

Understanding summertime peroxyacetyl nitrate (PAN) formation and its relation to aerosol pollution: Insights from high-resolution measurements and modeling

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Abstract: Peroxyacetyl nitrate (PAN), a key indicator of photochemical pollution, is generated similarly to ozone (O₃), through reactions involving specific volatile organic compounds (VOCs) and nitrogen oxides. Notably, PAN has been observed at unexpectedly high concentrations (maximum: 3.04 ppb) during the summertime. The average daily values of PAN showed a strong correlation with black carbon (BC) (R=0.77) and O₃ (R=0.77), suggesting a close connection between summertime haze and photochemical pollution. We addressed the puzzle of summertime PAN formation and its association with aerosol pollution under high O₃ conditions in Xiamen, a coastal city in southeastern China, by analyzing continuous high temporal resolution data utilizing box modeling in conjunction with the master chemical mechanism (MCM) model. The MCM model, with an index of agreement (IOA) value of 0.75, effectively investigate PAN formation, performing better during the clean period (R²: 0.68, slope K: 0.91) than haze one (R²: 0.47, slope K: 0.75). Using extreme gradient boosting (XGBoost), we identified NH₃, NO₃⁻, and PM_{2.5} as the primary factors for simulation bias. Moreover, the net production rate of PAN becomes negative with PAN constrained, suggesting

30 an unknown compensatory mechanism. Both relative incremental reactivity (RIR) and empirical kinetic modeling approach (EKMA)
31 analyses indicate that PAN formation is VOC-controlled. Controlling emissions of VOCs, particularly alkenes, C₅H₈, and aromatics,
32 would mitigate PAN pollution. PAN promotes OH and HO₂ while inhibiting the formation of O₃, RO₂, NO, and NO₂. This study
33 deepens our comprehension of PAN photochemistry while also offering scientific insights for guiding future PAN pollution control
34 strategies.

36 Introduction

37 PAN is a significant secondary gaseous pollutant commonly present in photochemical smog and poses risk to human health and
38 plant growth, being 1-2 magnitudes more phytotoxic than O₃ (Yukihiro et al., 2012; Taylor, 1969). Additionally, PAN's low aqueous
39 solubility, minimal reactivity with hydroxyl radicals (OH), and slow photolysis contribute to its capacity for long-range transport of
40 nitrogen oxides (NO_x) (Xu et al., 2018; Zhai et al., 2024; Marley et al., 2007b). Therefore, its formation in polluted areas holds
41 significant importance beyond local concerns. Similar to surface O₃, PAN is produced during the oxidation of VOCs in the presence
42 of NO_x (R1-R3) (Xu et al., 2021). PAN is formed when NO₂ reacts with peroxyacetyl (PA) radicals (CH₃C(O)OO·) (R2), but the
43 presence of NO consumes PA radicals, inhibiting PAN production (R3), which creates a comparable dependence of PAN and O₃ on
44 NO and NO₂ levels. Unlike O₃, however, PAN is influenced by only a limited number of oxygenated VOCs (OVOCs) that generate
45 PA radicals. These OVOCs, which are second-generation precursors of PAN, include acetaldehyde (CH₃CHO), acetone
46 (CH₃C(O)CH₃), methylglyoxal (MGLY, CH₃C(O)CHO), methyl vinyl ketone (MVK, CH₂CHC(O)CH₃), methyl ethyl ketone (MEK,
47 CH₃C(O)CH₂CH₃), methacrolein (MACR, CH₂C(CH₃)CHO), and biacetyl (CH₃C(O)C(O)CH₃). These compounds are typically
48 formed from the oxidation of alkenes, aromatics, and isoprene, which are the first-generation precursors of PAN (Xue et al., 2014;
49 Zhang et al., 2015). Identifying the dominant precursors is crucial for managing PAN pollution effectively. In the troposphere,
50 thermal decomposition (R4) is the primary process responsible for PAN loss.



55 In recent years, wintertime photochemical air pollution has increasingly garnered attention. At this time, the concentration of O₃ is
56 low due to the strong titration of NO, while the concentration of aerosol is high, and it is found that aerosol promotes PAN generation
57 (Xu et al., 2021). Surprisingly high concentrations of OH radical, particularly under hazy conditions, have been observed and are
58 largely attributed to HONO photolysis (Xu et al., 2021). Winter photochemical and haze pollution often exacerbate each other, with
59 photochemical trace gases supplying both oxidants and precursors for aerosol formation, and aerosols acting as mediums for

60 heterogeneous reactions that produce key oxidants such as HONO, H₂O₂, and OH radicals (Xu et al., 2021). The OH produced by
61 HONO photolysis can partially replace the UV action to promote PAN formation in winter in southeast coastal area of China when
62 particulate matter is high ($\geq 35 \mu\text{g}\cdot\text{m}^{-3}$) (Hu et al., 2020). Zhang et al. (2020) found the potential HONO sources significantly
63 improved the PAN simulations in wintertime heavy haze events with high concentrations of PAN. High concentrations of PAN are
64 a consequence of the increased levels of precursors and HONO observed during haze episodes (Liu et al., 2018). In conclusion,
65 most previous studies have studied the effect of aerosol on PAN generation in winter. Further research on PAN should determine
66 whether particulates significantly contribute to its formation during warmer seasons with elevated O₃ concentrations (Xu et al.,
67 2021). In Eastern China, photochemical air pollution often involves high concentrations of both O₃ and PAN, a persistent issue
68 during the warm season (April-September) for many years (Lu et al., 2020). The characteristics and formation pathways of PAN
69 during summer have been increasingly studied in regions such as the North China Plain (NCP), the Yangtze River Delta, the Pearl
70 River Delta, and southwestern China. These studies have generally shown consistent diurnal patterns and strong correlations
71 between PAN and O₃, identifying acetaldehyde—primarily derived from the degradation of aromatics and alkenes—as the key direct
72 precursor of PAN in the summer. However, there has been limited research on the formation of PAN and its relationship with aerosol
73 pollution during the summertime.

74 Xiamen is one of the fastest urbanizing regions in the southeast China and is also one of the cities with the best air quality in China,
75 where the air quality could represent the future of other Chinese urban regions. Between 2018 and 2023, Xiamen ranked among the
76 top 10 cities in China, achieving positions of 7th in 2018, 4th in both 2019 and 2020, 6th in 2021, 9th in 2022, and returning to 7th
77 in 2023 (mcc.gov.cn, last assessed October 30, 2014). Xiamen is located in a low-latitude coastal area, with abundant sunlight and
78 long daylight hours during the summer, resulting in strong solar radiation and rapid photochemical conversion rates. The city is
79 typically influenced by the East Asian monsoon and serves as a transport channel for atmospheric pollutants from both the Yangtze
80 River Delta and Pearl River Delta regions. Additionally, during the summer, Xiamen is often affected by complex meteorological
81 conditions such as typhoons and the West Pacific Subtropical High (WPSH). The WPSH creates weather conditions that promote
82 the formation and accumulation of photochemical pollutants and particulate matter (Wu et al., 2019). This setting provides an ideal
83 "laboratory" for investigating the complexities of summertime PAN formation and its relationship with aerosol pollution under high
84 O₃ concentrations. In summer, especially in July, high temperatures, high humidity, and intense radiation are likely to accelerate
85 both the formation and consumption rates of PAN. In this study, continuous measurements of trace gases, substances related to
86 aerosols, photolysis rate constants and meteorological parameters were performed at a suburban site in Xiamen from 10 to 31 July
87 2018. Firstly, we provide an overview of pollutant concentrations, meteorological parameters, and weather conditions during the
88 observation period. Secondly, we simulate PAN concentration with the aid of box modeling combined with master chemical
89 mechanism (MCM) model. Using machine learning with XGBoost, we identified the key factors that affect the observation-based
90 model (OBM) model's simulation results and clarified the mechanisms linking haze pollution to photochemical air pollution, as

91 indicated by PAN and O₃. Thirdly, the study identified the main precursors and oxidants responsible for summertime PAN production
92 in Xiamen and evaluated the influence of PAN on local atmospheric oxidation capacity. This study further emphasized the interplay
93 between haze and photochemical air pollution and highlighted significant implications for future research.

94 95 **2 Methodology**

96 **2.1 Field observations**

97 Trace gases (including PAN, O₃, HONO, HNO₃, HCl, NH₃, VOCs, NO_x, CO, and SO₂), substances related to aerosols (including
98 BC, PM₁, PM_{2.5}, PM₁₀, SO₄²⁻, NO₃⁻, NH₄⁺, Cl⁻, Na⁺, K⁺, Ca²⁺, Mg²⁺), photolysis rate constants (including JO¹D, JNO₂, JHONO,
99 JHCHO_M, JHCHO_R, JNO₃_M, JNO₃_R, JH₂O₂), and meteorological parameters (including temperature, relative humidity,
100 atmospheric pressure, wind speed, and wind direction) were continuously measured at an suburban site in Xiamen from 10 to 31
101 July 2018. All instruments were placed inside an air-conditioned container situated on the rooftop of a 20-story building at the
102 Institute of Urban Environment, Chinese Academy of Sciences (IUE: 118.06° E, 24.61° N) (Fig. S1(a)). When southerly winds
103 prevailed, Xiamen Island, characterized by dense population and traffic congestion, was located upwind of the IUE (Fig. S1(b)).
104 The IUE supersite is surrounded by Xinglin Bay, several universities and institutes, and major roadways with heavy traffic, such as
105 Jimei Road (< 200 m), Shenhai Expressway (870 m), and Xiasha Expressway (2300 m) (Fig. S1(c)).

106 PAN measurements were conducted using a PANs-1000 analyzer (Focused Photonics Inc., Hangzhou, China), which features an
107 automated system consists of a gas chromatograph, an electron capture detector, and a calibration unit. The analyzer provided PAN
108 readings every 5 minutes, with a detection limit of 50 ppt. The uncertainty and precision of the PAN measurements were ±10 % and
109 3 %, respectively. The PAN standard gas was produced through the reaction of acetone and NO under UV light. Calibration
110 procedures included monthly multi-point calibrations and weekly single-point calibrations. Detailed information about the PAN
111 detection system and calibration can be found in previous studies (Hu et al., 2020; Liu et al., 2022a). The VOC measurements were
112 conducted using a gas chromatography mass spectrometer (GC-FID/MS, TH-300B, Wuhan, China) at an hourly time resolution.
113 Detailed information regarding the VOC detection system and calibration procedures is available in our previous study (Liu et al.,
114 2022b). HONO measurements were conducted using a customized Incoherent BroadBand Cavity Enhanced Absorption
115 Spectroscopy (IBBCEAS) system developed by the Anhui Institute of Optics and Fine Mechanics (AIOFM), Chinese Academy of
116 Sciences. The HONO detection limit was 100 ppt, with a time resolution of 1 minute. The measurement principle and calibration
117 method of IBBCEAS can be found in the previous literature (Hu et al., 2022; Duan et al., 2018; Hu et al., 2024). The concentrations
118 of inorganic components in PM_{2.5} aerosols (including SO₄²⁻, NO₃⁻, NH₄⁺, Cl⁻, Na⁺, K⁺, Ca²⁺, Mg²⁺), as well as the concentrations of
119 gases such as NH₃, HCl, and HNO₃ were analyzed using a Monitor for AeRosols and Gases in ambient Air (MARGA, Model ADI
120 2080, Applikon Analytical B.V., the Netherlands) (Hu et al., 2022). The criteria air pollutants O₃, NO_x, CO, and SO₂ were measured
121 using different methods: ultraviolet (UV) absorption for O₃ (TEI model 49i), chemiluminescence with a molybdenum converter for

NO_x (TEI model 42i), non-dispersive infrared for CO (TEI model 48i), and pulsed UV fluorescence for SO₂ (TEI model 43i). A tapered element oscillating microbalance (TEOM1405, Thermo Scientific Corp., MA, USA) was used to continuously measure the mass concentrations of PM₁, PM_{2.5}, and PM₁₀ online. A photolysis spectrometer (PFS-100, Focused Photonics Inc., Hangzhou, China) was employed to measure the photolysis rate constants. An ultrasonic anemometer (150WX, Airmar, USA) was used to measure meteorological parameters.

2.2 Box modeling

This study employed a box model framework utilizing the Master Chemical Mechanism (MCMv3.3.1, <https://mcm.york.ac.uk/MCM/home.htm>) to investigate sensitivity and mechanisms of PAN formation. The model constraints were derived from observations of trace gases and meteorological parameters, which were averaged to 1-hour intervals. The reliability of model simulation results is often assessed using the index of agreement (IOA), which ranges from 0 to 1, with a higher IOA signifying greater alignment between observed and simulated values. Note that the model simulation values at this time are not constrained by PAN. For specific formulas, please refer to the supplementary information (Eq. S1). Other formulas, including PAN production rates (P(PAN)), net production of PAN (Net (PAN)), and the RIR, are provided in the supplementary information (Eq. S2- Eq. S4).

The MCM simulates the nonlinear interaction between PAN and its precursors by altering the VOCs-to-NO_x ratio across multiple scenarios, while keeping all other parameters fixed. In this study, a 20 % step size was applied, reducing VOCs and NO_x from 200 % down to 0 % to construct a scenario matrix. A total of 121 scenarios were generated to model the PAN production rate. The scenario representing the average VOCs and NO_x mixing ratio during the sampling period was designated as the base case, with the remaining 120 scenarios created by systematically adjusting the VOC-to-NO_x ratio. The output from these 121 simulations was used to construct isopleth diagrams depicting the relationship between VOCs, NO_x, and PAN.

2.3 Machine Learning Model

To identify the key factors influencing the performance of the model simulation, the Machine Learning (ML) model was applied to establish the prediction model of bias between simulation of OBM and observation. XGBoost is a supervised boosting algorithm that reduces the risk of over-fitting, captures the nonlinear relationships among predictor variables, and solves numerous data science problems in a rapid and accurate way (Li et al., 2024). It has demonstrated high performance in O₃ studies in China. As compared to other bagging tree models like random forest, XGBoost can handle more complex data while consuming fewer computing resources. To further improve the interpretability of the ML model, the feature importance of independent input variables in the XGBoost model is quantified using the Shaply Additive explanation (SHAP) approach (Lin et al., 2024). The SHAP calculates a value that represents the contribution of each feature to the model's outcome, which has been successfully applied in atmospheric environmental studies (Li et al., 2024; Lin et al., 2024). When the model was being adjusted, 90 % of the data was used as the

153 training data set, and 10 % of the data was used as the test data set. The hyperparameters were tuned using grid search and cross-
154 validation method. Specifically, for a single hyperparameter, grid search was used to obtain its more appropriate value range, and
155 for the combinations of hyperparameters, the whole training set was split into ten folds and then run a grid search over pre-adjusted
156 combinations of hyperparameters by training nine folds and predicting on the one fold in cross-validation procedure. For key
157 hyperparameters of XGBoost model, the number of trees was 100, learning rate was 0.1, max depth was 6. The model was trained
158 and tested on hourly data during the whole observation and the established model was examined by coefficient of determination (R^2)
159 value, the root-mean-squared error (RMSE) and mean absolute error (MAE). The formulas of RMSE and MAE are provided in the
160 supplementary information (Eq. S5 & Eq. S6). The performance of both models is illustrated in Fig. S3. The R^2 , MAE, and RMSE
161 for the training set are 0.90, 0.08, and 0.12, respectively, while the corresponding values for the test set are 0.77, 0.10, and 0.14,
162 respectively. These statistical metrics indicate that the XGBoost model is promising for further analysis.

164 3. Results and discussion

165 3.1 Overview of observation

166 The measured data of PAN, related trace gases and meteorological parameters at IUE over 10 to 31 July 2018 are documented in
167 Fig. 1. Combined with the synoptic situation shown in Fig. S4, the 8th typhoon of 2018, Typhoon Maria, made landfall on the
168 morning of the 11th at Huangqi Peninsula in Lianjiang County, Fujian. Due to the influence of the typhoon's outer spiral rain bands,
169 there was moderate to heavy rain on the 11 July 2018. Correspondingly, there was a noticeable decrease in ultraviolet radiation and
170 the temperatures. Starting from the 12 July 2018, a WPSH strengthened and extended westward, exerting control over Xiamen. In
171 the lower atmosphere, it was influenced by the eastward flow, resulting in predominantly cloudy weather. From the 16 to 18 July
172 2018, the area was affected by the outer periphery of Typhoon Shan Shen, which formed on the 17 July 2018, in the northeastern
173 part of the South China Sea and moved westward, making landfall along the coast of Wancheng Town, Wanning City, Hainan
174 Province in the early hours of the 17 July 2018. During this period, the city experienced strong winds with gusts reaching 5 to 6 on
175 the Beaufort scale in the urban areas. At the same time, the concentration of various pollutants reached their lowest levels, and the
176 daily variation patterns were less pronounced. From the 20 to 21 July 2018, Xiamen experienced the influence of the peripheral
177 descending airflow associated with Typhoon Ampil (which formed in the northwest Pacific Ocean around 8:00 PM local time (LT)
178 on the 18 July 2018, and moved northwest, making landfall along the coast of Chongming Island, Shanghai, around noon on the 22
179 July 2018). During this period, there were fewer clouds and higher temperatures. From the 22 to 24 July 2018, the city was
180 successively affected by the outer periphery of Typhoon Ampil and a tropical low-pressure system, resulting in occasional showers
181 or thunderstorms. From the 25 to 31 July 2018, a WPSH once again strengthened and controlled Xiamen. As a result, Xiamen
182 experienced stable meteorological conditions, with light winds ($w_s = 1.04 \text{ m s}^{-1}$), persistently high temperatures (maximum daily
183 average of $37.82 \text{ }^\circ\text{C}$), and high relative humidity (maximum daily average of 81.65 %). These factors created an environment that

184 favored the buildup of particulate matter and enhanced the photochemical formation of O₃ and PAN (Wu et al., 2019). The maximum
185 daily average of PM_{2.5}, O₃ and PAN were 49.26 μg.m⁻³, 93.62 ppb, and 1.37 ppb, respectively.

186 The phenomenon of simultaneous high levels of photochemical and particulate matter appears. Throughout the 22-days campaign,
187 12 days (including 11, 13, 21-23, and 25-31 July 2018) were observed with 1 h concentrations of PM_{2.5} exceeding 35 μg m⁻³; 13
188 days (including 11, 13, 15, 20-23, and 26-31 July 2018) were observed with 5-min concentrations of PAN exceeding 1 ppb. The
189 maximum concentration was recorded at 3.04 ppb (5-min data) at 11:09 LT of 13 July 2018. This concentration of PAN is comparable
190 to the levels recorded at downwind of Guangzhou, southern China (3.9 ppb, Wang et al., 2010), 2.51 ppb in Nashville, U.S (Roberts
191 et al., 2002). However, this value was significantly lower than heavily polluted areas in northern China in the summer, such as
192 Beijing (9.34 ppb, Xue et al., 2014), Lanzhou (9.12 ppb, Zhang et al., 2009), and Jinan (13.47 ppb, Liu et al., 2018). This is likely
193 because the higher summer temperatures in the southeastern coastal region are conducive to the thermal decomposition of PAN, and
194 the precursor concentrations of PAN, including NO₂ and VOCs, are significantly lower in the studied area compared to those in the
195 northern region. The concentration of alkanes is the highest, followed by alkenes, OVOCs and aromatics, while halogenated
196 hydrocarbons and C₅H₈ exhibit lower concentrations (Fig. S5). Furthermore, VOC concentrations for various species are elevated
197 during haze periods compared to clean periods (Fig. S5). Table S1 provides the detailed VOC concentrations observed during the
198 study period. Throughout the observation period, the variations in O₃ and PAN were almost identical, but the maximum
199 concentration of O₃ occurred at 3:00 PM LT on 29 July 2018 (114.12 ppb). The correlation between the average daily values of PAN
200 and both BC and O₃ is strong, with a correlation coefficient of 0.77 for each (Fig. S6), suggesting that summertime haze and
201 photochemical pollution were deeply connected.

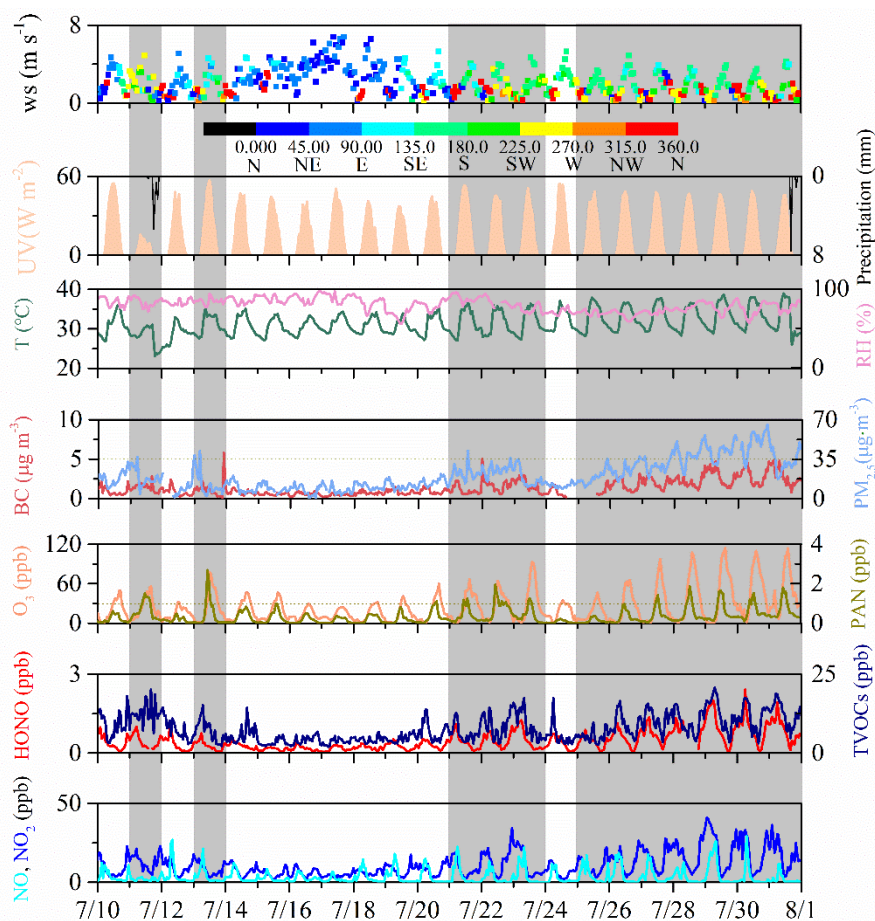


Figure 1. Time series of trace gases and meteorological parameters observed at IUE during 10-31 July 2018. The gray shading represents days when the PM_{2.5} hourly daily maximum value exceeded 35 $\mu\text{g m}^{-3}$.

We categorize it as "haze" and "clean" based on whether the PM_{2.5} hourly daily maximum value is greater than 35 $\mu\text{g m}^{-3}$. Specifically, "haze" includes 11, 13, 21-23, and 25-31 July 2018, while other days are categorized as 'clean'. To provide a quantitative perspective, the statistics for PAN and associated species were calculated and compiled in Table 1. PM_{2.5} concentrations during the haze period were significantly higher than those during the clean period, being 2.49 times those of the clean period. There was no significant difference of UV levels between clean and haze periods, while temperatures in the haze phase were notably higher than those in the clean phase. Therefore, without considering precursors, PAN concentrations should be lower during the haze phase due to higher thermal decomposition. In fact, PAN concentrations during the haze period were 2.35 times higher than those during the clean period. During the haze period, O₃ concentrations were also significantly higher than those during the clean period, being 2.04 times those of the clean period. These observations indicate that the atmospheric oxidation capacity is relatively strong during the haze period. Similar to PAN, HONO also exhibits higher concentrations during the haze phase (approximately 2.33 times that of clean conditions), which is consistent with current research findings that particles promote the generation of HONO (Ye et al., 2017). NO also experienced an increase from clean (3.28 ppb) to hazy (4.30 ppb) conditions, albeit less prominently than NO₂ (from 7.21 to 14.55 ppb). This observation further underscores that, during hazy periods, the

219 atmosphere demonstrates heightened oxidizing potential, facilitating the conversion of NO to NO₂. While the increased NO levels
 220 on hazy days reduced PA radicals and hindered PAN formation, this effect was offset by the concurrently higher concentrations of
 221 PAN precursors (NO₂ and VOCs) during those days. The TVOCs have increased to some extent, but in hazy conditions, they are
 222 only 1.34 times that of clean conditions. This is also because the strong oxidizing conditions during haze periods convert VOCs
 223 into secondary pollutants, such as O₃ and PAN. Although it is acknowledged that VOCs can also be converted into secondary
 224 organic aerosol (SOA), the discussion of SOA is beyond the scope of this study. The TVOC levels at this site are comparable to
 225 that at a rural site in a coastal city-Qingdao (7.6 ppb), significantly lower than inland sites, such as Wuhan (30.2 ppb, Liu et al.,
 226 2021a) and Chengdu (28.0 ppb, Yang et al., 2020) or economically more developed coastal cities, such as Shanghai (25.3 ppb, Zhu
 227 et al., 2020) and Hong Kong (26.9 ppb, Wang et al., 2018), and significantly higher than regional background locations like Mt.
 228 Wuyi (4.7 ppb, Hong et al., 2019) and Mt. Nanling (4.7 ppb, Wang et al., 2023), and global background station Mt. Waliguan (2.6
 229 ppb, Xue et al., 2013). The isoprene level during haze period was significantly higher than that during clean period probably due
 230 to haze period with higher temperature (Wang et al., 2023). The wind speed was very low during both the clean and haze periods,
 231 especially during the haze period with only 1.12 m s⁻¹. The relative humidity was high during both periods, and there was no
 232 significant difference between the clean and haze periods.

233 **Table 1.** Descriptive statistics of major trace gases (ppb), particulate matter (μg m⁻³) and meteorological parameters during 10-31
 234 July 2018.

Species	Clean (mean ± SD)	Haze (mean ± SD)
PAN	0.20 ± 0.23	0.47 ± 0.46***
O ₃	16.07 ± 12.73	32.79 ± 29.73***
HONO	0.27 ± 0.18	0.63 ± 0.43***
NO	3.28 ± 4.03	4.30 ± 8.39***
NO ₂	7.21 ± 3.87	14.55 ± 8.89***
TVOCs	6.13 ± 1.73	8.19 ± 2.55***
C ₅ H ₈	0.13 ± 0.04	0.17 ± 0.05***
PM ₁	10.13 ± 3.91	24.36 ± 10.77***
PM _{2.5}	11.21 ± 5.33	27.93 ± 13.16***
PM ₁₀	24.26 ± 9.45	47.28 ± 20.63***
UV (W m ⁻²)	14.29 ± 17.38	13.18 ± 17.40
T (°C)	30.68 ± 2.39	31.92 ± 3.36***
RH (%)	81.94 ± 8.60	77.18 ± 8.22
WS (m s ⁻¹)	1.64 ± 0.69	1.12 ± 0.61*

235 Note: *, **, and *** indicate that they passed the significance test at 0.05, 0.01 and 0.001 levels, respectively.

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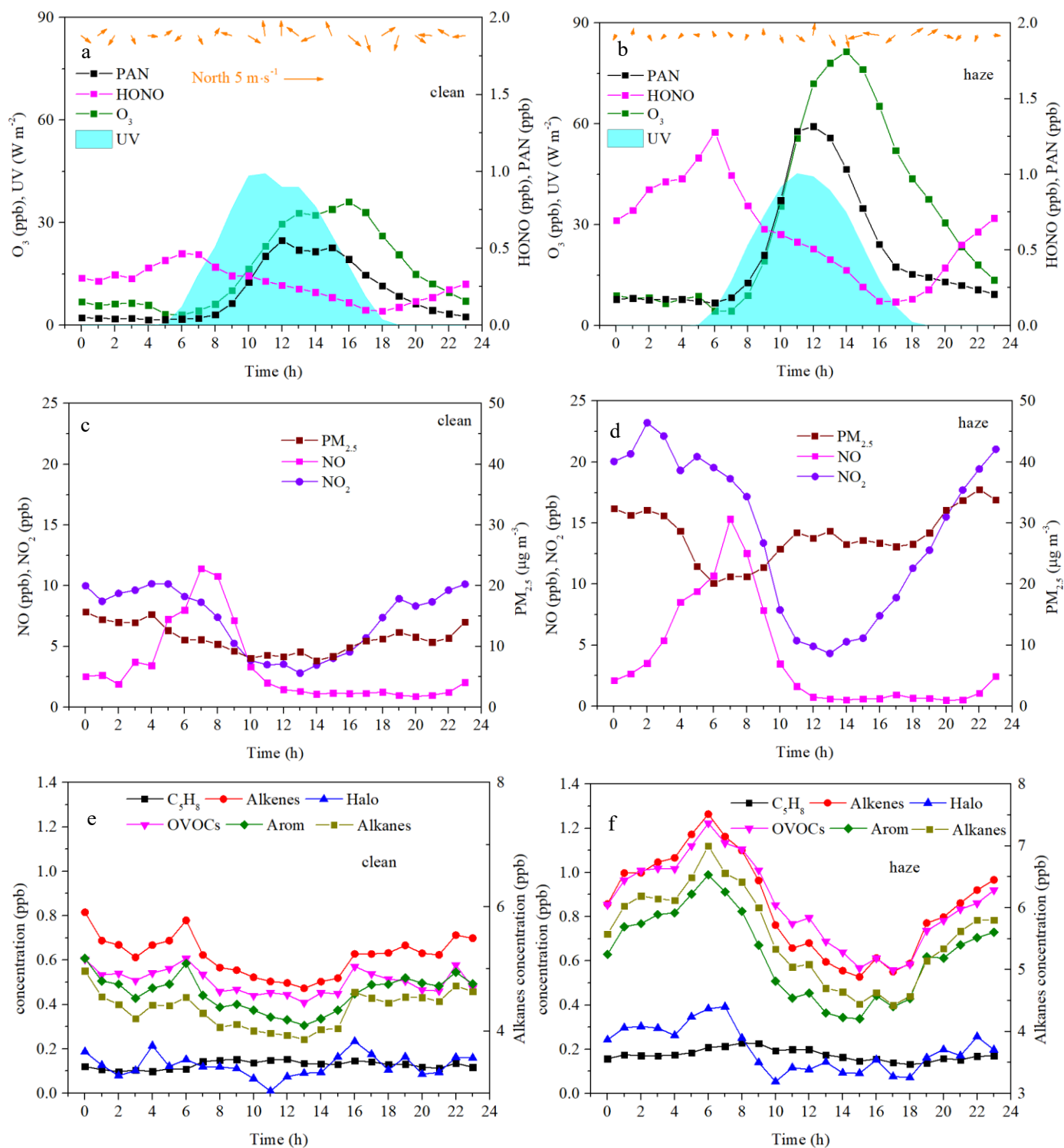
237 The average diurnal patterns of PAN and related variables have been averaged separately for clean and hazy conditions (Fig. 2).

238 The daily variation of PAN exhibits a clear unimodal pattern, with concentrations starting to rise after sunrise and decreasing after

239 12:00 LT caused by thermal decomposition of PAN at high temperatures (Fig. 2). Although PAN and O₃ exhibit a slight bimodal
240 pattern during the clean period, this is primarily due to the bimodal pattern of UV during this time. The peak occurring at noon
241 indicates that PAN primarily originates locally, as a delay of about 1-2 hours would be expected if it were influenced by
242 transportation (Liu et al., 2024). The daytime increment was much larger for hazy condition (1.17 ± 0.44 ppb) than for clean
243 condition (0.52 ± 0.21 ppb), indicating stronger photochemical production of PAN for hazy condition. The daily variation pattern
244 of O₃ is similar to PAN, except that O₃ reaches its peak relatively later compared to PAN, with the peak occurring at 16:00 during
245 the clean phase and 14:00 LT during the haze phase. Although PAN and O₃ are both products of photochemical reactions involving
246 NO_x and VOCs, their production efficiencies differ. PAN is specifically formed from VOCs that are precursors to the acetyl radical
247 (CH₃CO), whereas O₃ can be produced from the oxidation of any VOCs. Analyzing the correlation between PAN and O₃ can offer
248 insights into their respective photochemical production efficiencies. As shown in Fig. S7, the positive correlation between the daily
249 maximum values of PAN and O₃ for clean condition ($R^2=0.67$) was better than that for hazy condition ($R^2=0.15$). The slopes of
250 the linear regression were 0.021 ppb/ppb for clean conditions and 0.009 ppb/ppb for hazy conditions. This indicates that, on average,
251 approximately 2.1 ppb of PAN could be produced for each 100 ppb of O₃ formed under clean conditions, and about 0.9 ppb of
252 PAN for each 100 ppb of O₃ under hazy conditions in the air masses reaching IUE. The slope of linear regression for clean condition
253 is comparable to those determined in Hongkong (0.028, Xu et al., 2015), Mexico city (0.020, Marley et al., 2007a), and Nashville
254 (0.025, Roberts et al., 2002). The lower efficiency of PAN production relative to O₃ indicates that PAN precursors represent only
255 a small portion of the total VOCs, especially during hazy conditions. Additionally, the high temperatures in the southeast coastal
256 region likely contribute to the lower production efficiency of PAN. The average temperature during the entire observation period
257 was 31.39 °C, with an average temperature of 34.64 °C at 12:00 LT. This result is consistent with the result that RIR during the
258 cleaning period is higher than that during the haze period. As shown in Fig. S8, in the clean period, the correlation between PAN
259 and O₃ is the strongest ($R^2=0.70$), indicating that O₃ and PAN are both photochemical end products during clean periods. In contrast,
260 during hazy periods, the correlation between PAN and O₃×JO¹D is the strongest ($R^2=0.66$), suggesting that O₃ plays a more
261 significant role in promoting PAN formation through photolysis to generate OH during hazy periods.

262 Unlike the daily variation patterns of PAN and O₃, HONO exhibits a swift concentration decrease after sunrise in both clean and
263 hazy conditions, undergoing photolytic conversion into OH radicals. Subsequently, in clean conditions, HONO starts to increase
264 in concentration after sunset. In hazy conditions, however, the increase begins from 16:00 LT and not after sunrise. This suggests
265 a robust daytime net production or transport of HONO, where the rates surpass those of HONO photolysis and other sinks in the
266 afternoon in hazy conditions. The NO levels reach their peak at 7:00 LT during the morning rush hour, reflecting advection of fresh
267 urban plumes to the study site. The daily variation of NO₂ exhibits a 'U' shape, reaching its minimum value at 13:00 LT, mainly
268 owing to effects of emission, boundary layer height and photochemical reactions. In the clean period, the daily variation of PM_{2.5}
269 is similar to that of NO₂, both showing a 'U' shape, reaching their lowest values at noon. However, during the haze phase, the daily

270 variation pattern of $PM_{2.5}$ appears somewhat different. There is a noticeable trough in the early morning, remains stable during the
 271 day, and starts to rise after sunset. The diurnal variation of VOC concentrations for various species are not significant during clean
 272 periods (Fig. 2(e)), likely due to higher wind speeds that facilitate the dispersion of pollutants. In contrast, during haze periods, the
 273 daily variations are evident, with peaks occurring before sunrise, followed by a decline, and then an increase after sunset (Fig.
 274 2(f)). This is because the haze period is relatively stable at nighttime, which allows for the accumulation of pollutants, while during
 275 the daytime, sunlight converts VOCs into photochemical products like O_3 and PAN.



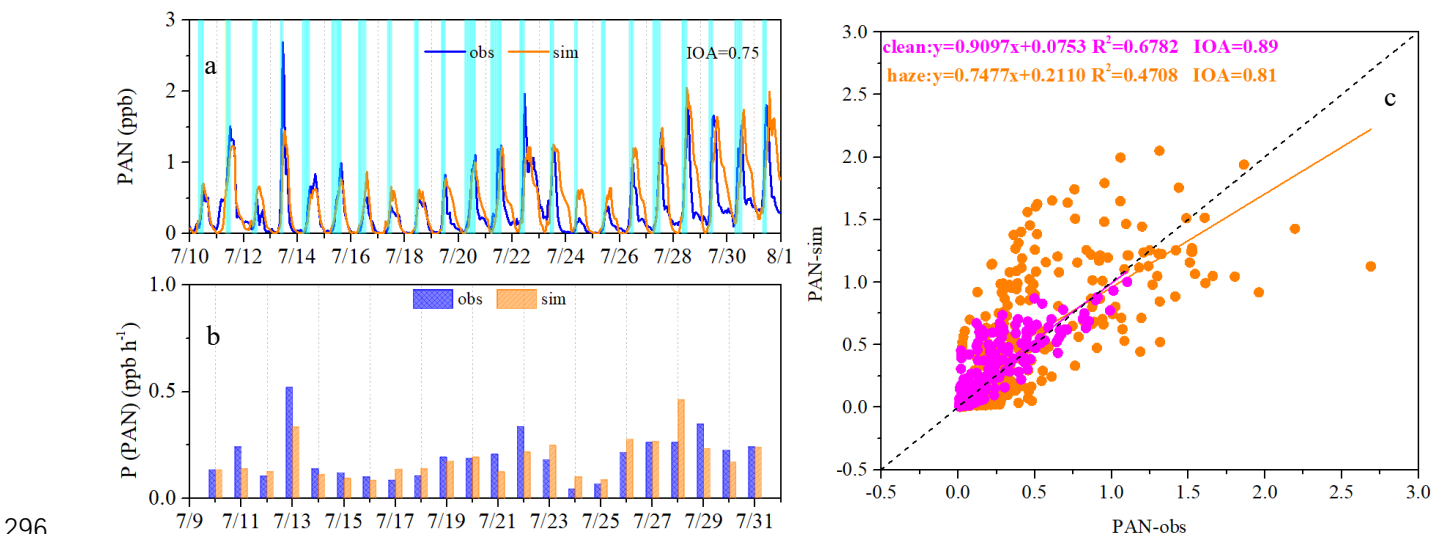
278
 279 **Figure 2.** The diurnal variations of PAN, HONO, O_3 , and UV during clean (a) and hazy (b) periods, the diurnal variations of $PM_{2.5}$,

280 NO, and NO₂ during clean (c) and hazy (d) periods, and the diurnal variations of isoprene (C₅H₈), Alkenes, halogenated
281 hydrocarbons (Halo), OVOCs, aromatics (Arom), and alkanes during clean (e) and hazy periods (f).

282

283 3.2 PAN formation: key factors and mechanisms

284 To investigate the key factors and mechanisms of PAN formation, PAN was simulated by constraining the MCM-based box model
285 with meteorological conditions and observed concentrations of precursor gases. The model successfully replicated the variations
286 in PAN, achieving an IOA of 0.75 (Fig. 3(a)), which was within the accepted range (0.66-0.87) in previous studies (Zeng et al.,
287 2019). The model captured its formation rate in general well, with observed rates varying from 0.04 to 0.52 ppb·h⁻¹ (average: 0.20
288 ppb·h⁻¹) and modeled rates ranging from 0.09 to 0.46 ppb·h⁻¹ (average: 0.19 ppb·h⁻¹) (see Fig. 3(b)). The similar result was found
289 in the North China Plain (NCP) region in the wintertime (Xu et al., 2021). When calculating the IOA separately for clean and hazy
290 periods, it was found that the IOA significantly increased to 0.89 and 0.81 (Fig. 3(c)), respectively. This phenomenon indicates a
291 substantial difference in the PAN production and destruction mechanisms between clean and hazy periods. Furthermore, the
292 simulated values are closer to the observed values during clean period, reflected in a higher R² value (R²=0.68) and a slope value
293 (K) closer to 1 (K=0.91) (Fig. 3(c)). In contrast, the R² value and the K value during hazy period are only 0.47 and 0.75, respectively
294 (Fig. 3(c)). This phenomenon suggests that some reactions related to PAN generation or destruction might be missing in the MCM
295 during the hazy period.



296

297 **Figure 3.** Comparisons of observed (obs) PAN and simulated (sim) PAN (daytime photochemical PAN production periods
298 indicated by cyan shading), (b) production rates, (c) correlation between PAN observations and simulated values.

299

300 To identify the key factors influencing the performance of the OBM model simulation, the bias (the model simulation minus the
301 observed value) as the target. The remaining variables, which were not input into the OBM model, such as NH₃, HNO₃, HCl
302 (alkaline and acidic gaseous pollutants), PM_{2.5} concentrations and their components, as well as physical process parameters like
303 wind speed and wind direction, were used as features. As shown in Fig. 4 (a), through XGBoost-SHAP machine learning, we found

304 that NH_3 is the most significant parameter affecting bias, contributing 19.68 % (Fig. S9). A scatter plot analysis of the SHAP values
 305 of NH_3 versus NH_3 concentrations revealed that as NH_3 concentrations increase (Fig. 4 (b)), the OBM model tends to overestimate
 306 PAN concentration more significantly. To date, there are very few studies that directly address the impact of NH_3 on PAN formation.
 307 Xu et al. (2021) suggested that NH_3 could promote the formation of HONO, which in turn affects PAN formation. However, since
 308 we included HONO as an input to constrain the model, the indirect influence of NH_3 on PAN formation through HONO can be
 309 excluded. NH_3 in the atmosphere can preferentially react with sulfuric acid (H_2SO_4) to form ammonium sulfate ($(\text{NH}_4)_2\text{SO}_4$)
 310 secondary inorganic aerosols (Behera et al., 2013), leading to the heterogeneous reaction removal of PAN by secondary inorganic
 311 aerosols (Pratap et al., 2021). This result is validated by the positive correlation between the SHAP values of NH_4^+ and SO_4^{2-} and
 312 their respective concentrations (Fig. S10). NO_3^- is the second most significant parameter influencing the bias between the two,
 313 contributing 11.33 % (Fig. S9). NO_3^- has a negative correlation with the bias (Fig. 4 (c)), indicating that higher NO_3^- levels lead to
 314 more significant underestimation by the model. Considering the significant positive correlation between PAN and NO_3^- at the 0.01
 315 level, with a correlation coefficient of 0.37, and the fact that both reach their peaks around noon (Fig. S11), it is likely that they
 316 have a common source. $\text{PM}_{2.5}$ is the third most significant parameter (Fig. 4 (a)), contributing 9.40 % (Fig. S9). $\text{PM}_{2.5}$ has a positive
 317 correlation with the bias (Fig. 4 (d)), indicating that higher $\text{PM}_{2.5}$ levels lead to more significant overestimation by the model,
 318 suggesting that PAN can undergo heterogeneous removal on the surface of $\text{PM}_{2.5}$ in the actual atmosphere (Sun et al., 2022).

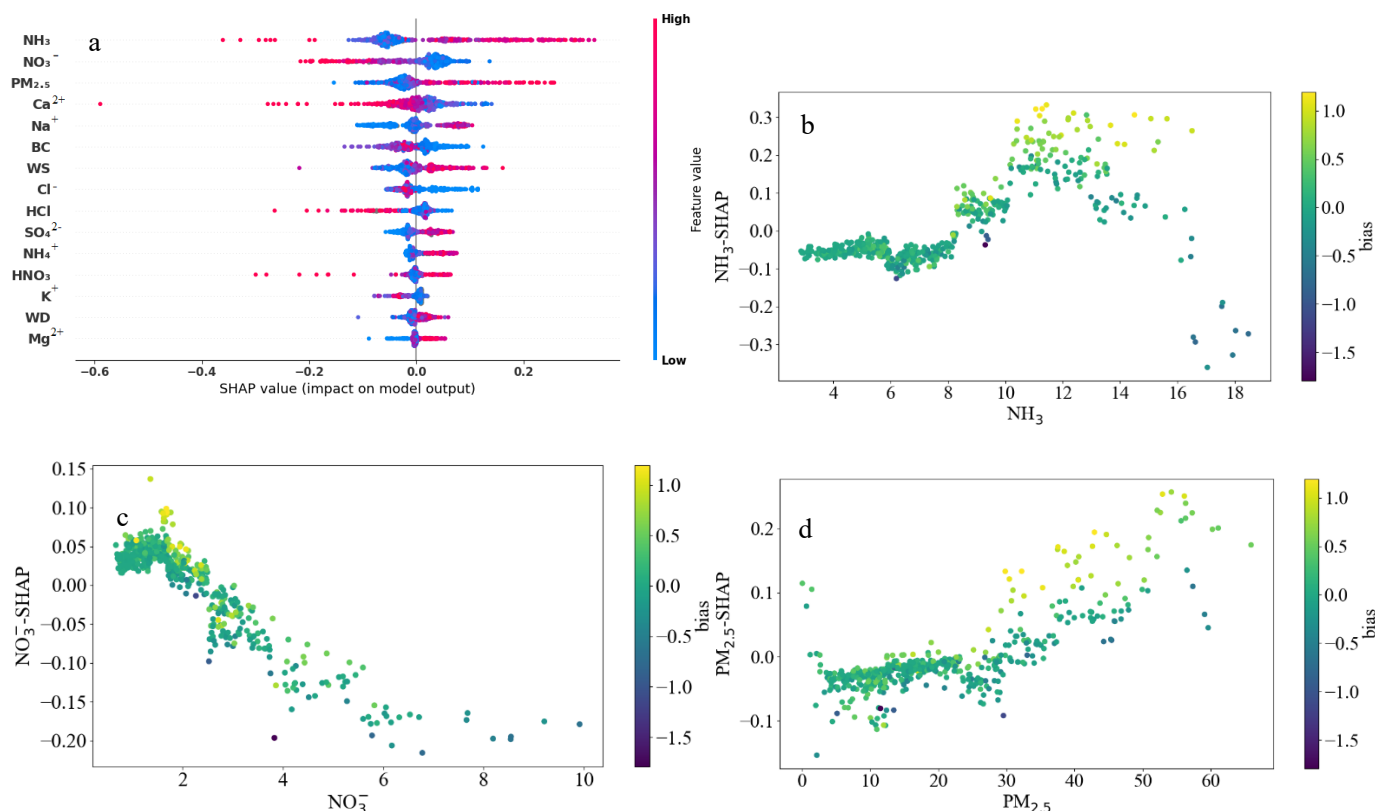


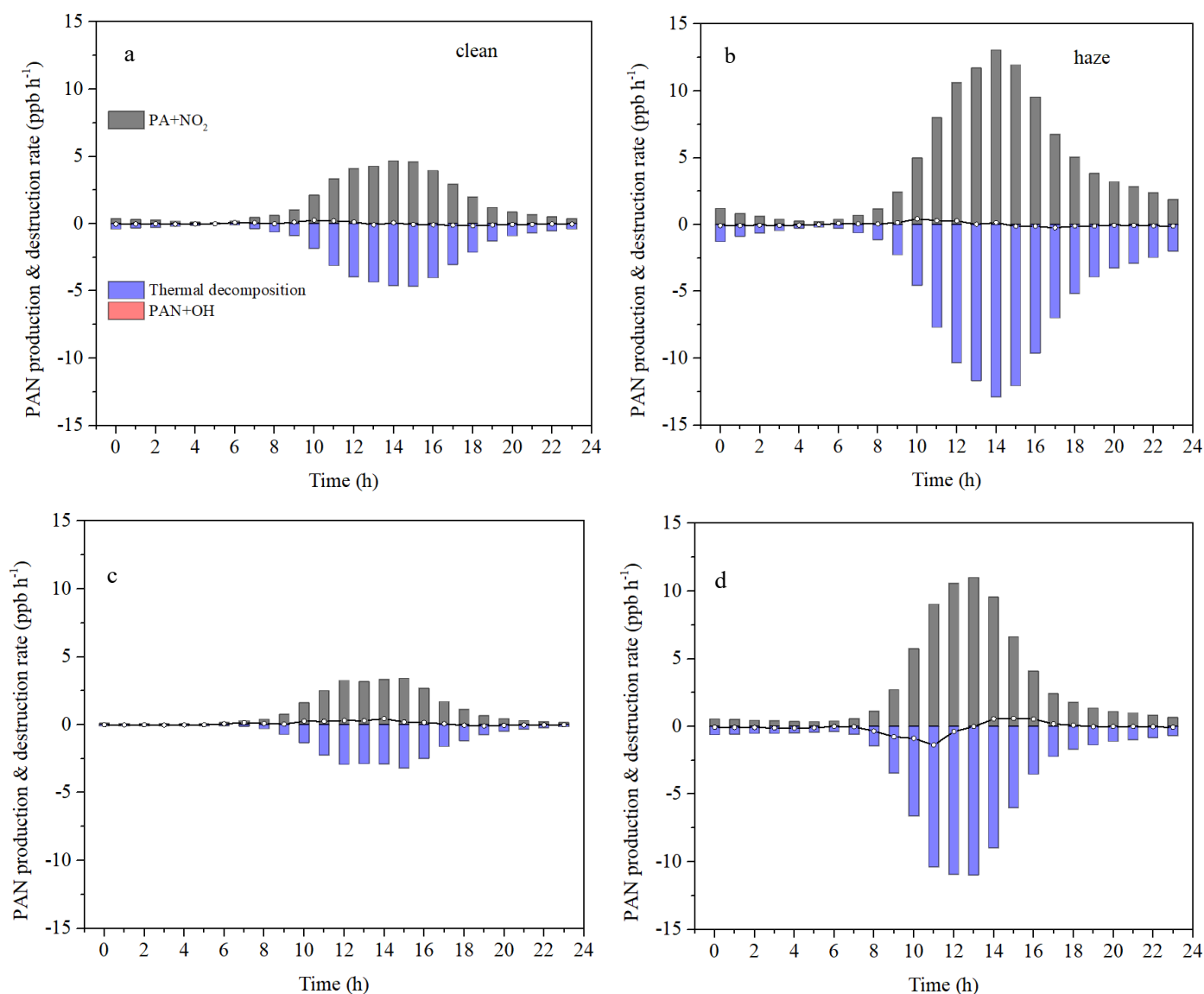
Figure 4. Feature importance was obtained by XGBoost-SHAP method (a). The scatter plots between concentration of top three important features and their SHAP values (b, c and d), and colored with the bias (the model simulation minus the observed value).

323

324 Figure 5 (a) and (b) show the average production and destruction rates of PAN during clean and haze periods, as simulated by
325 OBM without PAN constrained. During the haze period, both the production and destruction rates of PAN are significantly higher
326 than during the clean period. The higher production rate of PAN during the haze period is due to the higher concentration of PAN
327 precursors, while the higher destruction rate is because both the temperature and PAN concentration are higher. Regarding the net
328 production rate, it is also higher during the haze period than during the clean period, which corresponds to the previously observed
329 diurnal variation. From 6:00 to 12:00 LT during the haze period, the simulated net production rate of PAN is positive, with an
330 average value of 0.19 ppb h⁻¹. During the clean period, from 6:00 to 12:00 LT, the simulated net production rate of PAN is 0.12
331 ppb h⁻¹. The observed diurnal variation of PAN shows that from 6:00 to 12:00 LT, the average net production rates during the haze
332 and clean periods are 0.20 ppb h⁻¹ (Fig. 2(a)) and 0.09 ppb h⁻¹ (Fig. 2(b)), respectively. The model-simulated net production rate is
333 close to the observed net production rate, further indicating that the model can simulate PAN well, and also confirming that PAN
334 in summer mainly comes from local production. The net production rate of PAN during the haze period is similar to the summer
335 results in urban areas of the Pearl River Delta (PRD), which is 0.17 ppb h⁻¹, while the net production rate of PAN during the clean
336 period is similar to the summer results in rural areas of the PRD, which is 0.12 ppb h⁻¹ (Liu et al., 2024).

337 Figure 5 (c) and (d) show the average production and destruction rates of PAN during clean and haze periods, as simulated by OBM
338 with PAN constrained. The net production rate of PAN is approximately zero at night during both clean and haze periods, while
339 there is a significant difference in the net production rate during the day. During the clean period, the daytime net production rate of
340 PAN is greater than zero, with an average value of 0.19 ppb h⁻¹. In contrast, during the haze period, the net production rate of PAN
341 is negative from 6 AM to 1 PM LT, with an average value of -0.47 ppb h⁻¹, and positive from 2 PM to 5 PM LT, with an average
342 value of 0.47 ppb h⁻¹. Previous research has shown that an increase in temperature, an increase in PAN concentration, or a decrease
343 in PAN precursors (including VOCs and NO₂) can cause the net production rate of PAN to change from positive to negative (Liu et
344 al., 2024). We conducted a correlation analysis of the net production rate of PAN with temperature, PAN, VOCs, and NO₂
345 concentration and found that the net production rate of PAN had the best correlation with PAN concentration ($R^2=0.13$), showing a
346 significant negative correlation ($k=-0.53$, Fig. S12). Additionally, we also observed that when the net production rate of PAN is
347 negative, the PAN concentration is often very high (Fig. S12). As shown in Fig. 6, we conducted sensitivity experiments by reducing
348 the PAN concentration by 80 %, i.e., 0.2 times the observed value, and found that the simulated net production rate of PAN was
349 positive throughout the observation period. Conversely, when the PAN concentration was increased by 140 %, i.e., 2.4 times the
350 observed value, the simulated net production rate of PAN was found to be almost negative throughout the observation period.
351 Besides, we also conducted sensitivity experiments on temperature and found that when simulating winter temperatures, i.e., 0.4
352 times the observed value, with a temperature range of 9.25-15.29 °C, the simulated net production rate of PAN was positive
353 throughout the observation period. Similarly, when simulating spring and autumn temperatures, i.e., 0.6 times the observed value,

354 with a temperature range of 13.87-23.39 °C, the simulated net production rate of PAN was also positive throughout the observation
 355 period. In conclusion, the simulated net production rate of PAN becomes negative with PAN constrained, further suggesting the
 356 existence of an unknown compensatory mechanism.

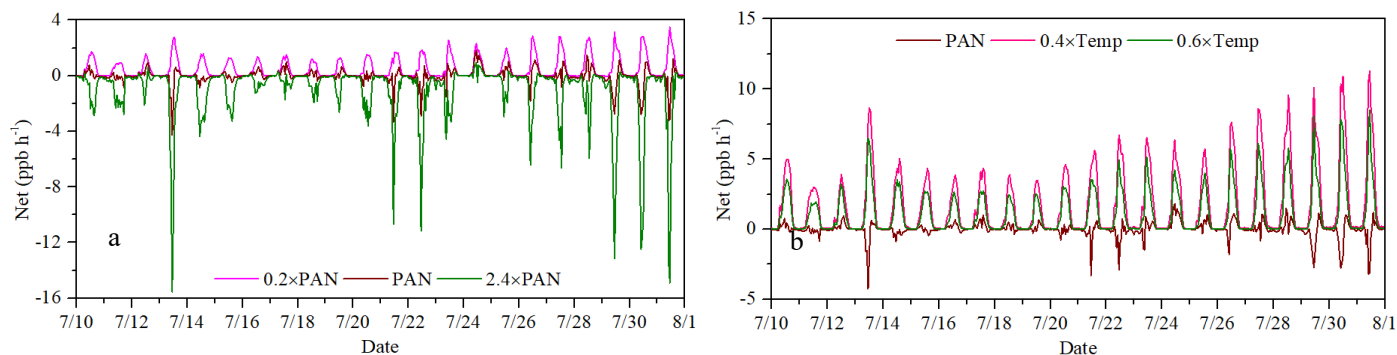


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358

359 **Figure 5.** Average diurnal variation of the OBM simulated production, destruction and net rates of PAN during clean (a) and haze
 360 days (b) without PAN constrained. And average diurnal variation of the OBM simulated production, destruction and net rates of
 361 PAN during clean (c) and haze days (d) with PAN constrained.

362



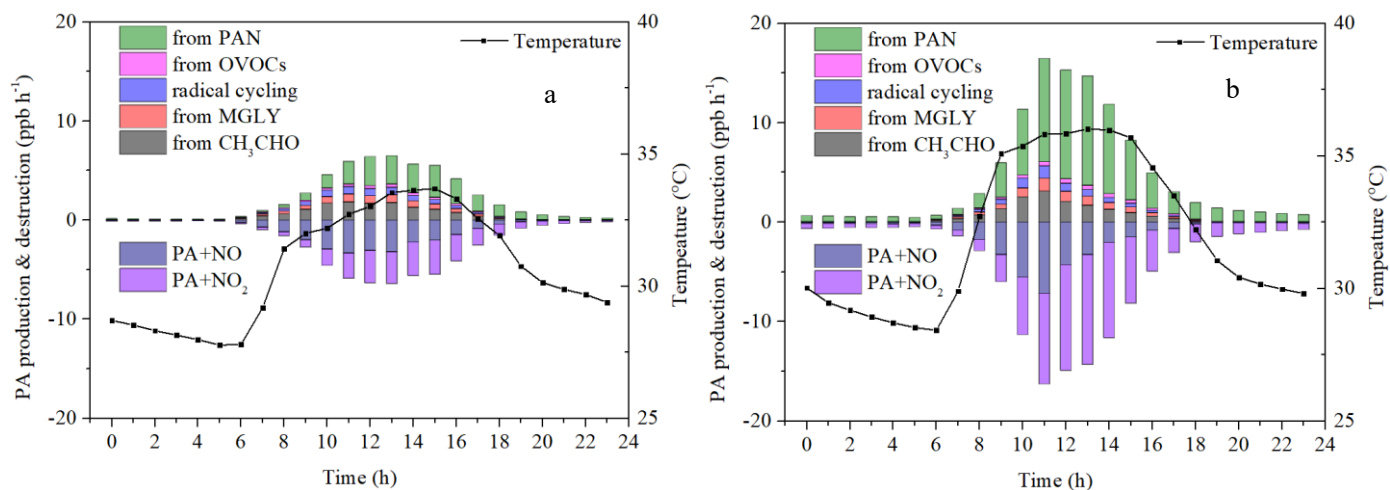
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364 **Figure 6.** Net PAN production rates simulated by OBM at different PAN concentrations (a) and different temperatures (b).

365
366 PAN is formed when the PA radical reacts with NO₂. Given the swift equilibrium between R2 and R4 at high temperatures, budget
367 analysis of PA's production and consumption pathways are frequently used to detail the mechanisms behind PAN formation (Sun et
368 al., 2020; Liu et al., 2022a; Liu et al., 2024). Figure 7 illustrates the diurnal patterns of the primary production and loss pathways
369 for the PA radical simulated by OBM across different periods. As shown in Fig. 7, during haze days, the rates of PA production and
370 destruction were twice as high as those on clean days. This indicates that radical cycling and photochemical formation were more
371 efficient during haze days, driven by higher temperatures and a greater abundance of precursors (Zeng et al., 2019). The PA radical
372 production rate from PAN thermal decomposition reached its peak at 15:00 LT (3.22 ppb h⁻¹) and 13:00 LT (10.99 ppb h⁻¹) for clean
373 and haze days, perfectly coinciding with the peak temperature time. In addition, the conversion of PAN into PA radical through
374 thermal decomposition had high exponential correlations with temperature during both haze (R²=0.95) and clean days (R²=0.91)
375 (Fig. S13). Previous laboratory experiments also indicated that the thermal decomposition of PAN is exponentially related to
376 temperature (Cox & Roffey 1977; Senum et al., 1986; Tuazon et al., 1990). The conversion of PAN into PA radical through thermal
377 decomposition during haze days was significantly higher than that during clean days, which was not only enhanced by higher
378 temperature but also maintained by higher PAN concentration during haze days. The thermal decomposition of PAN to PA radical
379 during the day (5:00-18:00 LT) accounted for 68.22 % and 45.59 % during haze and clean days, respectively. The pathways that did
380 not account for the transformation between PA and PAN reached their peak around noon (11:00 LT), coinciding with the highest
381 solar radiation and the most intense photochemical reactions, which has been observed in spring and autumn at the same site (Liu
382 et al., 2022a).

383 Production rates of PA from other pathways related to precursors, including OVOCs, radical cycling, MGLY, and CH₃CHO, showed
384 single-peak patterns around noon, which suggested that the PA radical generated from these pathways was primarily increased by
385 intense solar radiation at noontime (Sun et al., 2020). The average day PA radical production rates from CH₃CHO via reactions with
386 OH and NO₃ were 1.10 and 0.93 ppb h⁻¹, accounting for 48.85 % and 49.35 % (exclude PAN thermal decomposition sources) during
387 haze and clean days, respectively. These percentages were comparable to previous studies in Guangzhou (46 %, Yuan et al., 2018)
388 and Beijing (34.11-50.19 %, Xue et al., 2014), suburban site of Chongqing (47.72 %, Sun et al., 2020). The second production
389 pathway involved MGLY undergoing photolysis and oxidation through reactions with OH and NO₃ (haze: 0.50 ppb h⁻¹ and clean:
390 0.42 ppb h⁻¹), contributing to 22.27 % and 22.12 % for haze and clean days, respectively. Subsequently, radical cycling processes—
391 including the decomposition of RO radicals and the reactions of acyl peroxy radicals with NO—were also significant contributors
392 to PA production, accounting for 18.98 % on haze days and 19.54 % on clean days. PA from the other OVOCs (excluding CH₃CHO,
393 MGLY) via photolysis and oxidation reactions involving OH, NO₃, and O₃, accounted for 9.90 % and 8.99 % during haze (0.22 ppb
394 h⁻¹) and clean days (0.17 ppb h⁻¹). There were no notable differences in the proportions of individual pathways contributing to PA

395 haze and clean days, indicating comparable pollutant compositions in the atmospheric around IUE (Zeng et al., 2019). In
 396 summary, the thermal decomposition of PAN played the dominant role in boosting PA production rates during both clean and haze
 397 periods, followed by contributions from CH₃CHO, MGLY, radical cycling, and other OVOCs. The primary contributor to the PA
 398 destruction rate was the reaction between PA and NO₂, accounting for 67.72 % and 51.09 % during haze (4.74 ppb h⁻¹) and clean
 399 days (1.76 ppb h⁻¹), respectively, followed by PA+NO, contributing to 32.28 % and 48.91 % during haze (2.26 ppb h⁻¹) and clean
 400 days (1.69 ppb h⁻¹), respectively.



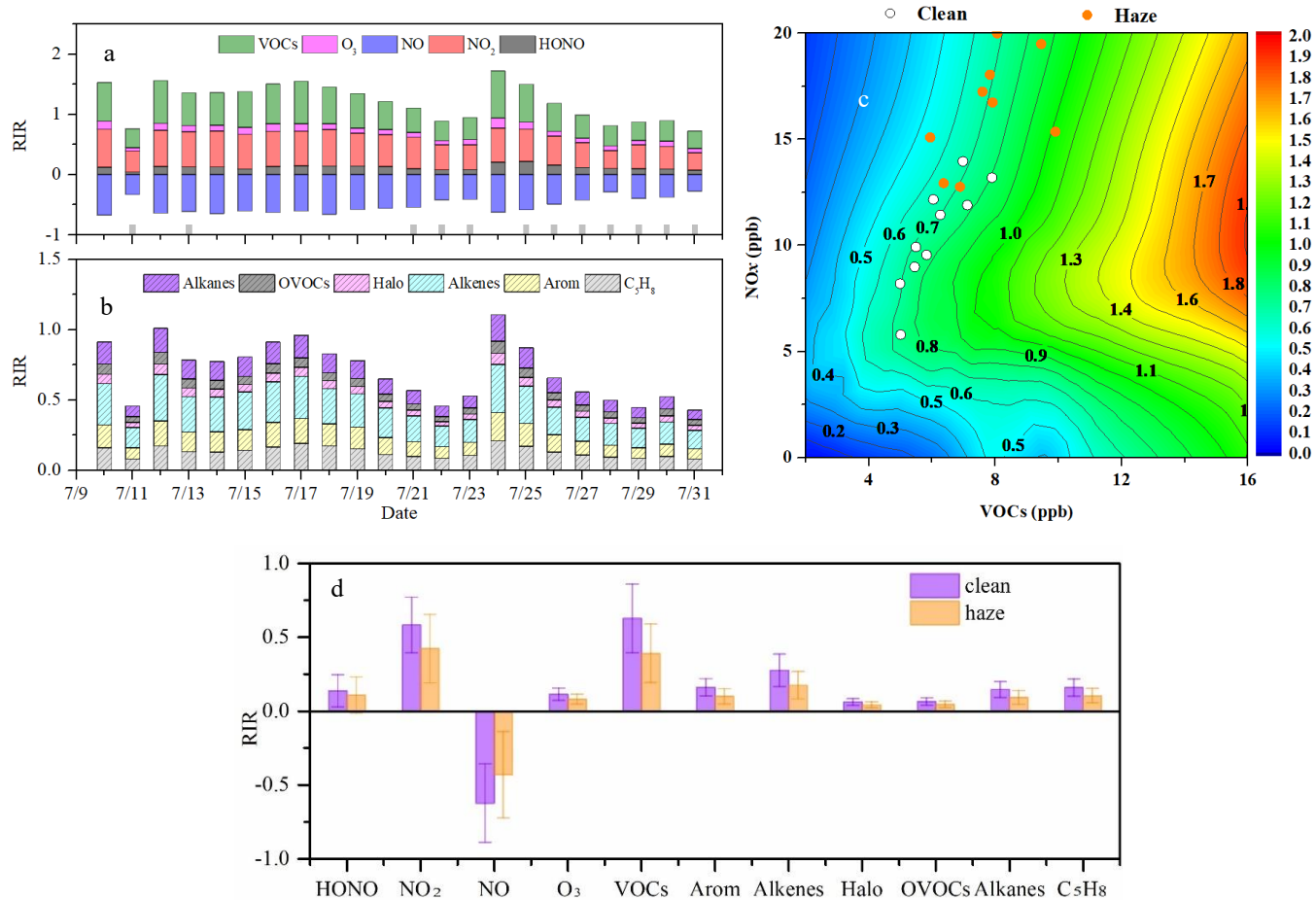
401
 402 **Figure 7.** PA radical production and destruction pathways simulated by OBM on (a) clean days and (b) haze days.

403
 404 **3.3 Sensitivity of PAN formation and its impact on the local atmosphere**

405 To determine the principal precursors influencing PAN formation, sensitivity modeling analyses were carried out to investigate
 406 how PAN relates to its precursors. The RIR reflects how sensitive PAN formation is to changes in its precursor levels. As shown
 407 in Fig.8 (a), the RIR of NO was negative, ranging from -0.67 to -0.27 (-0.52 ± 0.13) throughout the observation period. However,
 408 RIR is positive for other species, with NO₂ (0.50 ± 0.11) and VOCs (0.50 ± 0.15) having the highest RIR, followed by HONO
 409 (0.12 ± 0.04) and O₃ (0.10 ± 0.03). Around 50 types of VOCs were classified as alkanes, OVOCs, halogenated hydrocarbons
 410 (Halo), alkenes, aromatics (Arom), and isoprene (C₅H₈ representing biogenic hydrocarbons). Among these VOCs, the RIR of
 411 alkenes (0.22 ± 0.07) is the highest, followed by C₅H₈ (0.13 ± 0.04) and Arom (0.13 ± 0.04), while OVOCs (0.06 ± 0.01) and Halo
 412 (0.05 ± 0.01) have very low RIRs (Fig. 8 (b)). These phenomena indicated that increased NO level would inhibit the production
 413 of PAN while increased NO₂, VOCs (especially alkenes, C₅H₈, and Arom), HONO, and O₃ would promote the production of PAN.
 414 Because the values of NO and NO₂ RIR are approximately equal but with opposite signs, the RIR for NO_x is almost zero, indicating
 415 that the PAN generation at this site is not sensitive to NO_x. Zeng et al. (2019) also observed that NO₂ had a positive effect on PAN
 416 formation, while NO had a negative effect, in a suburban area of Hong Kong. This finding aligns with the fact that NO₂ directly
 417 contributes to PAN production, whereas NO reduces PA radicals, thereby inhibiting PAN formation. Based on the scenario analysis
 418 (Empirical Kinetic Modeling Approach (EKMA)), all data points for the 22 days fell above the ridge line (Fig. 8(c)). A reduction

419 in VOCs at these points resulted in lower PAN concentrations, indicating that PAN formation at IUE was influenced by VOCs and
 420 thus VOC-sensitive. Our previous research also found that in this coastal city, PAN generation is limited by VOCs during the
 421 spring and autumn seasons. The difference is that previous studies indicated that reducing NO₂, like reducing NO, also leads to an
 422 increase in PAN concentration in spring and autumn (Liu et al., 2022a). This is because the NO_x concentration in spring and autumn
 423 is significantly higher than in summer, which is consistent with that both NO₂ and NO inhibit the formation of PAN in regions
 424 with high NO_x concentrations (Liu et al., 2024).

425 We divided the RIRs for different species into haze and clean periods and found that the RIRs during clean periods were consistently
 426 higher than those during haze periods (Fig. 8(d)), which indicated that altering the concentrations of these species during clean
 427 periods had a greater impact on PAN formation. The rapid thermal decomposition of PAN at high temperatures is likely the primary
 428 reason. During the haze period, the main source of PA radical was PAN decomposition, which accounted for 68.22 %, and the other
 429 sources were smaller than that during the clean period (the source of the PA radical would be demonstrated in the following
 430 paragraph). Therefore, the sensitivity of PAN production to precursors and HONO & O₃ producing OH became lower during the
 431 haze period (Liu et al., 2021b; Liu et al., 2022a).



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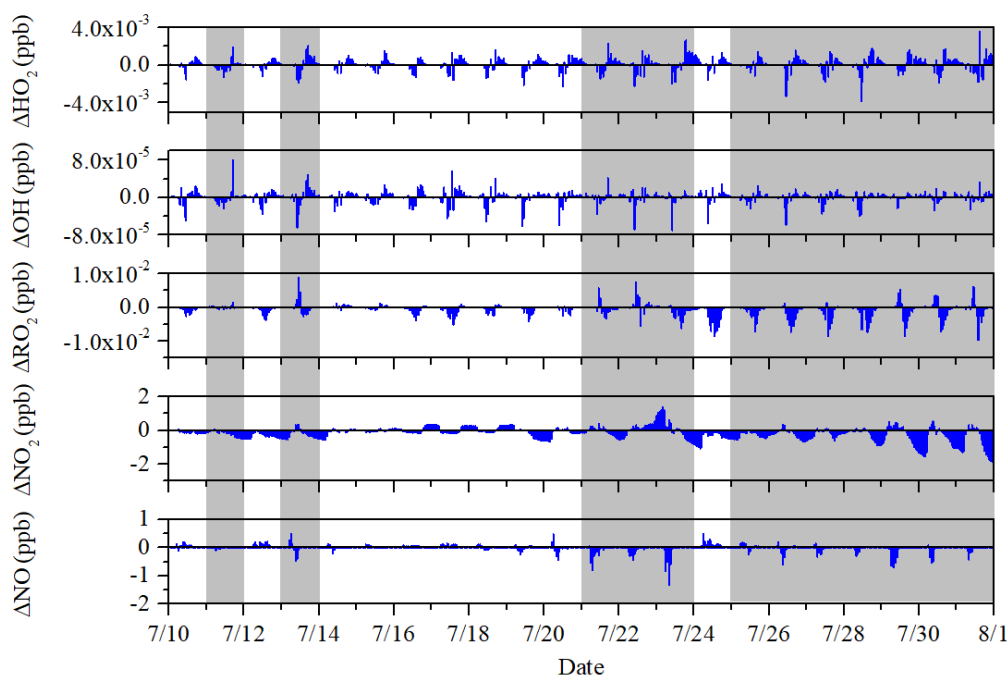
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Figure 8. These four figures illustrate the RIR of PAN formation to major precursors (a), the impact of different VOCs species (b), the isopleth diagrams of PAN formation (c), and a comparison of RIRs between clean and polluted periods (d).

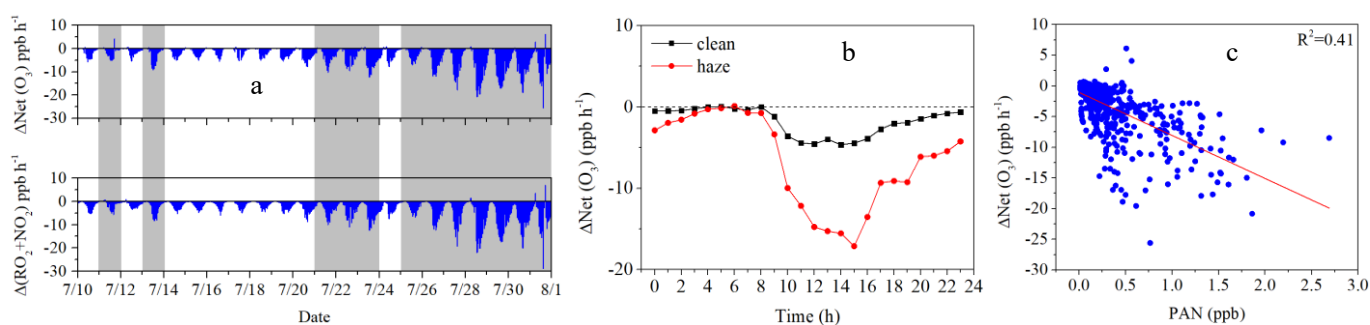
437 As shown in Fig. 9, ΔHO_2 and ΔOH are positive for most periods, accounting for 72.16 % and 70.83 %, respectively, indicating that
 438 the PAN mechanism promotes the generation of HO_2 and OH . Over the entire period, ΔHO_2 is $8.43 \cdot 10^{-5}$ ppb, with no significant
 439 difference between clean and hazy periods, being $8.18 \cdot 10^{-5}$ ppb and $8.64 \cdot 10^{-5}$ ppb respectively (Table S2). OH behaves similarly,
 440 with ΔOH being $4.55 \cdot 10^{-7}$ ppb over the entire period, and also showing no significant difference between clean and hazy periods,
 441 being $4.94 \cdot 10^{-7}$ ppb and $4.23 \cdot 10^{-7}$ ppb respectively (Table S2). The increase in simulated OH and HO_2 concentrations suggests that
 442 PAN photochemistry is in favor of radical formation and atmospheric oxidative capacity at this site (Liu et al., 2024). Unlike HO_2
 443 and OH , ΔRO_2 and ΔNO_2 are negative for most periods, accounting for 53.22 % and 67.23 %, respectively, because PAN formation
 444 uses up PA and NO_2 , the reduction in PA leads to a decrease in the amount of RO_2 . Over the entire period, ΔRO_2 is $-6.4 \cdot 10^{-4}$ ppb,
 445 with no significant difference between clean and hazy periods, being $-6.11 \cdot 10^{-4}$ ppb and $-6.55 \cdot 10^{-4}$ ppb respectively (Table S2). The
 446 average value of ΔNO_2 during the entire observation period is -0.17 ppb respectively, with significant differences between hazy and
 447 clean periods (Table S2). Specifically, ΔNO_2 is -0.22 during hazy periods and only -0.11 during clean periods, indicating that the
 448 PAN mechanism consumes more NO_2 during hazy periods. Although ΔNO is positive for most periods, accounting for 78.79 %, the
 449 overall mean is -0.01 , with significant differences between hazy and clean periods (Table S2). ΔNO is -0.05 during hazy periods,
 450 showing an inhibitory effect, while it is 0.03 during clean periods, showing a promoting effect.



451
 452 **Figure 9.** The time series of ΔHO_2 , ΔOH , ΔRO_2 , ΔNO_2 , and ΔNO . The ΔHO_2 , ΔOH , ΔRO_2 , ΔNO_2 , and ΔNO is calculated as the
 453 base scenario with the PAN mechanism minus the scenario without the PAN mechanism.

454
 455 As shown in Fig.10 (a), the PAN mechanism inhibited 85.80 % of net O_3 production during the entire observation period, with
 456 inhibition rates (the percentage of negative $\Delta\text{Net}(\text{O}_3)$) of 83.75 % and 87.50 % during clean and haze periods, respectively. This
 457 result is consistent with previous spring observations at the same site, where the inhibition rate was 83 % (Liu et al., 2022a). The

458 PAN mechanism mainly inhibits the net O₃ generation by increasing the RO₂+NO₂ reaction (Fig.10(a)), with negligible impact from
 459 other reactions (Fig. S14). As shown in Fig.10(b), the diurnal variation trend indicates that the PAN mechanism's inhibitory effect
 460 on O₃ is significantly greater during haze periods than during clean periods. Additionally, regardless of whether it is during haze
 461 periods or clean periods, the PAN mechanism's inhibitory effect on O₃ is significantly greater during the day than at night. These
 462 phenomena all indicate that the higher the PAN concentration, the more pronounced the inhibitory effect of the PAN mechanism on
 463 O₃ (Fig.10(c)). Under the condition of low precursors (including NO_x and VOCs) conditions, competition among these precursors
 464 may limit their secondary transformation, thus resulting in inhibition (Liu et al., 2024).



465
 466 **Figure 10.** (a) Time series plot of $\Delta\text{Net}(\text{O}_3)$ and the reaction of $\Delta(\text{RO}_2+\text{NO}_2)$, (b) Diurnal variation of $\Delta\text{Net}(\text{O}_3)$ during clean and
 467 hazy conditions, (c) Correlation between $\Delta\text{Net}(\text{O}_3)$ and PAN. $\Delta\text{Net}(\text{O}_3)$ and $\Delta(\text{RO}_2+\text{NO}_2)$ are calculated as the base scenario with
 468 the PAN mechanism minus the scenario without the PAN mechanism.

469
 470 **Conclusion**

471 This study thoroughly investigated the summertime PAN formation mechanism and established its connection to haze pollution. In
 472 addition to NO and TVOCs, the concentration of all pollutants during the haze period is above twice that during the clean period,
 473 indicating that the oxidation of NO and TVOCs during the haze period is stronger, which is conducive to the oxidation of NO and
 474 TVOCs into secondary pollutants, such as O₃ and PAN. The slopes of linear regression between the daily maximum values of PAN
 475 and O₃ were 0.021 ppb ppb⁻¹ and 0.009 ppb ppb⁻¹ for clean and hazy condition, respectively, implies that PAN precursors accounted
 476 for only a small fraction of the total VOCs, especially for hazy condition. High temperature should be another factor contributing
 477 to the lower production efficiency of PAN in the southeast coastal region. During the whole observation period, the IOA=0.75,
 478 indicating that the MCM model is well-suited for exploring the photochemical formation of PAN. During the clean period,
 479 simulation results were better than during the haze period (R^2 : 0.68 vs. 0.47, slope K: 0.91 vs. 0.75), indicating that some reactions
 480 related to PAN generation or destruction might be missing in the MCM during the hazy period. Additionally, the simulated net
 481 production rate of PAN becomes negative with PAN constrained. However, the observed increase in PAN concentrations indicates
 482 that the actual net production rate is positive, suggesting that there are additional sources contributing to PAN generation that are
 483 not considered in the MCM mechanism. Through XGBoost-SHAP machine learning, and given the significant positive correlation
 484 between PAN and NO₃⁻ ($R=0.37$) at the 0.01 level, and their peak around noon, they likely share a common source. Both RIR and

485 EKMA indicate that PAN formation in this region is VOC-controlled. Controlling emissions of VOCs, particularly alkenes, C₅H₈,
486 and aromatics, would be beneficial for mitigating PAN pollution. The RIR results also show that during the clean period, PAN is
487 more sensitive to changes in various pollutants than during the haze period, highlighting the significant importance of deep emission
488 reductions. PAN presented the promotion effects on OH and HO₂, while inhibited O₃ formation, RO₂, NO and NO₂. This study
489 improves our thorough understanding of PAN photochemistry and offers valuable scientific guidance for the future management of
490 PAN pollution.

491 **Data availability**

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

493 **Authorship Contribution Statement**

494 **Baoye Hu:** Methodology, Formal analysis, Investigation, Data curation, Writing – original draft. **Naihua Chen:** Software, Formal
495 analysis. **Rui Li:** Software, Formal analysis. **Mingqiang Huang:** Software. **Jinsheng Chen:** Funding acquisition, Supervision,
496 Writing - Review & Editing. **Youwei Hong:** Formal analysis. **Lingling Xu:** Investigation. Xiaolong Fan: Investigation. **Mengren**
497 **Li:** Investigation. **Lei Tong:** Investigation. **Qiuping Zheng:** Investigation. **Yuxiang Yang:** Writing - Review & Editing

498 **Competing interests**

499 The authors declare that they have no conflict of interest.

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506 (Institute of Urban Environment, Chinese Academy of Sciences).

507 **Supplementary information**

508 Attached please find supplementary information associated with this article.

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