Atmospheric oxidation capacity and ozone pollution mechanism in a

coastal city of Southeast China: Analysis of a typical photochemical

episode by Observation-Based Model

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Abstract:

A typical multi-day ozone (O₃) pollution event was chosen to explore the atmospheric oxidation 19 capacity (AOC), OH reactivity, radical chemistry, and O₃ pollution mechanism in a coastal city of 20 21 Southeast China, with an Observation-Based Model coupled to the Master Chemical Mechanism (OBM-MCM). The hydroxyl radical (OH) was the predominant oxidant (90±25%) for daytime AOC, while NO₃ 22 23 radical played an important role for AOC during the nighttime (72±9%). Oxygenated volatile organic 24 compounds (OVOCs, 30±8%), NO₂ (29±8%) and CO (25±5%) were the dominant contributors to OH reactivity, accelerating the production of O₃ and recycling of ROx radicals (ROx=OH+HO₂+RO₂). 25 Photolysis of nitrous acid (HONO, 33±14%), O₃ (25±13%), formaldehyde (HCHO, 20±5%), and other 26 OVOCs (17±2%) were major ROx sources, which played initiation roles in atmospheric oxidation 27 28 processes. Combined with regional transport analysis, the reasons for this O₃ episode were the accumulation of local photochemical production and regional transport. The results of sensitivity analysis 29 showed that VOCs were the limiting factor of radical recycling and O₃ formation, and the 5% reduction 30 of O₃ would be achieved by decreasing 20% anthropogenic VOCs, and controlling emissions of aromatics, 31 alkenes, and alkanes with >4 carbons were benefit for ozone pollution mitigation. The findings of this 32 study have significant guidance for emission reduction and regional collaboration on future 33 photochemical pollution control in the relatively clean coastal cities of China and similar countries. 34

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Keywords: Atmospheric oxidation capacity; Radical chemistry; O₃ formation mechanism; OH reactivity;

1 Introduction

Tropospheric ozone (O₃) is mainly produced by photochemical reactions of anthropogenic and natural emitted volatile organic compounds (VOCs) and nitrogen oxides (NOx), and is an important factor resulting in regional air pollution (Zhu et al., 2020; Lu et al., 2018). The elevated O₃ concentrations enhance the atmospheric oxidation capacity (AOC) and have harmful effects on global climate change, ecosystems, and human health (Liu et al., 2019a; Fowler et al., 2009). The formation mechanisms of O₃ pollution are extremely difficult to figure out, due to the complex types and sources of its precursors (Simon et al., 2015). O₃ formation is affected by multiple factors such as O₃ precursor speciation or level, atmospheric oxidation capacity, meteorological conditions and regional transport (Gong and Liao, 2019; Chang et al., 2019). To effectively control the tropospheric O₃ pollution, exploration of the photochemical mechanism and judgment on the controlling factors of O₃ formation become extremely important for scientific community (Chen et al., 2020; Li et al., 2018).

The atmospheric oxidation capacity reflects the essential driving force in tropospheric chemistry, and plays an important place in the loss rates of primary components and production rates of secondary pollutants, thus the key factors to quantify AOC are processes and rates of species being oxidized in the atmosphere (Elshorbany et al., 2009). The atmospheric conditions (such as photolysis rate, meteorology, pollutant concentrations and regional transport) together influence the AOC levels, and the AOC levels in the polluted urban regions are generally much higher than those at the background sites or remote regions due to the dominant limited factor for the significant differences of pollutant concentrations(Geyer et al., 2001; Xue et al., 2016). ROx radicals, including hydroxyl radical (OH), hydro peroxy radical (HO₂) and organic peroxy radical (RO₂), are very important indicators in atmospheric photochemistry and dominate the atmospheric oxidation capacity (Li et al., 2018). Meanwhile, radical chemistry drives the transformation and recycling of O₃ through initiating atmospheric oxidation processes (Wang et al., 2020). Among these radicals, the OH radical accounts for the majority of AOC over 90% during the daytime, thus the OH reactivity (i.e., OH loss) indicates the primary contribution of individual pollutants (Wang et al., 2018a; Mao et al., 2010). Hence, atmospheric oxidation capacity, OH reactivity, and radical chemistry are crucial aspects for understanding the complex atmospheric photochemistry processes (Li et al., 2018). For example, the major ROx sources are the photolysis reaction of O₃, formaldehyde (HCHO), other oxygenated volatile organic compounds (OVOCs), nitrous acid (HONO) and the reactions of O₃ with unsaturated VOCs (Volkamer et al., 2010). The dominant ROx sources at some rural sites were O₃

photolysis and O₃ reactions with VOCs (Li et al., 2018; Martinez et al., 2003), and those at many urban sites were HONO and OVOCs photolysis (Xue et al., 2016; Liu et al., 2012; Emmerson et al., 2005). For oil and gas field sites, there were highly abundant VOCs to promote the formations of O₃, and the contribution of OVOCs photolysis was 2-5 times higher than that in urban areas (Chen et al., 2020; Edwards et al., 2013, 2014). The HONO photolysis was a very important ROx source at the high-altitude or background sites. (Acker et al., 2001; Jiang et al., 2020).

Current studies of atmospheric O₃ photochemical pollution observations have been conducted at the urban, suburban, rural and remote sites around the world (Smith et al., 2006; Eisele et al., 1997; Kanaya et al., 2001; Hofzumahaus et al., 2009; George et al., 1999; Emmerson et al., 2005; Kanaya et al., 2007; Michoud et al., 2012). In China, O₃ photochemical pollution events have been reported in some megacities, such as Beijing, Shanghai, Guangzhou, and Chengdu (Liu et al., 2012; Tan et al., 2019; Zhu et al., 2020; Wang et al., 2020; Liu et al., 2019b; Ling et al., 2017). Few studies on O₃ photochemical pollution in cities with low O₃ precursor emissions have been reported, and the air quality in these areas usually depends on the change of meteorological conditions. In a coastal city of Southeast China, the concentrations of O₃ precursors were higher than those in remote sites and background, but lower than those in most urban and suburban areas, even lower than those in rural regions (Table S1). In a word, O₃ precursor emissions in our observation site were relatively low. Meanwhile, the southeast coastal region is influenced by the East Asian monsoon and acts as an important transport path between the Yangtze River Delta (YRD) and the Pearl River Delta (PRD) (Liu et al., 2020a; Liu et al., 2020b), which is a good 'laboratory' to further explore O₃ photochemical pollution and formation mechanism with relatively low O₃ precursors and complex meteorological conditions (Zhang et al., 2020b; Hu et al., 2020).

The Observation-Based Model (OBM) is widely used to investigate O₃-VOCs-NOx relationships and radical chemistry (Wang et al., 2018a; Tan et al., 2019). The O₃ sensitivity revealed the non-linear relationship between O₃ and its precursors (i.e., VOCs and NOx), which was conducted to investigate O₃ formation mechanism and control strategies (Wang et al., 2020). The OBM combined with the Master Chemical Mechanism (V3.3.1) (OBM-MCM) has been applied to explore the O₃ photochemical pollution mechanism in different environmental conditions (Chen et al., 2020; Li et al., 2018; Xue et al., 2016; Wang et al., 2018). In this study, we chose a typical multi-day O₃ pollution event in the coastal city Xiamen (Fig. S1), when Xiamen was affected by various meteorological conditions, such as typhoon and the West Pacific Subtropical High (WPSH) accompanied by temperature inversion phenomenon. Based on the OBM-MCM analyses, the study aims to clarify (1) the pollution characteristics of O₃ and its precursors, (2) the atmospheric oxidation capacity and radical chemistry, and (3) the O₃ formation mechanism and sensitivity analysis. The results are expected to enhance the understanding of the O₃ formation mechanism

with low O₃ precursor levels, and provide scientific evidence for O₃ pollution control in the coastal cities.

2 Materials and methods

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2.1 Study area and field observations

Xiamen is a coastal city in the southeast area of China, to the west coast of the Taiwan Strait. The field campaigns were carried out at the Atmospheric Environment Observation Supersite (24.61° N. 118.06° E) on the rooftop of around 70 m high building in the Institute of Urban Environment, Chinese Academy of Sciences. The supersite was equipped with complete monitoring instruments, including gas and aerosol species compositions, O₃ precursors, meteorological parameters, and photolysis rate. Criteria air pollutants of O₃, SO₂, NO-NO₂-NO_x, and CO were monitored by commercial instruments TEI 49i, 43i, 42i, and 48i (Thermo Fisher Scientific, USA), respectively. The meteorological parameters of wind speed (WS), wind direction (WD), air temperature (T), pressure (P), and relative humidity (RH) were measured by an ultrasonic atmospherium (150WX, Airmar, USA). HONO was measured with an analyzer for Monitoring Aerosols and Gases in Ambient Air (MARGA, ADI 2080, Applikon Analytical B.V., the Netherlands). A gas chromatography-mass spectrometer (GC-FID/MS, TH-300B, Wuhan, CN) was used for atmospheric VOCs concentrations monitoring, involving about 103 species of VOCs with a 1-hour time resolution. Photolysis frequencies were measured by a photolysis spectrometer (PFS-100, Focused Photonics Inc., Hangzhou, China). The photolysis rate constants include J(O¹D), J(NO₂), J(H₂O₂), J(HONO), J(HCHO), and $J(NO_3)$. Strict quality assurance and quality control were applied, and the detailed descriptions of the monitoring procedures were documented in our previous studies (Zhang et al., 2020b; Wu et al., 2020; Liu et al., 2020a; Liu et al., 2020b; Hu et al., 2020).

2.2 Observation-based chemical box model

In this study, the Observation-Based Model (OBM) combined with the latest version 3.3.1 of MCM (MCM v3.3.1; http://mcm.leeds.ac.uk/MCM/), involving 142 non-methane VOCs and more than 17000 elementary reactions of 6700 primary, secondary and radical species (Jenkin et al., 2003; Saunders et al., 2003), was used to explore the atmospheric oxidation processes and O3 formation mechanisms. The physical process of deposition within the boundary layer height (BLH), which varied from 300 m during nighttime to 1500 m during the daytime in autumn (Li et al., 2018), was considered in the model. Therefore, the dry deposition velocity was utilized to simulate the deposition loss of some reactants in the atmosphere and showed in Table S2, which avoided continuous accumulation of pollutant concentrations in the model (Zhang et al., 2003; Xue et al., 2016).

The observation parameters of the gaseous pollutants (i.e., O_3 , CO, NO, NO_2 , HONO, SO_2 , and VOCs), meteorological parameters (i.e., T, P, and RH), and photolysis rate constants ($J(O^1D)$, $J(NO_2)$, $J(H_2O_2)$, J(HONO), J(HCHO), and $J(NO_3)$) were input into the OBM-MCM model as constraints. The photolysis rates of other molecules such as OVOCs were parameterized by solar zenith angle and then scaled by the measured $J(NO_2)$ (Saunders et al., 2003). We pre-ran for 5 days before running the model to initialize the unmeasured compounds and radicals (Xue et al., 2014).

OBM-MCM is mainly used to simulate in situ atmospheric photochemical processes and quantify the O₃ production rate, AOC, OH reactivity, and ROx radical budgets. Among them, primary sources of ROx, including the photolysis reactions of O₃, HONO, formaldehyde (HCHO), and other OVOCs as well as reactions of VOCs with O₃ and NO₃ radicals, are important (Xue et al., 2016). The termination reactions of ROx are controlled by cross-reactions with NOx (under high NOx conditions) and ROx (under low NOx conditions) to form nitric acid, organic nitrates, and peroxides (Liu et al., 2012; Xue et al., 2016). Table 1 shows the production and destruction reactions and relevant reaction rates of O₃ in our study. The production rate of O₃ (P(O₃)) includes RO₂+NO (R1) and HO₂+NO reactions (R2, Eq. 1), and the destruction of O₃ (D(O₃)) involves reactions of O₃ photolysis (R3), O₃+OH (R4), O₃+HO₂ (R5), NO₂+OH (R6), O₃+VOCs (R7), and NO₃+VOCs (R8, Eq. 2). The net O₃ production rate (Pnet(O₃)) is calculated by P(O₃) minus D(O₃) as equation 3.

Table 1. Simulated production and destruction reactions and relevant reaction rates of O₃ in our study.

Reactions	Reaction rates	Number				
O ₃ production pathways-P(O ₃)						
$RO_2+NO \rightarrow RO+NO_2$ $2.7\times10^{-12}\times EXP(360/T)$		R1				
$HO_2 +NO \rightarrow OH +NO_2$	$3.45 \times 10^{-12} \times EXP(270/T)$	R2				
O ₃ loss pathways-D(O ₃)						
$O_3+hv \rightarrow O^1D+O_2$	JO_1D	R3a				
$O^1D+H_2O\longrightarrow OH$	2.14×10^{-10}	R3b				
$O_3+OH\rightarrow HO_2$	$1.70 \times 10^{-12} \times EXP(-940/T)$	R4				
$O_3+HO_2\longrightarrow OH$	$2.03\times10^{-16}\times(T/300)^{4.57}\times EXP(693/T)$	R5				
NO ₂ +OH→HNO ₃	$\begin{array}{c} 3.2\times10^{-30}\times9.7\times10^{18}\times\text{P/T}\times(\text{T/300})^{-4.5}\times3.0^{-11}\times10^{\log_{10}(0.41)}/(1+(\log(3.2^{-30}\times9.7\text{E}\times10^{18}\times\text{P/T}\times(\text{T/300})^{-4.5}\times3.0^{-11}/(0.75-1.27\times(\log_{10}(0.14))^2)/(3.2^{-30}\times9.7\text{E}\times10^{18}\times\text{P/T}\times(\text{T/300})^{-4.5}+3.0^{-11}) \end{array}$	R6				
O ₃ +VOCs→Carbonyls+Criegee biradical	Kcons.1	R7				
$NO_3+VOC_S \rightarrow RO_2$	Kcons.2	R8				

Note: The reaction rates of Kcons.1 and Kcons.2 were constant. There were around 700 reactions of VOCs+NO₃/O₃, and the relevant reaction rates were different, which can be obtained from this website http://mcm.leeds.ac.uk/MCM/.

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$$P(O_3) = k_1[HO_2][NO] + \sum (k_{2i}[RO_2][NO])$$
 (1)

156 $D(O_3) = k_3[O_1D][H_2O] + k_4[O_3][OH] + k_5[O_3][HO_2] + k_6[NO_2][OH] +$

$$\sum (k_{7i}[O_3][unsat.VOCs]) + 2\sum (k_{8i}[NO_3][unsat.VOCs])$$
 (2)

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$$Pnet(O_3) = P(O_3) - D(O_3)$$
 (3)

where *ki* is the related reaction rate constant. Detailed descriptions of the chemistry calculation can be found elsewhere (Chen et al., 2020; Wang et al., 2018a; Xue et al., 2014).

Relative incremental reactivity (RIR), an index to diagnose the sensitivity of O_3 formation to precursors, is defined as the ratio of the differences in O_3 production rate to the difference in precursor concentrations (Chen et al., 2020). Here, the $\Delta X/X$ in the OBM-MCM represents the percentage reduction in the input concentrations of each targeted O_3 precursor group and this value is adopted as 20% (Liu et al., 2020c).

$$RIR = \frac{\Delta P(O_3)/P(O_3)}{\Delta X/X} \tag{4}$$

2.3 Model performance

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The index of agreement (IOA) can be used to judge the reliability of the model simulation results, and its equation is (Liu et al., 2019b):

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$$IOA = 1 - \frac{\sum_{i=1}^{n} (o_i - S_i)^2}{\sum_{i=1}^{n} (|o_i - \bar{o}| - |S_i - \bar{o}|)^2}$$
 (5)

where Si is simulated value, Oi represent observed value, Oi the average observed values, and n is the sample number. The IOA range is 0-1, and the higher the IOA value is, the better agreement between simulated and observed values is. In many studies, when IOA ranges from 0.68 to 0.89 (Wang et al., 2018b), the simulation results are reasonable, and the IOA in our research is 0.80. Hence, the performance of the OBM-MCM model was reasonably acceptable.

2.4 Meteorological data and back trajectory calculation

The backward trajectories of air masses arriving at the observation site were calculated by the MeteoInfo during the episode (Wang et al., 2014). The backward trajectories with 72-h were run with the time resolution of 3 hours at 100 m height above ground level, and starting time was 0:00 LT and the ending Meteorological provided time was 23:00 LT. data were by NOAA ARL (ftp://arlftp.arlhq.noaa.gov/pub/archives/gdas1). The Final Operational Global Analysis data (FNL) is from the Global Data Assimilation System and analyzes results with the model which is also used by the National Center for Environmental Prediction (NCEP) in the Global Forecast System (GFS) (https://rda.ucar.edu/datasets/ds083.2/). The weather charts were conducted using Grid Analysis and Display System (GrADS) with the specific programmed script files. A detailed description of the synoptic information was shown in our previous study (Wu et al., 2019).

3 Results and discussion

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3.1 Overview of observations

The O₃ pollution events frequently appeared in the coastal city Xiamen during autumn time, related to the WPSH, carrying favorable photochemical reaction conditions (high temperature, low RH, and stagnant weather conditions) and encouraging the formation and accumulation of O₃ in the southeast coastal area (Wang et al., 2018a). The daily maximum 8-h-average O₃ concentrations (MDA8h O₃) from 20 to 29 Sep 2019 ranged from 53 to 85 ppby, partly exceeding the Grade II of China's National Ambient Air Quality Standard of 75 ppbv. The time series and descriptive statistics of air pollutants and meteorological parameters during this multi-day O₃ pollution event are shown in Fig. 1 and Table 2. During this period, the dominant wind direction was northeast, with an average wind speed of 1.8±0.9 m·s⁻¹. The maximum hourly temperature was as high as 35 °C, and the average RH was 56.4±12.6%. Our previous study showed that particulate pollution was slight in Xiamen, which could affect solar radiation by light-absorbing component, and the concentrations of particulate matter had not exceeded the National Ambient Air Quality Standard (Class II: 75 µg·m⁻³) for a whole year (Hu et al., 2021; Deng et al., 2020). Therefore, solar radiation intensity and J(NO₂) were strong, compared to those of the Yellow River Delta (Chen et al., 2020), Shanghai (Zhu et al., 2020) and Hong Kong (Xue et al., 2016). In general, these meteorological parameters were conducive to the production and accumulation of O₃. In addition, O₃ concentrations at nighttime kept relatively high (Fig.1), indicating the influence of regional transport and little NO titration (Zhang et al., 2020a; Wu et al., 2020). Figure S2 shows the 72 h back trajectories at the monitoring site. Among them, 80% of the air masses came from the Yellow Sea, and the other 20% air masses originated from the northeast China through long-range transport.

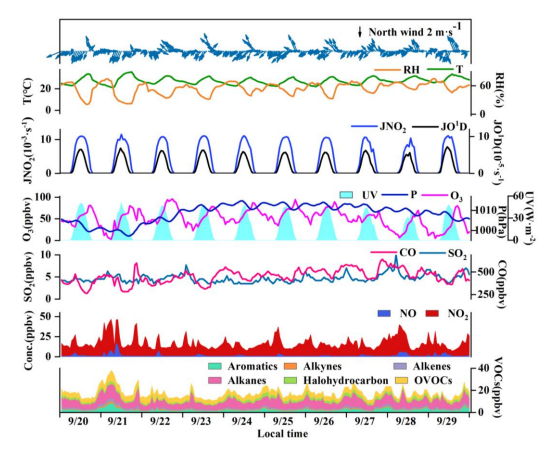


Figure 1. Time series of major trace gases, photolysis rate constants, and meteorological parameters during 20-29 Sep. 2019 in Xiamen.

Table 2. Descriptive statistics of major trace gases (ppbv) and meteorological parameters during 20-29 Sep. 2019.

Parameters	Mean±SD	Median	Max
MDA8h O ₃	67.4±17.2	52.6	89.3
TVOCs	17.2 ± 4.8	16.1	38.0
CO	452±77.3	454	641
NO	1.4±1.3	0.8	17.1
NO_2	15.4 ± 6.9	13.6	40.9
SO_2	4.7 ± 0.9	4.6	10.2
T (°C)	27.3 ± 3.21	26.9	35.6
RH (%)	56.4±12.6	56.6	75.0
P (hPa)	1008 ± 4.57	1010	1015
$UV (W \cdot m^{-2})$	46.4±1.12	0	51.1
Wind speed (m·s ⁻¹)	1.8 ± 0.9	1.6	3.8
Wind direction (°)	90.8±90.4	45.0	337

Table 3. Measured VOCs concentrations during 20-29 Sep. 2019 in Xiamen (Units: pptv), and the classification of VOCs were used and introduced in Section 3.3.

Chemicals	Classification	Mean±SD	Chemicals	Classification	Mean±SD
Aromatics		2131±1236	Alkanes		6970±2325
toluene	RAROM/AHC	995±632	ethane	LRHC/AHC	1552±342
m/p-xylene	RAROM/AHC	392±326	propane	LRHC/AHC	1546 ± 608
benzene	LRHC/AHC	236±95	iso-pentane	C4HC/AHC	930±316
o-xylene	RAROM/AHC	154±121	n-butane	C4HC/AHC	844±365
ethylbenzene	RAROM/AHC	138±94	n-dodecane	C4HC/AHC	618±101
styrene	RAROM/AHC	76 ± 65	iso-butane	C4HC/AHC	494 ± 201
1,2,4-trimethylbenzene	RAROM/AHC	75±37	n-pentane	C4HC/AHC	254±157

m-ethyltoluene	RAROM/AHC	16±11	n-hexane	C4HC/AHC	134±184
p-ethyltoluene	RAROM/AHC	10±6	3-methylhexane	C4HC/AHC	116±93
iso-propylbenzene	RAROM/AHC	5±3	n-heptane	C4HC/AHC	104 ± 78
1,3,5-trimethylbenzene	RAROM/AHC	8±6	3-methylpentane	C4HC/AHC	$82\pm\!48$
o-ethyltoluene	RAROM/AHC	8±5	2-methylhexane	C4HC/AHC	67±38
1,2,3-trimethylbenzene	RAROM/AHC	7±5	2-methylpentane	C4HC/AHC	56±46
n-propylbenzene	RAROM/AHC	7±4	2,3-dimethylbutane	C4HC/AHC	54±33
Halocarbons		1951±572	cyclohexane	C4HC/AHC	42±15
dichloromethane	AHC	998±392	n-undecane	C4HC/AHC	33±35
1,2-dichloroethane	AHC	499±210	n-octane	C4HC/AHC	24±15
chloromethane	AHC	294 ± 75	n-nonane	C4HC/AHC	15±13
1,2-dichloropropane	AHC	88±34	2,2-dimethylbutane	C4HC/AHC	15±7
bromomethane	AHC	47 ± 23	n-decane	C4HC/AHC	14±11
trichloroethene	AHC	15±6	Alkenes		1205±464
1,4-dichlorobenzene	AHC	9±3	ethene	Alkenes/AHC	671±361
OVOCs	AHC	4246±1263	propene	Alkenes/AHC	207±116
acetone	AHC	2802 ± 750	isoprene	BHC	171 ± 232
2-butanone	AHC	799±430	trans-2-pentene	Alkenes/AHC	105 ± 62
2-propanol	AHC	343 ± 283	1-butene	Alkenes/AHC	16±21
2-methoxy-2-methylpropane	AHC	169 ± 97	cis-2-butene	Alkenes/AHC	12±12
acrolein	AHC	66±22	1-pentene	Alkenes/AHC	10±7
4-methyl-2-pentanone	AHC	16±15	1,3-butadiene	Alkenes/AHC	8±7
2-hexanone	AHC	12±3	trans-2-butene	Alkenes/AHC	4 <u>±</u> 4
			Acetylene	LRHC/AHC	674±290

Table 3 lists the detailed VOCs concentrations during the observation period. Alkanes (6970±2325 pptv) were the predominant components of total VOCs, followed by OVOCs (4246±1263 pptv), aromatics (2131±1236 pptv), halocarbons (1951±572 pptv), alkenes (1205±464 pptv), and acetylene (674±290 pptv). The ratio of ethene/ethane (0.4±0.2) was significantly (p<0.05) lower than that in Hong Kong (0.7±0.1) with significant aged air masses, indicating that the high O₃ in Xiamen might be partially attributed to the aged air masses (e.g., transport of air from polluted regions or intense atmospheric oxidation) (Wang et al., 2018a). The concentration of TVOCs in Xiamen (17.2±4.8 ppbv) was much lower than that in the developed areas with large anthropogenic emissions (i.e., Beijing (44.2 ppbv), Lanzhou (45.3 ppbv), Wuhan (30.2 ppbv), Chengdu (36.0 ppbv), Hong Kong (26.9 ppbv), Los Angeles (41.3 ppbv) and Tokyo (43.4 ppbv), but was higher than that at the background and remote sites (i.e., Mt. Wuyi 4.7 ppbv and Mt. Waliguan 2.6 ppbv) (Table S1).

The O₃ formation process depends on its precursors and related environmental conditions, while the photochemical reactions during the daytime are the basis for O₃ changes. Figure 2 shows the diurnal patterns of major trace gases and meteorological parameters during 20-29 Sep. 2019. The O₃ concentration was maintained at relatively low levels from night to 07:00 LT, then rose and reached its maximum at around 17:00 LT. O₃ peak in the afternoon was related to the accumulation of both local

photochemical reaction and potential regional transport (including O_3 and its precursors in the upwind direction to the observation site), and the detailed analysis will be shown in Section 3.3.2. The reduction of observed O_3 (ΔO_3) in the early morning rush hour caused by NO titration did not appear, verifying the impacts of regional transport (Liu et al., 2019b; Zeren et al., 2019; Chen et al., 2020). Due to the photochemical reactions, the precursors of CO, NOx and VOCs were consumed during the daytime, and were accumulated during the nighttime with weak solar radiation. The diurnal patterns of VOCs, NOx and CO were similar, with the highest concentrations at around 08:00 LT and then decreasing during 9:00~16:00 LT and increasing at night, which is related to the human activity emissions (especially vehicle exhaust) and the variations of boundary layer (Elshorbany et al., 2009; Hu et al., 2020).

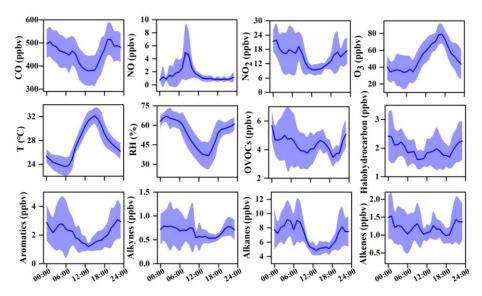


Figure 2. Average diurnal patterns of major trace gases and meteorological parameters during 20-29 Sep. 2019 in Xiamen. The error bar is the standard error.

3.2 Atmospheric oxidation and radical chemistry

3.2.1 Atmospheric oxidation capacity (AOC)

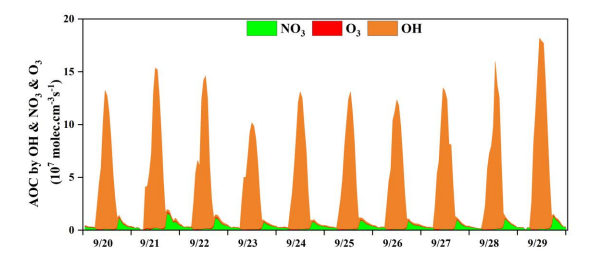


Figure 3. Time series of the model-calculated Atmospheric Oxidation Capacity (AOC) in Xiamen during 20-29 Sep. 2019.

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Figure 3 shows the time series of the model-calculated AOC during the O₃ pollution period. The AOC determines the removal rate of primary pollutants and the production rate of secondary pollutants, and was the basis for reflecting atmospheric photochemical pollution (Geyer et al., 2001). AOC is calculated as the sum of oxidation rates of various primary pollutants (CO, NOx, VOCs, etc.) by the major oxidants (i.e., OH, O₃, NO₃) (Chen et al., 2020; Xue et al., 2016; Xue et al., 2014). In this study, the average daytime AOC was 6.7×10^7 molecules cm⁻³ s⁻¹, and the daily maximum AOC was 1.3×10^8 molecules cm⁻³ s⁻¹, which was higher than those at rural sites with much low pollution emissions in Berlin $(1.4 \times 10^7 \text{ molecules cm}^{-3} \text{ s}^{-1})$ and a regional background in Hong Kong (6.2×10^7) , but lower than that in polluted cities, such as Santiago (3.2×10⁸ molecules cm⁻³ s⁻¹), due to the main limited factor of the significant differences of pollutant concentrations among different sites (Li et al., 2018; Xue et al., 2016; Geyer et al., 2001; Zhu et al., 2020). In some urban regions, the concentrations of air pollutants were higher than those in Xiamen, but their AOC levels (Hong Kong: 1.3×10⁸ molecules cm⁻³ s⁻¹; Shanghai: 1.0×108 molecules cm⁻³ s⁻¹) were comparable to or even lower compared with the AOC in Xiamen, which could be attributed to the relatively high solar radiation (Xue et al., 2016; Zhu et al., 2020) (Detailed descriptions showed in Section 3.1). The results of AOC characteristics in different regions were decided by the precursor concentrations/types and photochemical environment.

According to the diurnal patterns of the AOC contributed by OH, O₃, and NO₃, the predominant oxidant was OH (90±25%) during the daytime, followed by NO₃ (8±22%) and O₃ (2±3%). Meanwhile, the diurnal characteristics of AOC were consistent with the profile of the model-calculated OH (Fig. S3) and the observed photolysis rate constants (Fig.1) (Zhu et al., 2020). Meanwhile, NO₃ (72±9%) played the most important role in the oxidant capability during the nighttime, followed by OH (20±12%) and O₃

(8±1%). In particular, the contribution of NO₃ to AOC reached the maximum of 80% at around 18:00 LT, when the concentrations of O₃ and NO₂ were relatively high and accelerated the formation of NO₃ (Fig.2). In addition, solar radiation was weak during the nighttime, which resulted in the accumulation of NO₃ due to the cease of photolysis of NO₃ (Rollins et al., 2012; Chen et al., 2020). AOC contributed by O₃ was negligible, owing to the relatively low concentration of alkenes at the monitoring site (Fig.1 and Table 3), since O₃ contributed to the oxidation capacity through alkenes ozonolysis (Xue et al., 2016). In summary, the OH radical dominated the AOC, and it was necessary to further explore the partitioning of OH reactivity among different precursor groups.

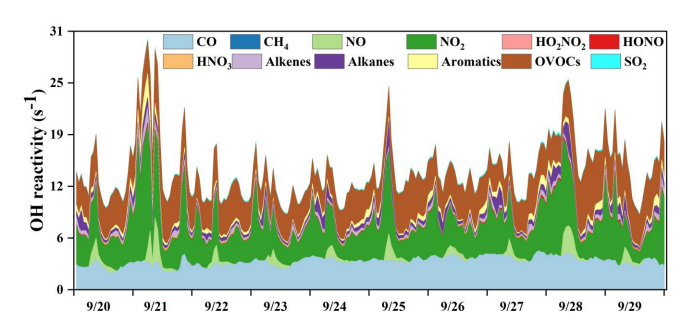


Figure 4. Time series of model-calculated OH reactivity and its partitioning to the major reactants in Xiamen during 20-29 Sep. 2019.

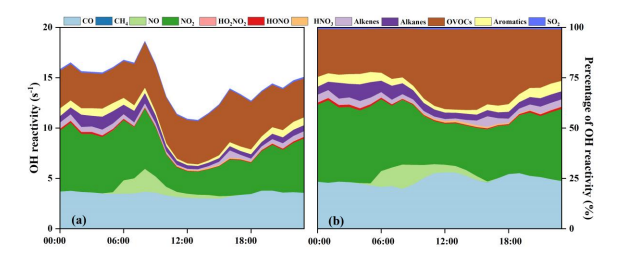


Figure 5. (a) Diurnal patterns and (b) percentage of model-calculated OH reactivity and its partitioning to the

major reactants.

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The OH reactivity is an indicator for the OH chemical loss frequency, computed as the reaction rates of OH with CO, NOx, SO₂, HONO, HNO₃, HO₂NO₂, and VOCs (Whalley et al., 2016; Chen et al., 2020). Time series and diurnal patterns of model-calculated OH reactivity as well as its partitioning to the major reactants during the episode are shown in Fig. 4 and Fig. 5. The OH reactivity reached the peak (18.6±4.8 s⁻¹) at around 8:00 LT, mainly caused by the reaction of OH with NOx, since vehicles exhausted large amounts of NOx during rush hours. The average daily OH reactivity was 14.4±3.83 s⁻¹, which was much lower than those in some polluted regions in Santiago (42 s⁻¹) and the PRD (50 s⁻¹), comparable to that at a rural site in Nashville (11 s⁻¹), but higher than that at a mountain site in Pennsylvania (6 s⁻¹) (Elshorbany et al., 2009; Lou et al., 2010a; Lou et al., 2010b; Kovacs et al., 2003; Ren et al., 2005). Figure 5 shows the diurnal variations and percentage of model-calculated OH reactivity to the major reactants during the episode. The OH reactivity exhibited a morning peak caused by the reactions of NO with OH, which should be ascribed to the freshly emitted urban plumes. Anymore, OVOCs showed high fractions at around 12:00-18:00 LT, which were mainly owing to the transport of the regional air masses containing the abundant OVOCs, as well as the oxidation effection by strong photochemical process. As shown in Fig. 5b, OVOCs (30±8 %), NO₂ (29±8%) and CO (25±5%), were the dominant contributors to OH reactivity, followed by alkanes ($5\pm3\%$), aromatics ($3\pm2\%$), alkenes ($3\pm1\%$), and NO ($2\pm4\%$). The high fraction of OVOCs and NO2 in OH reactivity indicated the high aged degree of air mass and the intensive NO_x emissions during the observation period, respectively (Li et al., 2018). However, the fraction of CO to OH reactivity at our observation site was higher than that at an urban site in Los Angeles (Hansen et al., 2021), a rural site in Hong Kong (Li et al., 2018), and a mountain site in Colorado (Nakashima et al., 2014), comparable to that at the urban site of Shanghai (Zhang et al., 2021a), which could be attributed to the abundant CO in our observation site. CO mainly comes from vehicle exhaust and the combustion of fossil fuels, and the observation site is a city with high density vehicles. Meanwhile, this pollution event was under the influence of the WPSH, which promoted the formation and accumulation of pollutants. The partitioning of OH reactivity elucidated the inherent photochemical processes and major reactants in Southeast China. High OH reactivity of OVOCs, NO2, and CO would promote the production of ROx radical. Therefore, the investigation of detailed chemical budget of the ROx, recycling, and termination reaction is meaningful to figure out the complex atmospheric photochemistry (Li et al., 2018; Lou et al., 2010b).

3.2.2 Radical chemistry

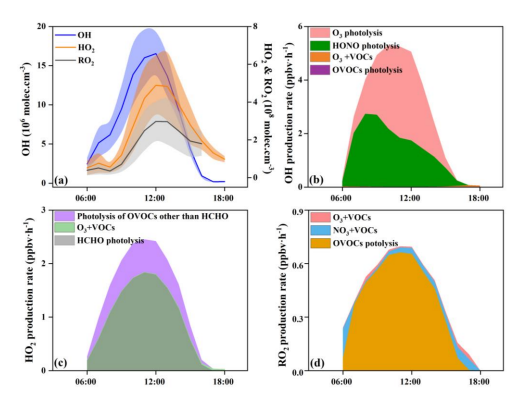


Figure 6. Model-simulated daytime average diurnal variations in (a) OH, HO₂, and RO₂ concentrations, and average primary production rates of (b) OH, (c) HO₂, and (d) RO₂ during 20-29 Sep. 2019 in Xiamen.

With the influence of NOx and VOCs, ROx radicals (OH, HO₂, and RO₂) undergo efficient recycling and produce secondary pollutants, such as O₃ and OVOCs (Sheehy et al., 2010). Figure 6 shows the model-simulated OH, HO₂, and RO₂ concentrations and their primary sources. The detailed time series of ROx concentrations and chemical budget are summarized in Fig. S3. Figure 6a shows the diurnal variations of the simulated OH, HO₂, and RO₂. The maximum daily values of OH, HO₂, and RO₂ concentrations were 2.4×10^7 , 7.9×10^8 and 4.7×10^8 molecules·cm⁻³, with the daytime average concentrations of 7.4×10^6 , 2.4×10^8 and 1.7×10^8 molecules·cm⁻³, respectively. Model-predicted concentrations of OH in Xiamen were higher than that in the Yellow River Delta (an oil field with high VOCs emission), while the concentrations of HO₂ and RO₂ showed reverse trends (Chen et al., 2020). The ROx recycling of OH \rightarrow RO₂ was mainly controlled by the reaction of OH+VOCs, and the RO₂ \rightarrow HO₂ and HO₂ \rightarrow OH depended on the reactions with NO (Fig.7). Combined with the ratio of VOCs/NOx (1.1±0.4), it was convinced that NOx would not be the limiting factor in the radical recycling processes. Hence, efficient conversions of radical propagation of RO₂+NO \rightarrow HO₂ and HO₂+NO \rightarrow OH were expected, and OH+VOCs \rightarrow RO₂ reaction was the rate-depended step of the radical recycling in our study. The detailed radical chemistry would be further discussed as follows.

Figure 6b shows the daytime average diurnal variations of primary OH sources. HONO photolysis

reached the maximum of 2.7 ppb h⁻¹ at around 8:00 LT, which occupied 56±19% of the total OH primary production rates. The second source of OH primary production was O₃ photolysis (42±21%), and the percentages of O₃+VOCs and OVOCs photolysis were minor. The highest HONO photolysis rate appeared in the morning rush hour, suggesting the influence of vehicle emissions and nocturnal accumulation of HONO (Hu et al., 2020). Considering the radical recycling, the reaction of HO₂+NO (8.0±6.2 ppb h⁻¹) dominated the total production of OH (Fig. S3a). Meanwhile, OH-initiated oxidations of VOCs (4.9±3.3 ppb h⁻¹) consumed OH most during the daytime, followed by OH+CO (2.6±1.9 ppb h⁻¹), OH+NO₂ (2.4±1.1 ppb h⁻¹), OH+NO (0.6±0.3 ppb h⁻¹), and OH+O₃ (0.2±0.1 ppb h⁻¹).

In this study, HCHO photolysis was identified as the most important source for HO₂ primary formation, with an average production rate of 1.1±0.6 ppb h⁻¹ (Fig.6c), followed by the other OVOCs photolysis (0.4±0.2 ppb h⁻¹). The rate of OVOCs photolysis in Xiamen was much lower than that in some megacities, such as Beijing (Liu et al., 2012) and Hong Kong (Xue et al., 2016). The reaction of OH+CO (2.6±2.2 ppb h⁻¹) and RO₂+NO (2.5±1.5 ppb h⁻¹) were also important sources of HO₂ (Fig. S3b). The main sink of HO₂ was HO₂+NO (7.9±6.2 ppb h⁻¹), while the loss rates of HO₂+HO₂ and HO₂+RO₂ were negligible.

In Fig. 6d, OVOCs photolysis contributed most to primary RO_2 production with a rate of 0.5 ± 0.2 ppb h^{-1} , accounting for $85\pm20\%$ of total RO_2 primary production. The reaction of unsaturated VOCs and NO_3 was the second important source, accounting for $11\pm18\%$ of the total primary RO_2 . The radical recycling rate of OH+VOCs was 8.4 times higher than the sum of RO_2 primary production. The consumption reaction of RO_2 was mainly caused by RO_2 +NO (3.7 ±2.9 ppb h^{-1}), and the cross-reactions by RO_3 themselves were limited.

The daytime average ROx budget and its recycling were also demonstrated (Fig. 7). For the ROx primary sources, the photolysis of HONO (33±14%), O₃ (25±13%), HCHO (20±5%) and other OVOCs (17±2%) were the major contributors. For ROx recycling, CO and VOCs reacted with OH producing HO₂ and RO₂ with the average rates of 4.0 and 4.4 ppbv h⁻¹, respectively. RO₂+NO and HO₂+NO enhanced the production of RO (3.6 ppbv h⁻¹) and OH (7.9 ppbv h⁻¹), with O₃ formed as a by-product. For the termination processes, the reaction rates of ROx and NOx were approximately 2-5 times faster than the cross-reaction rates of ROx.

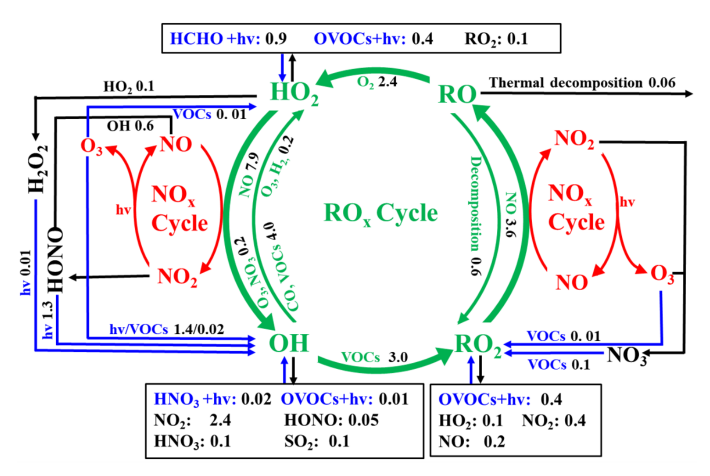


Figure 7. Daytime ROx budget during 20-29 Sep. 2019 in Xiamen. The unit is parts per billion per hour. The blue, black, and green lines and words indicate the production, destruction, recycling pathways of radicals, respectively.

3.3 O₃ formation mechanism

3.3.1 Chemical budget and sensitivity analysis of O₃ production

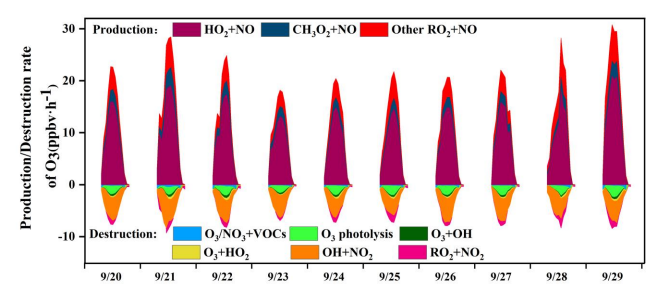


Figure 8. Time series of model-simulated O₃ chemical budgets during 20-29 Sep. 2019 in Xiamen.

The in situ O₃ production mechanism was examined, and the detailed reaction weights were shown

in Fig. 8. The daytime rate of HO_2+NO was 7.9 ± 6.2 ppb h^{-1} , accounting for $68\pm4\%$ of the total O_3 production. This result was consistent with that in section 3.2.2. The OH radical was the initiator of O_3 photochemical formation, and the source of OH from HO_2+NO was also the dominant pathway to produce O_3 (Liu et al., 2020c). The second pathway of O_3 production was RO_2+NO (3.6 ±2.0 ppb h^{-1}). The reaction of RO_2+NO contained more than 1000 types of RO_2 radicals, and the pathway of CH_3O_2+NO (34 $\pm6\%$) contributed mostly among them. In contrast, the contributors of O_3 destruction were O_3+NO_2 (61 $\pm18\%$), followed by O_3 photolysis (18 $\pm9\%$), RO_2+NO_2 (9 $\pm10\%$), O_3+HO_2 (4 $\pm4\%$), and O_3+OH (4 $\pm2\%$), while the other pathways of O_3+VOCs as well as NO_3+VOCs contributed limitedly. In addition, the net O_3 production (9.1 ±5.7 ppb h^{-1}) in Xiamen was ~2-5 times lower than that derived from the metropolis of Shanghai (26 ppb h^{-1}), Lanzhou (23 ppb h^{-1}) and Guangzhou (50 ppb h^{-1}), reflecting the influence of O_3 precursor emissions and photochemical conditions (Xue et al., 2014).

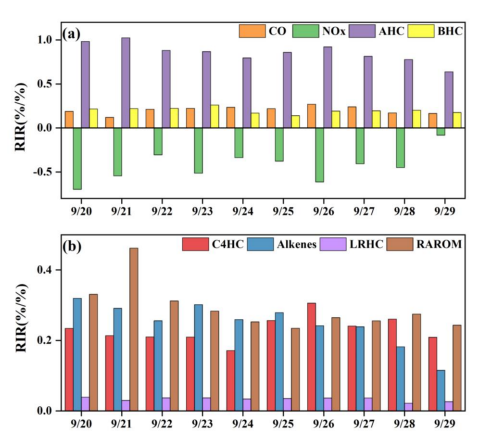


Figure 9. The model-calculated RIRs for (a) major O_3 precursor groups and (b) the AHC sub-groups during high O_3 daytime (06:00-18:00 LT) (AHC: anthropogenic hydrocarbons; BHC: biogenic hydrocarbons; RAROM: aromatics except for benzene; LRHC: low reactivity hydrocarbons; C4HC: alkenes, and alkanes with \geq 4 carbons).

In this study, we also calculated the relative incremental reactivity (RIR) to diagnose the sensitivity of O₃ formation to its precursors. Figure 9 shows the RIR values for major groups of O₃ precursors. Around 50 types of VOCs were classified as anthropogenic hydrocarbons (AHC), and the isoprene was categorized into biogenic hydrocarbons (BHC). Moreover, AHC further divided into four groups of

reactive aromatics (RAROM, including aromatics except for benzene), low reactivity hydrocarbons (LRHC, including ethane, acetylene, propane, and benzene), alkenes, and alkanes with \geq 4 carbons (C4HC). The in situ O₃ production was highly VOCs-sensitive, especially for AHC-sensitive (0.63–1.02 %/%) (Fig. 9a), followed by CO (0.17–0.27 %/%) and BHC (0.14–0.26 %/%), indicating the impacts from anthropogenic activities and flourishing vegetation emissions (Liu et al., 2020a; Lin et al., 2020). The RIRs were NOx-negative ranging from -0.70 to -0.08. As shown in Fig. 9b, the contributors of AHC sub-groups to RIRs were RAROM (0.24–0.46 %/%), C4HC (0.17–0.30 %/%), alkenes (0.11–0.32 %/%), and LRHC (0.03–0.04 %/%). Therefore, the reduction of aromatics, alkenes, and alkanes with \geq 4 carbons effectively decreased O₃ production, and the reduction of NOx might aggravate O₃ pollution.

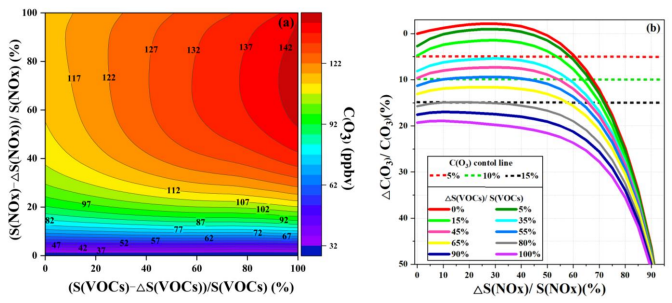


Figure 10. (a) Isopleth diagrams of modeled O_3 production potential $(C(O_3))$ on S(VOCs) and S(NOx) remaining percentages (i.e., (S(VOCs)-S(VOCs))/(S(VOCs)) and (S(NOx)-S(NOx))/(S(NOx)); (b) Relationship of $C(O_3)$ increment percentage $(\Delta C(O_3)/C(O_3))$ with S(NOx) and S(VOCs) reduction percentages $(\Delta S(NOx)/S(NOx))$ and $\Delta S(VOCs)/S(VOCs)$). Note: $C(O_3)$, S(NOx), and S(NOx) represent the concentrations of corresponding pollutants.

In order to investigate the O_3 control strategies during this multi-day O_3 pollution event, the scenario analysis with reduction by 0-100% at intervals of 5% for the reduction of anthropogenic VOCs $(\Delta S(VOCs)/S(VOCs))$ and NOx $(\Delta S(NOx)/S(NOx))$ were conducted using the OBM-MCM. According to the Empirical Kinetic Modeling Approach (EKMA) and scenario analysis, O_3 formation was in the NO-titration regime (Fig. 10), in accordance with those of RIR analysis, which meant VOCs should be reduced to effectively control O_3 during the O_3 pollution event. The maximum value of MDA8h O_3 during the monitoring period was 85 ppbv, exceeding the national air quality standard of 75 ppbv for O_3 by 13%. Hence, the O_3 reductions of 5%, 10%, and 15% were set to discuss the reduction schemes of

anthropogenic VOCs and NOx. As shown in Fig. 10b, achieving the 5% control target were 1) S(VOCs) is reduced by 15%, while S(NOx) remains unchanged; 2) S(VOCs) is reduced larger than 35%; 3) S(NOx) reduction is higher than 60%. The first scenario of just reducing VOCs emission was the most cost-efficient way for short-term or emergency control of O₃. However, NOx, as important precursors of PM_{2.5}, need to be reduced according to the long-term multi-pollutant control air quality improvement plan in China, thus the second scenario is a more practical and reasonable way to control air pollution. The 10% of O₃ control target was achieved by the 45% reduction of S(VOCs), and the S(NOx) keeps original emission. In view of the long-term control strategy of NOx and VOCs, S(VOCs) reduced by 55% and 80% could decrease 10% and 15% O₃ concentrations, respectively. Although VOCs and NOx control measures were drastically implemented, it is still challenging to achieve the 15% O₃ control goals in urban areas with relatively low precursor emissions. As the episode is a typical pollution process in the coastal region, the research results might act as reference for the policy makers. Meanwhile, as the O₃ sensitivity changed under the implementation of control measures, it is necessary to adjust timely the reduction of VOC and NOx policies.

3.3.2 O₃ from local photochemical production and regional transport

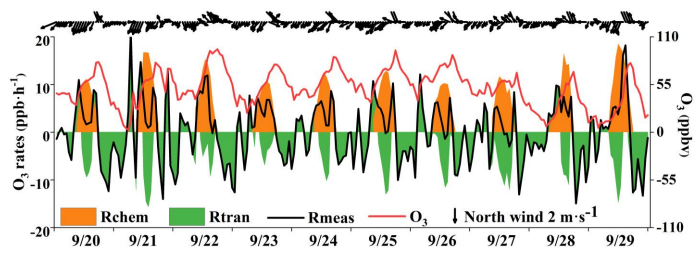


Figure 11. O₃ accumulation and contributions from local photochemical production and regional transport, and Rchem, Rtran, and Rmeas in figure caption represent local O₃ photochemical production, regional transport and observed O₃ formation rate, respectively.

Previous studies have found that the variation of O₃ mixing ratios was mainly influenced by chemical and physical processes (Xue et al., 2014; Tan et al., 2018). Figure 11 shows the time series of O₃ accumulation and contributions from local photochemical production and regional transport. The observed rate of change in O₃ (Rmeas) was calculated by the derivative of the observed O₃ concentrations (Rmeas=d(O₃)/dt). The local O₃ production (Rchem) was calculated by Equation 3, and computed hourly by the OBM as described in Section 2.2. The physical processes (Rtran) were calculated by the equation

of Rtran=Rmeas-Rchem, including horizontal and/or vertical transport, dry deposition dilution mixing, and so on. Many studies showed that the impacts of dry deposition were minor, thus the differences between observed O₃ changes and local O₃ production were mainly caused by the regional transport (note that the effect of atmospheric mixing was also included in this term), which could be treated as regional transport and could reasonably quantify the contributions of regional transport at our observation site (Zhang et al., 2021; Chen et al., 2020). The positive values of Rtran represented the O₃ import of regional transport, while the negative values indicated the O₃ export and deposition. We quantified the contributions of local photochemical formation and regional transport to the observed O₃, and figured out the reasons for the O₃ pollution process.

As shown in Fig. 11, two regular O₃ import phenomenon with positive values of Rtran were observed, and the curve of the Rmeas showed the "M" trend during the daytime. The first transient intense O₃ import happened in the early morning (at around 6:00-9:00), leading to a rapid increase in O₃ concentration, which was mainly attributed to the residual ozone from the day before. The O₃ export was remarkable at around 10:00-16:00, indicating the potential impacts on air quality in downwind areas. Generally, the maximum daily value of O₃ at this observation site appeared at around 15:00 LT without regional transport (Wu et al., 2019). In Figure 11, we found that the O₃ concentrations showed two peaks at around 15:00 and 17:00 LT, and O₃ concentrations rose slowly, or even decreased firstly and then increased between the two peaks. Under these circumstances, the local photochemical production kept producing O₃, but the decreased O₃ concentrations could be attributed to the favorable atmospheric conditions in diluting pollutants (O₃ export). When the near-surface wind direction changed from northeast to southeast, the second O₃ import phenomenon occurred in the afternoon (16:00-19:00 LT) in four days (20, 25 27 and 29 Sep.). Due to the persistence of Rtran in the afternoon, the daily maximum O₃ values appeared at around 17:00 LT. Under the conditions of southeast wind direction, downtown area with high density vehicles would make O₃ and its precursors transmitting to our observation site, consistent with the diurnal patterns of NO₂, OVOCs, alkanes, and aromatic in the early morning and afternoon (Fig. 2) to match with the "M" trend of Rmeas. This result indicated that the sudden changes of near-surface winds were corresponding to the variation in the transport of the urban plume.

According to the synoptic situations and meteorological parameters (Fig. 1, Fig. S4 and Fig.12), the environmental conditions also favored the O₃ pollution process during the observation periods. The contribution of Rchem (daily maximum: ranged from 10.2 to 19.1 ppb h⁻¹) during the daytime was observed (Fig.11). In Fig. S4abc, the monitoring site was continuously affected by the northerly airflow

with high O₃ and its precursors (from an industrial city adjacent to Xiamen of Quanzhou or polluted regions of Yangtze River Delta), due to the typhoon 'Tapah' from 20 to 22 Sep. 2019. The transport of O₃ import appeared on 21 Sep. (7.1±7.0 ppb h⁻¹), which resulted in the accumulation of O₃ (the MDA8h O₃: 85 ppbv) on 22 Sep. When the influence of typhoon disappeared, the direction of airflow turned from northerly into southwest with humid and warm at 500hPa (Fig. S4d), the surface wind on Sep. 23 was affected by the control of the cold northerly airflow (Fig. S4ef). Meteorological conditions including continental high pressure during 23 to 27 Sep. were favorable to the accumulation of air pollutants (Fig. 12). The isoline of 5880 gpm moving from north to the Yangtze River (Fig. 12a,b) indicated the strengthened subtropical high pressure during 23-27 Sep. 2019, which carried high temperature, low RH, and stagnant weather conditions, and the transport rate of O₃ export (5.4±3.4 ppb h⁻¹) on 24-26 Sep. was lower than that on other days (6.3±4.0 ppb h⁻¹). Favorable meteorological conditions significantly affected the formation and accumulation of O₃, and we chose five meteorological parameters (i.e. UV, T, RH, P and WS) to quantify the complex nonlinear relationships between O₃ and its influencing factors based on a generalized additive model (GAM) (Hua et al., 2021). Table S3 showed that the factors had significant non-linear impacts on O₃ concentration changes at the level of P-value<0.01 and degrees of freedom>1, indicating that each influencing factor has statistical significance as an explanatory variable. According to the F-values reflecting the importance of the influencing factors, the orders of the explanatory variables were RH (40.1) > WS (26.9) > T (10.9) > P (3.9) > UV (3.0). Response curves of O₃ concentration to explanatory factors are presented in Fig. 13. The O₃ concentration showed a remarkable upward trend until the UV increased to 17 W·m⁻², then changed little with the fluctuation of UV (Fig. 13a). In previous studies, UV had a significant positive correlation with O₃ concentrations (Ma et al., 2020), and these results showed the regional transport impacts on O₃ formation in our study. The RH and T had negative and positive correlations with O₃ concentrations, respectively (Fig. 13b and Fig. 13c). The increase of wind speed was favorable for O₃ regional transport (Fig. 13d). The influence of atmospheric pressure on O₃ seemed to be irregular and minor, which could be ignored (Fig. 13e). Hence, under the combined effects of favorable photochemical reaction conditions and strengthened WPSH, the MDA8h O₃ exceeded the standard of 75 ppbv during 24-26 Sep. Previous studies had found that severe multi-day O₃ pollution appeared under the WPSH control (Wang et al., 2018a). Overall, the results indicate that the three conditions of local photochemical production, synoptic situations and regional transport played very important roles in the pollution event.

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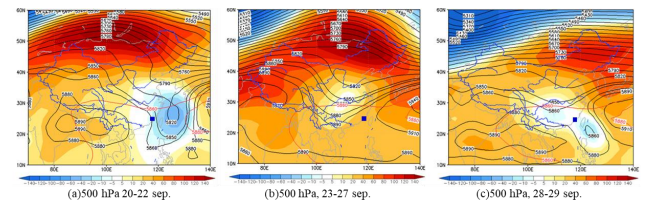


Figure 12. Synoptic situations of continental high pressure from 20 to 29 Sep. 2019. The gradient color area indicates the WPSH over the map and the contour line was from the characteristic isoline of 5880 gpm to the center isoline of 5920 gpm. The blue square is the study site.

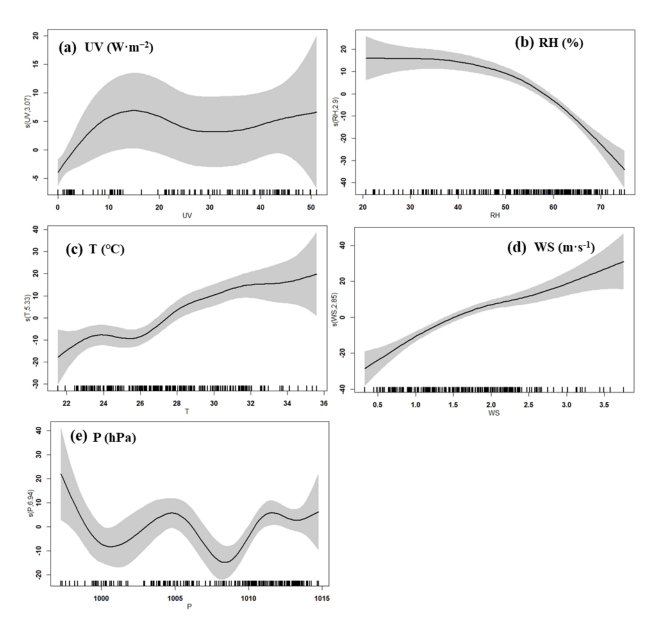


Figure 13. Response curves in GAM model of O₃ concentration to changes in (a) ultraviolet radiation (UV), (b) relative humidity (RH), (c) temperature (T), (d) wind speed (WS), and (e) pressure (P). The y-axis is the smoothing function values. The x-axis is the influencing factor; the vertical short lines represent the concentration distribution characteristics of the explanatory variables; the shaded area around the solid line indicates the 95%

confidence interval of O₃ concentration.

4 Conclusions

In the present study, we analyzed a typical high O_3 event during 20-29 Sep. 2019 in a coastal city of Southeast China. We clarified the characteristics of AOC, OH reactivity, and radical chemistry, as well as O_3 formation mechanisms using the OBM-MCM model. The predominant oxidant for AOC during the daytime and nighttime was the OH and NO_3 , respectively. During the period of O_3 pollution process, OVOCs, NO_2 , and CO consumed OH mostly. Meanwhile, the photolysis of HONO, O_3 , HCHO, and other OVOCs were major sources of ROx, which played the initiation roles in atmospheric oxidation processes. The radical termination reactions were governed by cross-reactions between ROx and NOx. The RIRs and EKMA results showed that the O_3 formation in autumn in the coastal city was VOCs-sensitive, and the VOCs were the limited factor of radical recycling and O_3 formation. The reduced emissions of aromatics, alkenes, and alkanes with \geq 4 carbons were benefit for ozone pollution control. The three conditions of local photochemical production, synoptic situations and regional transport played very important roles in the pollution event. Overall, the results clarified the O_3 pollution process with relatively low local precursor emissions, and implied the fact that O_3 pollution control in coastal cities needs to be further studied.

Code/Data availability

The observation data at this site are available from the authors upon request.

Authorship Contribution Statement

Taotao Liu and Youwei Hong contributed equally to this work. Jinsheng Chen and Likun Xue designed and revised the manuscript. Taotao Liu collected the data, contributed to the data analysis. Taotao Liu and Youwei Hong performed chemical modeling analyses of OBM-MCM and wrote the paper. Jinsheng Chen supported funding of observation and research. Lingling Xu, Mengren Li, Chen Yang, Yangbin Dan, Yingnan Zhang, and Min Zhao contributed to discussions of results. Zhi Huang and Hong Wang provided meteorological conditions in Xiamen.

Competing interests

The authors declare that they have no conflict of interest.

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Acknowledgments

- This study was funded by the Cultivating Project of Strategic Priority Research Program of Chinese
- Academy of Sciences (XDPB1903), the FJIRSM&IUE Joint Research Fund (RHZX-2019-006), the
- 555 Center for Excellence in Regional Atmospheric Environment, CAS (E0L1B20201), the Xiamen Youth
- 556 Innovation Fund Project (3502Z20206094), the foreign cooperation project of Fujian Province
- 557 (2020I0038) and Xiamen Atmospheric Environment Observation and Research Station of Fujian Province.

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Supplementary information

Attached please find supplementary information associated with this article.

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