

## Response to Reviewers

Comment on acp-2021-948

### RC1 Anonymous Referee #1

Comments on “Seasonal characteristics of atmospheric peroxyacetyl nitrate (PAN) in a coastal city 2 of Southeast China: Explanatory factors and photochemical effects” by Liu et al

The manuscript reports influencing factors to PAN pollution in China. The manuscript reports important results. It is suitable for the Journal Atmospheric Chemistry and Physics. I suggest authors incorporate the below suggestions before its publication.

**Response:** Thank you very much for your exploratory and constructive advice. Here, we have carefully revised the manuscript.

Major comments:

- In the abstract section, the authors state that the current paper reports the formation mechanism of PAN and its effect on ozone were identified. I suggest the authors explain it in brief.

**Response:** Thank you for your suggestions. PAN can be produced only through the reaction of PA+NO<sub>2</sub> (Liu et al., 2021; Xue et al., 2014). Hence, the production and sink of PA can represent that of the PAN mechanism indirectly. In the abstract section, we have explained the PA formation mechanism (hence PAN formation mechanism). The effect of PAN on ozone represented that PAN could promote or inhibit O<sub>3</sub> formation under high or low ROx levels, respectively. For better understanding, we revised the relevant expression, as follows:

“Model simulations revealed that acetaldehyde oxidation (46±4%) contributed to the dominant formation pathway of PA (hence PAN), followed by methylglyoxal oxidation (28±3%) and radical cycling (19±3%)”.

“The analysis of PAN formation mechanism and its positive or negative effect on ozone provided scientific insights into photochemical pollution mechanism under various pollution scenarios in coastal areas”.

- In section 2.2, some details of the box model should be added here, although it is explained in the previous study, e.g., details of computation of net production rate of O<sub>3</sub>.

**Response:** Thank you for your suggestions. More details about the box model have been added, as follows:

“The observed data with a time resolution of 1 h of pollutants (i.e., O<sub>3</sub>, CO, NO, NO<sub>2</sub>, HONO, SO<sub>2</sub>, and VOCs), meteorological parameters (i.e., T, P, and RH), and photolysis rate constants ( $J(\text{O}^1\text{D})$ ,  $J(\text{NO}_2)$ ,  $J(\text{H}_2\text{O}_2)$ ,  $J(\text{HONO})$ ,  $J(\text{HCHO})$ , and  $J(\text{NO}_3)$ ), which were mentioned in Section 2.1, were input into the OBM-MCM model as constraints”.

“The production pathways of O<sub>3</sub> include HO<sub>2</sub>+NO and RO<sub>2</sub>+NO reactions, and the destruction pathways of O<sub>3</sub> involve reactions of O<sub>3</sub> photolysis, O<sub>3</sub>+OH, O<sub>3</sub>+HO<sub>2</sub>, O<sub>3</sub>+VOCs, NO<sub>2</sub>+OH, and NO<sub>3</sub>+VOCs. The net O<sub>3</sub> production rate ( $P(\text{O}_3)$ ) is calculated by the difference of O<sub>3</sub> production rate and destruction rate”.

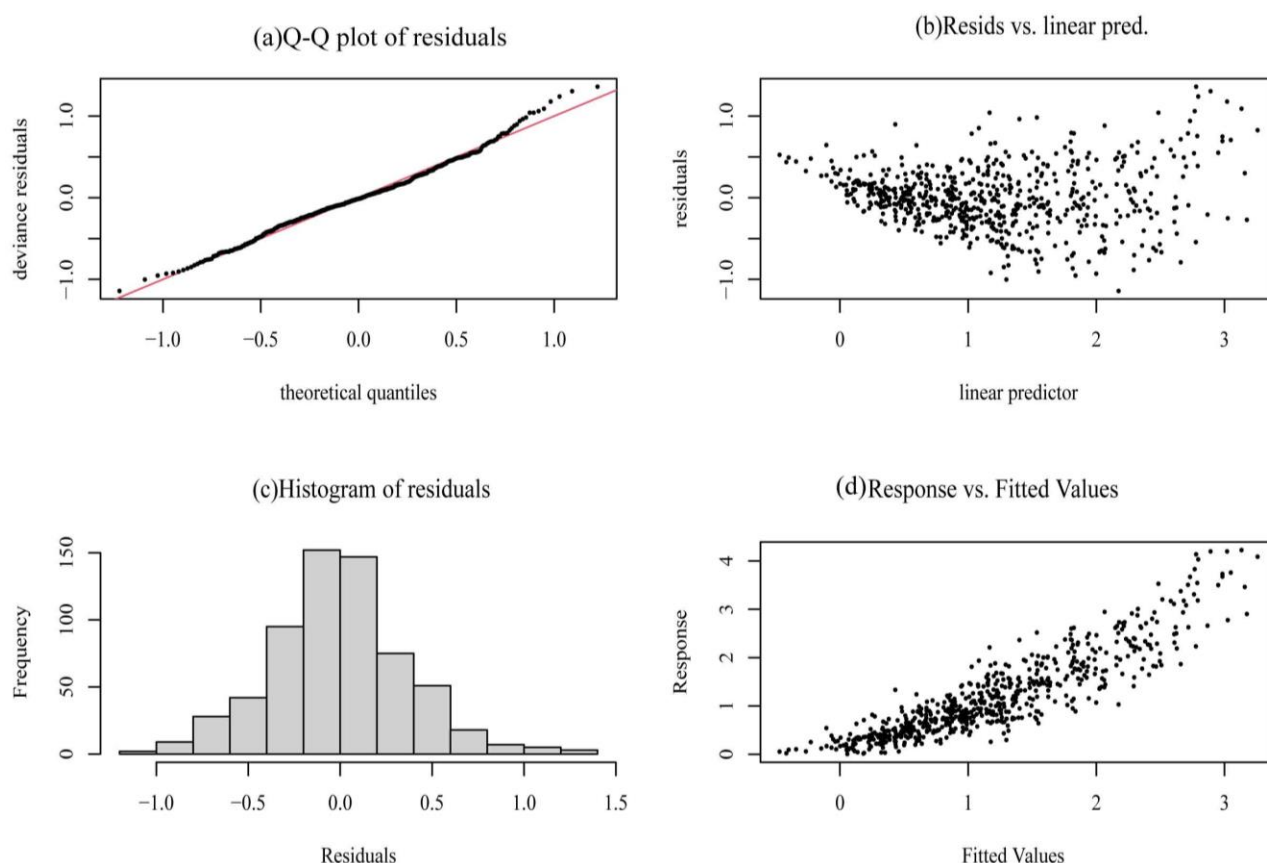
- The study used a non-parametric regression model. How good is it? Have you compared it with traditional chemistry models? Have you compared results with them?

**Response:** Thank you for your suggestions. We have added the validation of the GAM model in the supplementary material of Text 1, as follows.

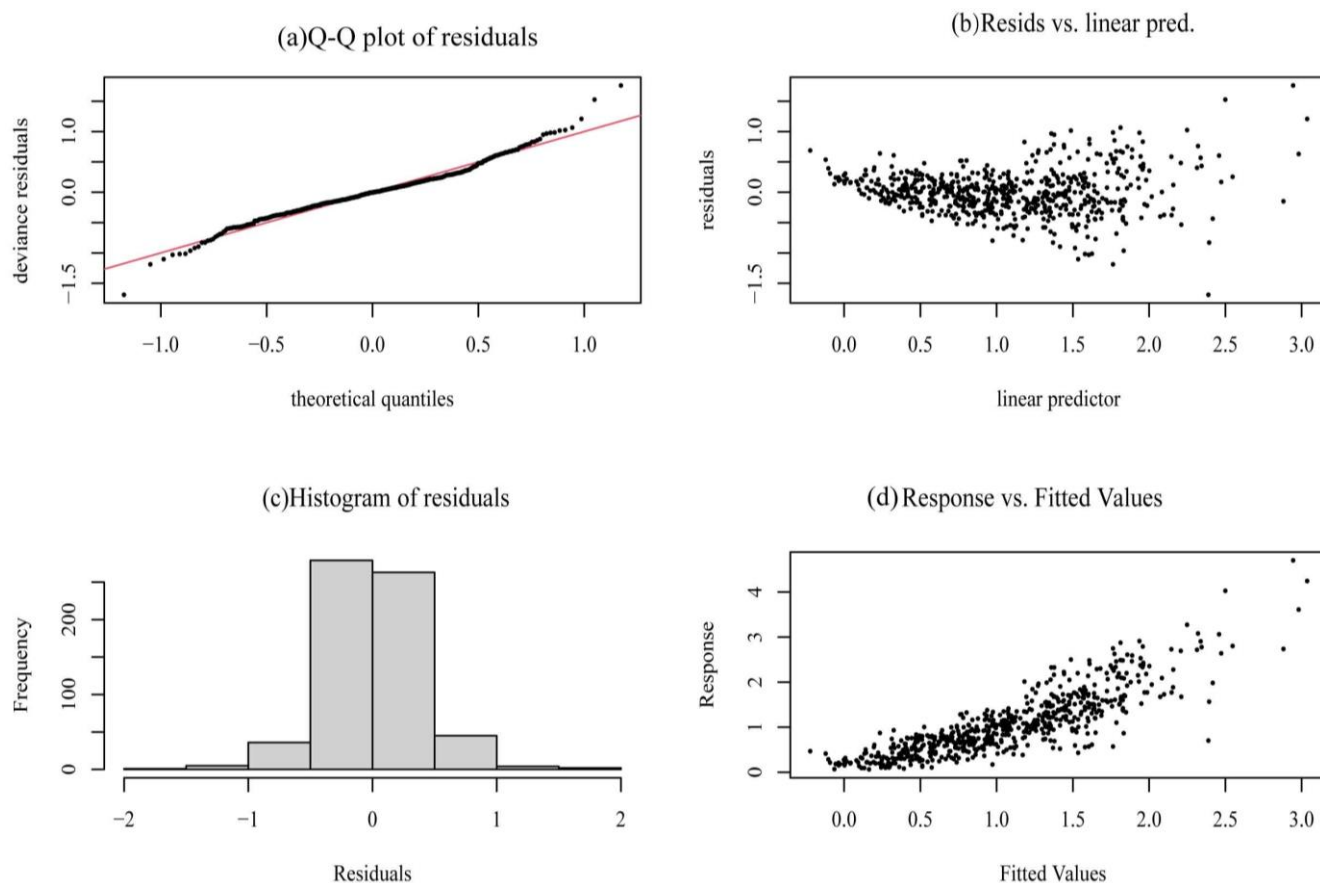
### Text 1 Model Validation.

Figure T1 and T2 show the residual test results of the Generalized Additive Model (GAM) in spring and autumn, respectively. From the residual Q-Q plots (Fig. T1 (a) and Fig. T2 (a)), the points were mostly on a straight line, indicating that the residuals conformed to a normal distribution. Meanwhile, the residual histogram of the model in Fig. T1 (c) and Fig. T2 (c) showed that the residuals were mainly concentrated around 0, which demonstrated the good fitting degree of the model. From the scatter plot of residuals and linear prediction values (Fig. T1 (b) and Fig. T2 (b)), the residuals were randomly distributed. From the scatter plot of the observed values and the fitted values (Fig. T1 (d) and Fig. T2 (d)), the response variables and the fitted values were well matched, and basically showed a “ $y = x$ ” distribution. Therefore, the fitting effect of this model was good.

The function of the Generalized Additive Models (GAM) is to analyze the correlation between the explanatory variables and the response variables. The GAM has been widely used in air pollution research and can effectively deal with the complex nonlinear relationship between air pollutants and influencing factors. Although GAM offers a flexible approach to calculating trends, the model is just a regression statistical model/method, which could not establish a connection with traditional chemistry models. Hence, comparing results with them might be very difficult and less significance.



**Fig. T1 Residual test results of the Generalized Additive Model (GAM) in spring.**



**Fig. T2 Residual test results of the Generalized Additive Model (GAM) in autumn.**

- Section 3.2, L226-229, statements are not clear. The statement "PAN pollution was mainly from local production" should be explained.

**Response:** Thank you for your suggestions, and we're sorry for the unclear expressions. We have revised the relevant contents in our manuscript.

“High PAN values in spring easily happened in the wind direction of the southeast with low wind speed ( $<3 \text{ m}\cdot\text{s}^{-1}$ ), showing the influence of urban plumes from the downtown of Xiamen island. High PAN values in autumn also appeared in the wind direction of the southeast, as well as the northeast with a relatively high wind speed (from Quanzhou city, an industrial city adjacent to Xiamen). Any more, PAN lifetimes in our observation site were relatively short due to the high ambient temperature, and the PAN lifetimes in autumn (2.02 hours) were significantly lower than that in spring (6.39 hours), which was not conducive to regional transport (Hu et al., 2020; Liu et al., 2018). Accordingly,  $\text{O}_3$  showed obvious characteristics of long-range transport, and PAN pollution was mainly from local production/accumulation in spring and autumn, but short-range transport from adjacent cities might contribute to the high PAN concentrations in autumn to a certain extent”.

- L230-235, it is not clear how local mixing is computed.

**Response:** Thank you for your suggestions. The mixing ratio in our study represents the volume concentration, which could be acquired directly from monitoring instruments. The mixing ratio of a gas component refers to

the occupied volume ratio of this gas component in dry air under the same temperature and pressure conditions. Mixing ratio values strictly speaking may not need a unit, but commonly ppb, nmol/mol or similar would be used. In our study, the units of the monitored pollutants (such as O<sub>3</sub>, PAN, and VOCs) were ppbv. Hence, we named their volume concentration as mixing ratio. For better understanding, we have changed the expression in this part. The detailed modifications are as follows:

“Based on the above analysis, we found that the photochemical reactions were still intense and even stronger under the low precursor levels. Although the precursor abundances of PAN and O<sub>3</sub> in spring were significantly higher than those in autumn ( $P < 0.01$ ), PAN values were comparable to and O<sub>3</sub> values were much higher in autumn than those in spring, respectively”.

- Fig. 3 and Fig. S4, nonlinear relations between variables (NO, UV, RH, T Ox) and PAN are well known. What is new in these figures should be explained.

**Response:** Thank you for your suggestions, we rewrote this part and added some new discussions in these figures. The details are as follows:

“The PAN in both seasons showed a downward trend with the increase of NO. PAN in spring was constant with NO fluctuation between 10 and 23 ppbv, and the confidence interval (CI) of NO concentration was relatively narrow. As we all know, the reaction of PA+NO is one of the most important loss pathways of PA, and the NO<sub>2</sub> production by NO oxidation in the O<sub>3</sub> formation cycle can react with PA radical to produce PAN, suggesting the fact that NO can consume and produce PAN indirectly (Liu et al., 2021). The consumption of NO to PAN was basically equal to the production when the NO levels were relatively high ( $>10$  ppbv), and the consumption of NO to PAN is greater than the production when the NO levels were low in spring. High values of NO mainly happened during rush hour traffic, thus controlling vehicle emissions can effectively alleviate PAN pollution. Ox had a positive correlation with PAN, representing the promotion effects of atmospheric oxidation capacity on PAN formation. The Ox levels  $<70$  ppbv (with narrow CI) played a significant promotion role in PAN formation (Fig. 3(b) and Fig. S4(b)). High Ox  $>70$  ppbv showed little influence on PAN, which could be explained as high Ox with relatively high air temperature leading to intense PAN thermal decomposition. When TVOCs were between 10 and 30 ppbv and PM<sub>2.5</sub> levels were  $<17 \mu\text{g}\cdot\text{m}^{-3}$ , PAN showed an upward trend with narrow CI. According to our previous study (Liu et al., 2022; Hu et al., 2020), the results of sensitivity analysis in Xiamen was VOCs-sensitive; the relatively low PM<sub>2.5</sub> concentrations in Xiamen showed limited influence on solar radiation through scattering and absorption, but promoted heterogeneous reactions producing radicals to a certain extent. UV and T had significant positive and negative nonlinear correlations with PAN, respectively. When UV changed between 0 and  $50 \text{ W}\cdot\text{m}^{-2}$  and T changed between 15 and  $35 \text{ W}\cdot\text{m}^{-2}$ , the CIs barely increased. In addition, when RH was more than 40%, the increase of RH was unfavorable for PAN production in both seasons. Some studies also found that high water vapor content could remove PAN and its precursors (Yan et al., 2018; Ma et al., 2020). Overall, the multiple-factor GAM analysis could better simulate the variations of PAN under real atmospheric conditions and evaluate the contributions of the influence factors to PAN formation”.

- L261-263: The solar radiations are stronger in spring than autumn; hence UV, T, and OX will be more effective in PAN formation during spring and vice-a-versa in autumn. Why should you mention it explicitly?

**Response:** Thank you for your suggestions, we agree that this summary was not accurate enough. In the previous question of Fig. 3 and Fig. S4, we rewrote this section and added some new information. Hence, we have rewritten this summary for a new version to avoid unnecessary misunderstandings.

“Overall, the multiple-factor GAM analysis could better simulate the variations of PAN under real atmospheric conditions and evaluate the contributions of the influence factors to PAN formation”.

- L292-302. These statements are huge. It is unclear whether they are supported by the model simulations or observations.

**Response:** We agree with your suggestions, and we rewrote these statements based on observations. The revisions in the manuscript are as follows.

“Diurnal variations of air pollutants during episodes and non-episodes are shown in Fig. 4, which could be explained by the evolution of the planetary boundary layer, local emissions, and atmospheric photochemistry. PAN reached a maximum value at 12:00-14:00, then decreased with weak solar radiation and reached the lowest in the early morning. Similar diurnal patterns of PAN and O<sub>3</sub> were observed, indicating the dominance of local photochemistry during the observation period (Zeng et al., 2019). CO, NO<sub>x</sub> and TVOCs showed highest values in the morning and the lowest values in the afternoon”.

- L325-328: All possible factors, meteorological conditions, accumulation of pollution, local transport, etc., are mentioned here. What is the most influencing factor?

**Response:** Thank you for your suggestions, we are sorry for the complicated and indirect expression of this sentence. We have revised the sentence as “These results indicated favorable meteorological condition was the dominant factor to produce PAN through accelerating its production rate and accumulation”.

Section 3.4 is lengthy but informative. It should be divided into two sections.

**Response:** Thank you for your suggestions, we have divided this section into two sections (Section 3.4.1 Inhibition and promotion effect of PAN on O<sub>3</sub> formation; Section 3.4.2 The influencing factors during inhibition and promotion stages).

Minor comments:

- The quality of figure 7 should be improved.

**Response:** Thank you for your suggestions, and we have improved the quality of figure 7.



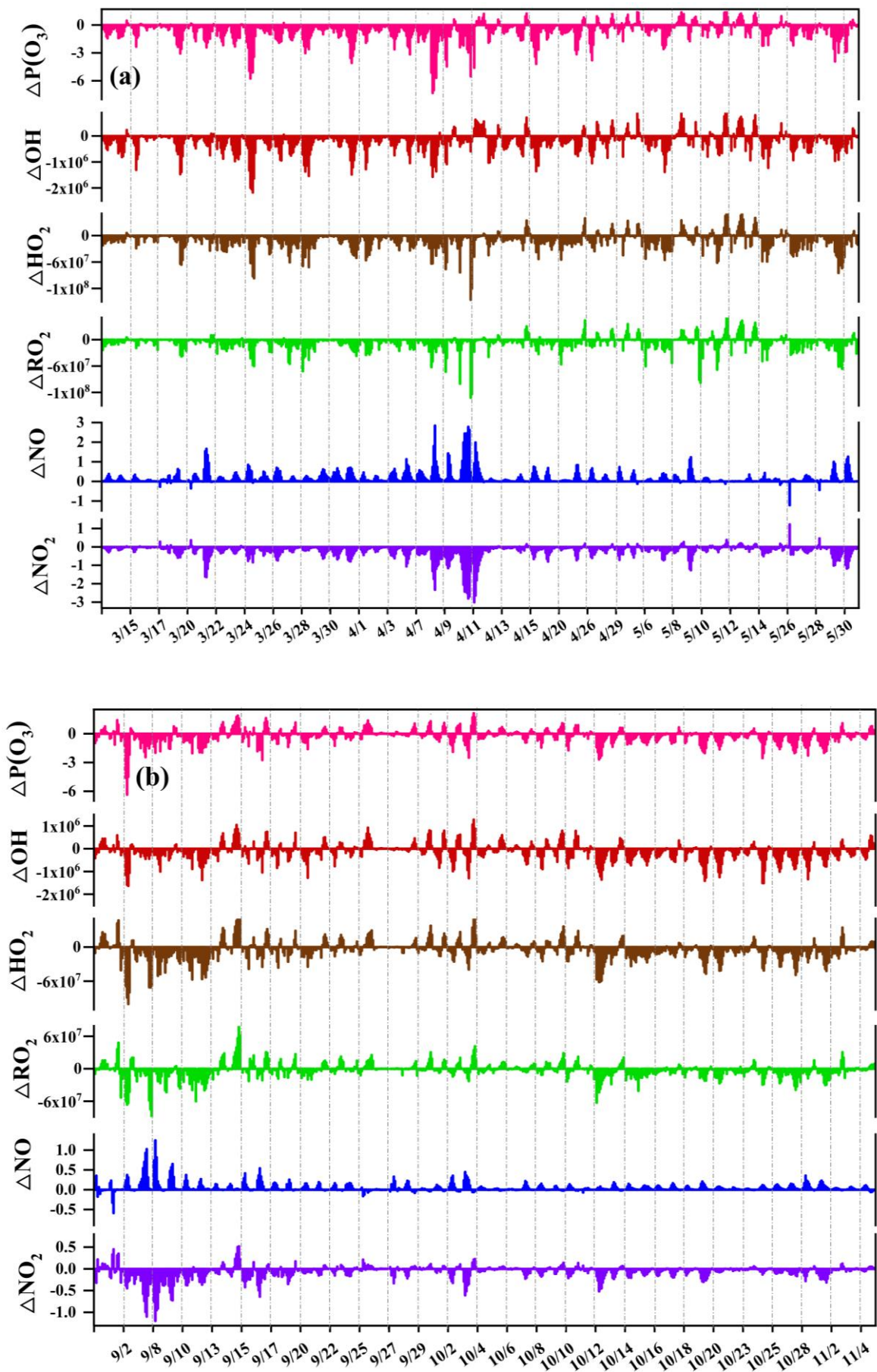
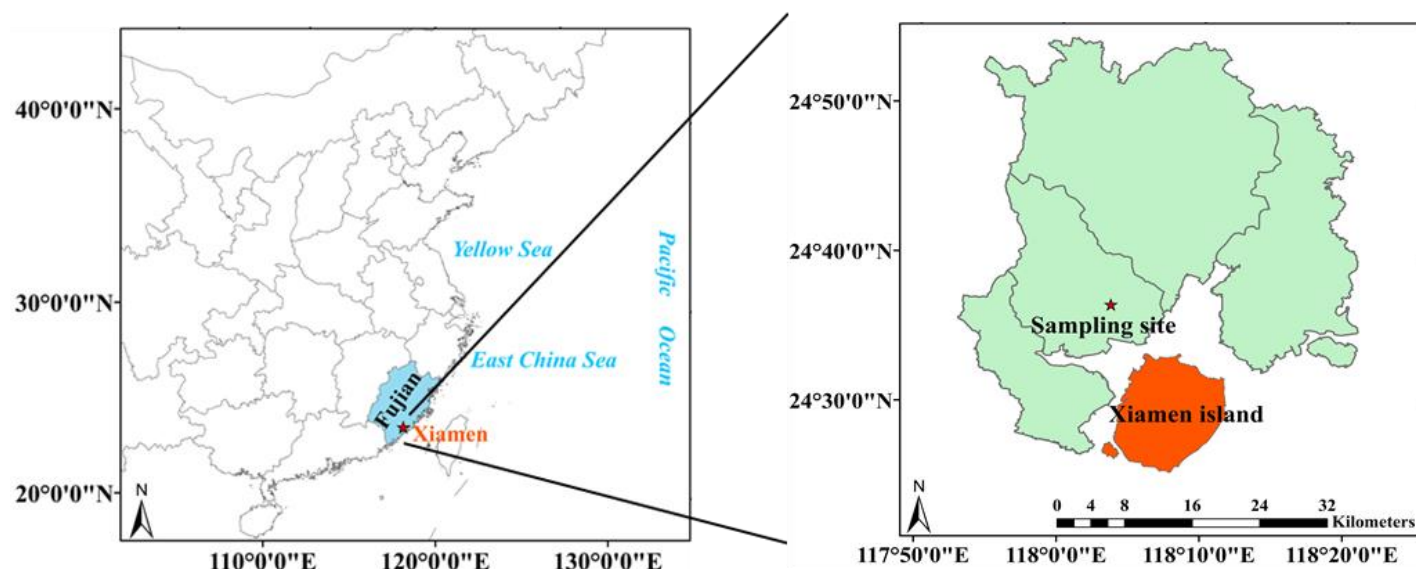


Fig. 7. The differences of O<sub>3</sub> net production  $\Delta P(O_3)$ ,  $\Delta OH$ ,  $\Delta HO_2$ ,  $\Delta RO_2$ ,  $\Delta NO$  and  $\Delta NO_2$  between the SC1 and the SC2 during the daytime (06:00-17:00) in (a) spring and (b) autumn (Unit: ppbv·h<sup>-1</sup> for  $\Delta P(O_3)$ ; ppbv for  $\Delta NO$  and  $\Delta NO_2$ ; molecules·cm<sup>-3</sup> for  $\Delta OH$ ,  $\Delta HO_2$  and  $\Delta RO_2$ ). The SC1 scenario was the base scenario putting all detected data (i.e. VOCs, trace gases, and meteorological parameters) into the model with all reaction pathways of the MCM mechanism, and the SC2 disabled the PAN chemistry, which is the only difference between SC1 and SC2.

- Section 2.1 can be moved to the supplementary material. I have difficulty in reading the x-axis labels in Figure 1.

**Response:** Thanks for your suggestion. We have changed a high resolution picture. Considering the importance of observation experiment details, we have simplified this part and divided them into two sections (Section 2.1 Observation site; Section 2.2 Measurement techniques). About Figure 1, the specific changes are as follows.

## 2.1 Observation site



**Fig. 1. Location of Xiamen and the observation site.**

Observations were carried out at the Atmospheric Environment Observation Supersite (AEOS, 24.61° N, 118.06° E; Fig. 1), located on the rooftop of around a 70 m high building in the Institute of Urban Environment, Chinese Academy of Sciences. The observations site is surrounded by highways, educational institutions, and residential buildings, which was characterized by rapidly urbanizing development area. When the prevailing wind direction was southerly winds, our observation site is downwind of the densely populated downtown (Xiamen island) (Hu et al., 2020; Liu et al., 2022). The field observations were continuously conducted from March 15 to November 4, 2020. The photochemical pollution events mainly appeared during spring and autumn in Xiamen, and we preferred to choose the periods with relatively high O<sub>3</sub> and PAN levels, then the measured data of 53 days in each season was chosen after excluding some special circumstances, such as extreme synoptic situations and instrument calibration.

## 2.2 Measurement techniques

PAN was monitored using a PAN analyzer (PANs-1000, Focused Photonics Inc., Hangzhou, CN) containing gas chromatography with electron capture detector (GC-ECD). During the observation period, multi-point standard curve calibration was conducted once a month, and single-point calibration was conducted every week, respectively. In the calibration mode of the PAN analyzer, the Mass Flow Controller (MFC) controls the flow rate of NO, acetone, and zero gas separately. The PAN standard gas is generated by the reaction of NO and acetone under ultraviolet light irradiation, and the sample is diluted to the required calibration mixing ratio for injection analysis. PAN was detected every 5 min and the detection limit was 50 pptv. The uncertainty and precision of PAN measurement were ±10% and 3%, respectively.

A gas chromatography-mass spectrometer (GC-FID/MS, TH-300B, Wuhan, CN) was used for monitoring the atmospheric VOCs with a 1-hour time resolution. The instrument conducted sampling with a

30 L/min sampling rate, then samples were pre-concentrated by cooling to -160 °C in a cryogenic trap followed by heating to 100 °C, and subsequently transferred to the secondary trap by high-purity helium (He). The flame ionization detector (FID) detected the low-carbon (C2-C5) hydrocarbons by a PLOT (Al<sub>2</sub>O<sub>3</sub>/KCl) column (15 m × 0.32 mm × 6.0 μm); the other species were quantified using a DB-624 column (60 m × 0.25 mm × 1.4 μm). The instrument system can quantitatively analyze 106 VOCs in the ambient atmosphere, including 29 alkanes, 11 alkenes, one alkyne, 17 aromatics, 35 halogenated hydrocarbons, and 13 OVOCs. The single-point calibration was performed every day at 23:00 with the standard mixtures of PAMS and TO15, and multi-point calibration was performed one month. The detection limits of the measured VOCs were in the range of 0.02 ppbv to 0.30 ppbv, and the measurement precision was ≤10%.

Criteria air pollutants of O<sub>3</sub>, CO, SO<sub>2</sub>, and NO<sub>x</sub>, were monitored by using Thermo Instruments TEI 49i, 48i, 43i, and 42i (Thermo Fisher Scientific, Waltham, MA, USA), respectively. HONO was monitored using an analyzer for Monitoring Aerosols and Gases in Ambient Air (MARGA, ADI 2080, Applikon Analytical B.V., the Netherlands). Particulate matters (PM<sub>2.5</sub>) were monitored by oscillating microbalance with tapered element (TEOM1405, Thermo Scientific Corp., MA, US), and the uncertainty of the PM<sub>2.5</sub> measurement was ±20%, respectively. The meteorological parameters (i.e. wind speed (WS), wind direction (WD), pressure (P), air temperature (T), and relative humidity (RH)) were measured by a weather station with sonic anemometer (150WX, Airmar, USA). Ultraviolet radiation (UV) was determined by a UV radiometer (KIPP & ZONEN, SUV5 Smart UV Radiometer). Photolysis frequencies including  $J(\text{O}^1\text{D})$ ,  $J(\text{NO}_2)$ ,  $J(\text{HONO})$ ,  $J(\text{NO}_3)$ ,  $J(\text{HCHO})$ , and  $J(\text{H}_2\text{O}_2)$  were analyzed by a photolysis spectrometer (PFS-100, Focused Photonics Inc., Hangzhou, China), and the uncertainty and detection limit of photolysis rates measurement were ±5% and around  $1 \times 10^{-5}$ , respectively.

Table S1 shows the detailed uncertainty and detection limit of instruments for trace gas observation. A schedule was applied to operate and inspect the AEOS monitoring station regularly and strictly to ensure the validity of the data. The detailed applications of the atmospheric monitoring procedure were shown in our previous studies (Wu et al., 2020; Liu et al., 2020a; Liu et al., 2020b; Hu et al., 2020).

## RC2 Anonymous Referee #2

The paper entitled “Seasonal characteristics of atmospheric peroxyacetyl nitrate (PAN) 1 in a coastal city 2 of Southeast China: Explanatory factors and photochemical effects” is generally well written, interesting and within the scope of ACP. However, the authors need to address a few minor comments, before the paper is suitable to be published in ACP.

**Response:** Thanks for your valuable comments. We have corrected this manuscript according to your suggestions.

Major comments:

I find the description of the VOC measurements a bit brief. Which classes of VOCs did the authors monitor with their GC and using which columns? It is not clear whether aldehydes and ketones were measured and if so which ones. Figure S5 doesn't discuss OVOCs as ozone precursors. While table S2 includes some OVOCs, it doesn't mention some important compounds like acetaldehyde, MVK, methylglyoxal and methacrolein. Where these compounds measured and if yes how? It appears to me that the authors are assuming that these are primarily secondary compounds that are derived from other primary emissions ethane, propene, isoprene and aromatics (line 57). If they are just model calculated and not measured then this could bias the results. While these compounds, no doubt have photochemical sources, there can also be substantial primary



emissions particularly from various types of biomass burning (Pallavi et al. <https://doi.org/10.5194/acp-19-15467-2019>), industrial sources and, in the case of acetaldehyde also vegetation (Sarkar et al. 2017 <https://doi.org/10.5194/acp-17-8129-2017>)

**Response:** Thank you for your suggestions. We have added more details of the description of the VOC measurements in the manuscript, as follows. Meanwhile, the detailed compounds of 106 VOCs were shown in Table R1.

“A gas chromatography-mass spectrometer (GC-FID/MS, TH-300B, Wuhan, CN) was used for monitoring the atmospheric VOCs with a 1-hour time resolution. The instrument conducted sampling with a 30 L/min sampling rate, then samples were pre-concentrated by cooling to -160 °C in a cryogenic trap followed by heating to 100 °C, and subsequently transferred to the secondary trap by high-purity helium (He). The flame ionization detector (FID) detected the low-carbon (C2-C5) hydrocarbons by a PLOT (Al<sub>2</sub>O<sub>3</sub>/KCl) column (15 m × 0.32 mm × 6.0 μm); the other compounds were quantified using a DB-624 column (60 m × 0.25 mm × 1.4 μm). The instrument system can quantitatively analyze 106 VOCs in the ambient atmosphere, including 29 alkanes, 11 alkenes, one alkyne, 17 aromatics, 35 halogenated hydrocarbons, and 13 OVOCs. The single-point calibration was performed every day at 23:00 with the standard mixtures of PAMS and TO15, and multi-point calibration was performed one month. The detection limits of the measured VOCs were in the range of 0.02 ppbv to 0.30 ppbv, and the measurement precision was ≤10%”.

Because the states of calibration gases of aldehydes and ketones are very unstable, thereby it is difficult to offer calibration gases during observation campaigns in our observation sites. Hence, the nine compounds can not be quantified (Acetaldehyde, Propanal, Crotonaldehyde, Methacrolein, n-butanal, Benzaldehyde, Valeraldehyde, m-Tolualdehyde, Hexanal). However, this phenomenon is common in many studies (Huang et al., 2021; Yang et al., 2019; Liu et al., 2021). Hence, the OVOCs that you mentioned can not be measured. All compounds of 106 VOCs detected by the gas chromatography-mass spectrometer are shown in Table R1. Meanwhile, the observation-based model has high requirements on the quality of the monitor data, and quantities of the pollutants inputting into the OBM model are limited, so VOCs with concentrations below the detection line were removed. Therefore, the VOCs in Table S2 were the all VOCs compounds that we used in the OBM model in our study.

The RIR values in Figure S5 were calculated by the observation data. As said above, some important aldehydes and ketones can not be considered, so the RIRs of OVOCs were underestimated. Hence, the discussion of RIRs of OVOCs is not included. In many relevant studies, their results of RIR also did not consider the OVOCs (Liu et al., 2021; Chen et al., 2020). In the future, we will calculate the RIR of OVOCs, because of our upgraded monitoring station with the OVOCs measurements.

Thank you for your suggestions. We agree with your opinions that the precursors of PAN of OVOCs have both primary and secondary sources (Sinha et al., 2019; Sarkar et al., 2017). When these OVOCs concentrations were not observed, the concentrations could be locally and reasonably calculated by the model. The three modules (initial value settings, observation data input, pre-ran for 2 days before running the OBM model) could constrain the unmeasured compounds reaching a steady-state and make the model localization. Some studies exploring the PAN formation mechanism based on OBM model also did not observe these OVOCs data (Chen et al., 2020, Liu et al., 2021; Xue et al., 2014). Meanwhile, we strongly agree with your idea and realized the importance of OVOCs, and some OVOCs of model calculation without measuring could bias the results. Hence, our team improved the monitoring of OVOCs in October 2021. A more optimized and complete monitoring system is also the future optimization goal of our model. Anymore, the index of agreement (IOA) judging the reliability of the model simulation results showed the performance of the OBM-MCM model was reasonably acceptable, and its detailed calculation is shown in the next question.

**Table R1 The detailed compounds of 106 VOCs and the detectors.**

Compounds	Detector	Compounds	Detector
ethene	FID	p-ethyltoluene	MS
ethane	FID	1,2,4-trimethylbenzene	MS
propane	FID	1,2,3-trimethylbenzene	MS
iso-butane	FID	m-diethylbenzene	MS
n-butane	FID	p-diethylbenzene	MS
iso-pentane	FID	naphthalene	MS
n-pentane	FID	dichlorodifluoromethane	MS
cyclopentane	FID	chloromethane	MS
propene	FID	1,1,2,2-tetrachloro-Ethane	MS
1-butene	FID	vinylchloride	MS
cis-2-butene	FID	bromomethane	MS
trans-2-butene	FID	chloroethane	MS
1-pentene	FID	trichlorofluoromethane	MS
trans-2-pentene	FID	1,1,2-trichloro-1,2,2-trifluoroethane	MS
cis-2-pentene	FID	carbondisulfide	MS
acetylene	FID	dichloromethane	MS
isoprene	FID	cis-1,2-dichloroethene	MS
2,2-dimethylbutane	MS	1,1-dichloroethane	MS
2,3-dimethylbutane	MS	trans-1,2-dichloroethene	MS
2-methylpentane	MS	trichloromethane	MS
3-methylpentane	MS	1,1,1-trichloroethane	MS
n-hexane	MS	1,2-dichloroethane	MS
2,4-dimethylpentane	MS	carbon tetrachloride	MS
methylcyclopentane	MS	trichloroethene	MS
cyclohexane	MS	1,2-dichloropropane	MS
2-methylhexane	MS	dichlorobromomethane	MS
2,3-dimethylpentane	MS	cis-1,3-dichloropropene	MS
3-methylhexane	MS	trans-1,3-dichloropropene	MS
2,2,4-trimethylpentane	MS	1,1,2-trichloroethane	MS
n-heptane	MS	dibromochloromethane	MS
methylcyclohexane	MS	tetrachloroethene	MS
2,3,4-trimethylpentane	MS	1,2-ethylenedibromide	MS
2-methylheptane	MS	chlorobenzene	MS
3-methylheptane	MS	tribromomethane	MS
n-octane	MS	Ethane, 1,1,2,2-tetrachloro-	MS
n-nonane	MS	1,3-dichlorobenzene	MS
n-decane	MS	chlorotoluene	MS
n-undecane	MS	1,4-dichlorobenzene	MS
n-dodecane	MS	1,2-dichlorobenzene	MS
1-hexene	MS	1,2,4-trichlorobenzene	MS
1,3-butadiene	MS	1,1,2,3,4,4-hexachloro-1,3-butadiene	MS
1,1-dichloroethene	MS	acrolein	MS
benzene	MS	acetone	MS
toluene	MS	2-butanone	MS
m/p-xylene	MS	2-propanol	MS
ethylbenzene	MS	2-methoxy-2-methylpropane	MS
styrene	MS	vinylacetate	MS

o-xylene	MS	ethylacetate	MS
iso-propylbenzene	MS	tetrahydrofuran	MS
n-propylbenzene	MS	methyl methacrylate	MS
o-ethyltoluene	MS	1,4-dioxane	MS
m-ethyltoluene	MS	4-methyl-2-pentanone	MS
1,2,3-trimethylbenzene	MS	2-hexanone	MS

The authors should include a more detailed description of their measurements different VOC classes and if some VOCs were only calculated as secondary products in the observation-based model (OBM) then this should be mentioned and the corresponding uncertainties should be discussed. If they were directly measured, it would be better to include OVOCS in the analysis of Figure 6.

**Response:** Thank you for your suggestions. We have added more details of the description of the VOC measurements in the manuscript, as the above question shown. The instrument system can quantitatively analyze 106 VOCs in the ambient atmosphere as shown in Table R1, but the observation-based model has high requirements on the quality of monitor data, so VOCs with concentrations below the detection line were removed. Therefore, the VOCs in Table S2 were the all VOCs compounds that we used in the OBM model in our study, and these valid data contained key species used in most relevant studies, which could constrain the model and realize the localization of the model well (Liu et al., 2021; Chen et al., 2020; Xue et al., 2014).

About Figure 6: Firstly, it is RIRs of first-generation precursors for PAN. Secondly, the RIRs of measured OVOCS were very minor, which could be ignored in Figure 6. And the detailed information of some VOCs was added in the manuscript, as follows:

“The second-generation precursors of PAN of CH<sub>3</sub>CHO and MGLY have both primary and secondary sources, and the other OVOCS are mainly oxidation products of hydrocarbons (Sinha et al., 2019; Sarkar et al., 2017). Consequently, the contribution and importance of first-generation precursors of PAN are necessary to identify to better control photochemical pollution, which will be discussed in the next section”.

“During these simulations (except for NO and NO<sub>2</sub>), the model was not constrained by the OVOC measurements considering that these first-generation precursors contribute to PAN production through formation of OVOCS”.

About the uncertainties of the model simulation results, the index of agreement (IOA) can be used to judge the reliability of the model simulation results, and the model validation results were added in the supplementary material of Text 1.

### Text 1 Model validation

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

$$IOA = 1 - \frac{\sum_{i=1}^n (O_i - S_i)^2}{\sum_{i=1}^n (|O_i - \bar{O}| + |S_i - \bar{O}|)^2} \quad (4)$$

where  $S_i$  is simulated value,  $O_i$  represents observed value,  $\bar{O}$  is 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 0.90 (Wang et al., 2018), the simulation results are reasonable, and the IOA in our research is 0.88. Hence, the performance of the OBM-MCM model was reasonably acceptable.

**Table S2. Descriptive statistics of measured VOCs mixing ratios (Units: ppbv).**

Chemicals	Spring		Autumn		Chemicals	Spring		Autumn	
	Mean	SD	Mean	SD		Mean	SD	Mean	SD
<b>Alkanes</b>	<b>9.41</b>	<b>5.30</b>	<b>5.47</b>	<b>2.88</b>	<b>Alkyne</b>	<b>1.00</b>	<b>0.55</b>	<b>0.63</b>	<b>0.34</b>
Ethane	2.39	1.02	1.31	0.43	<b>Aromatics</b>	<b>2.71</b>	<b>2.33</b>	<b>1.62</b>	<b>1.15</b>
Propane	2.31	1.29	1.19	0.58	Benzene	0.27	0.14	0.16	0.09
iso-Butane	0.87	0.57	0.52	0.37	Toluene	1.37	1.21	0.85	0.84
n-Butane	1.30	0.94	0.77	0.59	m/p-Xylene	0.53	0.63	0.39	0.32
iso-Pentane	1.15	1.27	0.52	0.44	Ethylbenzene	0.18	0.18	0.09	0.10
n-Pentane	0.44	0.42	0.24	0.21	Styrene	0.09	0.16	0.02	0.04
2,2-Dimethylbutane	0.02	0.02	0.02	0.01	o-Xylene	0.19	0.23	0.04	0.09
2,3-Dimethylbutane	0.05	0.06	0.05	0.05	m-Ethyltoluene	0.02	0.02	0.01	0.01
2-Methylpentane	0.08	0.09	0.05	0.04	1,3,5-Trimethylbenzene	0.01	0.01	0.01	0.01
3-Methylpentane	0.14	0.15	0.06	0.06	p-Ethyltoluene	0.01	0.01	0.01	0.005
n-Hexane	0.20	0.25	0.10	0.20	1,2,4-Trimethylbenzene	0.03	0.05	0.01	0.02
Cyclohexane	0.04	0.04	0.02	0.02	1,2,3-Trimethylbenzene	0.01	0.01	0.01	0.004
2-Methylhexane	0.05	0.06	0.04	0.05	<b>Isoprene (BHC)</b>	<b>0.08</b>	<b>0.14</b>	<b>0.10</b>	<b>0.17</b>
3-Methylhexane	0.08	0.09	0.05	0.08	<b>Halocarbons</b>	<b>2.54</b>	<b>1.27</b>	<b>1.95</b>	<b>0.90</b>
n-Heptane	0.07	0.08	0.05	0.06	Chloromethane	0.51	0.23	0.46	0.18
n-Octane	0.04	0.06	0.09	0.06	Bromomethane	0.06	0.03	0.04	0.02
n-Nonane	0.02	0.01	0.01	0.005	Dichloromethane	1.19	0.81	0.87	0.50
n-Decane	0.01	0.01	0.01	0.01	Trichloromethane	0.07	0.03	0.05	0.02
n-Undecane	0.02	0.02	0.03	0.03	1,2-Dichloroethane	0.51	0.34	0.36	0.22
n-Dodecane	0.12	0.29	0.36	0.84	Trichloroethene	0.02	0.02	0.02	0.01
<b>Alkenes</b>	<b>1.30</b>	<b>0.89</b>	<b>0.85</b>	<b>0.48</b>	1,2-Dichloropropane	0.12	0.13	0.10	0.08
Ethene	0.90	0.65	0.51	0.34	Tetrachloroethene	0.05	0.05	0.04	0.05
Propene	0.20	0.14	0.19	0.11	<b>OVOCs</b>	<b>4.49</b>	<b>1.83</b>	<b>4.17</b>	<b>2.57</b>
1-Butene	0.04	0.03	0.03	0.02	Acrolein	0.06	0.03	0.04	0.02
cis-2-Butene	0.05	0.06	0.03	0.03	Acetone	2.22	0.94	2.21	0.91
trans-2-Butene	0.03	0.06	0.03	0.02	2-Butanone	0.67	0.45	0.50	0.44
1-Pentene	0.02	0.02	0.01	0.01	2-Propanol	0.24	0.31	0.12	0.12
trans-2-Pentene	0.04	0.04	0.04	0.02	2-Methoxy-2-methylpropane	0.24	0.32	0.09	0.09
1,3-Butadiene	0.01	0.02	0.01	0.01	Ethylacetate	1.07	0.83	1.20	1.31

Minor comments:

Line 28: I could not quite understand what the authors intend to say in the following sentence “Without considering the transformation of peroxyacetyl radical (PA) and PAN, acetaldehyde contributed to the dominant production of PA (46±4%), followed by methylglyoxal (28±3%) and radical cycling (19±3%).”

**Response:** Thank you for your suggestions, and we are sorry for the confused expressions. The thermal decomposition of PAN can produce NO<sub>2</sub> and PA radicals (Eq. R1), and PA can quickly react with NO<sub>2</sub> producing PAN (Eq. R2). The ratio of reaction rates of K<sub>1</sub> and K<sub>2</sub> was close to 1, thus the contribution of PAN thermal decomposition to PA was minor. For better understanding, we revised the sentence as “Model simulations revealed that acetaldehyde oxidation (46±4%) contributed to the dominant formation pathway of PA (hence PAN), followed by methylglyoxal oxidation (28±3%) and radical cycling (19±3%)”.





Line 30: Grammar needs to be improved, maybe the following will convey the intended meaning better: “The PAN formation was highly VOC-sensitive, as surplus NO<sub>x</sub> (compared with VOCs abundance) prevented NO<sub>x</sub> from being the limiting factor photochemical formation of secondary pollution.”

**Response:** Thank you for your suggestions, we strongly agree with your descriptions, and we have revised this sentence as you said.

“The PAN formation was highly VOC-sensitive, as surplus NO<sub>x</sub> (compared with VOCs abundance) prevented NO<sub>x</sub> from being the limiting factor photochemical formation of secondary pollution”.

Line 32: This sounds like a generic statement “PAN could promote or inhibit O<sub>3</sub> formation under high or low RO<sub>x</sub> levels, respectively.”. It may be more appropriate to target this at the results of the present study “At our site, PAN promoted and inhibited O<sub>3</sub> formation under high and low RO<sub>x</sub> levels, respectively.”

**Response:** Thank you for your suggestions, your expression was appropriate, and we revised the sentence accordingly.

“At our site, PAN promoted and inhibited O<sub>3</sub> formation under high and low RO<sub>x</sub> levels, respectively”.

Line 36: The authors could be a bit more assertive and specific in highlighting the contribution of their study to the scientific understanding. “Might be helpful” doesn’t sound very convincing to me and doesn’t specify the main contribution.

**Response:** Thank you for your suggestions. Your suggestions are pretty good, and we have replaced some words and emphasized the main contribution.

“The analysis of PAN formation mechanism and its positive or negative effect on ozone in our study provided scientific insights into photochemical pollution mechanism under various pollution scenarios in coastal areas”.

Line 52: Grammar: “is the only formation pathway” instead of “is solely formation pathway”

**Response:** Thank you for your suggestions, we revised the sentence accordingly.

Line 72 Language: “were the most significant contributors” instead of “offered the highest contribution”

**Response:** Thank you for your suggestions, we have revised the expressions accordingly.



Line 74: “Recently, negative and positive impacts of PAN photochemistry on O<sub>3</sub> production were captured under the low and high NO<sub>x</sub> conditions, respectively.” This statement should include a reference to the corresponding study.

**Response:** Thank you for your suggestions, we have added references to the corresponding study.

*Zeng, L., Fan, G. J., Lyu, X., Guo, H., Wang, J. L., and Yao, D.: Atmospheric fate of peroxyacetyl nitrate in suburban Hong Kong and its impact on local ozone pollution, Environ Pollut, 10.1016/j.envpol.2019.06.004, 2019.*

*Liu, Y., Shen, H., Mu, J., Li, H., Chen, T., Yang, J., Jiang, Y., Zhu, Y., Meng, H., Dong, C., Wang, W., and Xue, L.: Formation of peroxyacetyl nitrate (PAN) and its impact on ozone production in the coastal atmosphere of Qingdao, North China, Sci Total Environ, 778, 146265, 10.1016/j.scitotenv.2021.146265, 2021.*

Line 99: “was attributed to the downwind region of the downtown (Xiamen island) with densely population ” can be simplified to “was downwind of the densely populated downtown region (Xiamen island)”

**Response:** Thank you for your suggestions, we have simplified the description.

Line 100 in my opinion field observation should be plural “field observations were” not singular

**Response:** Thank you for your suggestions, we agree with your opinion and changed its format into plural.

Line 122 I have never come across the term “ultrasonic atmospherium” before. I believe the correct name would be "weather station with 2D sonic anemometer".

**Response:** Thank you for your suggestions. We are sorry for the wrong usage of the term, and the correct name you mentioned was right. Hence, we changed the name to “weather station with sonic anemometer”.

Line 209 please avoid colloquial language “The wind directions in late spring and early autumn were messy due to the season switch.” More scientific “During the transition from spring to summer the wind direction fluctuated between ... and ... while during the transition from summer to autumn the wind direction fluctuated from ... to ...”

**Response:** Thank you for your suggestions. We have revised this description as “During the transition from spring to summer the wind direction fluctuated between northwest and southeast while during the transition from summer to autumn the wind direction fluctuated from southeast to northeast”.

Line 210, respectively missing at the end of the sentence “The wind rose charts showed that the wind direction frequencies with relatively high wind speed in spring and autumn were southeast wind and northeast wind (Fig. S2), respectively.” Also define “high” by inserting a number “(>... m/s)” in brackets.

**Response:** Thank you for your suggestions. We have revised this sentence as “The wind rose charts showed that the wind direction frequencies with relatively high wind speed ( $>3 \text{ m}\cdot\text{s}^{-1}$ ) in spring and autumn were southeast wind and northeast wind (Fig. S2), respectively”.

In general, the manuscript should be run through a grammar check software before resubmission. It is better to avoid colloquial language and indirect phrases. It is OK to be direct and use simple sentences.

**Response:** We have performed the grammar corrections of our manuscript through grammar check software, and we also invited native speakers in related fields to polish the manuscript. Thanks for your suggestion, simple sentences, and direct phrases make my point clear.

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