

1 **Extreme Weather exacerbates Ozone Pollution in the Pearl**  
2 **River Delta, China: Role of Natural Processes**

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## Abstract

25

26 Ozone (O<sub>3</sub>) pollution research and management in China have mainly  
27 focused on anthropogenic emissions, while the importance of natural  
28 processes is often overlooked. With the increasing frequency of extreme  
29 weather events, the role of natural processes in exacerbating O<sub>3</sub> pollution  
30 is gaining attention. In September 2022, the Pearl River Delta (PRD) in  
31 South China experienced an extended period (25 days) of regional O<sub>3</sub>  
32 exceedances and high temperatures (2<sup>nd</sup> highest over last 2 decades) due to  
33 extreme weather conditions influenced by the Subtropical High and  
34 typhoon peripheries. Employing an integrated approach involving field  
35 measurements, machine learning, and numerical model simulations, we  
36 investigated the impact of weather-induced natural processes on O<sub>3</sub>  
37 pollution by considering meteorological factors, natural emissions,  
38 chemistry pathways, and atmospheric transport. The hot weather  
39 intensified the emission of biogenic volatile organic compounds (BVOC)  
40 by ~10%. **Isoprene and biogenic formaldehyde accounted for 47% of the**  
41 **in-situ O<sub>3</sub> production, underscoring the predominant role of BVOC**  
42 **emissions in natural processes.** The chemical pathway of isoprene  
43 contributing to O<sub>3</sub> formation was further explored, with O<sub>3</sub> production  
44 more attributable to the further degradation of early generation isoprene  
45 oxidation products (**contributed 64.5%**) than the direct isoprene oxidation  
46 itself (**contributed 35.5%**). Besides, it was found that the hot weather  
47 significantly promoted regional photochemical reactions, with  
48 meteorological factors contributing to an additional 10.8 ppb of O<sub>3</sub> levels  
49 compared to normal conditions. Temperature was identified as the  
50 dominant meteorological factor. Furthermore, the typhoon nearing landfall  
51 significantly enhanced the cross-regional transport of O<sub>3</sub> from northern to  
52 southern China through stratosphere-to-troposphere exchange (STE). The  
53 CAM-Chem model simulations revealed that the STE-induced O<sub>3</sub> on the  
54 PRD surface could reach a maximum of ~ 8 ppb, highlighting the non-  
55 negligible impact of STE. This study highlights the importance of natural  
56 processes exacerbated by extreme weather events in O<sub>3</sub> pollution and  
57 provides valuable insights for O<sub>3</sub> pollution control under global warming.

## 58 **1 Introduction**

59 Ground-level ozone (O<sub>3</sub>) is a secondary air pollutant with adverse effects  
60 on human health, vegetation, crop yields, and climate (Knowlton et al.,  
61 2004; Ashmore, 2005; Eyring et al., 2013). The formation of tropospheric  
62 O<sub>3</sub> is a result of **sunlight-driven** photochemical reactions involving  
63 nitrogen oxides (NO<sub>x</sub>), volatile organic compounds (VOC), and other  
64 pollutants (Derwent et al., 1998; Jacob, 2000). The relationship between  
65 O<sub>3</sub> and its precursors exhibits a non-linear pattern that varies across  
66 different regions (Jenkin and Clemitshaw, 2000). O<sub>3</sub> and its precursors  
67 originate from both anthropogenic activities and natural processes such as  
68 fossil fuel combustion, **biogenic volatile compound (BVOC)** emissions and  
69 stratosphere-troposphere exchange (STE). Meteorological conditions also  
70 play a crucial role in influencing O<sub>3</sub> pollution, adding complexity to  
71 mitigation efforts (Wang et al., 2017).

72 Since the industrial revolution, the northern hemisphere has experienced a  
73 significant increase in O<sub>3</sub> pollution, particularly in mid-latitude cities with  
74 large populations and industries (Gaudel et al., 2018). In recent decades,  
75 Europe and the United States have made notable progress in mitigating O<sub>3</sub>  
76 pollution through emission control efforts. However, eastern Asia, notably  
77 China, continues to face a severe O<sub>3</sub> pollution problem (Lyu et al., 2023).  
78 Despite the implementation of strict emission control measures, such as the  
79 Air Pollution Prevention and Control Action Plan and the reduction in fine  
80 particulate matter concentrations, O<sub>3</sub> levels in China have continued to rise  
81 (Wang et al., 2019). Lu et al. (2018) reported that O<sub>3</sub> pollution days in  
82 China are 93 to 575% higher compared to other developed countries,  
83 indicating the significant public concern surrounding this issue.

84 Despite human activities being recognized as major contributors to severe  
85 O<sub>3</sub> pollution, it is also important to acknowledge the role of meteorological  
86 conditions on the dynamics of tropospheric O<sub>3</sub> concentrations. For example,  
87 temperature has a direct impact on chemical reaction rates involved in O<sub>3</sub>  
88 formation as well as the emissions of **BVOC** from vegetation (Lu et al.,  
89 2019). Atmospheric water vapor, on the other hand, plays a crucial role by  
90 providing **hydrogen oxide (HO<sub>x</sub>)** radicals and directly influencing O<sub>3</sub>  
91 photochemistry (Camalier et al., 2007). Additionally, wind patterns

92 contribute to the transport and dispersion of pollutants, thereby influencing  
93 the spatial distribution of O<sub>3</sub> and its precursors (Wang et al., 2022).  
94 Nonetheless, the local meteorological parameters are controlled by  
95 synoptic weather system. Generally, the role of weather systems manifests  
96 in two aspects, one is via the influence on the regional transport of  
97 pollutants, and the other is modulating the aggregation and dispersion of  
98 local air pollutants (Ding et al., 2017). Extensive research conducted in  
99 eastern China has shed light on the importance of weather patterns and  
100 addressed on the impact of extreme weather contributing to O<sub>3</sub> pollution.  
101 Notably, anticyclones (such as high-pressure systems) and the periphery of  
102 typhoons have emerged as prominent factors (Chan and Chan, 2000; Han  
103 et al., 2020; Gao et al., 2021). Besides, **natural processes**, including the  
104 natural sources, chemistry and atmospheric transport of O<sub>3</sub>, are highly  
105 meteorology-sensitive and might further aggravate O<sub>3</sub> pollution under  
106 extreme weather. For example, the wildfire caused by hot and dry weather  
107 could emit large amount of **carbon monoxide (CO)**, NO<sub>x</sub> and VOCs and  
108 exacerbate O<sub>3</sub> pollution (Westerling et al., 2006; Yue and Unger, 2018; Lei  
109 et al., 2022); **vegetation-released** BVOC emissions are sensitive to  
110 temperature and have been proven to increase during hot season and thus  
111 accelerate urban O<sub>3</sub> formation (Pusede et al., 2015; Wang et al., 2022); **an**  
112 **STE** event would bring stratospheric O<sub>3</sub> to the troposphere under a **large-**  
113 **scale**/mesoscale process such as tropopause folds, gravity wave breaking,  
114 and deep convections (Stohl et al., 2003; Wang et al., 2020). As global  
115 warming progresses, there is an increase in the frequency of extreme  
116 weather events which further impact surface O<sub>3</sub> (Banerjee et al., 2016; Lu  
117 et al., 2019). These impacts may undermine or offset the efforts by  
118 anthropogenic emission reductions, posing risks to the ecological  
119 environment. Therefore, understanding the influence of natural process on  
120 the formation of ground-level O<sub>3</sub> is essential for gaining insights into the  
121 dynamics of O<sub>3</sub> pollution and developing effective strategies for managing  
122 air quality.

123 The Pearl River Delta (PRD) region, known for its high levels of  
124 anthropogenic emissions and surrounded by significant vegetation cover,  
125 frequently suffers from extreme weather events such as heat waves and

126 tropical cyclones. The PRD region has emerged as a typical hotspot  
127 witnessing an increase in O<sub>3</sub> pollution, making it an ideal location to  
128 investigate the impact of extreme weather on O<sub>3</sub> pollution. In September  
129 2022, the PRD endured a prolonged period of hot weather, leading to more  
130 than 20 days of regional O<sub>3</sub> exceedance. Here, by integrating simultaneous  
131 measurements, machine learning, and numerical model simulations, we  
132 aim to improve the understanding of how natural processes induced by  
133 extreme weather events affecting O<sub>3</sub> pollution and provide new insights for  
134 future O<sub>3</sub> pollution control efforts.

## 135 **2 Methods and Materials**

### 136 **2.1 Data Source**

137 In-situ observations were conducted at the Guangzhou Haizhu Urban  
138 Ecological Meteorological Comprehensive Observation Base (HZ Base,  
139 23°05'N, 113°22'E), which is located in the North area of the Guangzhou  
140 Haizhu District National Wetland Park (as shown in Fig. 1(a)). The  
141 observation base is surrounded by basic farmland protection areas and  
142 represents a typical composite wetland system consisting of river channels,  
143 creeks, fruit orchards, and the Jiangxinzhou Island. It also encompasses  
144 commercial streets, residential areas, and major transportation routes,  
145 providing a representation of wetland climate and human activities.  
146 Synchronous online observations of O<sub>3</sub>, NO<sub>x</sub>, SO<sub>2</sub>, CO, components of  
147 VOCs, and meteorological parameters (surface winds, temperature,  
148 relatively humidity, precipitation and solar radiation) were carried out at  
149 this observation base. All the data are collected at the HZ Base from Sept.  
150 1<sup>st</sup> to Sept. 30<sup>th</sup>, with a time resolution of 1 hour. Detailed information about  
151 the data used, including the monitoring instruments, data coverage, and  
152 access method, was summarized in Table S1. Briefly, ambient  
153 concentrations of O<sub>3</sub>, NO<sub>x</sub>, CO and SO<sub>2</sub> were routinely measured using  
154 instruments produced by Thermo Scientific (49i-D1NAA, 42i-  
155 DNMSDAA, 48i-DNSAA and 43i-DNSAA, respectively). The species of  
156 VOC components were monitored by GC5000 analysis systems coupled  
157 with flame ionization detectors (FID) from AMA Instruments GmbH  
158 (AMA, Germany). The target compounds of the instrument were the 56  
159 VOCs designated as photochemical precursors by the US Environmental

160 Protection Agency (EPA). The gas standards utilized were identical to  
161 those employed by the US EPA Photochemical Assessment Monitoring  
162 Stations (PAMS). More details were documented in our previous paper  
163 (Zou et al., 2015). All instruments were regularly calibrated and maintained  
164 for different durations. Meteorological data, including temperature, solar  
165 radiation, precipitation, relative humidity and winds, were obtained at the  
166 same site from the China Meteorological Administration. The in-situ  
167 measurements were mainly used to drive a photochemical box model as  
168 described below. Additionally, the air quality monitoring network  
169 established by the Ministry of Ecology and Environment of China was  
170 utilized to identify O<sub>3</sub> pollution event in PRD. There were 56 monitoring  
171 sites distributed in the whole region (Fig. 1(a)) and the 90th percentile of  
172 the maximum daily 8-hour average (MDA8-90) O<sub>3</sub> concentration was  
173 employed to assess the regional degree of O<sub>3</sub> pollution. A regional O<sub>3</sub>  
174 exceedance occurs when the MDA8-90 exceeds the China's Grade II  
175 standard (i.e., 160 µg/m<sup>3</sup>).

176 In addition to the in-situ data, ancillary data from the fifth generation of the  
177 European Centre for Medium-Range Weather Forecasts atmospheric  
178 reanalysis (ERA5) were utilized. The ERA5 data, at a resolution of 0.25°  
179 × 0.25° grid, corresponded to the same time period and interval as the  
180 observed data and provided information such as boundary layer height,  
181 potential vorticity, vertical velocity, geopotential height, specific humidity  
182 and O<sub>3</sub> mass mixing ratio. Besides, typhoon tracks occurred in West Pacific  
183 Ocean were collected from China Meteorology Administration Tropical  
184 Cyclone Data Center (<https://tcdata.typhoon.org.cn>).

## 185 **2.2 Stepwise Regression Analysis**

186 The stepwise regression analysis, as a common method of machine  
187 learning, was used to simulate the dynamics of O<sub>3</sub> concentrations and  
188 quantitatively assess the influence of various meteorological factors on  
189 pollutant variations. The method was designed to construct an optimal  
190 equation by iteratively selecting significant factors while eliminating non-  
191 significant ones to address autocorrelation concerns (Johnsson, 1992). In  
192 this study, simple regressions were firstly performed for each explanatory  
193 variable, and the variable with the most significant contribution was

194 identified. Subsequently, additional variables were gradually introduced,  
195 and the F-test and t-test were employed to evaluate their significance. Non-  
196 significant variables were progressively eliminated until we obtained a  
197 final set of critical explanatory variables. Following variable selection, a  
198 multivariate linear regression equation was established to capture the  
199 variation of O<sub>3</sub> concentrations:

$$200 \quad O_3(t) = \alpha \text{VAR}_1(t) + \beta \text{VAR}_2(t) + \gamma \text{VAR}_3(t) + \dots + \eta \text{VAR}_n(t) + R(t) \quad (1)$$

201 In Eq. (1), O<sub>3</sub>(t) represents the temporal changes in O<sub>3</sub> concentration at  
202 hour t. The coefficients ( $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\eta$ ) for each variable (VAR) were  
203 determined during the stepwise regression process, while R(t) represents  
204 the residual error term. This approach enables us to effectively analyze the  
205 intricate relationship between meteorological factors and O<sub>3</sub>  
206 concentrations, providing valuable insights into the dynamics of O<sub>3</sub>  
207 pollution. This utilized approach has proven effective in our previous  
208 pollutant simulations (Chen et al., 2022). In this study, the model's  
209 performance was validated through Fig. S1, demonstrating a strong  
210 correlation (Pearson correlation coefficient (R) = 0.84 and p-Value (from  
211 two-tailed t-test) < 0.01) between the observed and simulated O<sub>3</sub>  
212 concentrations.

### 213 **2.3 Lagrangian Dispersion Modeling**

214 We conducted backward Lagrangian particulate dispersion modeling  
215 (LPDM) using the Hybrid Single-Particulate Lagrangian Integrated  
216 Trajectory model (HYSPLIT, <https://www.arl.noaa.gov/hysplit/>, last  
217 access: Nov. 15, 2023) to identify the dominant air flow impacting the  
218 receptor area. The meteorology fields were from the Global Data  
219 Assimilation System (GDAS) data. In this study, we released 3000  
220 particulates at 100 m above sea level (a.s.l) over the site (HZ Base) and  
221 tracked their backward movement for 48 hours with a time resolution of 1  
222 hour. The positions of the particulates were determined using both vertical  
223 and horizontal calculations, considering mean wind and turbulence  
224 transport. The model finally identified the "retroplume" footprint  
225 representing the spatial residence time of the particulates and could be  
226 regarded as the distribution of the simulated air mass's surface probability  
227 or residence time.

## 2.4 MEGAN model

The Model of Emissions of Gases and Aerosols from Nature (MEGAN, version 2.1, available at: <https://bai.ess.uci.edu/megan/>, last access Nov. 15, 2023) was used to estimate BVOC emissions from terrestrial ecosystems (Guenther et al., 2012). The model could calculate 147 individual biogenic compounds and lump them into the appropriate VOC mechanisms such as CB05, RACM, SAPRC99 and etc. Herein, the CB05 mechanism was adopted for VOC treatment. Due to the versatility and compatibility, the MEGAN model could be incorporated into many widely used chemical transport models with horizontal resolution ranging from a few kilometers to several hundred kilometers. In this study, the plant function type (PFT) data was from Moderate-Resolution Imaging Spectroradiometer (MODIS) MCD12Q1 product and the leaf area index (LAI) data was from MODIS MCD15A2H product. The input meteorology including temperature, solar radiation, and relative humidity was obtained from Weather Research Forecast-Community Multi Scale Air Quality (WRF-CMAQ) simulations with a horizontal resolution of 12km×12km. The configuration of WRF-CMAQ was summarized in Table S2. In particular, to emphasize the influence of extreme weather, comparisons of BVOC emissions from parallel simulations were conducted. In detail, we employed two sets of meteorological fields to drive the MEGAN model, respectively. One set was based on the WRF-simulated meteorology from September 2022, while the other utilized the average meteorological fields from the preceding three years (2019-2021).

## 2.5 In-situ photochemistry modeling

In-situ O<sub>3</sub> formation was simulated using the Framework for 0-D Atmospheric Modeling (F0AM) incorporating Master Chemical Mechanism v3.3.1 (<https://sites.google.com/site/wolfegm/models>, last access: Nov. 15, 2023). The application method was roughly in line with that in Lyu et al. (2022). Briefly, the model was constrained by observations including O<sub>3</sub>, NO<sub>x</sub>, CO, sulfur dioxide (SO<sub>2</sub>), VOCs species and meteorological parameters collected at HZ Base at hourly resolution. Specifically, HCHO was not measured and was constrained by the WRF-CMAQ simulation results (The validation of WRF-CMAQ was



262 summarized in Table S3). A ‘family conservation’ that set the total  $\text{NO}_x$  to  
263 the observed value every hour and allowed **nitrogen monoxide (NO) and**  
264 **nitrogen dioxide ( $\text{NO}_2$ )** to evolve over time was applied. Photolysis  
265 frequency of  $\text{NO}_2$  observed at HZ Base was input and used to correct the  
266 photolysis frequencies of other species. Net  $\text{O}_3$  production rate (OPR) was  
267 calculated as the difference between  $\text{O}_3$  production rate ( $\text{NO}_2$  production  
268 through NO oxidation by peroxy radicals) and destruction rate ( $\text{NO}_2$   
269 reacting with OH,  $\text{O}_3$  photolysis, ozonolysis of alkenes, and  $\text{O}_3$   
270 consumption by OH and  $\text{HO}_2$ ), in line with the way adopted in previous  
271 studies (Lyu et al., 2019; Wang et al., 2017). The reaction rates for a total  
272 of 17,224 reactions were **extracted** to diagnose the main isoprene-related  
273 pathways leading to  $\text{O}_3$  formation.

## 274 **2.6 CAM-Chem model**

275 The CAM-Chem (Chemical Lagrangian Model of the Stratosphere), a  
276 component of the **National Center for Atmospheric Research (NCAR)**  
277 Community Earth System Model (CESM) version 2.2.0, was utilized to  
278 simulate the impact of **STE** transported  $\text{O}_3$  on the troposphere. The  
279 meteorological fields were obtained from Model-Era Retrospective  
280 analysis for Research and Applications-Version 2 (MERRA-2) and  
281 regrided to a 32-vertical layer with a horizontal resolution of  $0.9 \times 1.25^\circ$ .  
282 The chemistry mechanism employed was the **Model for Ozone and Related**  
283 **Chemical Tracers (MOZART)**. Further information about the model can be  
284 found in the CAM-Chem Wiki (<https://wiki.ucar.edu/display/camchem>).  
285 To validate the model's performance, we compared the distribution of the  
286 simulated  $\text{O}_3$  with  $\text{O}_3$  from AIRS (Atmospheric Infrared Sounder). Both  
287 the monthly averaged distribution and an STE-induced  $\text{O}_3$  intrusion case  
288 were compared (**Fig. S2**). Although the CAM-Chem simulations showed  
289 slightly higher  $\text{O}_3$  levels in southern China, **the correlation coefficient in**  
290 **Eastern China was 0.79 (passed a 95% significance test), indicating the**  
291 **CAM-Chem model relatively well produced  $\text{O}_3$  at higher levels (Fig. S3).**  
292 Besides, it was worth noting that the satellite retrievals themselves contain  
293 uncertainties, mainly from the impact of clouds, aerosols, surface albedo  
294 and the inversion algorithms (Briegleb et al., 1986; De Smedt et al., 2010;  
295 Povey and Grainger, 2015). Overall, the simulated  $\text{O}_3$  showed good

296 agreement with the AIRS data in terms of magnitude and spatial pattern,  
297 indicating satisfactory model performances. Furthermore, a comparison of  
298 the CAM-Chem simulated ground-level O<sub>3</sub> with the surface network  
299 monitoring was also conducted (Fig. S4). The daily magnitude and  
300 variation trend were successfully captured in Guangzhou, with a mean bias  
301 error (MBE) of -7.9 ppb and a root mean square error (RMSE) of 16.3 ppb.  
302 This demonstrated a good reproduction of surface O<sub>3</sub> concentrations.  
303 Indeed, our previous paper **has** also shown the good performance of the  
304 CAM-Chem model application in eastern China (Wang et al., 2023).

### 305 **3 Results and discussion**

#### 306 **3.1 Exacerbation of O<sub>3</sub> Pollution due to Extreme Weather** 307 **Conditions**

308 In September 2022, the PRD region experienced continuous extreme  
309 weather, resulting in a prolonged period of hot weather conditions. As a  
310 consequence, the region encountered 25 consecutive days of regional O<sub>3</sub>  
311 pollution (Fig. 1(b)). Monthly O<sub>3</sub> **concentrations** (MDA8-90 O<sub>3</sub>) were  
312 situated in high levels reaching up to 92 ppb, approximately 20 ppb higher  
313 than the average of the same period during past three years. Meanwhile,  
314 the monthly average daily-maximum temperature soared to 32°C, making  
315 it the second-highest temperature recorded in September over the past 2  
316 decades (Fig. S5). The extremely high temperature appeared to be a  
317 significant driver of O<sub>3</sub> pollution, as evidenced by a high correlation  
318 coefficient of 0.70 (p<0.05) between O<sub>3</sub> levels and air temperature (Fig.  
319 1(b)). From a synoptic weather perspective, the occurrence of hot weather  
320 was the combined effect of multiple tropical cyclones and Western Pacific  
321 Subtropical High (WPSH) (Fig. 1(c)). It was recorded that there were four  
322 tropical cyclones within the one-month time (2022 September) influenced  
323 PRD (described in Table S4). As shown in Fig. 1(c), the combined effects  
324 of tropical cyclones and WPSH resulted in the splitting of the subtropical  
325 high into two parts. One part lingered over the western Pacific Ocean,  
326 while the other remained over the southern region of China, leaving the  
327 PRD under the fully control of the mainland high-pressure system.  
328 Affected by the sinking air flow under the mainland high, the PRD region  
329 experienced conditions characterized by high temperatures, intense solar

330 radiation, low humidity, and reduced precipitation, creating a favorable  
331 environment for photochemical pollution (Fig. 1(d)).

332 To assess the influence of meteorological parameters on O<sub>3</sub> concentrations,  
333 we developed a stepwise regression model to simulate regional O<sub>3</sub>  
334 concentrations. By incorporating an extensive range of input variables,  
335 including surface and 850hPa meteorological factors, we rigorously tested  
336 and identified ten significant factors through T-test analysis. These factors  
337 comprised the following: 2m temperature (T2), boundary layer height  
338 (BLH), surface relative humidity (RH), surface wind speed (WS), 10m U-  
339 component of wind (U10), vertical wind speed (W), 850hPa U-component  
340 of wind (U850), total cloud coverage (TCC), 10m V-component of wind  
341 (V10), and 850hPa V-component of wind (V850). As illustrated in Fig. 2,  
342 meteorological parameters exerted a crucial influence on O<sub>3</sub> concentrations  
343 in 2022, surpassing levels in previous years (2019-2021). This underscored  
344 the profound impact of meteorology on O<sub>3</sub> pollution. Notably, the factor  
345 associated with photochemistry, such as T2, BLH, and RH, played a  
346 substantial role, contributing 43.1 ppb, 35.7 ppb, and -9.3 ppb, respectively,  
347 to the overall O<sub>3</sub> concentration. In particular, the average daily-maximum  
348 air temperature in September maintained at a typically high level (32 °C),  
349 which not only accelerated the rates of photochemical reactions, but also  
350 stimulated the emission of BVOC from vegetation, thereby exacerbating  
351 O<sub>3</sub> concentrations. Furthermore, the increase in BLH and WS compared to  
352 previous years indicated relatively favorable ventilation conditions, which  
353 facilitated the transport of local and upstream pollutants. Subsequent  
354 investigations unveiled that air pollutants from northern regions could be  
355 transported to the PRD, contributing to the observed O<sub>3</sub> concentrations  
356 (refer to Section 3.3). Additionally, September exhibited relatively dry  
357 conditions with lower relative humidity (RH) and less precipitation. Our  
358 model revealed a negative correlation between O<sub>3</sub> and RH, suggesting that  
359 the presence of water vapor contributed to the photochemical removal of  
360 O<sub>3</sub> concentrations (e.g., through HO<sub>x</sub> reactions). The reduced RH in  
361 September also likely facilitated the persistence of O<sub>3</sub> pollution in the  
362 region.

363

### 3.2 Weather-boosted BVOC emissions aggravating O<sub>3</sub> production

364 As one important precursor of O<sub>3</sub> formation, BVOC emissions are sensitive  
365 to ambient temperature and solar radiation. Here, we utilized MEGAN  
366 model to calculate the regional BVOC emissions. Parallel simulations  
367 driven by different meteorological inputs, i.e., meteorological fields in  
368 September 2022 and the average meteorological fields in September of the  
369 previous three years, were conducted, respectively. It was found that the  
370 hot weather in September 2022 led to an increase in BVOC emissions in  
371 the PRD region by approximately 10%, relative to that in the same period  
372 in the past (Fig. 3(a)). Besides, the in-situ observed isoprene exhibited a  
373 significant concentration difference between day and night, i.e., 0.52 – 1.25  
374 ppb during 6:00 – 17:00 and an average of 0.10 ppb at other times (Fig.  
375 S6). Not surprisingly, isoprene contributed 7.77 ppb h<sup>-1</sup> (~ 40%) to the in-  
376 situ net O<sub>3</sub> production rate (OPR) in daytime (Fig. 3(b)). Nevertheless, this  
377 was likely a conservative estimate of the biogenic contributions, due to lack  
378 of consideration of other biogenic VOC. HCHO, as an important O<sub>3</sub>  
379 precursor, is of both anthropogenic and biogenic origin. Here, we utilized  
380 WRF-CMAQ simulated biogenic HCHO as input to examine its impact on  
381 O<sub>3</sub> formation with F0AM. It was found that biogenic HCHO at an average  
382 concentration of 2.46 ppb elevated the OPR by 1.29 ppb h<sup>-1</sup>. This increased  
383 the contribution to OPR of biogenic emissions to 47%. Overall, the  
384 modeling results underlined the crucial role of biogenic emissions in  
385 building up O<sub>3</sub> levels in September 2022.

387 Next, we explored the detailed mechanisms of O<sub>3</sub> formation enhancement  
388 induced by the rise in isoprene levels due to hot weather (Fig. 4).  
389 Simulations were performed for a base case with observations in 2022 and  
390 a hypothetical case of lower isoprene levels. We used the ratio of isoprene  
391 emissions between 2022 and previous years to scale the observed isoprene  
392 in September 2022. So, isoprene in the base case was 10% higher than that  
393 in the hypothetical case. HCHO was constrained by the same profile in  
394 both cases. It was simulated that the 10% increase in isoprene would lead  
395 to an additional O<sub>3</sub> production of 7.5 ppb (OPR of 1.00 ppb h<sup>-1</sup> at 12:00).  
396 While there was little change in the O<sub>3</sub> destruction pathways, the  
397 production of NO<sub>2</sub> through RO<sub>2</sub> + NO and HO<sub>2</sub> + NO (pathways leading

398 to O<sub>3</sub> formation following NO<sub>2</sub> photolysis) increased by 0.63 ppb h<sup>-1</sup> and  
399 0.38 ppb h<sup>-1</sup>, respectively. As shown in Fig. 4, this overall effect was caused  
400 by multiple reactions involving several generations of isoprene oxidation  
401 products/intermediates. The direct oxidation of isoprene by OH and the  
402 following transformation from RO<sub>2</sub> through RO to HO<sub>2</sub> only accounted for  
403 30.3% and 42.8% of the increase in total rate of RO<sub>2</sub> + NO and HO<sub>2</sub>  
404 production, respectively. The rest was contributed by the degradation of  
405 methyl vinyl ketone (MVK) and methacrolein (MACR), two typical  
406 isoprene oxidation products (Pierotti et al., 1990). In particular, the  
407 formation of peroxyacetyl radical (CH<sub>3</sub>CO<sub>3</sub>) was enhanced by 0.16 ppb h<sup>-1</sup>  
408 <sup>1</sup>, which further accelerated the rate of RO<sub>2</sub> oxidizing NO by 0.30 ppb h<sup>-1</sup>  
409 (45.6%) and HO<sub>2</sub> production rate by 0.15 ppb h<sup>-1</sup> (32.5%). Methylglyoxal  
410 (MGLYOX) and CH<sub>3</sub>CO<sub>3</sub> were the key intermediates in photochemical  
411 degradation of MVK and MACR that largely enhanced O<sub>3</sub> formation. The  
412 effect of MVK was much more significant than MACR, which was  
413 reasonable, due to the presence of a more reactive vinyl group in the MVK  
414 molecule.

415 It is well documented that isoprene emitted from vegetation is highly  
416 reactive in the troposphere and is therefore not prone to transport over long  
417 distances. Here, we show that the primary oxidation products of isoprene  
418 that may be formed during air mass transport (Wang et al., 2022),  
419 especially MVK and MACR, make significant contributions to O<sub>3</sub>  
420 formation. This presents to be an important mechanism of isoprene  
421 contributing to O<sub>3</sub> formation. Hence, the impacts of BVOC oxidation  
422 intermediates on downwind air quality warrant more attention (Dreyfus et  
423 al., 2002; Lee et al., 2014).

### 424 **3.3 O<sub>3</sub> enhancement by STE and cross-regional transport**

425 In addition to its influence on meteorological factors and natural emissions,  
426 extreme weather can also impact atmospheric transport, modulating the  
427 regional air quality. For instance, the STE process is a significant natural  
428 process that facilitates the exchange of O<sub>3</sub>-rich air from the stratosphere to  
429 the troposphere, impacting O<sub>3</sub> levels in the lower atmosphere (Wang et al.,  
430 2023). STE often occurs in association with synoptic weather systems such  
431 as cyclones, westerly jet stream, frontal activities and troughs of low

432 pressure (Banerjee et al., 2016). Being affected by the combined influence  
433 of the Subtropical High and typhoons, we diagnosed a continuous STE  
434 event occurring from September 13, 2022, to September 16, 2022.  
435 Initially, on September 13, a trough of low pressure extended from  
436 northwest Inner Mongolia to central China, affecting a large portion of  
437 mainland China (Fig. S7). Concurrently, Typhoon "Muifa" developed near  
438 the coastline in the western Pacific Ocean, leading to the gradual  
439 development of this trough towards the southeastern part of China. On the  
440 15th, the typhoon made landfall in the Yangtze River Delta region. The  
441 combined influence of the typhoon's low-pressure center and the external  
442 strong anticyclone further extended the trough of low pressure southward  
443 (Fig. S7). The dynamic evolution of the weather system facilitated the  
444 favorable conditions for cross-regional transport from higher latitudes of  
445 China to the lower latitudes, such as the PRD region.  
446 Here, we utilized multiple methods to illustrate the impact of the STE-  
447 induced O<sub>3</sub>. First of all, we employed potential vorticity (PV) at 300 hPa  
448 to distinguish between stratospheric and tropospheric air masses,  
449 considering a threshold of 2 potential vorticity units (PVU) as the  
450 dynamical tropopause (Li et al., 2023; Wang et al., 2020). According to Fig.  
451 5(a), a notable high value of PV was observed in eastern China, specifically  
452 spanning from the North China Plain (NCP) area to southern China on  
453 September 14, 2022. This extensive cross-regional transport area is closely  
454 associated with typhoon "Muifa" (as depicted in Fig. S5). The presence of  
455 a strong anticyclone on the outer periphery of the typhoon further  
456 intensified the cross-regional transport in eastern China. This was true with  
457 the LPDM simulation, as it revealed that the PRD region was  
458 predominantly influenced by northerly air flow originating from central  
459 China (Fig. S8). As a result, the potential impact of stratospheric O<sub>3</sub>  
460 intrusion on the troposphere formed a distinct and extensive band that  
461 stretched from the north to the south over eastern China. The subsequent  
462 investigations further supported this finding, as we found similar patterns,  
463 including notable high O<sub>3</sub> distribution at 300 hPa (Fig. 5(b)), low specific  
464 humidity (Fig. 5(c)), and low geopotential height (Fig. 5(d)) along the high  
465 PV area. These patterns suggested that the stratospheric intrusion did

466 transport both dry and O<sub>3</sub>-rich air masses to the troposphere. Meanwhile,  
467 the transported region exhibited a prevailing downward airflow with  
468 positive vertical velocity (Fig. 5(e)), and a distinct high O<sub>3</sub> area was also  
469 observed along the transported band at 700 hPa (Fig. 5(f)), indicating that  
470 the O<sub>3</sub> induced by STE could impact the lower troposphere. Similar  
471 patterns were consistently observed on other days between September 13-  
472 16, 2022 (Fig. S9-S11), confirming the continuous nature of the STE event.  
473 The CAM-Chem model was further adopted to quantify the impact of STE-  
474 induced O<sub>3</sub>. In this model, we introduced a tracer, O<sub>3</sub>S, to represent the  
475 concentration of O<sub>3</sub> from stratosphere. Fig. 6 convinced again the previous  
476 analyses that the transport of O<sub>3</sub>-rich air from the stratosphere to the  
477 troposphere, spanning from the northern to the southern China. The cross-  
478 regional transport of O<sub>3</sub>S was notable at higher levels (between 500hPa and  
479 300hPa) in the troposphere with substantial contributions exceeding 50 ppb.  
480 Though the influence reduced in the lower of the troposphere, the impacted  
481 contribution was still high. The simulated maxima of O<sub>3</sub>S at the surface  
482 level could reach up to approximately 8 ppb, indicating a non-negligible  
483 impact of STE.

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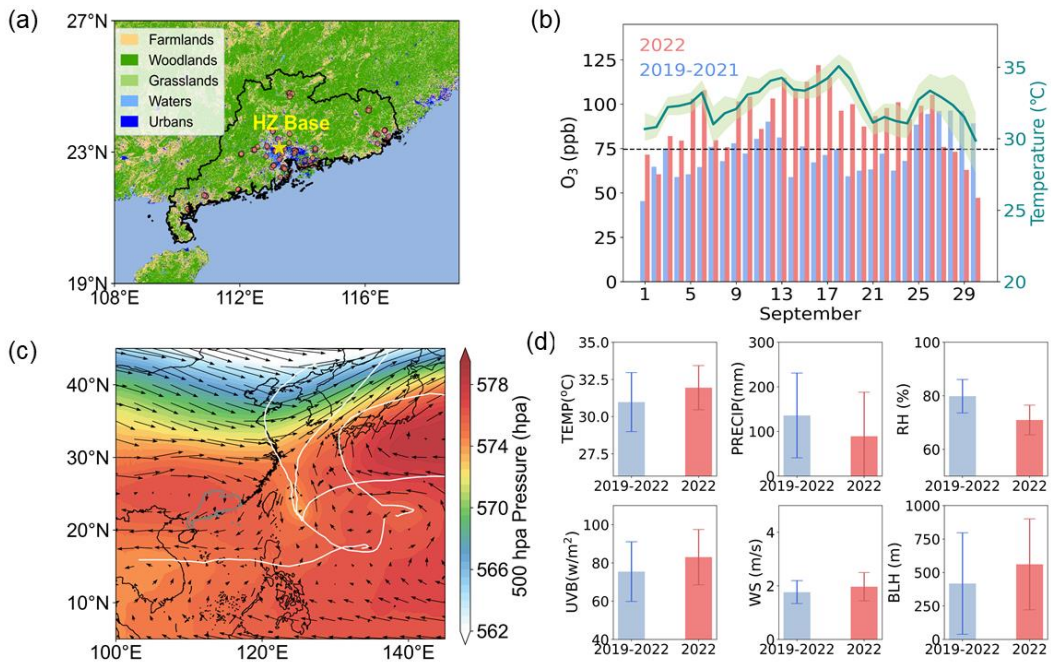
#### 485 **4 Conclusion and implication**

486 This study adopted an integrated methodology, utilizing concurrent  
487 observations, machine learning techniques, and numerical simulations, to  
488 probe how natural processes triggered by extreme hot weather conditions  
489 (the continuous combined influence of the Subtropical High and typhoon  
490 peripheries) on O<sub>3</sub> pollution. Various natural processes, including  
491 meteorological factors, natural emissions, chemistry pathway and  
492 atmospheric transport, were investigated and summarized in Fig. 7. Firstly,  
493 we found that meteorological conditions during extreme weather events,  
494 characterized by high temperatures, high pressure, and low humidity,  
495 greatly facilitated regional photochemical reaction. Through the  
496 application of machine learning techniques, we identified that  
497 meteorological factors contribute an additional 10.8 ppb to O<sub>3</sub> levels  
498 compared to the same period in previous years, with surface temperature  
499 exerting the most prominent influence. Furthermore, our investigation  
500 revealed that the hot weather stimulated BVOC emissions (increased by

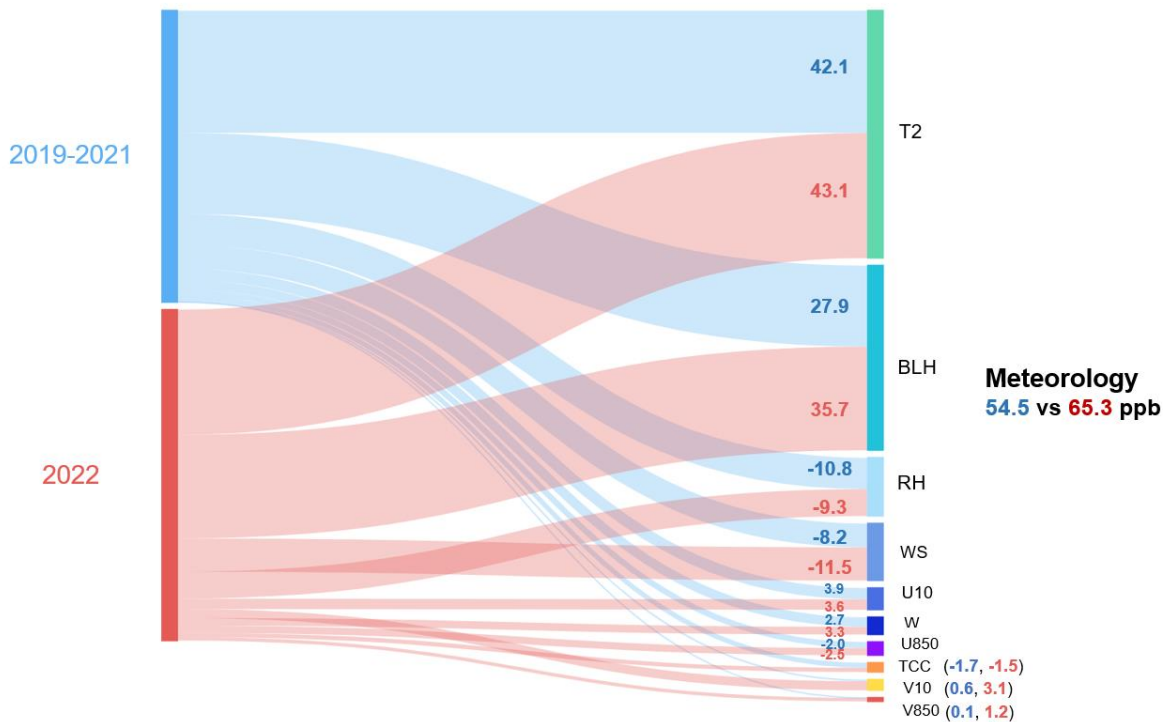
501 10%). Due to the typical high NO<sub>x</sub> environment (mainly from  
502 anthropogenic emission) in the PRD region, BVOC emissions aggravated  
503 photochemical reaction and contributed nearly half of in-situ O<sub>3</sub> production.  
504 The chemical transformation pathways of isoprene and its intermediate  
505 products were further explored, it was found that the further degradation of  
506 initial oxidation products of isoprene was responsible for a large fraction  
507 of isoprene contributions to O<sub>3</sub> formation. This could be an important  
508 mechanism of isoprene affecting downwind air quality. In addition, the  
509 impact of extreme weather on atmospheric transport was also investigated.  
510 The phenomenon of STE usually takes place in high latitudes. Interestingly,  
511 we discovered that the outer periphery of a typhoon, aggravated the cross-  
512 regional transport of STE-induced O<sub>3</sub>, spanning from the northern China  
513 to southern China. This process resulted in a non-negligible contributor to  
514 the surface levels in downwind area (such as the PRD, reached a maximum  
515 of ~ 8 ppb).  
516 Our study underscores the importance of natural processes induced by  
517 extreme weather events in O<sub>3</sub> pollution and provides valuable insights for  
518 future endeavors in O<sub>3</sub> pollution control. Given the impact of climate  
519 change, many regions around the world are experiencing an increase in the  
520 frequency of extreme weather events, thereby intensifying natural  
521 processes. This trend is particularly notable in developed regions with high  
522 levels of anthropogenic emissions, such as eastern China, southeastern  
523 America and northern India. The interaction between natural process and  
524 human activities might further exacerbate air pollution. Future pollution  
525 control and prevention efforts should not solely focus on reducing  
526 anthropogenic emissions. Instead, a comprehensive consideration of both  
527 anthropogenic impact and natural impact should be taken into account, and  
528 a coordinated cross-regional joint emission control is highly recommended.  
529



530 **List of Figures**

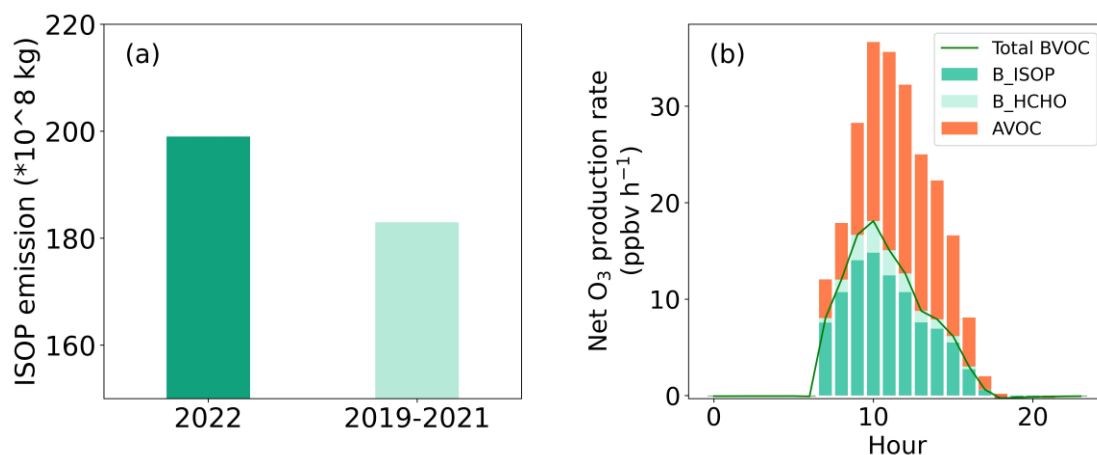


531  
 532 Figure 1 (a) Map showing the geographical distribution of PRD. The dots show the air quality monitoring network and the yellow star shows the in-situ site at Guangzhou Haizhu Urban Ecological Meteorological Comprehensive  
 533 Observation Base (HZ Base); (b) Variation of MDA8-90 O<sub>3</sub> concentrations and regional daily max temperature (The green line shows the average, and the upper and lower shade indicate the 75<sup>th</sup> and 25<sup>th</sup> percentile, respectively); (c)  
 534 Distribution of 500 hPa pressure and winds of September 2022. The white line shows the typhoon track; (d)  
 535 Comparisons of meteorological parameters (temperature (TEMP), precipitation (PRECIP), relative humidity (RH),  
 536 ultraviolet radiation b (UVB), wind speed (WS) and boundary layer height (BLH)) between 2022 and 2019-2021  
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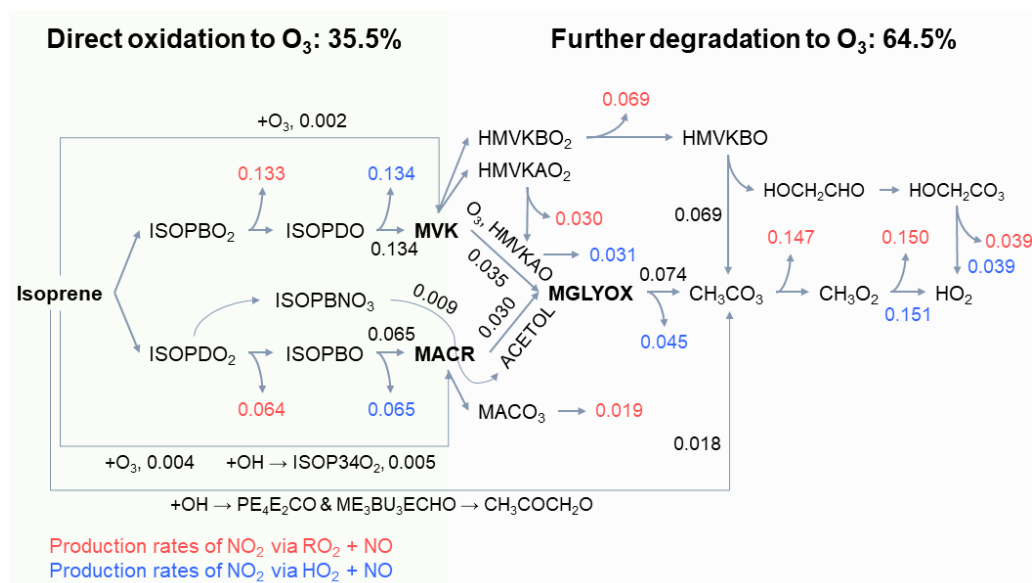


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 540 Figure 2 Contributions of multi-meteorological factors (2m temperature (T2), boundary layer height (BLH),  
 541 relative humidity (RH), wind speed (WS), 10 m u-component of wind (U10), w (vertical wind speed), 850 hPa u-

542 component of wind (U850), total cloud coverage (TCC), 10m V-component of wind (V10), and 850hPa V-  
 543 component of wind (V850)) to O<sub>3</sub> in the September of 2022 and 2019-2021.

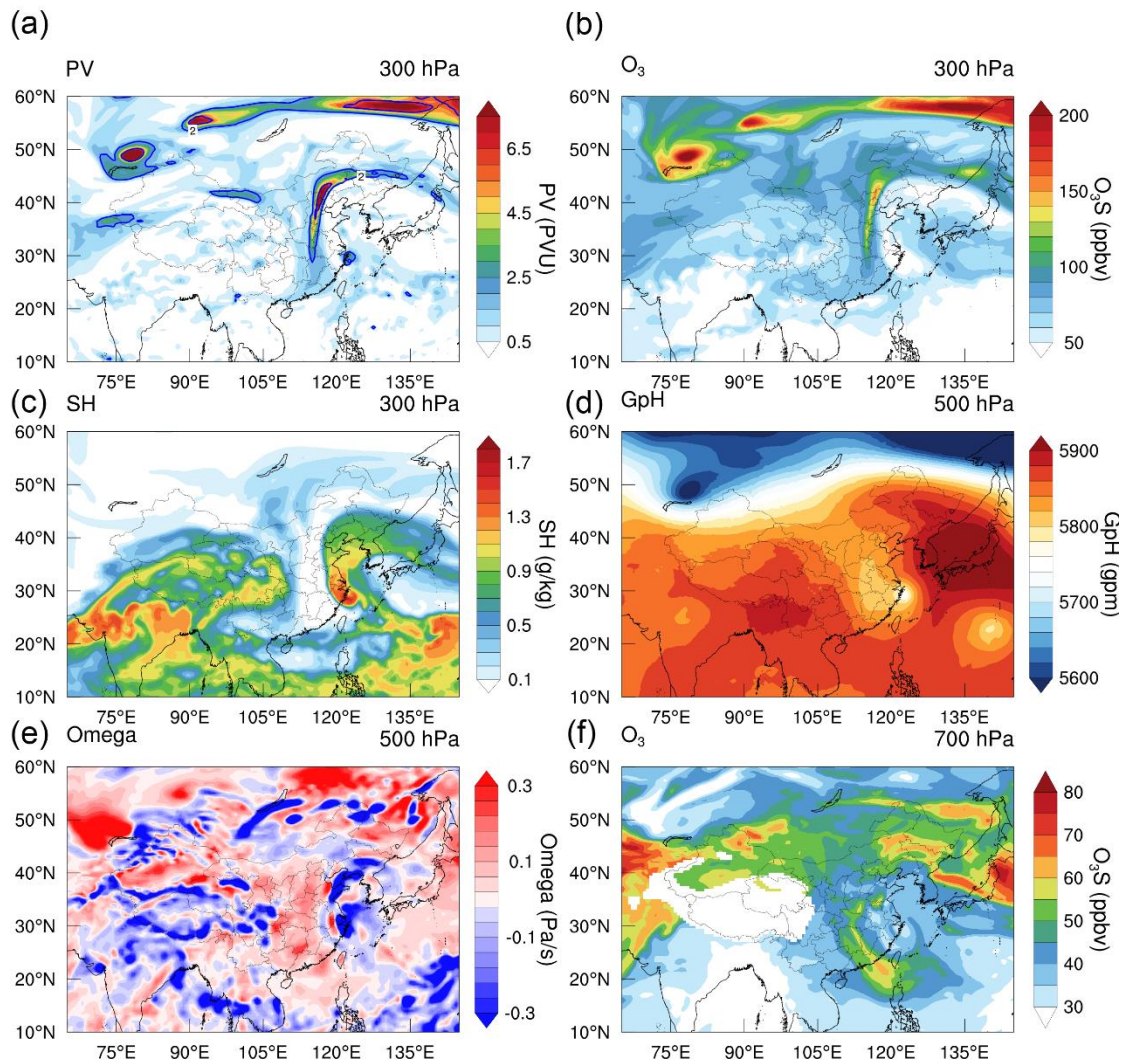


544  
 545 Figure 3 (a) Isoprene emissions in PRD in September in 2022 and 2019 – 2021; (b) Net OPR attributed to biogenic  
 546 isoprene (B\_ISOP), biogenic HCHO (B\_HCHO), total BVOC and anthropogenic VOC (AVOC) in September 2022.  
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549  
 550 Figure 4 Changes in the rates (numbers; unit: ppbv h<sup>-1</sup>) of major reactions leading to O<sub>3</sub> formation at 12:00 induced  
 551 by 10% increase in isoprene concentrations. Red and blue fonts indicate the production rates of NO<sub>2</sub> via RO<sub>2</sub> + NO  
 552 and HO<sub>2</sub> + NO, respectively. Abbreviations of the species conform to the MCM naming convention  
 553 (<http://chmlin9.leeds.ac.uk/MCMv3.3.1/home.htm>).

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557  
 558 Figure 5 Evidence illustrating STE O<sub>3</sub> intrusion on September 14, 2022. (a) Spatial distribution of potential vorticity  
 559 (PV) at 300hPa over China (The blue solid line indicates the dynamical tropopause of 2PVU, 1 PVU=10<sup>-6</sup> m<sup>2</sup> s<sup>-1</sup> K  
 560 kg<sup>-1</sup>); (b-e) The distribution of O<sub>3</sub> concentration (at 300 hPa), specific humidity (at 300 hPa, SH), geopotential height  
 561 (at 500 hPa, GpH), vertical velocity (at 500 hPa, Omega), and O<sub>3</sub> concentration (at 700 hPa), respectively. All the  
 562 data were identified based on ERA5 database.

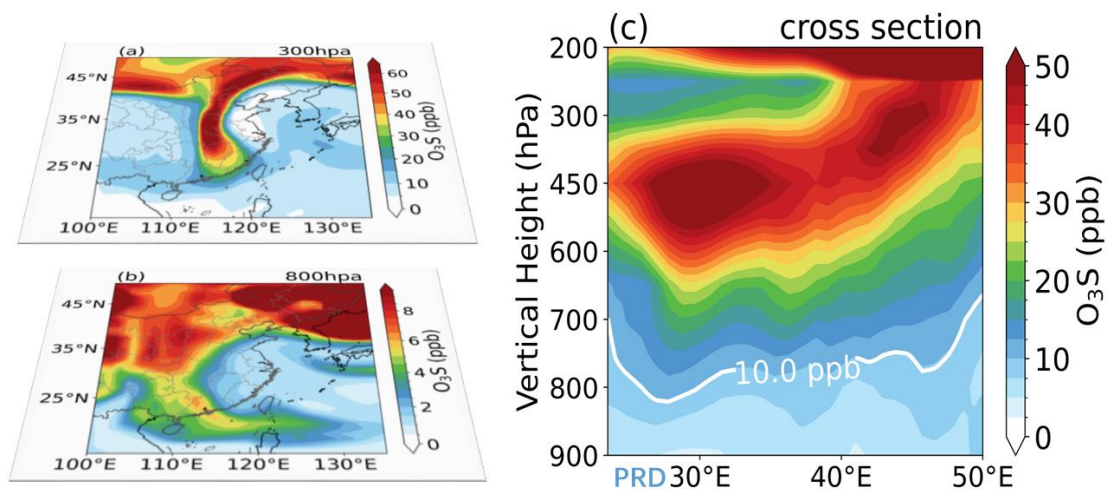
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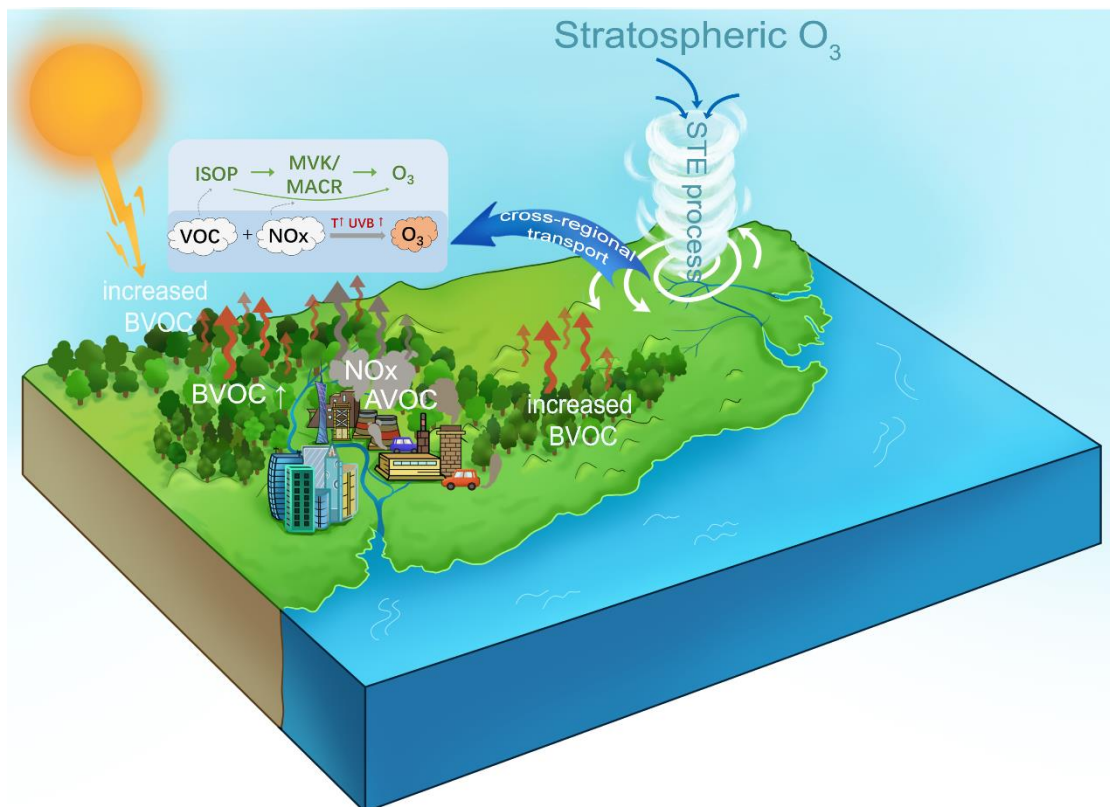


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569 Figure 6 Distribution of CAM-chem simulated O<sub>3</sub>S. (a) O<sub>3</sub>S distribution at 300 hPa; (b) same as (a) but at 800 hPa;  
 570 (c) Vertical transection of O<sub>3</sub>S along the 113°E.

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574 Figure 7 Conceptual scheme illustrating how extreme weather induced natural processes affecting O<sub>3</sub> in PRD. ISOP,  
 575 MVK and MACR refer to isoprene, methyl vinyl ketone and methacrolein, respectively.

576

577 **Associated Content (Supporting Information)**

578 Validation of the stepwise regression model(Fig. S1); Synoptic weather  
579 distribution (Fig. S2); LPDM simulated 48h retroplume (footprint  
580 residence time) (Fig. S3); Evidence illustrating STE O3 intrusion (Fig. S4-  
581 S6); Introduction of monitoring instruments (Table S1); Introduction of the  
582 recorded tropical cyclones (Table S2)

583 **Author Contributions**

584 N.W. designed the research. N.W. and X.L. wrote the manuscript. N.W. ,  
585 X.L., H.W., X.C. and F.Y. contributed to the interpretation of the results.  
586 All authors provided critical feedback and helped shape the research,  
587 analysis, and manuscript.

588 **Competing Interests**

589 The contact author has declared that none of the authors has any competing  
590 interests.

591 **Code/Data availability**

592 The code and data used in this study are available upon request from Nan  
593 Wang ([nan.wang@scu.edu.cn](mailto:nan.wang@scu.edu.cn)) and Xiaopu Lyu ([xiaopu\\_lyu@hkbu.edu.hk](mailto:xiaopu_lyu@hkbu.edu.hk))

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