in the GBA. In contrast, the  $I_{dis}$  values of EM20, EM23, and EM28 were higher than those of the other EMs, indicating that they performed worse. Consequently, we classified EM05, EM16, and EM17 as good members, while EM20, EM23, and EM28 were classified as bad members. As shown in Fig. 3b, the comparison between the mean results of good EMs and the observations clearly

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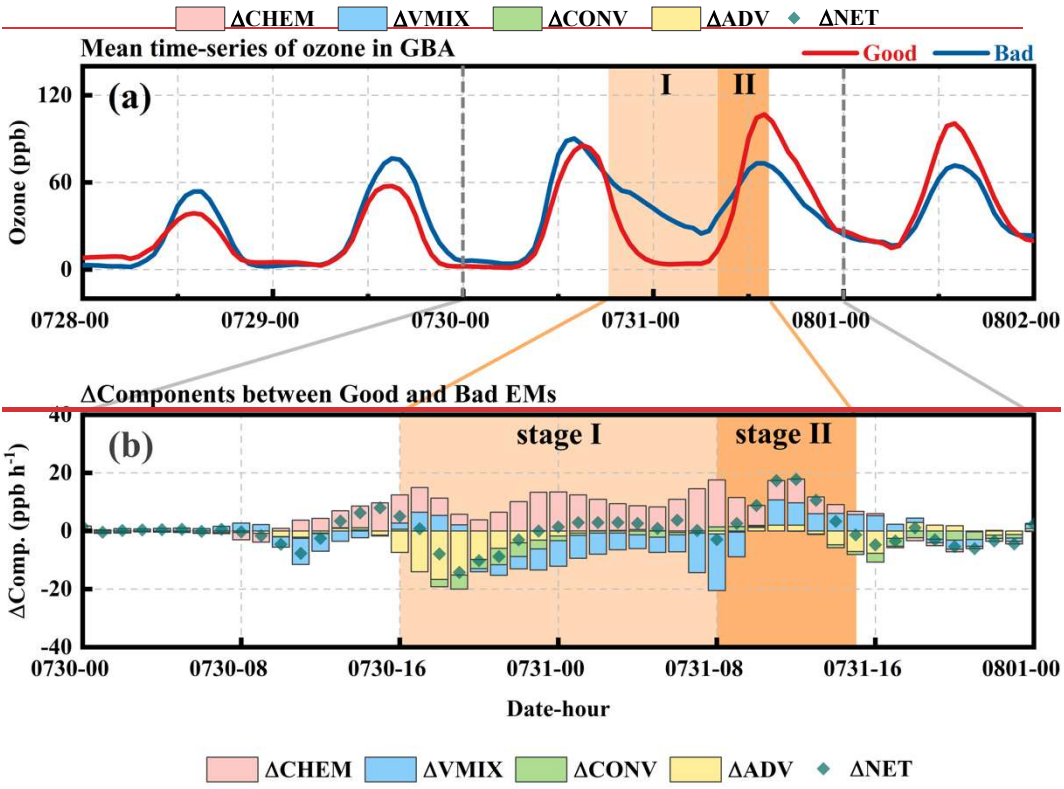
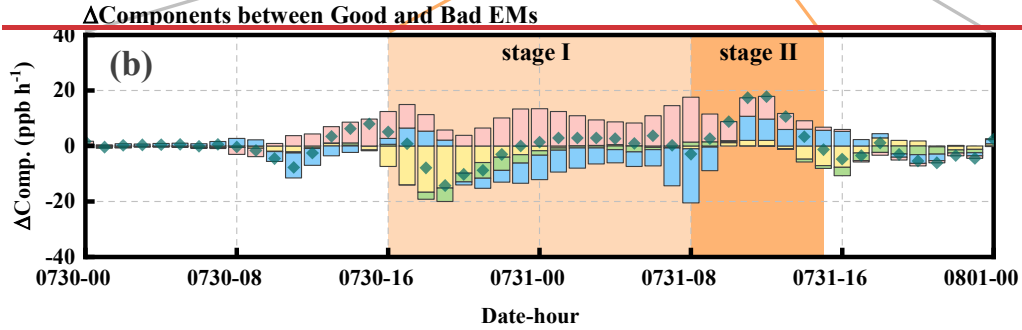
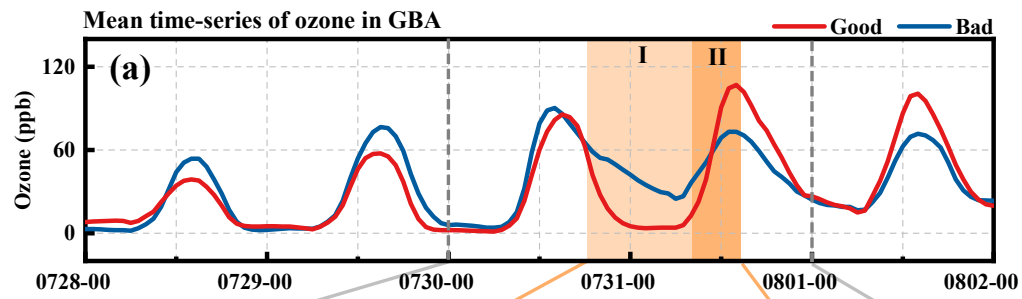


revealed that the ozone time series of the good EMs agreed well with the observations and basically reproduced the variation pattern of ozone in the GBA during this episode, particularly the maximum concentration that occurred on July 31. For the bad EMs (Fig. 3c), ~~time~~the time series of surface ozone exhibited an acceptable diurnal variation. However, they failed to capture the maximum ozone concentration on July 31. In addition, for both the good and bad EMs, we also did the model validations on meteorological factors (temperature at 2 m, T2; wind speed, WS; and wind direction, WD) and NO<sub>2</sub> from hundreds of stations located in middle and southern China (Table S3). ~~The statistical~~Statistical metrics revealed that, both inside and outside the GBA, the good EMs demonstrated better model performance than the bad EMs. For example, all the meteorological factors of the good EMs exhibited higher indices of agreement (IOAs) than did those of the bad EMs. Most variables of the good EMs exhibited lower biases than those of the bad EMs in all regions, except T2 outside the GBA and WD in all regions. The good EMs exhibited a satisfactory model performance and could effectively reproduce the high-ozone episode in the GBA.

### 3.2 Physical and chemical causes of the extremely high ozone concentration on July 31

#### 3.2.1 Process analysis of the average time series of surface ozone in the GBA

For surface ozone in the GBA, the average time series of the good and bad EMs (abbreviated as ozone\_good and ozone\_bad, respectively) are shown in Fig. 4a. The time series exhibited similar diurnal variations and concentrations from July 28 to the afternoon of July 30. The two ozone time series then began to vary differently, and the differences increased until the afternoon of July 31. Specifically, compared with that of ozone\_bad, the time series of ozone\_good declined more sharply in the afternoon of July 30, and the concentration remained relatively low until the early morning of July 31 (Stage I; light orange shaded area). From 08:00 LT on July 31, the time series of both ozone\_good and ozone\_bad began to increase, and the trends persisted until the afternoon, when high ozone concentrations occurred (Stage II; dark orange shaded area). Notably, ozone\_good increased much more than did that of ozone\_bad, resulting in the maximum concentration during this high-ozone episode in the GBA. The variation pattern was also consistent with that in the observations. As mentioned above, the relatively high maximum concentration of ozone\_good may be attributed to the significant differences between ozone\_good and ozone\_bad from the afternoon of July 30 to the afternoon of July 31.



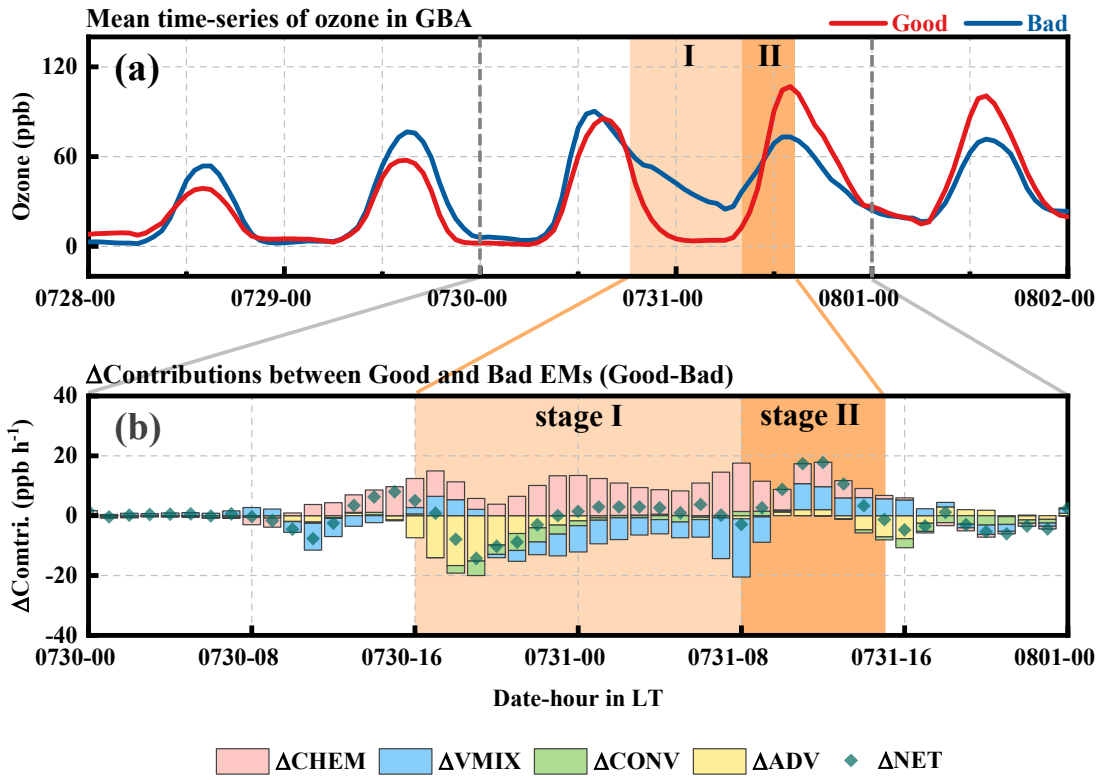


Figure 4: (a) Averaged time series of ozone<sub>good</sub> and ozone<sub>bad</sub> in the GBA<sub>-</sub> and (b) the differences in each ozone **component contribution** between the good and bad EMs (**good-bad**) from July 30 to 31. CHEM = chemistry, VMIX = vertical mixing, CONV = convection, ADV = advection, NET = net contribution.

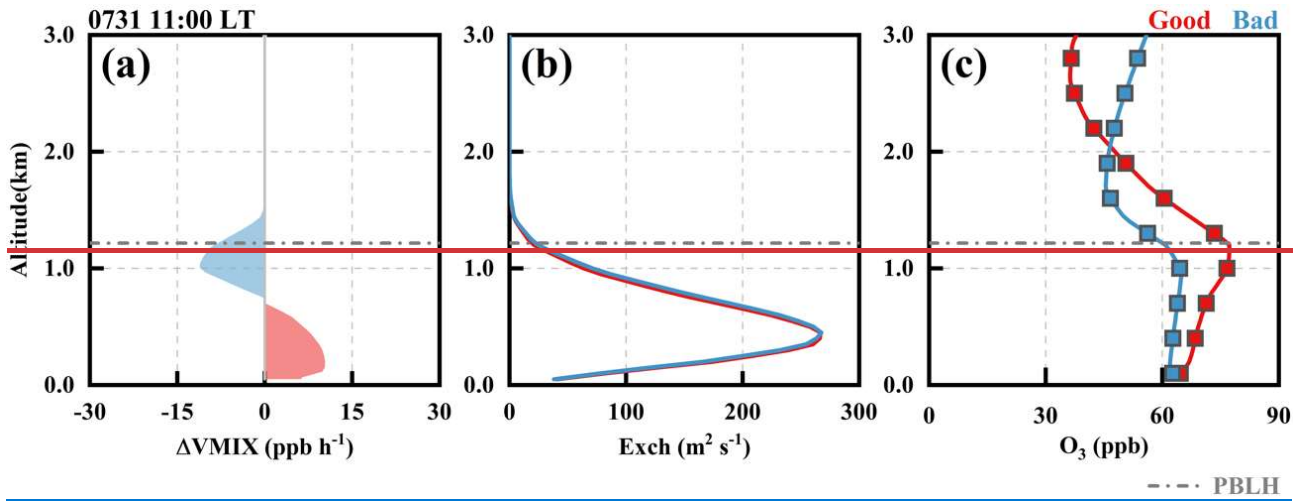
The changes in ozone **components-contributions between good and bad EMs (Good-Bad)** (**good-bad**; Fig. 4b) could help us clarify the chemical and physical causes of the extremely high ozone concentration on July 31 (Fig. 4b) (Fig. 4b).  $\Delta$ NET is calculated as the sum of all differences from each process between the good and bad EMs, which can represent the relative changes in ozone<sub>good</sub> relative to ozone<sub>bad</sub>. For example, when both ozone<sub>good</sub> and ozone<sub>bad</sub> increased at Stage II, the positive  $\Delta$ NET value indicated a more significant increase in ozone<sub>good</sub>. At these two stages,  $\Delta$ CHEM,  $\Delta$ ADV and  $\Delta$ VMIX significantly contributed to  $\Delta$ NET. At Stage I, although both ozone<sub>good</sub> and ozone<sub>bad</sub> decreased, the negative values of  $\Delta$ NET during 18:00~21:00 on July 30 suggested that ozone<sub>good</sub> decreased more significantly, which was attributed primarily to  $\Delta$ ADV. At Stage II, ozone<sub>good</sub> increased more than ozone<sub>bad</sub> did since  $\Delta$ NET became positive. The budget of  $\Delta$ NET suggested that the significant increase in ozone<sub>good</sub> resulted from the combined effects of  $\Delta$ CHEM and  $\Delta$ VMIX. Hence, it could be concluded that the extremely high concentration of ozone<sub>good</sub> that occurred in the afternoon of July 31 could be directly attributed to the ozone contributions from chemical and vertical mixing processes. In addition,  $\Delta$ ADV during the evening of the previous day also contributed to the extremely high concentration. In the

following section, we will discuss the causes of the extremely high ozone concentration from physical and chemical perspectives.

### 3.2.2 Physical cause of the extremely high ozone concentration

Among the physical processes, VMIX was the direct contributor to the high concentration of ozone\_good [from 10:00 to 15:00 \(LT\) on July 31](#). ~~Here we took~~[taking the situation at 11:00 as an example \(Fig. 5\)](#). By comparing the VMIX of ozone\_good with that of ozone\_bad ( $\Delta$ VMIX in Fig. 5a4), the negative values aloft and positive values near the surface suggested that more ozone from ozone\_good was entrained downward to the surface, leading to a greater increase in ozone at ground level. As the two key factors of VMIX (Gao et al., 2018), the differences in the vertical exchange coefficient and the vertical ozone profile between the good and bad EMs could explain the significant contribution of VMIX in good EMs. In the morning of July 31, there were no obvious differences between the good and bad EMs (Fig. 5ba2). However, the vertical ozone profiles (Fig. 5a35c) revealed that ozone\_good exhibited higher vertical gradients from the surface to the top of the PBL. Under these conditions, compared with the bad EMs, the good EMs showed more transport of ozone from the top of PBL to surface, despite similar vertical exchange intensities.





**Figure 5: The vertical profiles of the changes in VMIX between good and bad EMs (good-bad) (a), exchange coefficients (b) and ozone concentrations of good and bad EMs at 11:00 (LT) on July 31.**

**Figure 5: (a) Profiles of (1)  $\Delta VMIX$ , (2) exchange coefficient and (3) ozone at 10:00 and 11:00 on July 31; (b) vertical distributions of (1) ozone good and the changes in ozone [(2)  $\Delta Ozone$ ] and NET [(3)  $\Delta NET$ ]; (c) vertical distributions of the changes in the ozone contribution of each process [(1)  $\Delta ADV$ ; (2)  $\Delta VMIX$ ; (3)  $\Delta CHEM$ ; (4)  $\Delta CONV$ ].**

The vertical distributions of ozone<sub>good</sub> (Fig. 5b1) revealed that the vertical gradient persisted from the night of July 30 to the next morning, which is consistent with the vertical structure of ozone concentrations reported in other urban areas (Xu et al., 2018; Gao et al., 2020). The greater vertical gradients of ozone from good EMs occurred not only in the morning of July 31 but also in the previous evening. As shown in Fig. 6a, the vertical distribution of the difference in ozone between good and bad EMs ( $\Delta Ozone$ ) showed positive values aloft and negative values in the lower layers from the evening of July 30 to the morning of July 31. this indicated that good EMs exhibited greater vertical gradients of ozone during this period. The difference between ozone<sub>good</sub> and ozone<sub>bad</sub> ( $\Delta Ozone$  in Fig. 5b2) showed that the vertical gradient of ozone<sub>good</sub> was even greater, as positive  $\Delta Ozone$  values occurred aloft and since negative values occurred in the lower layers during this period. Notably, the positive  $\Delta Ozone$  values aloft were even greater during the morning of July 31, which could further increase the vertical gradient of ozone.  $\Delta NET$  (Fig. 5b3b) revealed a negative zone (blue square) at altitudes of 0–750 m from 18:00 to 23:00 LT on July 30 and a positive zone (red square) at altitudes of 1000–1500 m from 05:00 to 11:00 LT on July 31. These two changes were the direct causes of the vertical structure. By separating  $\Delta NET$  into changes in physical and chemical each ozone component contributions (Fig. 5e1–46c–6f), it could be observed that  $\Delta ADV$  exhibited the same features (the negative and positive zones in Fig. 5e16b) as did  $\Delta NET$  at the same altitude during the same period, suggesting that the significant vertical gradients of ozone<sub>good</sub> were primarily caused by distinct  $\Delta ADV$  values at different levels during various periods.



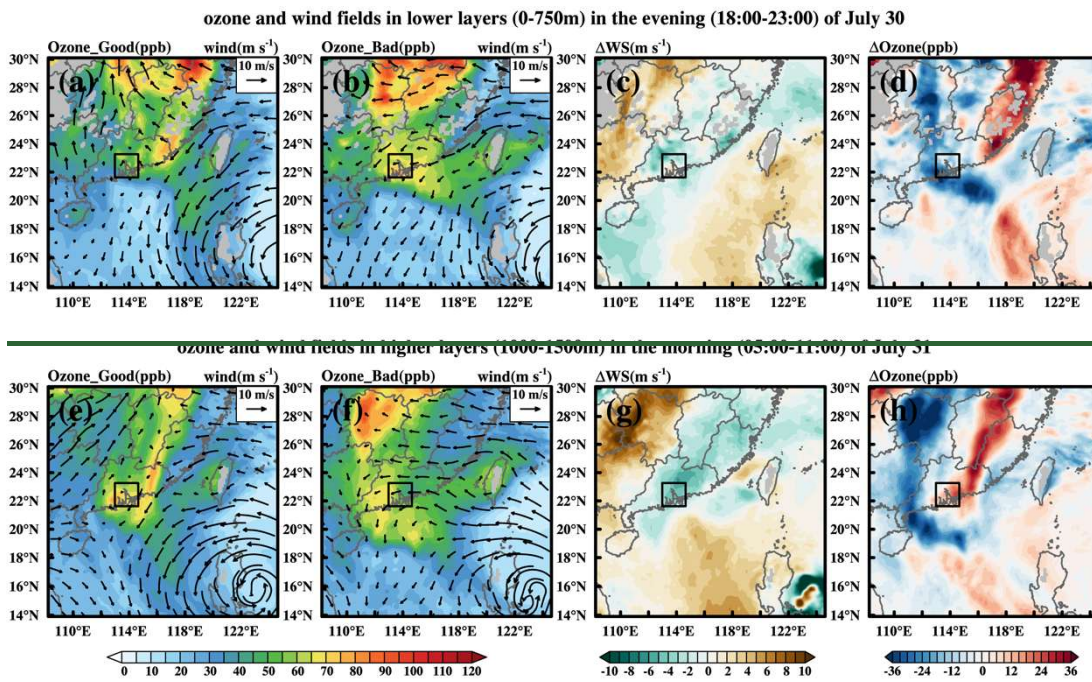


Figure 6: Mean distributions of the ozone concentration and wind fields and their respective differences between the good and bad EMs in the lower (a-d) and higher layers (e-h).

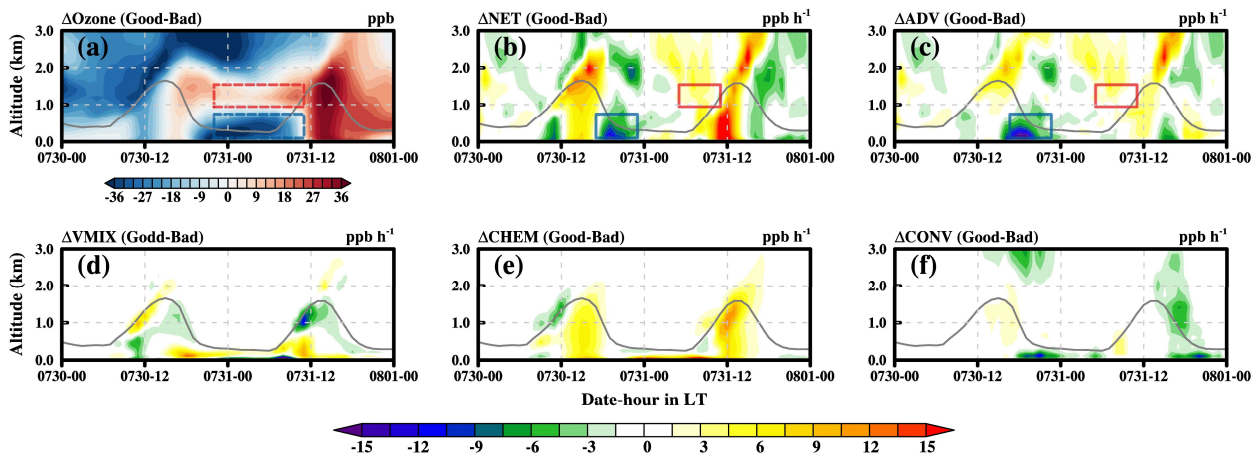


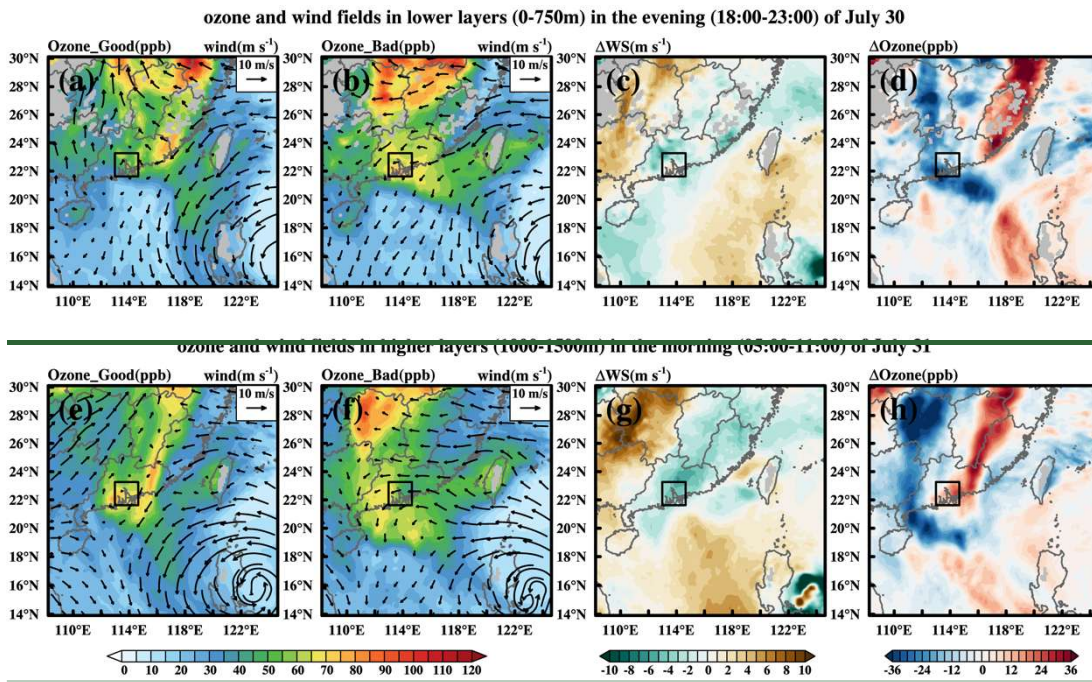
Figure 6: The vertical distributions of the changes in ozone (a) and its contributions (b-f) between good and bad EMs (good-bad).

The contribution of ADV is influenced by wind fields and ozone spatial distributions (Gao et al., 2017). The mean distributions of the ozone concentration and wind fields are shown in Fig. 67. During the evening of July 30, the good EMs indicated southwest winds in the GBA in the lower layers (Fig. 6a7a). With the transport of southwesterly winds, the low



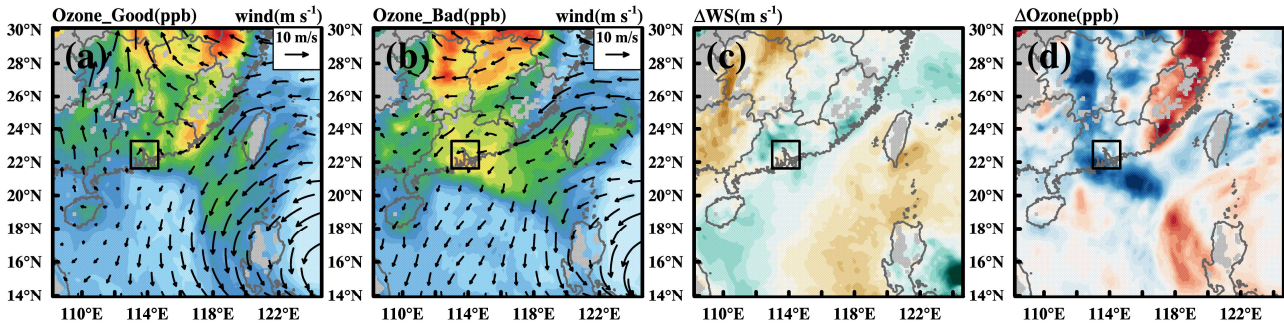
concentrations of ozone over sea areas (upwind regions) could be transported to the GBA, leading to a decrease in ozone. However, the bad EMs showed that the GBA was controlled by easterly winds (Fig. 6b7b). The ozone in the upwind region was approximately equal to or even higher than that in the GBA, which did not cause a significant decrease in ozone levels. Thus, the concentrations of ozone\_good were lower than those of ozone\_bad in the lower layers of the GBA in the evening of July 30 (Fig. 6d7d). The significant vertical ozone gradient was more closely associated with the increase in ozone aloft in the morning of July 31 (Fig. 6e7e-6h7h). In the higher layers, the good EMs exhibited high ozone concentrations and approximate static winds across the GBA, which was favorable for ozone retention (Fig. 6e7e). However, the bad EMs exhibited stronger east winds but lower ozone concentrations in upwind regions, which might have led to a decrease in ozone over the GBA (Fig. 6f7f). Thus, the concentrations of ozone\_good were higher than those of ozone\_bad over the GBA during the morning of July 31 (Fig. 6h7h). In addition, the good EMs showed weaker winds than the bad EMs did at the two stages (Fig. 6e-7c and 6g7g, respectively). Especially in the higher layers, the weak winds played an important role in the increased  $\Delta\text{Ozone}$  value.

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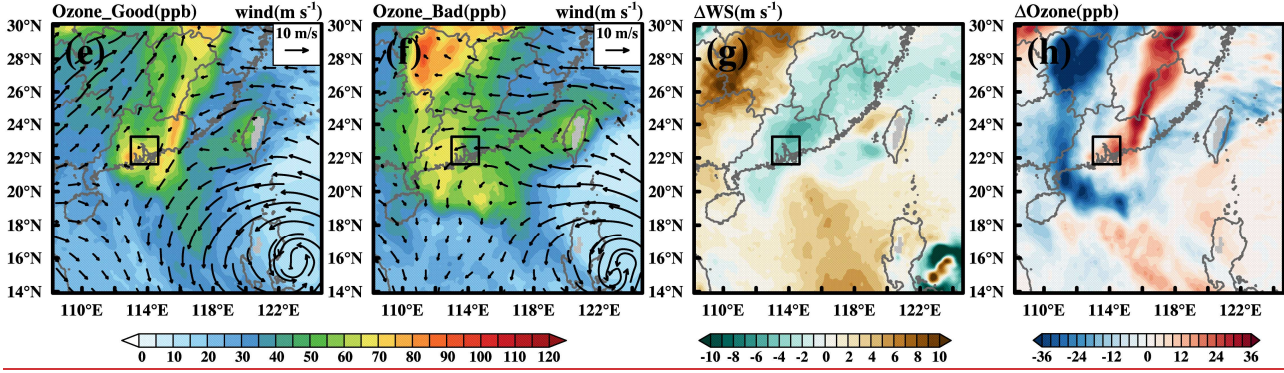


**Figure 6: Mean distributions of the ozone concentration and wind fields and their respective differences between the good and bad EMs in the lower (a-d) and higher layers (e-h).**

### Ozone and wind fields in lower layers (0-750m) in the evening (18:00-23:00 LT) of July 30



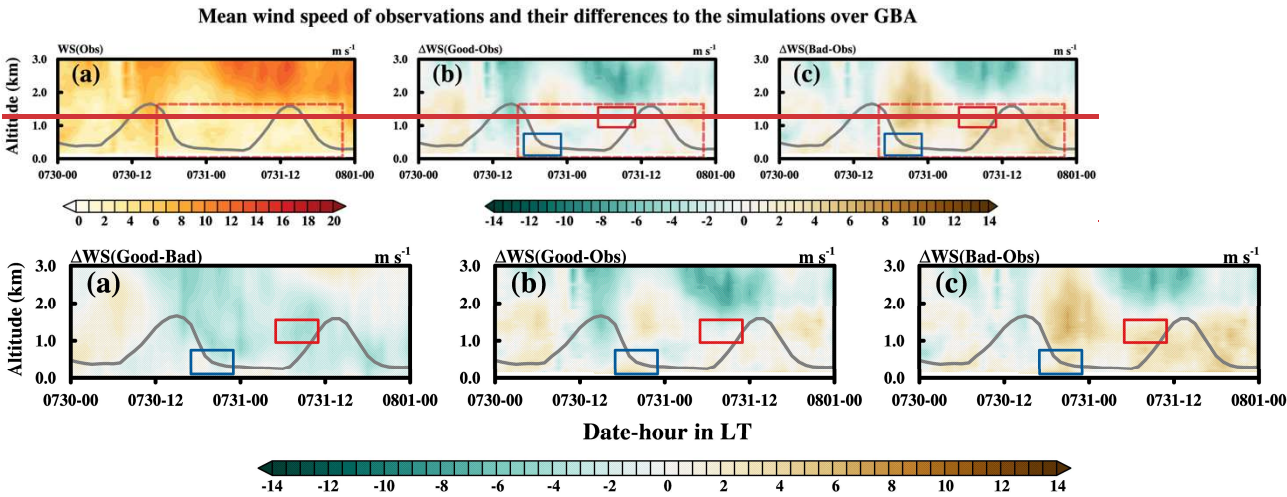
### Ozone and wind fields in higher layers (1000-1500m) in the morning (05:00-11:00 LT) of July 31



**Figure 7: Mean distributions of the ozone concentration and wind fields and their respective differences between the good and bad EMs (good-bad) in the lower (a-d) and higher layers (e-h).**

The changes in WS between good and bad EMs (Good-Bad; Fig. 8a) showed that the WS of good EMs was weaker than that of bad EMs from the surface up to nearly 1700 m. The weak wind could confine ozone in the residual layer from the evening of July 30 to the early morning of July 31. To evaluate the model performance in WS, we collected the vertical sounding observations of wind speed vertical WS from four stations in the GBA (the station codes and locations are listed in Table S2 in the Supplementary supplementary Informationinformation). As shown in Fig. 7a, the observations revealed a low WS value at heights ranging from the surface to nearly 1700 m (dashed red square), which could confine ozone in the residual layer from the evening of July 30 to the early morning of July 31. By comparing the observations with the simulations, it can be concluded that Comparison of the simulations and observations revealed that the vertical wind of the WS of the good EMs agreed better with the observations, since-as their differences were relatively small in the lowest range of 2 km (Fig. 7b8b). In contrast, the bad EMs performed worse. Most of the time, the WS value-of of the bad EMs was greater than that of the observations, especially during the evening of July 30 and the morning of July 31 (Fig. 7e8c). By comparing the differences (Fig. 7b-8b and 7e8c), especially in areas (solid squares) where ADV significantly differed (mentioned in Fig. 5e16c), the WS value-of of the good EMs was closer to the observations, which not only indicates the better model performance

of the good EMs for the vertical ~~wind~~ features of WS but also indicates the important contribution of static winds to the formation of high ozone levels in the residual layer.

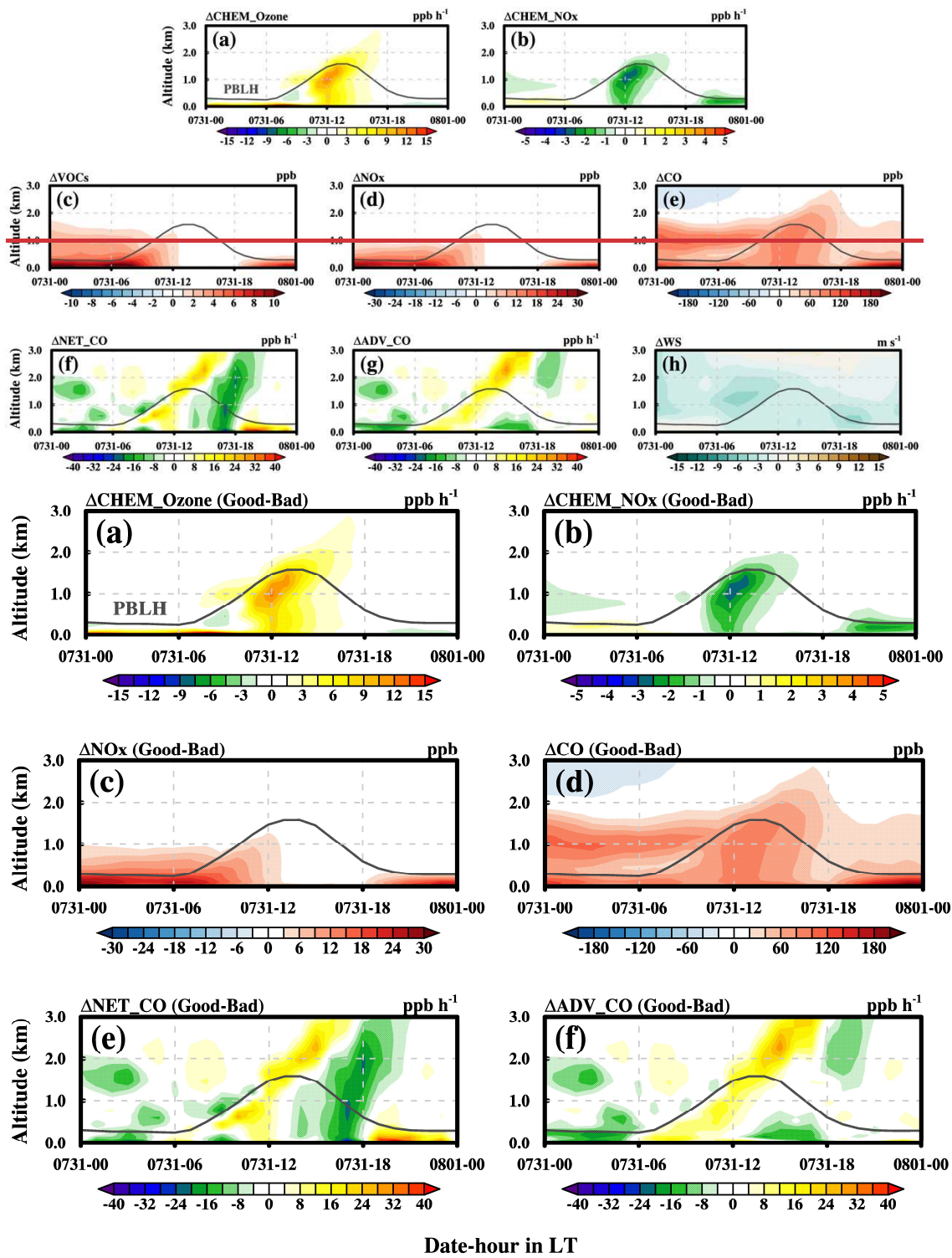


**Figure 78:** Vertical distributions of ~~the three differences of WS. (a) Good-Bad; (b) Good-Obs; (c) Bad-Obs~~WS observations in the GBA (a) and differences of WS between the simulations and observations [(b) for good EMs and (c) for bad EMs].

### 3.2.3 Chemical cause of the extremely high ozone concentration

Enhanced ozone contribution of chemistry (CHEM\_Ozone) ~~-CHEM~~ was another key contributor to the extremely high ozone concentration. The changes in CHEM Ozone-of ozone ( $\Delta$ CHEM\_Ozone) were positive within the PBL from 11:00 to 16:00 LT (Fig. 8a9a), suggesting that the photochemistry in the good EMs was significantly greater than that in the bad EMs. Notably, enhanced photochemistry was more significant in the middle and upper layers of the PBL. Correspondingly, the changes in the NOx contribution of chemistry ( $\Delta$ CHEM\_NOx) showed negative values, which ~~-CHEM-of NOx ( $\Delta$ CHEM\_NOx) in the good EMs decreased significantly, as indicated by the negative values - $\Delta$ CHEM\_NOx (Fig. 8b), which~~ also indicated that enhanced ozone photochemistry occurred in the good EMs.





**Figure 89: Vertical distributions of the changes in CHEM of ozone and the inducing causes on July 31. and the induced factors on July 31. (a) Changes in CHEM of ozone; (b) changes in CHEM of NOx; (c) changes in the concentrations of VOCs; (d) changes in the concentrations of NOx; (e) changes in the concentrations of CO; (f) changes in NET of CO; (g) changes in ADV of CO; and (h) changes in the WS.**

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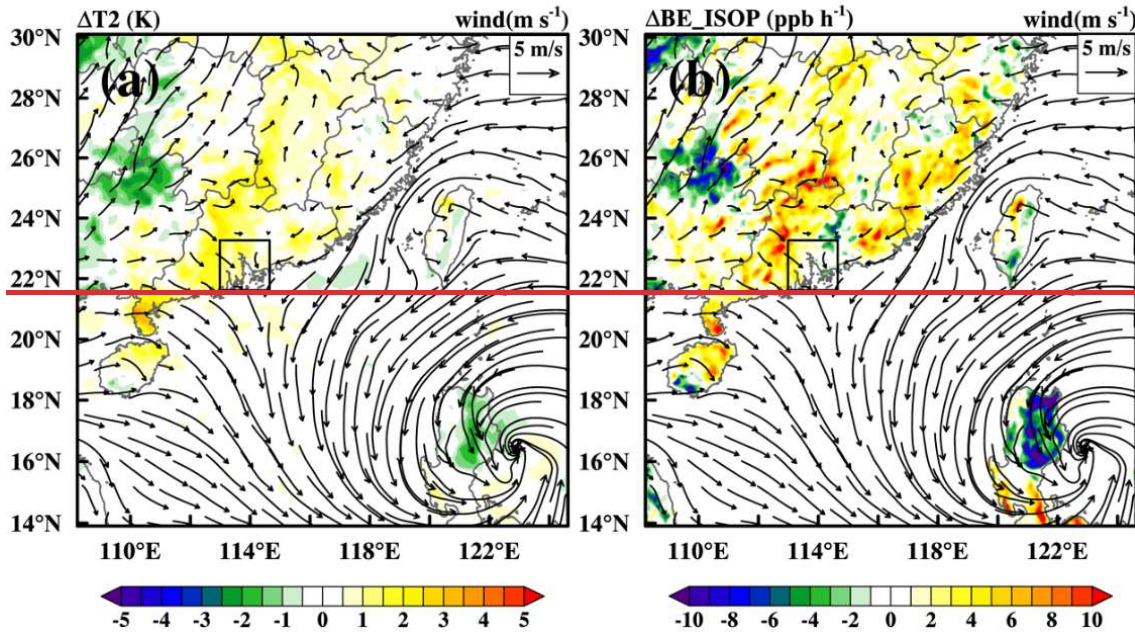
The photochemical production of ozone primarily depends on ozone precursors and chemical reaction constants (Monks et al., 2015). Although the meteorological factors differed significantly between the good and bad EMs, the relevant photochemical reaction constants, such as the photolysis rates (Fig. S2), did not show significant differences in the PBL during daytime. Thus, the enhanced photochemical production of ozone in the good EMs could be attributed primarily to the increase in ozone precursors. Taking NOx as an example, As shown in Fig. 8e and 8d, the changes in NOx ( $\Delta\text{NOx}$ ) showed positive values in the morning which and VOCs ( $\Delta\text{VOCs}$ ) suggested that the good EMs contained more ozone precursors within the PBL and in the morning of July 31, which could enhance ozone photochemistry. Interestingly,  $\Delta\text{NOx}$  and  $\Delta\text{VOCs}$  could approach 0 ppb in the afternoon, which did not align with the significant increase in  $\Delta\text{CHEM\_Ozone}$  during this period. This discrepancy arises because photochemical reactions are rapid, leading to the swift consumption of ozone precursors during their accumulation. Thus, although the meteorological conditions favored the accumulation of ozone precursors, they were consumed quickly by photochemical reactions.

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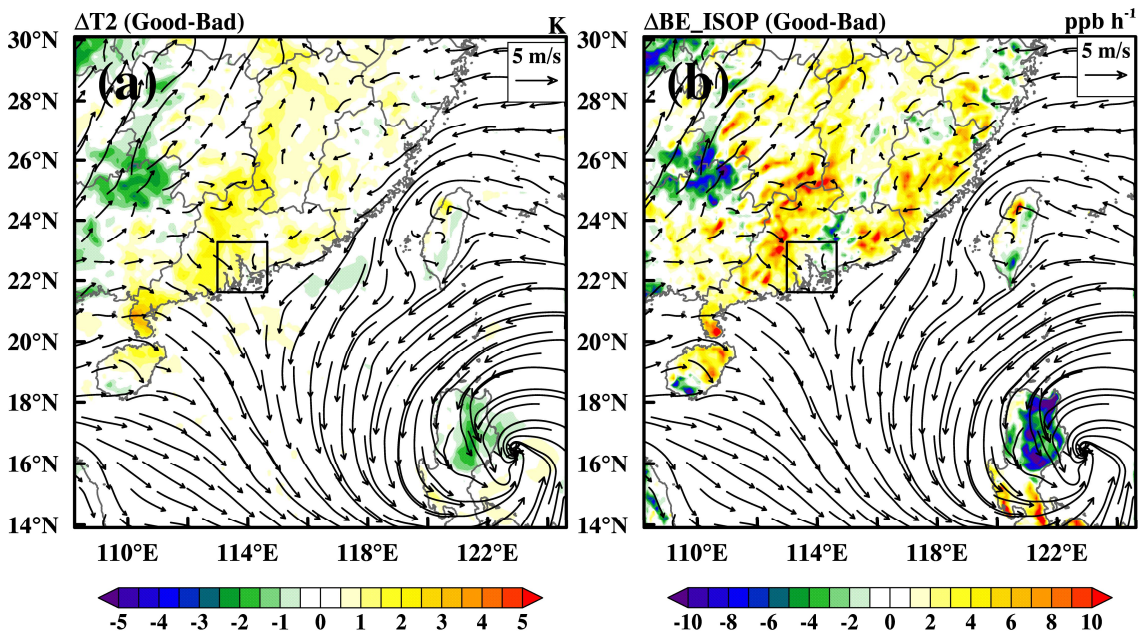
To better characterize the accumulation of ozone precursors, we applied CO as a substitution. Since CO changes induced by chemistry are much lower than those induced by physical processes, the distributions of CO better reflect the accumulation of gaseous pollutants (Ding et al., 2013b). As shown in Fig. 8e9d, the changes in CO showed positive values during the whole day, which indicated the concentrations of CO in the good EMs were much higher than those in the bad EMs within the PBL not only in the morning but also in the afternoon. By comparing the changes in NET of CO (Fig. 8f9e) with those in ADV of CO (Fig. 8g9f), it could be concluded that the increased CO was primarily attributed to the advection process within the PBL during daytime. The enhanced CO contribution of ADV of CO was also due to weak winds. As shown in Fig. 8h8a, the WS in the good EMs was lower than that in the bad EMs within the lowest 2 km during daytime. Lower WS favored the accumulation of gaseous pollutants, particularly ozone precursors, which could subsequently enhance the ozone photochemistry.

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# Mean $\Delta T2$ and $\Delta BE\_ISOP$ in the morning of July 31



# Mean $\Delta T2$ and $\Delta BE\_ISOP$ in the morning of July 31



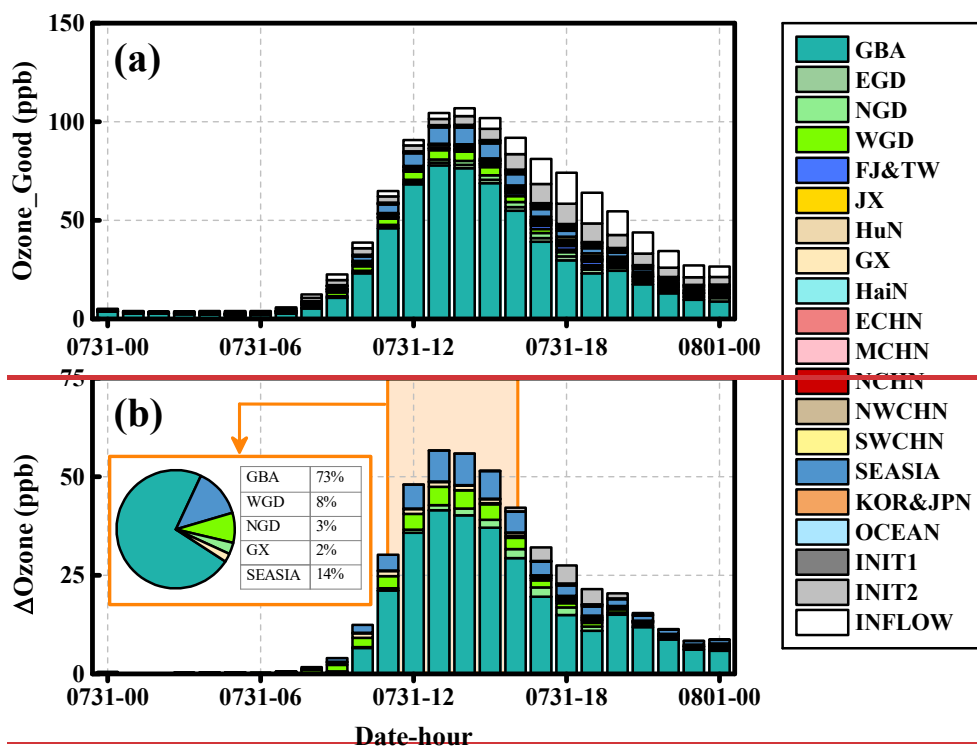
375 Figure 910: Mean distributions of  $\Delta T2$  (a) and  $\Delta BE\_ISOP$  (b) between good and bad EMs (Good-Bad) during the morning (09:00~12:00 LT) of July 31.

In addition, high temperature is a key factor in enhancing biogenic emissions, which can increase the concentrations of VOCs in the atmosphere. Comparative analysis showed that the temperatures in the good EMs were higher in most parts of Guangdong (GD) and Jiangxi (JX) Provinces (Fig. 9a10a). Consequently, increased temperature could lead to the enlargement of leaf stomata, thereby emitting more biogenic VOCs (BVOCs) into the atmosphere (Guenther et al., 2006; 2012). Adopting isoprene as an example (Fig. 9b10b), the changes in the isoprene induced by biogenic emissions ( $\Delta BE\_ISOP$ ) closely matched the changes in T2 ( $\Delta T2$ ), indicating that the higher temperature in the good EMs resulted in increased biogenic VOCs in the atmosphere, especially in the western and northern parts of GD. Northwest winds facilitated the transport of more VOCs to the GBA, promoting the photochemical production of ozone, which is another important chemical cause of the extremely high ozone concentration on July 31. Our results support the findings of a previous study that highlighted the impact of BVOCs on ozone pollution in South China during typhoon approaches (Wang et al., 2022).

### 3.3 Source apportionment of extremely high ozone concentrations

With the implementation of the ozone source apportionment method, the contributions of ozone from various geographical source regions on July 31 were quantified. As illustrated in Fig. 10a11a, the local area made the greatest contribution to surface in the GBA during the daytime. Especially during the high ozone period (11:00~16:00 LT), the mean local contribution could reach 65.4 ppb. In addition to the GBA, the surrounding areas (i.e., WGD) and the remote source region (i.e., SEASIA) showed obvious ozone contributions (with mean contribution of 4.0 ppb and 6.8 ppb, respectively) during this period. In addition, INIT2 and INFLOW, which derived from the initial and boundary conditions of chemistry, also contributed to the surface ozone in the GBA.





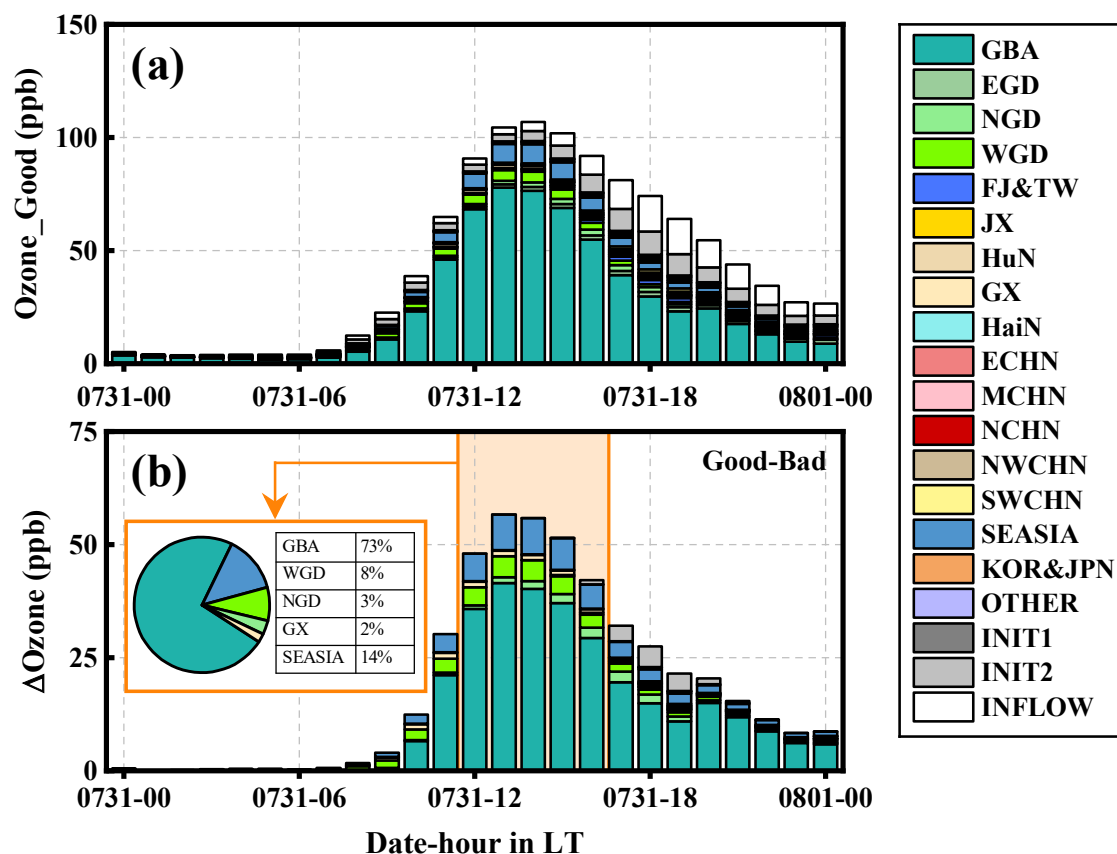
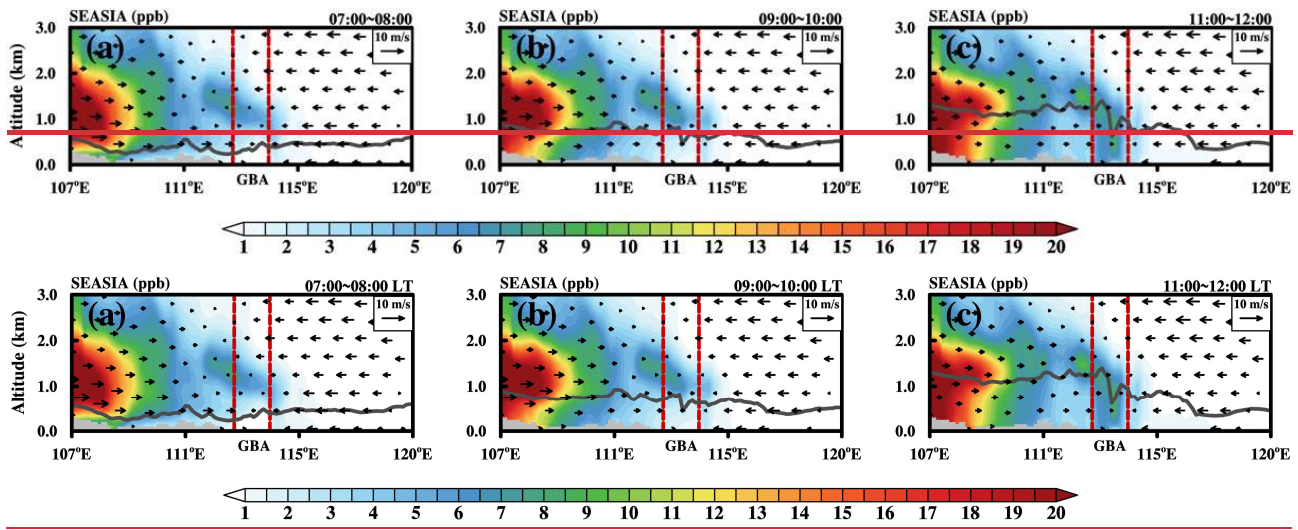


Figure 1011: Mean surface ozone contributions of good EMs (a) and mean changes in surface ozone contributions between the good and bad EMs (b) in the GBA on July 31 (LT).

By comparing with the bad EMs (Fig. 1011b), the changes in ozone contributions were quantified, which could be used to determine the primary contributors to the increase in ozone during the daytime on July 31. It's clear that ozone contributions from the local region (GBA), surrounding regions (WGD, NGD and GX), and remote regions to the west (SEASIA) increased on July 31. During the period from 11:00 to 16:00 LT, their respective proportions in the ozone increase accounted for 73%, 8%, 3%, 2%, and 14%, respectively. In terms of local contribution, the GBA was controlled by stagnant weather, which was not favorable for the diffusion of atmospheric pollutants. As a result, ozone precursors accumulated locally and produced more ozone through photochemical reactions. As shown in Fig. 9b10b, WGD, NGD and GX were controlled by westerly winds during the morning of July 31. Considering the greater amount of BVOCs emitted into the atmosphere over these regions, more ozone and its precursors might be transported and contribute to the increase in surface ozone in the GBA. Our results are consistent with those of previous studies in this region (Li et al., 2012; 2022, Wang et al., 2022).



**Figure 11:** Mean vertical cross sections of the ozone contribution from SEASIA along the west-east direction during the morning on July 31. (a) 07:00~08:00 LT; (b) 09:00~10:00 LT; (c) 11:00~12:00 LT.

As a remote source region, the ozone contribution from SEASIA also significantly increased on July 31. However, unlike the local and surrounding regions, ozone and its precursors from SEASIA are challenging to transport to the GBA at ground level or low altitudes due to dry/wet deposition and chemical consumptions, suggesting a reliance on higher transport pathways. Based on the vertical cross section of the ozone contribution from SEASIA along the west-east direction (Fig. 11), the high-concentration zone was located to the west of the GBA. Under the control of westerly winds, ozone from SEASIA could be transported to high altitudes in the area. Notably, due to the low wind speeds over the GBA, high ozone concentrations from SEASIA were confined to the residual layer in the early morning (Fig. 11a). When the PBL developed, the ozone aloft began to be entrained downward (Fig. 11b). With the continuous rise of the PBL, ozone aloft was kept on being transported to surface through vertical mixing, ultimately contributing to increase surface ozone (Fig. 11c). This finding aligns with the physical cause of the high ozone concentration we discussed in section 3.2.2.

## 4 Conclusion

In the summer of 2016, an extremely high-ozone episode occurred in the GBA when Typhoon NIDA approached. The observations revealed that the maximum concentration reached as high as  $366 \mu\text{g m}^{-3}$  in the afternoon of July 31. Such extremely high concentration poses a challenge for model simulations. To more accurately capture the pollution characteristics of this ozone episode, we conducted an ensemble simulation using the WRF-Chem-O3tag model. By using R and MNB, we introduced an index,  $I_{dis}$ , to select good and bad EMs based on their model performance on ozone. By comparing the good and bad EMs, we quantitatively examined the physical and chemical causes of the extremely high concentrations and the ozone contributions from various geographical source regions in this study.

The extremely high ozone concentration could be explained from both physical and chemical perspectives. From physical perspective, the GBA experienced weak winds as Typhoon NIDA approached. Weak winds facilitated a more significant accumulation of ozone in the residual layer during the evening and early morning. When the PBL developed in the morning, ozone aloft was continuously entrained downward to the surface and then significantly contributed to surface ozone. From chemical perspective, the approach of typhoon provided favorable meteorological conditions for ozone photochemistry, such as increased temperature and more solar irradiance. However, our results revealed that the photochemical reaction constants (i.e., photolysis rate) changed only slightly during the high ozone period. More importantly, weak winds contributed to the accumulation of ozone precursors in the GBA. The increased ozone precursors were the primary reason for the dramatic enhancement of photochemistry. In addition, the increased temperature could increase the concentration of biogenic VOCs by intensifying biogenic emissions from vegetation. The increase in biogenic VOCs could also significantly contribute to the increase in ozone precursors.

The ozone source apportionment results in the good EMs revealed that ozone from the local source region (GBA) contributed most significantly to the high ozone episode. Furthermore, the surrounding regions (i.e., WGD) and remote regions (i.e., SEASIA) also made significant contributions. During the high-concentration period on July 31, the mean ozone contributions from the GBA, WGD, and SEASIA were 65.4 ppb, 4.0 ppb, and 6.8 ppb, respectively. Compared to the bad EMs, the ozone contributions from the GBA, WGD, NGD and GX significantly increased, which was primarily caused by stagnant weather in the GBA and westerly transport effects from outside the area. In addition, ozone from the western and remote region (SEASIA) could be transported to the GBA via westerly winds at high altitudes. When the PBL developed in the morning, ozone aloft was entrained downward, significantly contributing to ~~surfae~~the surface ozone in the GBA.

**Data availability.** We have uploaded the data used in this paper to Zenodo. Please check it via the website <https://doi.org/10.5281/zenodo.13868062>

**Author contributions.** GJ and XH designed the research. XH prepared the input files for the ensemble simulation. GJ ran the model and analyzed the results. GJ wrote the paper. GJ and XH revised the paper.

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

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