Atmospheric Measurements at the Foot and the Summit of Mt. Tai - Part II: HONO Budget and Radical ($RO_x + NO_3$) Chemistry in the Lower Boundary Layer

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Abstract

In the summer of 2018, a comprehensive field campaign, with measurements on HONO and related parameters, was conducted

- 30 at the foot (150 m a.s.l.) and the summit of Mt. Tai (1534 m a.s.l.) in the central North China Plain (NCP). With the implementation of a 0-D box model, the HONO budget with six additional sources and its role in radical chemistry at the foot station were explored. We found that the model default source, NO + OH, could only reproduce 13% of the observed HONO by 13%, leading to a strong unknown source strength of up to 3 ppby h^{-1} . Among the additional sources, the NO₂ uptake on the ground surface dominated (~70%) night-time HONO formation, and its photo-enhanced reaction dominated (~80%)
- daytime HONO formation. Their contributions were sensitive to the mixing layer height (MLH) used for the parameterizations, 35 highlighting the importance of a reasonable MLH for exploring ground-level HONO formation in 0-D models and the necessity of gradient measurements. A Δ HONO/ Δ NO_x ratio of 0.7% for the direct emissions from vehicle exhaust was inferred and a new method to quantify its contribution to the observations was proposed and discussed. Aerosol-derived sources, including the NO₂ uptake on the aerosol surface and the particulate nitrate photolysis, did not lead to significant HONO formation, with
- 40 their contributions lower than NO + OH.

HONO photolysis in the early morning initialized the daytime photochemistry at both the foot and the summit stations, and also It was also a substantial radical source throughout the daytime, with contributions higher than or about one quarter of O_3 photolysis to OH initiation at the foot and the summit stations, respectively. Moreover, we found that OH dominated the atmospheric oxidizing capacity in the daytime, while modeled NO₃ appeared to be significant at night. Peaks of modeled NO₃

- time series and average diurnal variation reached 22 and 9 ppty, respectively. NO₃ induced reactions contribute 18% of nitrate 45 formation potential ($P(HNO_3)$) and 11% of the isoprene (C_5H_8) oxidation throughout the whole day. At night, NO₃ chemistry led to 51% or and 44% of P(HNO₃) or the C₅H₈ oxidation, respectively. NO₃-chemistry may significantly affect night-time secondary organic and inorganic aerosol formation in this high O₃ region, implying that NO₃ chemistry could significantly affect night-time secondary organic and inorganic aerosol formation in this high-O₃ region. Considering the severe O₃ pollution
- 50 in the NCP and the very limited NO₃ measurements, we suggest that besides direct measurements of HO_x and primary HO_x precursors (O_3 , HONO, alkenes, etc.), NO₃ measurements should be conducted to understand the atmospheric oxidizing capacity and air pollution formation in this and similar regions.

1 Introduction

Numerous field campaigns coupled with model simulations have been conducted worldwide to understand the summertime 55 atmospheric chemistry as it is linked to the regional air quality and global climate (Alicke et al., 2003; Elshorbany et al., 2012; Heard et al., 2004; Kanaya et al., 2009, 2013; Michoud et al., 2012; Ren et al., 2003; Rohrer and Berresheim, 2006; Tan et al., 2017; Travis et al., 2020). One of the key issues is the level and the production/loss paths of the atmospheric oxidizing capacity governed by radicals (OH, NO₃, etc.). This is also essential in converting primary to secondary pollutants and the removal of greenhouse gases (Lu et al., 2019; Seinfeld and Pandis, 2016).

On a global scale, OH controls atmospheric oxidation. As the detergent in the troposphere, OH can oxidize most trace gases, including inorganic (SO₂, NO₂, etc.) and organic compounds (VOCs, etc.), and determines the lifetime of greenhouse gases (e.g., CH₄). Besides the fast conversion of HO₂ to OH (R-1) as part of the radical propagation cycle, primary OH (radical initiation) mainly originates from photolysis reactions, including O₃ ((R-2) to (R-4)), HONO (R-5), HCHO ((R-6) to (R-9), and (R-1)), and H₂O₂ (R-10), and the ozonolysis of alkenes (not shown in detail here). In particular, HONO photolysis is reported to be an important or even the major OH source in the lower atmosphere of polluted regions, with a contribution of 20 – 90% (Alicke et al., 2003; Elshorbany et al., 2009; Kleffmann et al., 2005; Platt et al., 1980; Slater et al., 2020; Whalley et al., 2021; Xue et al., 2020). However, this process still needs more global quantifications due to the incomplete understanding of HONO formation and its vertical distribution in the atmosphere (Kleffmann, 2007). A state-of-art summary of the reported HONO sources can be found in our recent study (Xue et al., 2020).

7	Λ
1	v

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 $HO_2 + NO \rightarrow OH + NO_2 \tag{R-1}$

$\boldsymbol{0}_3 + \boldsymbol{h} \boldsymbol{v} \rightarrow \boldsymbol{O}(^1 \boldsymbol{D}) + \boldsymbol{O}_2$	(λ<320 nm)	(R-2)
$O(^1D) + H_2O \rightarrow 2OH$		(R-3)
$O(^1D) + M \rightarrow O + M$	(M=N ₂ or O ₂)	(R-4)
$HONO + hv \rightarrow NO + OH$	(λ<400 nm)	(R-5)
$HCHO + hv \rightarrow H + HCO$	(λ<340 nm)	(R-6)
$HCHO + hv \rightarrow H_2 + CO$	(λ<365 nm)	(R-7)
$H + O_2 \rightarrow HO_2$		(R-8)
$HCO + O_2 \rightarrow CO + HO_2$		(R-9)

$$H_2O_2 + h\nu \rightarrow 20H$$
 (λ <362 nm) (R-10)

Besides, other oxidants can also be of importance on a regional scale. For example, <u>the_NO3</u> radical could be a major oxidant in forests (vegetation shadows slow down its photolysis) or in the nocturnal boundary layer at high O₃ regions (Brown and Stutz, 2012). Formed by the reaction of NO₂ + O₃ (R-11), high NO₃ levels usually occur at night, concerning its very rapid photolysis during the daytime (R-12) and (R-13). Moreover, high NO₃ concentrations are only observed for high O₃ and medium NO_x concentrations in the absence of significant levels of NO caused by reaction (R-15). Like OH, NO₃ also has high reactivity with various trace gases (Brown and Stutz, 2012; Mellouki et al., 2021). For example, NO₃ reacts with NO₂ to form N₂O₅ (R-14), which can undergo hydrolysis on wet surfaces or clouds to produce HNO₃ (or NO₃⁻) (R-16) or decomposition back to NO₃ + NO₂ (R-17). NO₃ can also react with various organic compounds to form secondary organic aerosol (SOA). For instance, NO₃ reacts with isoprene (C₅H₈), leading to significant organic nitrates (e.g., alkyl nitrates) production (Rollins et al., 2009).

 $NO_2 + O_3 \rightarrow NO_3 + O_2 \tag{R-11}$

$$NO_3 + hv \rightarrow NO + O_2$$
 ($\lambda < 690 \text{ nm}$) (R-12)

$$NO_3 + hv \rightarrow NO_2 + 0 \qquad (\lambda < 690 \text{ nm}) \tag{R-13}$$

$$NO_2 + NO_3 \rightarrow N_2O_5 \tag{R-14}$$

$$NO + NO_3 \rightarrow 2NO_2$$
 (R-15)

$$N_2 O_5 + H_2 O \xrightarrow{\text{surface}} 2HNO_3 (or 2NO_3^{-})$$
(R-16)

$$N_2 O_5 \rightarrow N O_2 + N O_3 \tag{R-17}$$

In the past decade, particle pollution, such as PM_{2.5}, is going down while O₃ pollution is increasing in many cities of China (Han et al., 2020; Li et al., 2019; Sun et al., 2016, 2019)., especially-Especially in the North China Plain (NCP), where there exists a large population (>330 million) and air pollution in this region becomes a major environmental risk for the public health of >330 million people living in this region. This raises efforts in exploring the NO_x-VOCs-O₃ chemistry. Meanwhile,

- 100 health <u>of >330 million people living in this region</u>. This raises efforts in exploring the NO_x-VOCs-O₃ chemistry. Meanwhile, high O₃ indicates an enhanced atmospheric oxidizing capacity; that is, elevated OH and NO₃ levels are expected. However, <u>very few observations have been reported of OH and NO₃ levels</u>, as well as their production (e.g., HONO photolysis or NO₂ + O₃) or loss (e.g., to oxidize primary pollutants), <u>have been reported</u> in the high-O₃ region of the NCP, by far, are very few <u>reported</u> (Lu et al., 2019; Suhail et al., 2019). Herein, we provided the first HONO measurements at the foot of Mt. Tai (in
- 105 Tai'an city, a typical urban site), followed by measurements at the summit of Mt. Tai in the summer of 2018. Data from the summit station <u>was is</u> presented in the companion paper, in which daytime HONO formation and its role in the atmospheric oxidizing capacity at the summit level <u>were are</u> studied. In this paper, coupled with the box model, the HONO budget and the radical chemistry at the ground level <u>were are</u> explored and discussed.

2 Methods

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110 2.1 Field Campaign

2.1.1 Measurement Sites

In the summer (from late May to July) of 2018, a comprehensive field campaign was conducted to understand the atmospheric oxidizing capacity and O₃ pollution in Tai'an, a city in the middle of the NCP. <u>Tai'an is located nearly in the middle between</u> <u>Beijing and Shanghai. <u>Tai'an</u>The city has a population of about 5.6 million and is about 60 km south of Jinan city (the capital</u>

- 115 <u>city of Shandong province, population: ~8.7 million).</u> Measurements were conducted both at the ground level (the foot of Mt. Tai, 150 m a.s.l.) and the summit level (the summit of Mt. Tai, 1534 m a.s.l., 36.23°N, 117.11°E). The foot station was inside Shandong College of Electric Power (36.18°N, 117.11°E), in the Taishan district of Tai'an city. There are no industrial activities around this site, which is surrounded by the campus, residential area, and a business district. The 801st province road is in the northeast of this typical urban site. which represents a typical urban site. Inside the campus (about 50 ha) frequent traffic was not observed, but it sometimes occurred on the urban roads nearby. Tai'an city has a population of about 5.6 million
 - and is about 60 km south of Jinan city (the capital city of Shandong province, population: -8.7 million).-Mt. Tai is locateds in

the north part of Tai'an city. Locations of these two stations on the map <u>could_can</u> be found in the companion ACP paper (entitled "Atmospheric Measurements at the Foot and the Summit of Mt. Tai - Part I: HONO Formation and Its Role in the Oxidizing Capacity of the Upper Boundary Layer". Xue et al. (2021a).

125 Measurements at these two stations allow us to study HONO formation and its role in the atmospheric oxidizing capacity of the lower (the foot study) and the upper boundary layer (the summit study). Briefly, in the summit study, we found remarkably high daytime HONO levels as well as high unknown HONO source strength, which was mainly caused by rapid vertical transport from the ground to the summit levels driven by mountain winds.

2.1.2 Instrumentation

- HONO mixing ratios were continuously measured by the LOPAP technique (LOPAP-03, QUMA GmbH, Germany) (Heland et al., 2001; Kleffmann et al., 2006) from 29th May to 8th July 2017-2018 at the foot station, followed by measurements at the summit station from 9th to 31st July 2017. At the foot station, NO-NO₂-NO_x, O₃, CO, and SO₂ were online measured by a series of Thermo Fisher Scientific instruments (42i, 49i, 48i, and 43i, respectively). Because chemiluminescence techniques using molybdenum converters were reported to overestimate the NO₂ level caused by other NO_y-interferences of other NO_y species,
- 135 we furtherly corrected the measured NO₂ was corrected with a family constraint in a model run (see Section 3.1.2). VOCs (56 species) and OVOCs (15 species) were measured by a homemade GC-FID instrument (Liu et al., 2016) and the USEPA DNPH-HPLC method (Wang et al., 2020), respectively. Gas-phase H₂O₂ was measured by a monitor based on the wet chemical method (AL2021, Aerolaser GmbH, Germany), and details about of which the used instrument can be found in Ye et al. (2018). Water-soluble inorganic particle ions-species (i.e., NO₃⁻, SO₄²⁻, Cl⁻, Na⁺, K⁺, Ca²⁺, etc.) of PM_{2.5} were collected on Teflon
- 140 filters every two hours at a sampling flow of 100 L min⁻¹ and analyzed by an ion chromatograph (Liu et al., 2020). Meteorological parameters, including atmospheric temperature (T), pressure (p), relative humidity (RH), wind direction (WD), wind speed (WS), and solar irradiance (Ra) were continuously measured by an auto meteorological station. J(NO₂) was measured by a 4-π filter radiometer (Metcon GmbH, Germany). 10-min and hourly-average data (except for PM_{2.5}) were used for the following-analysis (of time series and statistic descriptions of the data)-and-In contrast, hourly data were used for
- 145 model simulations, respectively. $PM_{2.5}$ measurements werewas obtained from the Tai'an monitoring station (200 m east of the foot station), and only hourly-average data was-were available. Other J-values used in this study, including J(HONO), J(O(¹D)), J(H₂O₂), J(HCHO)_{rad}, and J(HNO₃), are calculated by the box model based on trigonometric SZA functions (MCM default photolysis frequency calculation, see Jenkin et al. (1997)) and scaled by the measured J(NO₂). For instance, J(HONO) = J(HONO)_{model}*J(NO₂)_{measured}/J(NO₂)_{model}.

150 2.2 Model Description

2.2.1 Box Model and Constraints

The Framework for **0**-D Atmospheric Modeling, F0AM v4.0 (available at <u>https://github.com/AirChem/F0AM</u>) developed by Wolfe et al. (2016) was used to explore the HONO budget and the radical chemistry. The <u>used_MCM v3.3.1</u> chemical mechanism was <u>MCM v3.3.1</u>, which could be obtained from <u>http://mcm.leeds.ac.uk/MCMv3.3.1/home.htt</u>. Note that the

155 present F0AM model could also be run with family constraints (see details in Wolfe et al. (2016)), such as the NO_y and family, Cl_y familiesy, etc., and hHence, it allows us to correct for interferences of in the NO₂ measurements by the chemiluminescence method (see Section 3.1.2).

The model was constrained by the measured $J(NO_2)$, T, RH, P, VOCs, OVOCs, and all the other measured inorganic species, including the corrected NO₂ by the family constraint. Continuous VOCs measurements was were available from 12th June to

160 6^{th} July, and hence box model simulations were performed during this period. While J(NO₂) measurement was available from 16th June, J(NO₂) from 12th to 16th June was estimated through the high quadratic correlation (R² = 0.96, Figure S1) between J(NO₂) and solar irradiance.

2.2.2 Model Scenarios

Table 1 shows the description of different model scenarios. A base case (Sce-0) with all the measured parameters as constraints
was run to simulate radicals² concentrations and their production/loss rates. The family constraint was used in this scenario to correct for interferences of NO₂ measurements (Section 3.1.2). Meanwhile, the role of HONO in radical chemistry was also explored by several model sensitivity tests with reducing or increasing the constrained HONO.

With the simulated OH and the corrected NO_2 from the base case, we could further explore the HONO budget. Three model scenarios were conducted to assess the potential contributions of different HONO sources, including one with only the default

- 170 model source (Sce-1), and one with all the six additional sources, including direct emission, the dark and the photo-enhanced NO₂ uptake on the aerosol and ground surfaces and nitrate photolysis (Sce-2). In Sce-3, photo-enhanced NO₂ uptake on the ground surface was reduced by a factor of 10, aerosol-derived sources (NO₂ uptake on the aerosol surface or particulate nitrate photolysis) were significantly enhanced (see Section 3.2.3) to test whether the aerosol-derived sources could well explain the observations.
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Table 1: Description of different model scenarios.

Scenarios	Constraints	Objectives
Sce-0	All measurements; NOz family	NO ₂ correction; radical concentration
Sce-0	constraint	and chemistry
Sce-1	All measurements + corrected NO ₂	HONO simulation with NO+OH
Sce-1	and simulated radicals from Sce-0	HONO SIIIUIAUOII WIUI NO+OH

Sce-2

Same as Sce-1

HONO simulation with additional sources

Sce-3

Same as Sce-1, but with reduced ground NO₂ uptake and enhanced aerosol-derived sources

Testing the performance of aerosolderived sources

3. Results and Discussion

3.1 Overview of the Measurements and Potential Interferences on the Measurements

3.1.1 Overview of the Measurements

Figure 1 shows the meteorological parameters measured at the foot station. During the campaign, it was generally sunny except
slightly rainy (<10 mm) on 9th, 10th, 13th, and 28th and heavy rainy (~100 mm) at night of 25th/26th June. Ambient temperature was normally around 25 °C at night and around 30 °C during the daytime, except for rainy days when the temperature was relatively low. The relative atmospheric humidity (RH) was high (up to 80%) on those rainy or cloudy days and low (around 40%) on other days. Campaign-averaged temperature and RH were 27.5 °C and 46.6%, respectively (Table 2). Air masses observed at this site was originated from multiple directions, including west, south, and east, which can be obtained from are shown in the wind rose plot in (Figure S2A). Wind speed was generally low, with an average of about 2 m s⁻¹. In addition, the wind rose results generally agree well with HYSPLIT trajectory results in Figures S2B and S2C.

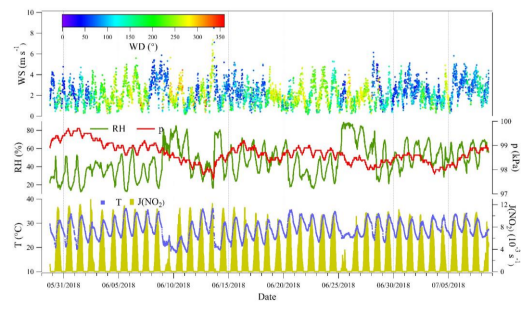


Figure 1: Meteorological parameters measured at the foot of Mt. Tai during the campaign.

Table 2: Statistic summary of meteorological parameters and the measured species. Note that the observation data point number(190)(Obs) of hourly PM2.5 is about 1/6 of others (10 min time resolution), and. The measured rather than the corrected NO2 was used
here. SD: standard deviation; Min: minimum; Max: maximum.

Parameters	Mean	SD	Median	Min	Max	Obs
WS (m s ⁻¹)	2.2	1.1	2.1	0.2	9.7	5749
RH (%)	46.6	17.5	44.9	12.2	88.7	5749
P (kPa)	98.7	0.4	98.6	97.6	99.7	5749
T (°C)	27.5	3.8	27.4	17.9	36.1	5749
$J(NO_2) (s^{-1})$	3.2E-03	3.7E-03	1.0E-03	0	1.1E-02	3183
O ₃ (ppbv)	63	31	62	0.1	145	5727
PM _{2.5} (µg m ⁻³)	29	12	28	10	66	959
CO (ppmv)	0.28	0.25	0.20	0.01	2.08	5717
SO ₂ (ppbv)	3.6	4.0	2.2	0	36.2	5648
NO (ppbv)	2.0	8.3	0.3	0	126.0	5749
NO ₂ (ppbv)	15.2	10.8	12.3	0	78.8	5601
HONO (ppbv)	0.62	0.42	0.52	0.05	2.97	5423

In Figure 2 and Figure S3 the measured HONO and related species at the foot station and their diurnal variation profiles are presented, respectively. The measured HONO showed a typical diurnal variation with accumulation after sunset and decay after sunrise. Mixing ratios of the measured HONO varied from 0.05 to about 3 ppbv, with an average of 0.62 ± 0.42 ppbv.

- 195 The measured NO₂ showed a very similar variation to HONO, and their correlation was high (R = 0.73), indicating a potential role of NO₂ in HONO formation. Severe O₃ pollution (maximum: 145 ppbv) was observed at this site, with O₃ levels frequently exceeding the Class-1 limit value (1-h 160 µg m⁻³, equivalent to 82 ppbv at 298 K and 101 kPa) of the National Ambient Air Quality Standard of China (GB3095-2012), while the NO_x level was typically lower than O₃. Consequently, a relatively low NO was frequently found, whose concentration was generally lower than 1 ppbv, except for some fresh plumes with higher
- NO concentrations inside. The two primary pollutants, CO and SO₂, were generally lower than 0.5 ppmv and 5 ppbv, respectively, except for several polluted events, within which CO and SO₂ reached around 2 ppmv and around 35 ppbv, respectively. However, all the primary pollutants, including NO, CO, and SO₂, showed poor correlations with HONO (R = 0.49, 0.44, and 0.13, respectively), implying the minor role of direct emission in HONO formation. The measured hourly PM_{2.5} varied from 10 to 66 µg m⁻³, with an average of 29 µg m⁻³. The correlation of PM_{2.5} and HONO was very low (R = 0.06), suggesting a minor role of aerosol-derived sources in HONO formation. More discussion on the HONO budget is presented in Section 3.2.

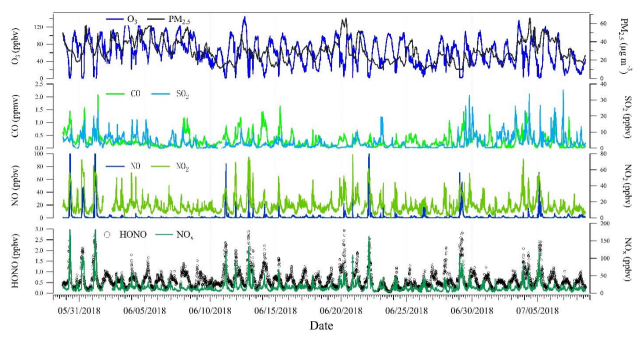


Figure 2: HONO and related species measured during the campaign.

- Compared to other previous summertime measurements worldwide (Table 3), the measured HONO level at this site is similar
 to some measurements in China, such as Beijing in 2007 (Hendrick et al., 2014), Beijing in 2008 and 2017 (Crilley et al., 2019; Hendrick et al., 2014) and Guangzhou in 2006 (Yang et al., 2014); in Europe, such as Milan in 1998 (Alicke et al., 2002) and Rome in 2001 (Acker et al., 2006); and in North America, such as New York in 2001 (Ren et al., 2003) and Colorado in 2011 (Vandenboer et al., 2013). Besides, it is lower than measurements in cities during polluted periods, such as Jinan in 2016 (Li et al., 2018), Santiago de Chile in 2005 (Elshorbany et al., 2009), Santiago de Chile in 2009 (Villena et al., 2011), and Mexico
 in 2003 (Volkamer et al., 2010), but higher than recent measurements near European cities, including Forschungszentrum Karlsruhe (Kleffmann et al., 2003), Forschungszentrum Jülich (Elshorbany et al., 2012), suburban Paris (Michoud et al., 2014), and Cyprus (Meusel et al., 2016). It is noteworthy that the measured HONO at the foot station is significantly higher than that observed at the summit station in the same summer, indicating possibly different roles and formation paths of HONO at these two stations.
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Table 3: Examples of v	worldwide HONO	measurements at the	e ground level.
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Location	Period	Techniques	Mean (pptv)	Range (pptv)	HONO/NO _x %	Reference
Europe						
Milan	May-Jun 1998	DOAS	920ª/140b	<4400		(Alicke et al., 2002)
Pabstthum	Jul-Aug 1998	DOAS	$330^{a}/70^{b}$	<1200	$1.4^{a}/1.0^{b}$	(Alicke et al., 2003)
Rome	May-Jun 2001	LP-DOAS, DNPH- HPLC, WEDD	580	<2000	3*	(Acker et al., 2006)
Forschungszentrum Karlsruhe	Oct 2001	LOPAP	400	180-1100	1-6 ^a	(Kleffmann et al., 2003)
Forschungszentrum Jülich	Jun-Jul 2005	LOPAP	220	50-1100	2 (0.6-12)	(Elshorbany et al., 2012)

Suburban Paris	Jul 2009	Ni-troMAC	~150	10-500		(Michoud et al., 2014)
Cyprus	Jul-Aug 2014	LOPAP	35	<300	33	(Meusel et al., 2016)
South America						
Santiago de Chile	Mar 2005	LOPAP	2500ª/2300b	670-7100	3.9 (1.3-9.2)	(Elshorbany et al., 2009)
Santiago de Chile	Nov 2009	LOPAP	1500/1100**	220-3800/150- 4600**	2.0 (0.7-5.9)	(Villena et al., 2011)
North America						
California	Aug-Sep 1979	DOAS	$1090^{a^*}/\!\!<\!\!280^{b}$	<4100		(Platt et al., 1980)
New York	Jul-Aug 2001	HPLC	660	400-1400		(Ren et al., 2003)
Colorado	Feb-Mar 2011	NI-PT-CIMS	$500^{a^*}/100^{b}$	<2000	3.5-7.6 ^a	(Vandenboer et al., 2013)
Mexico	Mar-May 2003	LP-DOAS	1200^{*}	<3000		(Volkamer et al., 2010)
China						
Beijing	Aug-Sep 2004	DOAS		<6100	8.4 ^c	(Qin et al., 2006)
Guangzhou	Jun 2006	LOPAP	950 ^a /240 ^b	10-5000	4.3ª/4.5 ^b	(Li et al., 2012)
Yufa	Jul-Aug 2006	LOPAP	890 ^a /430 ^b	30-3600	$4.6^{a}/4.8^{b}$	(Yang et al., 2014)
Beijing	Aug 2007	DOAS	1450	440-2900	5°	(Spataro et al., 2013)
Beijing	Jul 2008	DOAS	180 ^d	100-800	0.8 ^{cd}	(Hendrick et al., 2014)
Xianghe	Jun 2012	DOAS	90 ^d	100-700	1.7 ^{cd}	(Hendrick et al., 2014)
Jinan	Jun-Aug 2016	LOPAP	$1200^{a}/1010^{b}$	<6000	6ª/5 ^b	(Li et al., 2018)
<u>Wangdu</u>	<u>Aug 2016</u>	<u>SCIC</u>	<u>230</u>	<u>30-1140</u>	<u>2.1^b</u>	(Xue et al., 2021c)
<u>Wangdu</u>	Jun 2017	LOPAP	360e/720f/1360g	<u>260-860^e/320-</u> <u>1490^f/400-3130^g</u>	$2.6^{e}/5.4^{f}/11.8^{g}$	(Xue et al., 2021c)
<u>Beijing</u>	<u>May-Jun 2017</u>	Intercomp.h		<u>~100-10000</u>		(Crilley et al., 2019)
Tai'an	May-Jul 2018	LOPAP	620	50-2970	4.2	This study
Mt. Tai Summit	Jul 2018	LOPAP	133	1880	6.4	This study

^a: night-time, ^b: daytime, ^c: HONO/NO₂, ^d: noontime, ^e: non-fertilization period, ^f: pre-fertilization period, ^g:

intensive fertilization period, Intercomph: intercomparison of 5 instruments.

*: half of the diurnal maximum.

**: 3rd and 21st floors, respectively.

225 3.1.2 NO₂ Interference and Correction

As the most important HONO precursor, accurate measurement of NO_2 plays a key role in analyzing HONO formation. The NO_x monitor used in this study could specifically detect NO. To measure NO_2 , a molybdenum converter is used to convert NO_2 to NO. NO_2 measured by the chemiluminescence methodHowever, this chemical conversion process suffers from the interference of other NO_y species (Villena et al., 2012), primarily including inorganic species such as (measured) HONO, (non-

230 measured) HNO₃, HNO₄, N₂O₅, and NO₃, peroxyacyl nitrates (PANs, RC(O)OONO₂), organic nitrates (RONO₂), and peroxy nitrates (ROONO₂), etc. The sum of the latter two was<u>In this study, we</u> defined the sum of RONO₂ and ROONO₂ here as organic nitrates^{*}. Hence, the measured NO₂ is the sum of real NO₂ and those interferingence species. HONO was measured and subtracted from the measured NO₂, and we defined NO₂^{*} = the measured NO₂ – HONO. As NO₂ is the most important HONO precursor, we used the family constraint (NO₂^{*} = NO₂ + HNO₄ + 2N₂O₅ + NO₃ + PANs + organic nitrates^{*}) in the base case (Sce-0) to separate each species from NO₂^{*}. In the term of To describe the PANs class, PAN, PPN, and MPAN (MCM names, see their structures at http://mcm.leeds.ac.uk/MCMv3.3.1/) were considered. In the elass of organic nitrates class, CH3NO3, C2H5NO3, NC3H7NO3, IC3H7NO3, TC4H9NO3, NOA, ISOP34NO3, ISOPANO3, ISOPDNO3, ISOPCNO3, and ISOPBNO3 (MCM names) were considered included. Considering that HNO₃ is very sticky, we expect HNO₃ was mostly absorbed by the filter and/or sampling tubes before the converter rather than being converted to NO by the converter. Therefore, HNO₃ was generally not included in the family constraint and only considered for the uncertainty analysis.

Figure 3 shows the model results of the relative contribution of each NO_2^* species to NO_2^* . At night with the absence of photochemistry, the real NO₂ dominated NO₂^{*} components, with a contribution of >95%, suggesting a small interference on the NO₂ measurement. However, the contribution of real NO₂ was found to decrease during the daytime due to the increasing interference. For example, at 11:00, the real NO₂ contributed 82% of the $NO_2NO_2^*$, which means indicating that the interferences could be as high as +22% (calculated from 18%/82%). In particular, at 1113:00, PANs alone caused the most 245

highest interference by +21% (calculated from 17%/81%).

The variations of the simulated PANs and NO₃ and their ratios to NO₂ were similar to previous observations (Brown and Stutz, 2012; Roberts et al., 1998; Su et al., 2008; Villena et al., 2012; Xue et al., 2011), indicating that the uncertainty of the method is small. For the following model simulations and analysis, only the corrected NO₂ was used. Besides, Figure S43 exhibits the

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parallel test results, in which HNO₃ was included in the family constraint. It can be found that the interference became more significant; for instance, the interference could be as high as +75% (calculated from 43%/57%, at 11:00). This represents the upper limits of the interferences, if the sampling tubes and the inlet filter are heated so that HNO₃ could have reached the converter.

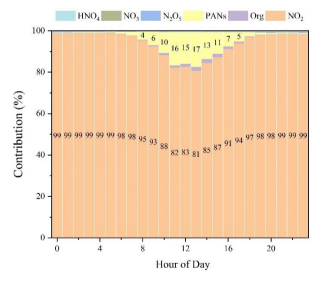


Figure 3: Relative contribution of each NO₂* species. PANs = PAN + PPN + MPAN, and. Org represents organic nitrates* (RONO₂ 255 + ROONO₂).

Additionally, as shown in Figure S5, the simulated HNO₄ showed 1) a different diurnal variation from, 2) generally 1 - 2orders of magnitude lower concentrations than, and 3) a very poor correlation ($R^2 = 0.06$) with the observed HONO, indicating its negligible interference on the HONO measurement by the LOPAP technique (Legrand et al., 2014). It is worth noting that

260 for the description of O_3 formation in the polluted atmosphere, accurate measurements of $\frac{\text{VOCs an}}{\text{NOx}}$ and $\frac{\text{HONO}}{\text{are}}$ are necessary.

3.2 HONO Sources and Budget

3.2.1 Model Default Source (NO + OH) and Unknown Source Strength

The homogeneous reaction of NO and OH has been adopted as the default HONO source in atmospheric chemistry models, including MCM. Model results from Sce-1 that only contains the homogeneous source with the modeled OH from Sce-0 are shown in Figure 4. Apparently, the source of NO + OH is too small to explain the observed HONO as the simulated one is almost one order of magnitude lower than observations. Its contributions to the measured daytime <u>or-and</u> night-time HONO are 15% and 12%, respectively.

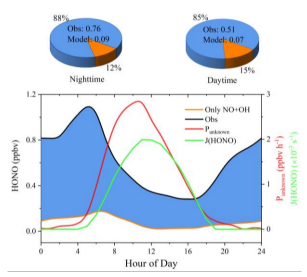


Figure 4: Simulated HONO by the default mechanism (Sce-1, left axis) compared with the observations (Obs, left axis), unknown source strength (P_{unknown}, right axis), <u>HONO photolysis frequency (J(HONO), right axis), and the The</u> relative contributions of NO + OH to the observations at night (19:00 - 4:50, left pie chart) and day (5:00 - 18:50, right pie chart) were shown in left and the right pie charts, respectively. The shaded area in blue represents the difference between the observation and modeled values.

Then we calculated the unknown source strength (Punknown) based on the following equation (Sörgel et al., 2011; Su et al., 2008).

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$$P_{unknown} = \frac{\Delta HONO}{\Delta t} + L(HONO)_{pho} + L(HONO)_{HONO+OH} - P(HONO)_{NO+OH}$$
(Eq-1)

where t<u>T</u>he HONO loss rates through photolysis (L(HONO)_{pho}) and reaction with OH (L(HONO)_{HONO+OH}) and production rate from NO + OH were obtained from the base model scenario (Sce-0 with a constraint of the measured HONO). HONO mixing ratio differences within a one-hour interval, $\frac{\Delta HONO}{\Delta t}$, was were calculated by the measurement, and its comparison are compared with P_{unknown} was shown in Figure S<u>6</u>5. A typical unknown HONO strength variation, with high values at noontime, was obtained (Figure 4). As shown in Figure 4, P_{unknown} rapidly increased in the morning and peaked <u>at</u> nearly 3 ppbv h⁻¹ at 11:00, followed by a decrease, revealing a photo-enhanced source. Note that the profile of $P_{unknown}$ was asymmetric around 11:00, indicating the unknown source is not simply photolytic but also includes its precursors that also have an asymmetric distribution (e.g., NO₂, Figure S3). The possible additional HONO sources that are responsible for $P_{unknown}$ are discussed in the following section.

285 3.2.2 Additional Sources vs. Punknown

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3.2.2.1 Direct Emission: AHONO/ANO_x Ratio

The Δ HONO/ Δ NO_x ratio for the direct emission was determined from fresh plumes, which reached the following requirements: 1) at night when photolysis was absent, 2) rapid NO increase by >10 ppbv within 10 min. Only 17 cases were obtained throughout the campaign due to the persistent high O₃ and the fast NO-to-NO₂ conversion₂₇ If NO is completely titrated by O₃, there is a risk that the plume may not be fresh and that for which the inferred Δ HONO/ Δ NO_x might be overestimated due to heterogeneous secondary HONO sources. In Table 4 the obtained Δ HONO/ Δ NO_x was shown, varying from 0.18% to 1.86%,

- with an average of 0.98% and a median of 0.90%. The inferred value might be larger than the real one as <u>potential</u> NO₂-to-HONO conversion leads to an <u>positive interferenceoverestimation</u>, which This is consistent with the observation that the inferred Δ HONO/ Δ NO_x is generally higher in high RH conditions (in favor of due to the potential NO₂-to-HONO conversion)
- (Figure 5). <u>AlsoMore importantly</u>, we found that the observed HONO/NO_x is convergent as NO/NO₂ increases (Figure 5), which allows a further correction on <u>of</u> ΔHONO/ΔNO_x. The reported NO/NO₂ ratios from <u>the</u> combustion process<u>es</u> vary from <u>digits to hundredsshow significant variations</u>, e.g., 6.7 in Wuppertal (Kurtenbach et al., 2012), ~18 in Denver (Wild et al., 2017), 5 30 in London (Carslaw and Beevers, 2005), and 13 43 from China IV/V vehicles (He et al., 2020). Furthermore, in the emission inventory, the NO/NO₂ emission ratio in the NCP is about 9 (Zhang et al., 2009). However, the measured
- 300 night-time NO/NO₂ ratios were less than 3 (Figure 5), much lower than that from on road<u>compared to direct emission</u> measurements, indicating that the obtained observed plumes were not fresh enough to directly obtain the primary $\Delta HONO/\Delta NO_x$ emission ratio. By using a typical NO/NO₂ ratio of 10 from car exhaust, the calculated $\Delta HONO/\Delta NO_x$ through the convergent function (Figure 5) is 0.7%, similar to that obtained from laboratory or tunnel experiments (Kirchstetter et al., 1996; Kramer et al., 2020; Kurtenbach et al., 2001; Liu et al., 2017a).
- 305 Considering that HONO from Caused by its fast daytime photolysis, direct HONO emissions (HONO_{emi}) is are likely significantly overestimated with when a constant Δ HONO/ Δ NO_x ratio is considered, as done in former studies (Kramer et al., 2020; Liu et al., 2017b). because of different lifetimes Due to the much shorter lifetime of HONO (τ (HONO)) and compared to NO_x (τ (NO_x)), a much smaller fraction of the emitted HONO compared to NO_x will survive in the daytime atmosphere. in the daytime (also see Section 3.1.1 where very poor correlations of HONO with primary pollutants were presented and the
- 310 minor role of direct emission in HONO formation was inferred). Then Accordingly, for the first time, we calculated τ (HONO) and τ (NO_x) and considered them in the parameterization of HONO_{emi} (see the method in Section 1 of the Supporting Information). As shown in Figure S76A, daytime τ (NO_x) was typically one order of magnitude longer-higher than τ (HONO)

(Figure S₂₆A), indicating the <u>a</u> remarkable overestimation of the contribution of HONO_{emi} to the measured HONO when using a constant Δ HONO/ Δ NO_x emission ratio (Figure S₂₆B). Hence, HONO_{emi} was quantified by the following equations:

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$$HONO_{emi} = 0.7\% \times [NO_x]$$
 (night-time) (Eq-2)

$$HONO_{emi} = 0.7\% \times [NO_x] \times \frac{\tau(HONO)}{\tau(NO_x)} \text{ (daytime)}$$
(Eq-3)

In summary, direct emission contributed about 1 - 26% of the measured HONO, with an average of 13%. Moreover, the new method developed here may <u>still</u> have <u>some</u> uncertainties but largely <u>reduced</u> the significant overestimation of <u>the</u> <u>contribution of</u> HONO_{emi} to the observations <u>in theduring</u> daytime compared to using <u>only</u> a constant Δ HONO/ Δ NO_x (Figure S<u>7</u>6B).

Table 4: △HONO/△NO_x ratios determined from night-time (19:00 – 4:50) data of the campaign. <u>Concentrations (ppbv) of</u> <u>HONO/NO/NO_x at the start and the end of each plume of fresh emissions are also shown.</u>

Date	Period HONO		NO	NO		NO _x		$\Delta HONO/\Delta NO_x$
1 st June	2:20-2:30	1.58	1.77	3.1	13.5	64.4	75.8	1.67%
	4:20-4:30	2.67	2.74	0.6	38.2	36.0	74.2	0.18%
11 th June	3:00-3:10	1.32	1.71	0.2	18.9	24.9	50.0	1.55%
	3:50-4:20	1.68	2.21	15.7	71.7	51.3	107.0	0.95%
	4:30-4:40	1.91	2.31	41.2	72.4	75.3	107.3	1.25%
	22:00-22:10	1.41	1.69	1.0	16.6	41.1	78.4	0.75%
	23:30-23:40	1.74	1.99	1.7	17.0	53.2	71.1	1.40%
12 th June	22:10-22:30	2.64	2.78	0.8	59.0	68.7	132.9	0.22%
14 th June	3:00-3:10	1.43	1.76	6.6	21.2	38.8	56.5	1.86%
20th June	20:50-21:10	0.94	1.65	0.9	29.7	30.0	108.5	0.90%
22 nd June	2:10-2:20	1.29	1.98	4.6	95.1	51.8	147.2	0.72%
	2:40-3:00	1.58	2.20	38.3	120.1	83.1	163.7	0.77%
29th June	1:00-1:20	0.96	2.26	3.1	70.5	24.3	109.4	1.53%
	2:20-2:40	0.34	0.40	7.0	21.1	45.6	59.6	0.43%
	4:20-4:30	1.79	1.97	11.7	42.2	44.0	74.0	0.60%
5 th July	1:10-1:30	0.77	1.38	0.1	18.2	33.8	70.6	1.66%
	2:10-2:30	1.09	1.26	6.4	71.0	56.2	124.6	0.25%
Mean								0.98%

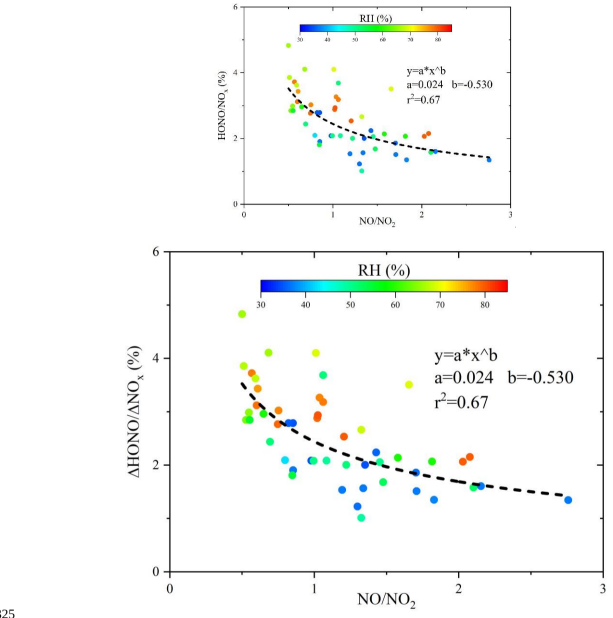




Figure 5: The inferred direct emission ratio (AHONO/ANOx) and NO/NO2 colored by RH. Only data with NO/NO2 >0.5 was shown as lower ones indicate much aged rather than fresh plumes.

3.2.2.2 NO₂ Uptake on the Aerosol Surface

Parameterizations of HONO formation from the NO₂ uptake on the aerosol surface without (P(HONO)_{a dark}) and with 330 (P(HONO)_a) photo-enhanced effects (P(HONO)_{a dark} and P(HONO)_a) are described by (Eq-4) and (Eq-5), respectively. In (Eq-4) and (Eq-5), HONO yields of 50% and 100% were considered for the dark and the photo-enhanced NO₂ conversion, respectively (Finlayson-Pitts et al., 2003; George et al., 2005). A relatively large NO₂ uptake coefficient γ_{a_dark} of 1×10^{-5} was used here to represent its upper limit. Its overestimation should not cause significant uncertainties as P(HONO)_{a_dark} was negligible to HONO formation (see the following discussion). For the photo-enhanced_NO₂ uptake coefficient γ_a values of 1.3×10^{-4} (overestimated oneupper limit derived from the summit measurement, see (Xue et al., (2021b)) and 2×10^{-5} (popularly used_more realistic one-value derived from laboratory experiments, see Stemmler et al. (2006 and 2007)) were used in (Eq-5) to constrain the upper limit and general onea more realistic value of P(HONO)_a.

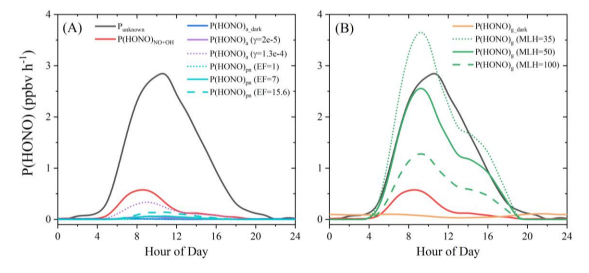
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$$P(HONO)_{a_dark} = \frac{v(NO_2) \times S_a \times [NO_2]}{8} \times \gamma_{a_dark},$$
(Eq-4)

$$P(HONO)_{a} = \frac{\nu(NO_{2}) \times S_{a} \times [NO_{2}]}{4} \times [\gamma_{a} \times \frac{J(NO_{2})_{measured}}{0.005 \, s^{-1}}],$$
(Eq-5)

340 where-v(NO₂), S_a, [NO₂], and J(NO₂)_{measured} denote the average NO₂ molecular speed (m s⁻¹), aerosol surface density (m⁻¹), NO₂ concentration (ppbv), and the measured NO₂ photolysis frequency (s⁻¹), respectively. As aerosol size distribution measurements was-were not available at the foot station, we estimated S_a based on the measured PM_{2.5} concentrations because as they were highly correlated. For instance, measurements at the summit station during this campaign and other sites in the NCP found high correlations between PM_{2.5} and S_a (derived from particle size distribution measurement) with a S_a/PM_{2.5} ratio of about 8×10⁻⁶ – 1.3×10⁻⁵ m² µg⁻¹ (Wu et al., 2008; Xue et al., 2020). Here a S_a/PM_{2.5} ratio of 1.0×10⁻⁵ m² µg⁻¹ was used, and its uncertainty will not cause significant changes in the HONO simulations because of its small contribution (see the following discussion).

Diurnal variations of P(HONO)_{a_dark} and P(HONO)_a, in comparison with P_{unknown} and P(HONO)_{NO+OH}, are shown in Figure 6A. Clearly, both P(HONO)_{a_dark} and P(HONO)_a ($\gamma_{a} = 2 \times 10^{-5}$) were negligible compared to daytime P_{unknown}. P(HONO)_a increased with γ_{a} , but even when using an extremely highthe upper limit of $\gamma_{a} = 1.3 \times 10^{-4}$, it was still too small to be comparable to P(HONO)_{NO+OH} and far from explaining P_{unknown}. This observation revealsing minor impacts of P(HONO)_{a_dark} and P(HONO)_a heterogeneousin HONO formation on particle surfaces to the missing HONO source, particularly during the daytime.



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Figure 6: HONO production rates from different ((A): aerosol-derived, (B): ground-derived) sources and unknown HONO source strength.

3.2.2.3 pNO₃ Photolysis

Parameterization of HONO formation from particulate nitrate photolysis (P(HONO)_{pn}) is presented in (Eq-6). Recent studies found that EF values were generally lower than one magnitude 10, for instance, a value of 7 from a recent field study (Romer et al. 2018). and ~1 from In laboratory studies even lower values of ~1 were observed (Laufs and Kleffmann, 2016; Shi et al., 2021; Wang et al., 2021). Hence an EF value of 7 was used in the P(HONO)_{nn} calculation, and values of 1 and 15.6 (overestimated one derived from the summit measurement) were also used to test the sensitivities (the upper limit value was

derived from the summit measurement, see Xue et al. (2021a)).

$$P(HONO)_{pn} = pNO_3 * J(HNO_3) * EF,$$
(Eq-6)

where pNO₃ and J(HNO₃) represent the measured particulate nitrate (with unit converted from $\mu g m^{-3}$ to ppbv) and the photolysis frequency of gas-phase HNO₃ (s⁻¹), respectively. 365

Diurnal variations of P(HONO)_{pn} with different EF values are shown in Figure 6A. With EF varying from 1 to 7, P(HONO)_{pn} was 1-2 orders of magnitude lower than P_{unknown}. Even using a high EF = 15.6, P(HONO)_{nitrateon} was still significantly less than half of $P(HONO)_{NO+OH}$. Therefore, model results constrained by field measurements and recent kinetics suggested that the two aerosol-derived sources (NO₂ conversion and nitrate photolysis) may not have significant impacts on daytime HONO formation, with their contributions significantly lower than half of $P(HONO)_{NO+OH}$.

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3.2.2.4 NO₂ Uptake on the Ground Surface

Parameterizations of HONO production from the NO₂ uptake on the ground surface without (P(HONO)₂ dark) and with $(P(HONO)_g)$ photo-enhanced effects $(P(HONO)_g \text{ dark} \text{ and } P(HONO)_g)$ are demonstrated in (Eq-7) and (Eq-8), respectively. NO₂ uptake coefficients of $\gamma_{g_{dark}}$ and γ_{g} were set to 1.6×10⁻⁶ and 2×10⁻⁵ (Han et al., 2016; Stemmler et al., 2006, 2007), respectively. The photo-enhancement effect was reflected described by $\frac{J(NO_2)_{measured}}{0.005 \text{ s}^{-1}}$ (Vogel et al., 2003; Wong et al., 2013; Xue et al., 2020).

$$P(HONO)_{g_dark} = \frac{v(NO_2) \times [NO_2]}{8 \times MLH} \times \gamma_{g_dark},$$

$$P(HONO)_g = \frac{v(NO_2) \times [NO_2]}{4 \times MLH} \times \gamma_g \times \frac{J(NO_2)_{measured}}{0.005 \, s^{-1}},$$
(Eq-8)

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It can be found that As shown above, one of the most important parameters for calculating ground HONO formation in a box model is the mixing layer height (MLH) as it is part of the denominators in both (Eq-7) and (Eq-8). MLH for HONO should be significantly lower than the boundary layer height (BLH) due to its formation on the ground level and its short lifetime, which could be confirmed by the gradient measurements (Kleffmann et al., 2003; Meng et al., 2020; Vogel et al., 2003; Wang et al., 2019; Wong et al., 2012; Xing et al., 2021; Ye et al., 2018b). For instance, a recent gradient HONO measurement study by the MAX-DOAS technique in southwest China found a very rapid-steep HONO decrease as increasing altitude gradient

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- 385 from 0 to 4 km (Xing et al., 2021). When considering their measurement at 17:00 (UTC+8) as an example, HONO levels rapidly decreased from 4.8 ppbv at the ground level (~4 m above the ground surface) to 1.6, 0.7, 0.3, 0.2, and 0.1 ppbv averaged in height ranges the layers of 0 - 100, 100 - 200, 200 - 300, 300 - 400, and 400 - 500 m above the ground level, respectively. In contrast, both NO₂ and aerosol extinction remarkably increased from the ground level to about 200 m above the ground level and then decreased with altitude (>200 m). These observations indicate $\frac{1}{2}$ that 1) ground-derived sources dominated
- 390 daytime HONO formation; 2) the MLH for HONO was much less than 100 m, and 3) significant overestimation, i.e., by a factor of >3 in Xing et al. (2021), could be expected if using measurements on the ground surface to represent the average HONO within an MLH higher than 100 m. A similar phenomenon could also be found in Brown et al. (2013) and Vandenboer et al. (2013). Therefore in the present study, the MLH for HONO was set at a constant height of 50 m, were used to scale the MLH with sensitivity tests performed with the MLH set atom 35 and 100 m. Similar values (25 - 100 m) were also used in
- 395 previous box model studies (Lee et al., 2016; Xue et al., 2020, 2021c). It should be highlighted that a box model as used in the present study is not an ideal tool for studying a ground source when comparing with near-near-ground surface measurements in the atmosphere. For the future, gradient measurements are recommended, which should be compared with 1-D model simulations.
- Diurnal variations of P(HONO)_g dark and P(HONO)_g, in comparison with P_{unknown} and P(HONO)_{NO+OH}, are shown in Figure 6B. 400 P(HONO)_{g dark} was is the largest HONO source during the night-time, while it was negligible during the daytime, which is consistent with many previous studies (Li et al., 2010; Liu et al., 2019; Vogel et al., 2003; Xue et al., 2020; Zhang et al., 2019b, 2019a, 2016). With Considering the photo-enhanced effectformation, P(HONO)_g showed shows a similar shape and a similar level to daytime Punknown, indicating the potential dominance of P(HONO)_g in the daytime HONO formation. When changing MLH to 100 (or 35) m, the level of P(HONO)_g became becomes much lower (or higher) than Punknown. Due to this, for which 405 they were discussed here as sensitivity tests on MLH were conducted but-not used in Sce-2 a constant MLH of 50 m was used.

Small differences in the shapes of measured and modeled results may be also caused by the variable MLH induced by variable vertical mixing in the atmosphere and the variable photolytic lifetime of HONO during the daytime.

3.2.3 HONO Budget

Along with the previous discussion, we conducted a model run (Sce-2) with all the discussed HONO sources. As shown in

- 410 Figure 7, the this model with present HONO source parametrizations showed magnificent performanceperformed well in predicting HONO, indicating reasonable parameterizations of the HONO sources. Here,-as the time series of the modeled HONO was were very consistent with those of the observations in both for the variations and the absolute levels, except Exception was only the heavy rain during the period from 25th to 28th June (because of heavy rain, see the next section),. indicating reasonable parameterizations of the HONO sources. In particular, the model exhibited very high performance in
- predicting noontime (10:00 16:00) HONO as the modeled HONO was very close to the observed HONO (Figure 7B). 415 Moreover, in Sce-3 we reduced γ_g by a factor of 10 and enlarged γ_a from 2×10^{-5} to 1.2×10^{-3} or EF from 7 to 400. We found While that the <u>modified</u> model could also generally predict the observed HONO levels (Figures S_{2}^{87} A and S_{2}^{98} A) but <u>it largely failed</u>

to reproduce the noontime observations in levels and variations (Figures S<u>8</u>7B and S<u>98B) including its asymmetry as</u> mentioned in Section 3.2.1., This observation reinforcesing our conclusion that the non-dominated roles of aerosol-derived sources play only a minor role sources in the daytime HONO formation.

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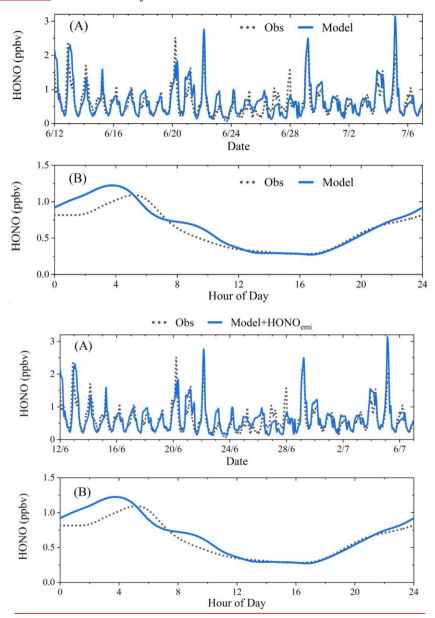
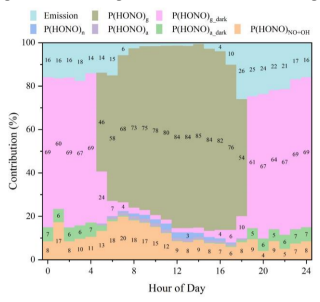


Figure 7: Modeled HONO mixing ratios (Model, in blue) in comparison with observations (Obs, in black). (A): time series; (B): average diurnal variations. Note that Model+HONO_{emi} was included in represents the sum of the modeled HONO and HONO_{emi}.

425 Figure 8 displays the <u>diurnal</u> relative contributions of different HONO sources-<u>at different hours</u>. It clearly shows that dark NO₂ uptake on the ground surface dominated (~70%) night-time HONO formation while photo-enhanced NO₂ uptake on the

ground surface dominated (~80%) daytime HONO formation. P(HONO)_{NO+OH} played a moderate role throughout the whole day, with a contribution of 5 - 15% except for a relatively larger contribution (~20%) in the early morning due to high NO levels. Direct emissions made moderate contributions of 15 - 25% at night but negligible ones during daytime. Contributions of P(HONO)_{a_dark}, P(HONO)_a, and P(HONO)_{pn} were always lower than 10%, and their contributions could be even smaller when using smaller-realistic kinetic parameters derived in recent studies. Therefore, aerosol-derived HONO sources may not

significantly contribute to HONO formation at this site for near-ground surface measurements (Chen et al., 2019; Neuman et al., 2016; Sarwar et al., 2008; Vogel et al., 2003; Wong et al., 2013; Xue et al., 2020; Zhang et al., 2016, 2019b).



435 Figure 8: Relative contributions of different HONO sources. Note that the contribution from direct emission was calculated based on the ratio of HONO_{emi} to the observed HONO.

3.2.4 Other Potential Sources

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As discussed before, the model (Sce-2) could generally well reproduce most observations except for the period from 25th to 27th June. A significant overestimation occurred from midday of 25th to the morning of 26th, which was caused by the enhanced wet/dry deposition due to the heavy rain (>100 mm, Figure S109) on-during the night of 24th/25th. In contrast, from midday of 26th to the night of 27th/28th, a significant underestimation by the model was obtained. Besides, an elevation of HONO/NO_x was found during this period (Figure S109). This might be caused by 1) the enhanced HONO emission from urban soil or 2) the enhanced NO₂ uptake on the ground surface. The former oneHONO emissions from soils may occur through biological processes observed in the laboratory experiments or field measurements over the agricultural fields (Oswald et al., 2013; Scharko et al., 2015; Tang et al., 2019; Xue et al., 2019), while evidence for its occurrence on the urban soil surfaces after the rain was is still not sufficientuncertain. At 13:00 on 26th or and 27th June, the model predicted lower HONO by almost a factor of 2 – 4 (observation: 0.45 or 0.45 ppbv on both days; model: 0.13 or and 0.21 ppbv), which needs an enhancement of at least 2 – 4 in γ_g if using NO₂ uptake on the ground surface to explain the underestimation. Current laboratory experiments studies

have studied the enhancement effect of atmospheric RH (in the range of 10 - 70%) on the NO₂ uptake coefficient on the humic

- 450 acid surface of target substances and the with enhancement factors was of less than 3 (Han et al., 2016; Stemmler et al., 2006, 2007). Campaign averages of the measured NO₂ and RH at 13:00 were 7.4 ppbv and 35.5%, respectively. At 13:00 on 26th (or 27th) June, the measured NO₂ of 7.9 (or 4.3) ppbv was similar to (or lower than) the campaign average, but the RH of 67.6% (or 53.1%) was significantly higher than the campaign average. Bbut the RH was still in the range (10-70%) where for which RH showed an small enhancement (less than 3) effect on γ_g (less than 3). Hence, after rain, the an enhanced NO₂ uptake was
- likely tomay be responsible for the underestimation. MeanwhileIn addition, soil HONO emission may co-exist, but more 455 evidence for urban regions was needed. However, the impact of direct water addition to those substances highly humid surfaces (e.g., rainwater on the ground surface) on different HONO sources and sinks was still not-unclear. i.e., it may enhance NO2 uptake and/or soil emission to produce HONO or deposition to consume HONO. Further studies may explore on the impact of rain on urban soil surface processes are necessary, such as field studies of the soil HONO emission fluxes and laboratory studies of NO₂ uptake kinetics on relevant surfaces. 460
 - **3.3 Radical Chemistry**

Comprehensive field measurements in comparison to model studies allow studying the role of HONO in the radical chemistry of the atmosphere. HONO is expected to strongly impact OH levels in the lower atmosphere due to strong daytime HONO sources and due to its fast photolysis. In addition, considering high O_3 levels at the present field site, NO₃ chemistry could also be important particularly during night-time, which will also be discussed in this section.

3.3.1 Role of HONO in Radical Concentrations

Figure 9 shows the simulated radical concentrations in different model scenarios, in which-where their sensitivities to the constrained HONO were tested. It can be obtained that RO_x radicals (OH, HO₂, and RO₂) were significantly affected by the constrained HONO, implying the vital role of HONO in the RO_x budget. For instance, the peak OH concentration in the base 470 case was 0.42 pptv (equivalent to 1.0×10^7 molecules cm⁻³). It decreased to 0.37 (or 0.32) pptv when HONO was reduced by 50% (or 100%) and increased to 0.46 (or 0.51) pptv when HONO was enlarged by 50% (or 100%). In contrast, modeled NO_3 concentrations showed very small variations whether HONO was reduced or enlarged., which is because NO₃ concentrations is are mainly governed by the levels of $O_3 + NO_2$ during night-time, when HONO has no impact on radical levels, caused by due to its-the missing photolysis. Nevertheless, the The almost same radical concentrations in the model cases NO + OH

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and case -100% indicate the minor role of NO + OH in the radical budget as this OH sink is exactly compensated by the OH production through (R-5).

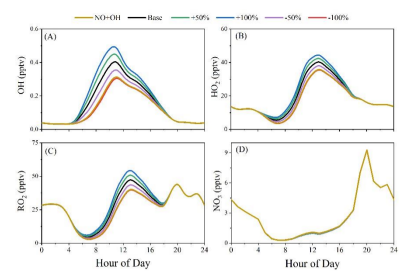
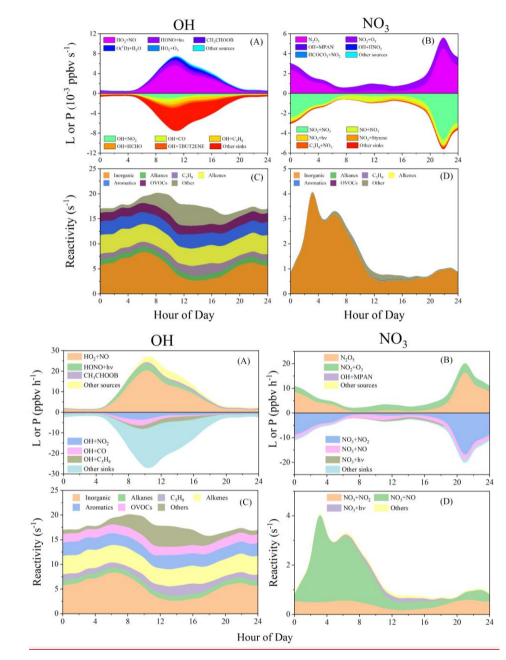


Figure 9: Simulated concentrations of (A): OH, (B): HO₂, (C): RO₂, and (D): NO₃. <u>Different colored lines show results</u> in different scenarios: (NO + OH: only with the homogeneous source; Base: constrained by the observed HONO; +50%: constrained by the observed HONO×1.5; +100%: constrained by the observed HONO×2; -50%: constrained by the observed HONO×0.5; -100%: constrained by HONO = 0].

3.3.2 Radical Production/Loss Rates and Reactivity

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- Figure 10A and Figure 10B illustrate the production/loss rates of OH and NO₃, respectively. The total production rates of these radicals were similar to their loss rates due to their short lifetimes and high reactivities. For OH (Figure 10A), its largest source was the reaction of HO₂ + NO <u>(average contribution: 70%)</u>, which is part of the propagation cycle and which is not a radical initiation source (Elshorbany et al., 2010). HONO photolysis was the second-largest OH source <u>(average contribution: 11%)</u>, and it is expected to be the largest primary OH source after subtracting OH loss through HONO + OH and NO + OH (see Section 3.3.4). Reactions with NO₂, CO, and C₅H₈ acted as the top three OH sinks but did not dominate OH loss due to high OH reactivity caused by various other reactions, particularly those with other VOCs (see below).
- Figure 10C and S1<u>1</u>0A show the OH reactivity with different classes of pollutants and their relative contributions, respectively. Total OH reactivity showed a small peak of 20 s⁻¹ in the morning and then kept almost constant around 17 s⁻¹. Among different classes of pollutants, the measured inorganics (including – ordered by OH reactivity contribution; – NO₂ > CO > NO > O₃ > HONO > SO₂ > H₂O₂) contributed the largest to the OH reactivity with values in the range of 2.6 – 8.4 s⁻¹. Their total contribution was larger in the morning (43%) due to high NO, NO₂, and CO levels (Figure 2) and decreased to 15% at noontime. Reactivities with the measured alkanes, alkenes, aromatics, and OVOCs were 0.95 – 1.2 s⁻¹, 3.3 – 3.9 s⁻¹, 2.2 – 2.9 s⁻¹, and 1.65 – 1.9 s⁻¹, leading to relative contributions of around 5 – 7%, 18 – 22%, 11 – 17%, and 9 – 12% throughout the whole day, respectively. Likewise, C₅H₈ alone contributed 4% of the OH reactivity in the early morning (0.85 s⁻¹), and its contribution increased to 12% at noontime (2.1 s⁻¹) as a result of high levels of C₅H₈ and OH at noontime.



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Figure 10: Production <u>rates</u> (P)<u>and</u>_loss<u>(L)</u> rates (L) and reactivities of <u>radicals</u>. (A): <u>L</u> and <u>P</u> of OH<u>and</u>; (B): <u>L</u> and <u>P</u> of NO₃; and reactivities of (C): <u>Reactivities of</u> OH and (D): <u>Reactivities of NO₃ with measured species</u>. In (A) and (B), the top-five-<u>3</u> sources or sinks are shown, and all the others are summarized in "Other sources" or "Other sinks". In (C), and (D), OH reactivities with <u>different families of the measured species are shown and reactivities with</u> all the unmeasured species are summarized in "Others<u>"</u>. In (D), NO₃ reactivities from top-3 reactions are shown and all the others are summarized in "Others".

Figure 10D and S11 θ B show NO₃ reactivity with different pollutant classes and their relative contributions, respectively. Compared with the total OH reactivity, the total NO₃ reactivity exhibited lower values and a different variation profile. It showed a minimum of 1 s⁻¹ at noontime and increased to around 4 s⁻¹ at 2:00. In addition to the N₂O₅-decomposition (R-17), The reaction NO₂ + O₃ (R-11) is the most important <u>net NO₃ source</u>, which is also, in fact, the most important net NO₃-source,

510 In contrast, NO₃ formation by the N₂O₅ deposition (R-17) considering is almost compensated by the same amount of NO₃ loss during N₂O₅ production through (R-14). NO₃ loss was dominated by its photolysis and its reactions with NO during the daytime and reactions with NO₂ at night. More discussion on the NO₃ chemistry is presented in the following section.

3.3.3 NO₃ Chemistry

As shown in Figure 9D, high NO₃ levels (diurnal peak: 9.3 pptv, time-series peak: 22 pptv) were simulated by the model. High
515 NO₃ concentrations, as well as its-high NO₃ reactivitiesy (Figures 10D), generally appeared at night (18:00 to 4:00 in the next day) when OH was very low and NO₃ was not lost by photolysis, indicating that the NO₃-initialized chemistry may play an important role in night-time chemistry at this site. To verify this assumptionimplication, we compared the C₅H₈ oxidation and nitrate formation through NO₃-induced reactions with other pathsreactions.

3.3.3.1 C₅H₈ Oxidation

- Figure 11 shows (A): the C₃H₈ loss rates (L(C₃H₈)) through different oxidation paths and (B): their relative contributions. L(C₃H₈) through O₃ was generally in the range of 1.0 3.2×10⁻⁵0.04-0.12 ppbv sh⁻¹. L(C₃H₈) through OH showed high values in the daytime and low ones in theduring night-time. On the contrary to OH, low-loss rats L(C₅H₈) through NO₃ occurred-were low in the daytime and high one occurred in the night timeduring night. On average, L(C₃H₈) through OH, O₃, and NO₃ oxidation were 3.6×10⁻⁴1.3, 2.0×10⁻⁵0.07, and 4.5×10⁻⁵0.16 ppbv sh⁻¹, with relative contributions of 84%, 5%, and 11%, respectively. During the daytime, L(C₃H₈) through OH oxidation was generally one order of magnitude higher than those through NO₃ or O₃ oxidation, leading to a dominated C₃H₈ loss contribution of generally >90% through OH oxidation (Figure 11B). However, at night, OH was much lower and NO₃ was higher due to the absence of photochemistry, resulting in an increasing contribution of L(C₅H₈) through NO₃ oxidation (Figure 11B). Average L(C₃H₈) through night-time NO₃ oxidation increased to 8.4×10⁻⁵0.30 ppbv sh⁻¹, but L(C₅H₈) through OH oxidation decreased to 9.2×10⁻⁵0.33 ppbv sh⁻¹, resulting in a
- ⁵³⁰ relatively high contribution of NO₃ oxidation (32 57%). NO₃ oxidation contributed to 44% of the night-time C₅H₈ loss, which is comparable to OH oxidation (48%) and much higher than O₃ oxidation (8%). Considering that C₅H₈ is an important common hemiterpene emitted from multitudinous by many species of vegetations and its oxidation plays a key role in secondary organic aerosol (SOA) formation₂₇. The results of the present study show that daytime OH-induced C₅H₈ oxidation was highlighted is the most important, while NO₃-induced oxidation of C₅H₈ may also significantly affect the SOA formation during the night-
- time (Brown and Stutz, 2012; Mellouki et al., 2021).

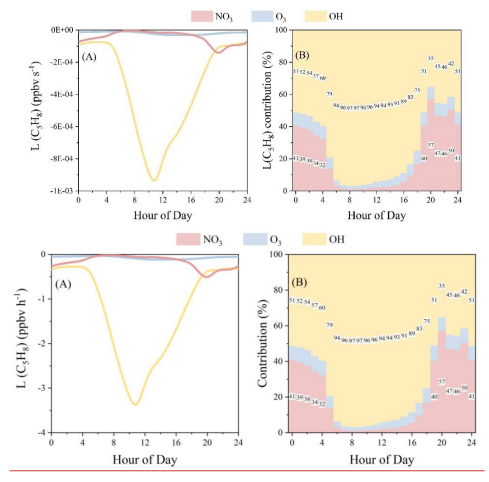


Figure 11: (A): <u>Diurnal Lloss</u> rate of C₅H₈ through each path<u>by NO₃, O₃ and OH oxidation</u> and (B): their <u>diurnal</u> relative contributions at each hour of the day.

540 3.3.3.1-2 HNO₃ Formation

As an important component of particulate matter, inorganic nitrate (pNO₃) was-is produced through by the partitioning of gas-phase_HNO₃ and by heterogeneous uptake of N₂O₅. HNO₃ formation from OH+NO₂ and from N₂O₅ hydrolysis on the aqueous aerosol surface are also the major daytime and night-time sinks for removing NO_x from the atmosphere. Hence, the production of HNO₃, defined as P(HNO₃) = P(HNO₃)_{OH} + P(HNO₃)_{NO₃}, represents the upper limits of pNO₃ production. P(HNO₃)_{OH}
545 denotes the HNO₃ production through (R-18) in the model (Sce-0). For calculation of P(HNO₃)_{NO₃}-calculation, both HNO₃ formation through N₂O₅ heterogeneous uptake on the aerosol surface (R-16) and other NO₃-induced reactions were considered (the former iswas the dominated dominant one). In the model, parameterization for the heterogeneous N₂O₅-uptake is presented in (Eq.9).

$$NO_2 + OH \rightarrow HNO_3$$

(**R-18**)

550 In the model, parameterization for the heterogeneous N₂O₅ uptake is described by presented in (Eq-9).

$$P(HNO_3)_{N_2O_5} = \frac{\nu(N_2O_5) \times S_a \times [N_2O_5]}{4} \times \gamma_{N_2O_5} \times 2,$$
(Eq-9)

where v(N₂O₅), [N₂O₅], and γ<sub>N₂O₅ represent the molecular speed, concentration, and heterogeneous uptake coefficient of N₂O₅, respectively. γ<sub>N₂O₅ was typically set as 0.1 reported in previous studies (Brown and Stutz, 2012; Wang et al., 2017). As shown in Figure 12, the overall P(HNO₃) was high during the daytime and low during the night-time. During the daytime, P(HNO₃)_{NO₃} was generally much lower than P(HNO₃)_{OH}, leading to high contributions of P(HNO₃)_{OH} (>90%). However, during the night-time, P(HNO₃)_{OH} and its relative contribution to P(HNO₃)_{NO₃} to the sum P(HNO₃). On average throughout all day, P(HNO₃)_{NO₃} contributed 18%, significantly lower than P(HNO₃)_{OH} (82%). However, at night, P(HNO₃)_{NO₃} contribution increased to 51%, slightly higher than P(HNO₃)_{OH} (49%). Model results may have uncertainties but shed light on the atmospheric chemistry in this polluted region. By far, very few NO₃ measurements are available in China (Lu et al., 2019; Suhail et al., 2019), while its high concentration and important role in chemical oxidation presented in this study shed light on the summertime O₃ levels areis substantially increasing (Han et al., 2020; Li et al., 2019; Sun et al., 2016, 2019).
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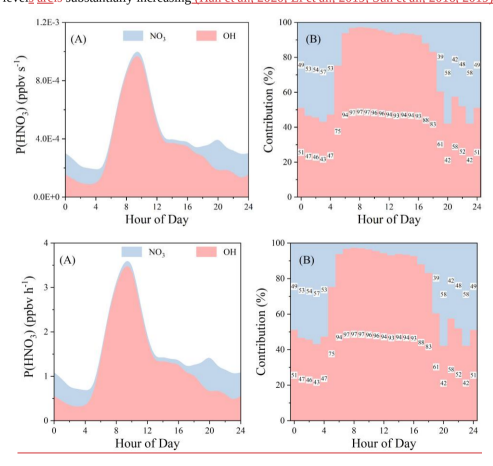
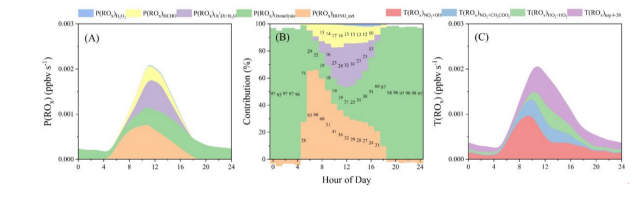


Figure 12: (A): HNO₃ production (P(HNO₃)) from NO₃- or OH- induced reactions and (B): their relative contribution at each hour of the day. NO₃-induced reactions include heterogeneous uptake of N₂O₅ on the aerosol surface and all the other NO₃ reactions that produce HNO₃.

3.3.4 Radical Initiation vs. Termination

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- Measurements on of other radical precursors, such as H₂O₂ (through photolysis to produce OH), HCHO (through photolysis to produce HO₂), and alkenes (through ozonolysis via Criegee intermediate to produce OH₂ and HO₂, and RO₂), were available, which allows a comparison of radical initiation (primary production) and termination (T(RO_x)). As shown in Figure 13, the overall radical initiation and termination showed similar variations and levels. Both, the sum of the RO_x initiation and termination showed peaks of about 2.1×10⁻³7.6 ppbv sh⁻¹ at noon, which are in the range of 0.7 3.4×10⁻³2.5-12.2 ppbv sh⁻¹
 reported in previous studies (Elshorbany et al., 2010, 2012; Hofzumahaus et al., 2009; Kukui et al., 2014; Liu et al., 2012; Ren et al., 2003). During the daytime, it is evident that HONO photolysis (P(RO_x)_{HONO_net}) made the largest contribution (20 70%, Figure 13B) to RO_x initiation, with an average of 37% (or 32% for all-day, Figure S124), followed by ozonolysis (29%), O₃ photolysis (21%), HCHO photolysis (13%), and H₂O₂ photolysis (1%). In particular, RO_x production from the ozonolysis of alkenes was significantly lower than that from HONO during 6:00 14:00 until later after 17:00 when it started to dominate
- 580 RO_x production. At night with the absence of photochemistry, ozonolysis was the major source for primary RO_x. <u>Due to the</u> <u>also high production of radicals during the daytime, and exhibited similar levels to itself during the daytime, leading</u> <u>toozonolysis played anits</u> important role in primary RO_x production (39% for all day). <u>Besides, The termination of radicals</u> T(RO_x) was dominated by NO₂ + OH \rightarrow HNO₃, NO₂ + CH₃COO₂ \rightarrow PAN, and HO₂ + HO₂ \rightarrow H₂O₂ (Elshorbany et al., 2010, 2012; Hofzumahaus et al., 2009; Kukui et al., 2014; Liu et al., 2012; Stone et al., 2012).



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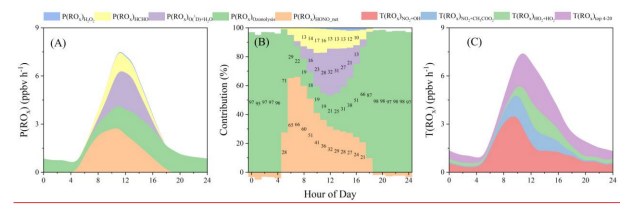


Figure 13: (A):-Primary RO_x production and net RO_x loss. (A): Production from different sources and (B): their relative contributions at different hours of the day, and (C): the top-20 RO_x loss rates. Note that: 1) due to an integration problem, the top-20 net radical loss paths were summarized here and i. It could represent the majority of total $T(RO_x)$ as others (<1×10⁻⁵0.03 ppbv sh⁻¹) were at least 2 orders of magnitude lower than the sum of top-20, 2). Nnight-time $P(RO_x)_{HONO_net}$ was negative (a net sink for OH) so that its contribution was also negative at night. and 3) tThe same amounts of radical loss or production from equilibrium reactions (e.g., HO₂ + NO₂ \leftrightarrow HNO₄; CH₃COO₂ + NO₂ \leftrightarrow PAN) was excluded from radical initiation or termination. $T(RO_x)_{NO_2+CH_3COO_2}$ represents the net PAN formation.

3.3.5 Role of HONO in OH Production at the Foot and the Summit Stations

595 Although measurements at the foot and the summit stations were conducted during two consecutive periods rather than the same one in summer 2018, it still allows a reasonable comparison of HONO contribution to OH formation at the foot station (lower boundary layer) and the summit station (upper boundary layer). Because of limited data available at the summit station, we only compared HONO with O₃ in primary OH formation. As reported in the companion paper, rapid vertical transport maintains the high HONO level at the summit station, promising HONO an important role in integrated OH production with a contribution of 26% to the sum of OH production (P(OH)_{O₃+HONO}) considering only HONO and O₃-photolysis. If OH loss through HONO + OH and NO + OH was subtracted from P(OH)_{HONO} (then it becomes P(OH)_{HONO} ner), its contribution

decreased to 18%, about one quarter of $P(OH)_{\Theta_2}$.

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Then net OH production from HONO (P(OH)_{HONO_net}) and O₃-(P(OH)_{O₃}) photolysis at the foot and summit stations were also summarized and compared. As shown in Figure 14, it is apparent that HONO photolysis initialized the daytime photo-

- 605 chemistry at both the foot and the summit stations as P(OH)_{HONO_net} dominated OH production in the early morning. Average P(OH)_{HONO_net} and P(OH)_{O₃} at the foot station are 2.4×10⁻⁴ and 1.4×10⁻⁴ ppbv s⁻¹, respectively, both of which are significantly higher than those (1.7×10⁻⁵ and 7.7×10⁻⁵ ppbv s⁻¹) at the summit station as a result of relatively lower HONO and O₃ concentrations and lower solar photolysis frequencies observed at the summit station. The latter was caused by frequent cloud formation neat the summit. Nevertheless, the considerable contributions of P(OH)_{HONO_net} to P(OH)_{O₃+HONO} at the foot (64%)
- 610 and the summit (18%) stations indicate the essential role of HONO in the atmospheric oxidizing capacity at both the ground (lower boundary layer) and the summit (upper boundary layer) levels in mountainous regions.

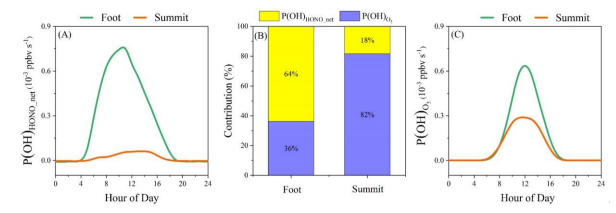


Figure 14: (A): Net OH production from HONO (P(OH)_{HONO_net}), (C): net OH production from O₃ (P(OH)_{O3}) photolysis, and (B): their relative contributions at the foot and the summit of Mt. Tai.

615 4. Summary

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Atmospheric HONO and related parameters (VOCs, NO_x, PM_{2.5}, J(NO₂), etc.) were measured at the foot and the summit of Mt. Tai in the summer of 2018. The present study was conducted mainly based on measurements at the foot station. The observed HONO varied from 0.05 to about 3 ppbv, with an average of 0.62 ± 0.42 ppbv. With the implementation of a 0-D box model (F0AM) coupled with the Master Chemical Mechanism (MCM v3.3.1), the HONO budget and the radical (RO_x +

620 NO₃) chemistry were explored.

The main conclusions are summarized as follows:

- The default HONO source, NO + OH, significantly underestimated the observed HONO <u>concentrations</u>. This reaction <u>could only account for 13% of the observed HONO by 87%</u>, revealing a strong unknown source (P_{unknown}). The diurnal profile of P_{unknown} rapidly increased in the morning and peaked nearly 3 ppbv h⁻¹ at <u>ca. 1 hour before</u> noon, suggesting additional photo-enhanced HONO formation processes.
- 2. A HONO/NO_x ratio of 0.7% was derived for direct emission, and its contribution (15 25% at night but negligible during the daytime) was furtherly-quantified by a new method developed in this study. Based on the constraints on the aerosol-derived HONO sources (NO₂ uptake on the aerosol surface and nitrate photolysis) obtained from the summit measurement (see the companion paper) and from recent laboratory studies, we found that the aerosol-derived HONO sources may make moderate or small contributionsnot significantly contribute to HONO formation at the summit level and the ground level₂, respectively, but their Their contributions to HONO formation were not higher always smaller than NO + OH. Heterogeneous NO₂ conversion on the ground surface made the largest contribution to P_{unknown}, but it was sensitive to the MLH used for its parameterization. This addressed the importance of a reasonable MLH for exploring ground-level HONO formation in 0-D models and the necessity of vertical measurements.
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3. HONO played an important role in RO_x but a negligible role in NO₃ concentrations. OH dominated the atmospheric oxidizing capacity in the daytime, while NO₃ appeared to be significant at night. Peaks of NO₃ time series and diurnal variation reached 22 and 9 pptv, respectively. NO₃ induced reactions contribute 18% of nitrate formation potential and 11% of the C₅H₈ oxidation throughout the whole day. While at night, NO₃ chemistry led to 51% or and 44% of the nitrate formation potential or and the C₅H₈ oxidation, respectively. NO₃ chemistry may significantly affect night-time secondary organic and inorganic aerosol formation in this high O₃ region. Hence, the direct measurement of NO₃ (along with HO_x, N₂O₅, ClNO₂, etc.) in this region should be conducted.

4. A comparison of HONO contributions to primary OH at the summit and the ground levels was conducted and it was confirmed that HONO photolysis initialized daytime photo chemistry at both sites in the early morning. On average,
 645 HONO made contributions of 64% and 18% to P(OH)_{O₃+HONO} at the summit and the ground levels, respectively. As HONO observed at the summit level was mainly transported from the ground level, it addressed the role of HONO in the atmospheric oxidizing capacity in both the lower and the upper boundary layer over mountainous regions.

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Author Contribution: C.X., C.Y., W.Z., X.H., P.L., C.Z., X.Z., C.L., Z.M., J.L., and J.W. performed the field measurements. C.X. analyzed the observation data, performed model simulations, and wrote the paper with input from all co-authors. C.Y. and J.K. also contributed by fruitful discussions and comments on model simulations and writing. J.K., C.Y., K.L., V.C., J.K., A.M., and Y.M. revised the manuscript.

660 **Competing Interests**: The authors declare no competing financial interest.

Data Availability: All the data used in this study is available upon request from the corresponding authors.

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