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Atmospheric nitrous acid (HONO) at a rural coastal site in North China: Seasonal variations and effects of biomass burning

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HIGHLIGHTS

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- · Significant seasonal variations of HONO
- pollution and chemistry were found. Source strength of HONO can be enhanced by solar radiation and
- biomass burning. · Biomass burning enhanced the role of aerosol in heterogeneous HONO formation.

GRAPHICAL ABSTRACT



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ABSTRACT

Nitrous acid (HONO) plays a significant role in atmospheric chemistry due to its contribution to hydroxyl radical (OH). However, no scientific consensus has been achieved about the daytime HONO formation mechanisms. To identify the seasonal variations of HONO chemistry and the impacts of biomass burning (BB), we performed a twophased field study in winter-spring and summer (covering a harvest season) in 2017 at a rural coastal site in North China. Though the mean HONO concentration in winter-spring (0.26 \pm 0.28 ppbv) was higher than in summer (0.17 \pm \pm 0.19 ppbv), the maximum HONO concentrations were comparable (~2 ppbv) in the two campaigns. Both the HONO/NO_x ratio and nocturnal heterogeneous conversion efficiency of HONO (C_{HONO}) in summer were over twice of that in winter-spring. The daytime budget analysis also revealed that the strength of Pother (i.e., the HONO sources apart from the reaction of OH + NO) in summer was double of that in winter-spring. BB affected the HONO concentration by enhancing the contribution of heterogeneous HONO production on the aerosol surface but weakening the role of photo-related HONO formation. HONO photolysis was a significant source of OH in both winter-spring and summer, and its contribution could be further enhanced during the BB episode in summer. Our study demonstrates the significant seasonal variations of HONO and the effects of BB, and suggests needs for more multiseason observations and considerations of BB, especially during the harvest time, in HONO research.

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1. Introduction

Atmospheric nitrous acid (HONO) has attracted great attention in recent decades due to its contribution to the hydroxyl radical (OH) and participation in the NO_x cycle (Lammel and Cape, 1996; Ren et al., 2003; Ye et al., 2016b). OH is the major atmospheric oxidant initiating the diurnal photochemistry that can affect air quality and climate (Finlay-son-Pitts and Pitts, 2000). Numerous studies have shown that the photolysis of HONO can make a significant contribution to OH not only in the early morning but also throughout the rest of the day (Acker et al., 2006b; Fu et al., 2019). Therefore, investigating the pollution characteristics and chemistry of HONO is essential for a better understanding of atmospheric chemistry and the causes of regional air pollution.

Despite the relatively good understanding of the sinks (R1, R2, and dry deposition), sources of HONO are still under debate. Tropospheric HONO sources include chemical formation and direct emissions.

$$HONO + h\nu \rightarrow NO + OH \quad (320 \text{ nm} < \lambda < 400 \text{ nm}) \tag{R1}$$

$$HONO + OH \rightarrow NO_2 + H_2O$$
 (R2)

The chemical reactions producing HONO include homogeneous gasphase reactions and heterogeneous reactions on various surfaces. So far, the proposed gas-phase reaction sources of HONO include reactions R3 to R7. Among them, R4 is of minor importance due to a low HONO yield (Sörgel et al., 2011; Ye et al., 2015). R5 is still under debate due to the unconfirmed reaction rate coefficient (Carr et al., 2009; Li et al., 2008). R6 is proposed to be a non-negligible HONO source in the urban atmosphere but should be verified through more laboratory studies (Bejan et al., 2006). R7 is only proposed by density functional theory calculations, and further experimental demonstrations are needed (Zhang and Tao, 2010). The proposed heterogeneous reactions producing HONO include reactions R8 to R13. R8 is the earliest identified heterogeneous source of HONO, but it generally produces insignificant HONO because of the small reaction rate constant (Finlayson-Pitts et al., 2003). R9 is the alternative reaction of R8 and is demonstrated to be an important HONO source, but its contribution has not been accurately quantified due to the complex influences of NO2 concentrations, moisture content, and surface characteristics on the NO2 uptake and HONO product yield (Finlayson-Pitts et al., 2003; Sakamaki et al., 1983). R10, the heterogeneous reactions of NO₂ on the surfaces of reducing matters such as soot, humic acid, mineral dust, and other organic substrates, can produce HONO and the production rate has been proved to be photo-enhanced (Han et al., 2017; Monge et al., 2010; Stemmler et al., 2006). In R10, 'ads' means surface-adsorbed, and 'red' and 'ox' denote the reducing matters and oxidizing substances, respectively. R11 is proposed as a HONO source in an environment surrounding with H₂O molecules but later believed to have little contribution to the ambient HONO (Kleffmann et al., 2004; Saliba et al., 2001). R12 is the photolysis and remobilization of HNO₃ and nitrate on surfaces, and it has been recognized as the dominant HONO source in the low-NO_x situation near the ground surface in daylight (Yang et al., 2018; Ye et al., 2016a; Zhou et al., 2003). Besides, the photolysis of particulate nitrate (R13) has been proposed to be a HONO source with the photolysis frequency ranging from 10^{-6} to 10^{-4} s^{-1} , which is affected by nitrate loading, organic matters, aerosol pH, and relative humidity (Ye et al., 2017; Gen et al., 2019).

 $OH + NO + M \rightarrow HONO + M$ (R3)

 $HO_2 \cdot H_2O + NO_2 \rightarrow HONO + O_2 + H_2O$ (R4)

 $NO_2^* + H_2O \rightarrow HONO + OH$ (R5)

ortho – nitrophenol + $h\nu \rightarrow$ HONO (R6)

$$2NO_2(g) + H_2O(g) + NH_3(g) \rightarrow HONO(g) + NH_4NO_3(s)$$
(R7)

$$NO + NO_2 + H_2O \xrightarrow{surface}{\rightarrow} 2HONO$$
 (R8)

$$2NO_2 + H_2O \xrightarrow{\text{surface}} HONO + HNO_3$$
(R9)

$$NO_2 + red_{(ads)} \rightarrow HONO + ox_{(ads)}$$
 (R10)

$$HNO_{3(ads)} + NO_{(g)} \rightarrow HONO + NO_2$$
(R11)

$$HNO_{3(ads)} / nitrate_{(ads)} + h\nu \rightarrow HONO$$
(R12)

$$pNO_3 + h\nu \rightarrow HONO + NO_2$$
 (R13)

Direct emission sources of HONO include fossil fuel combustion (Liu et al., 2017), microbial activities in soil (Oswald et al., 2013), and biomass burning (BB) (Burling et al., 2010). While fossil fuel combustion such as vehicle exhaust is a non-negligible HONO source in urban areas (Liang et al., 2017; Yun et al., 2017), soil emissions can make a considerable contribution to ambient HONO in the areas that retain a soil cover (Oswald et al., 2013; Zhang et al., 2016). Besides, both laboratory and field studies have shown that BB could emit HONO directly and produce HONO indirectly through heterogeneous reactions of NO₂ on soot particles (Burling et al., 2010; Yokelson et al., 2009). In a field study, direct BB emission was estimated to contribute up to 17% to the observed HONO, and heterogeneous conversion of NO₂ contributed over 80% in a nighttime BB plume (Nie et al., 2015). Despite being a significant HONO source during the harvest seasons, the BB emission of HONO has been relatively less studied so far.

In recent decades, numerous HONO field studies have been conducted around the world. Most of the observations were short-term and mono-season, and only a few long-term or multi-season measurements have been carried out. Wang et al. (2013) observed weak seasonal variations of HONO and HONO/NO₂ at an urban site in Shanghai, China. In contrast, significant monthly and seasonal differences of HONO and HONO/NO₂ were observed in some other regions. Xu et al. (2015) and Li et al. (2018) reported the maximum HONO concentrations in winter and elevated HONO/NO₂ ratio in summer at an urban site in the Pearl River Delta (PRD) and the North China Plain (NCP), respectively. In comparison, the highest HONO concentration was observed in autumn with the lowest in winter at an urban site in Beijing (Wang et al., 2017). In view of the diverse temporal variations of HONO, more multi-season field studies are needed for the comprehensive understanding of HONO pollution, sources, and impacts on regional air quality.

To investigate the seasonal variations of HONO and the influence of BB, we carried out a two-phased field study during the winter-spring and summer (covering a harvest season) in 2017 at a rural coastal site in the Yellow River Delta (YelRD) in North China. This site is located in the warm temperate monsoon zone with four distinct seasons and limited anthropogenic activities. We present the results to show the significant seasonal variations of HONO pollution characteristics, nocturnal heterogeneous conversion coefficient (C_{HONO}), diurnal sources and sinks, and impacts on atmospheric oxidation capacity. We also analyze BB cases to highlight its important contribution to HONO during the harvest time.

2. Experiments

2.1. Study site

The field study was carried out at the Yellow River Delta Ecology Research Station of Coastal Wetland, Chinese Academy of Science (118.98°E, 37.76°N), which is located in the YelRD region in North China (see Fig. 1). The station is surrounded by wide-open land with limited population or industrial activities within a radius of 10 km. The three closest populated areas are a town 15 km southwest, a town 20 km northwest, and a town 23 km north-northwest with resident populations of about 23,000, 46,000, and 46,000, respectively. The nearest city of Dongying is about 40 km southwest, with a population of over two million. Besides, the site is 45 km to the northeast of the Shengli Oilfield and 15 km to the southwest of the Gudong Oilfield. As the North China Plain is a major winter-wheat growing area, the air quality in this site may be affected by the wheat straw burning during the harvest season in June (Yin et al., 2019). To sum up, the station is characterized as a typical rural coastal site with distinct seasons and limited local anthropogenic emissions, but could be influenced by potential complicated transported regional air masses. More descriptions of this site were given by Zhang et al. (2019) and Zheng et al. (2019).

2.2. Measurements

The two-phased field study was performed in the winter-spring (from February 8th to March 24th) and summer (from June 1st to July 10th) of 2017. The instruments were housed in an air-conditioned container, and the sampling inlets were mounted on the roof-top of the container, which was about 3 m above the ground.

HONO was measured by an online commercial LOng Path Absorption Photometry instrument (LOPAP-03, QUMA, Germany) using photometric detection method (Heland et al., 2001). The collection efficiency was more than 99%, and the detection limit was 5 pptv at a time resolution of 30 s. Instrument background signals were automatically detected by sampling high purity nitrogen for 30 min every 12 h. Single-point calibrations were carried out using diluted NO₂ standard (Nitrite Standard for IC, 1000 mg/L, Sigma-Aldrich) at a time interval of 4 days. NO and NOv were detected with a commercial chemiluminescence analyzer (T-API Model T200U, USA), which was equipped with an externally-mounted molybdenum oxide converter. NO2 was measured with a Cavity Attenuated Phase Shift (CAPS) detector operating as an optical absorption spectrometer (T-API Model T500U, USA). O3 was measured by a standard UV photometric detector (TEI Model 49C, USA). CO was detected using an IR absorption analyzer equipped with a gas filter correlation wheel (T-API Model T300U, USA). $PM_{2.5}$ was measured using a synchronized hybrid ambient real-time particulate monitor (Thermo Scientific Model 5030 SHARP, USA). Meanwhile, PM_{2.5} samples were collected on the quartz filter membrane (08:00-19:30 for daytime samples and 20:00-07:30 for nighttime samples) using a medium volume sampler (TH-150, Tianhong, China). Inorganic water-soluble ions (K^+ , NO_3^- , NH_4^+ , etc.) in the PM_{2.5} samples were then analysed by an ion chromatography (ICs-90, Dionex, USA).

Photolysis frequencies of HONO, NO₂, and O₃ (j(HONO), j(NO₂), and j (O¹D)) were measured with a spectro radiometer (Metcon, GmbH, Germany). Meteorological parameters (temperature, relative humidity (RH), and wind direction and speed) were monitored by a portable meteorological station (PC4, China). The time resolution of all the measurements above was 1 min except for HONO at 30 s, meteorological parameters at 5 min, PM_{2.5} at 30 min, and water-soluble ions in fine particles at 12 h. More details about their detection limits, precisions, and quality assurance and quality control procedures were provided by Xue et al. (2016) and Zhang et al. (2019).

3. Results and discussion

3.1. Seasonal and diurnal variations

Fig. 2 presents the overview of the observed HONO, related species and meteorological parameters, and the calculated OH throughout the two-phased field study. The OH concentration ([OH]) was calculated using equation (E1) with measured $j(O^{1}D)$ and the coefficients of a = 2.4×10^{6} cm⁻³, b = 1, c = 0.13×10^{6} cm⁻³, which were proposed by Rohrer and Berresheim (2006) for rural areas. The calculated OH concentrations match well with the simulated results by the MCM model (Zhang et al., 2019). The descriptive statistics of NO, NO₂, NO_y, HONO, temperature, and RH are listed in Table 1.

$$[OH] = a \times (j(O^{1}D)/10^{-5} s^{-1})^{b} + c$$
(E1)

Generally, concentrations of air pollutants (except for O₃ and K⁺ in fine particles) were higher in winter-spring than in summer. The average HONO concentrations were 0.26 \pm 0.28 ppbv in winter-spring and 0.17 \pm 0.20 ppbv in summer, with the maximum values of 2.07 and 2.00 ppbv, respectively. The mean HONO concentration at this rural coastal site was lower than that in other polluted rural sites in the Yangtze River Delta (YRD) (Nie et al., 2015), PRD (Li et al., 2012), and Beijing-Tianjin-Hebei region (Yang et al., 2014) in China. However, it was much higher than that in most European sites (Acker et al., 2006a; Alicke et al., 2002; Meusel et al., 2016). NOx and NOy, which are significant precursors of HONO, showed the mixing ratios ranging from several to tens of ppbv with much higher concentrations in winter-spring than in summer. PM2.5 also showed a similar seasonal variation pattern with NO_x and NO_y . As a tracer of biomass burning, K^+ in fine particles showed elevated concentrations in two episodes during June 16-17 and June 29-30 in summer with the daily average mixing ratio of more than



Fig. 1. Maps showing (a) NCP and anthropogenic NO_x emission in China, and (b) the observation site in the YelRD and fire points in the surrounding regions. The anthropogenic NO_x emission data in 2016 in China was obtained from the MEIC (the Multi-resolution Emission Inventory for China, http://www.meicm odel.org/). The fire map of June 2017 was provided by NASA, and fire points from VIIRS, MODIS/Aqua, and MODIS/Terra were colored with red, purple, and orange, respectively (https://firms.modaps.eosdis.nasa.gov/map/). NCP and YelRD represent the regions of the North China Plain and Yellow River Delta, respectively. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)



Fig. 2. Time series of the measured HONO, NO, NO_x, NO_y, O₃, PM_{2.5}, K^+ in fine particles, j(HONO), meteorological parameters, and the calculated OH during the two-phased field study in winter-spring and summer.

Table 1 Statistics of the measured HONO, NO, NO₂, NO_y, and meteorological parameters during winter-spring and summer.

	Winter-Spring						Summer						
	Min.	Max.	Med.	Mean \pm SD			Min.	Max.	Med.	Mean \pm SD			
				All Day	Daytime	Nighttime				All Day	Daytime	Nighttime	
HONO (ppbv)	BDL	2.07	0.18	0.26 ± 0.28	0.22 ± 0.24	0.30 ± 0.28	BDL	2.00	0.11	0.17 ± 0.20	0.16 ± 0.20	0.20 ± 0.18	
NO (ppbv)	BDL	46.92	0.17	1.49 ± 3.28	2.91 ± 3.71	0.33 ± 1.95	BDL	19.03	0.17	0.32 ± 0.50	0.43 ± 0.61	0.16 ± 0.20	
NO ₂ (ppbv)	0.32	55.34	7.86	10.41 ± 9.11	$\textbf{9.85} \pm \textbf{8.10}$	10.50 ± 9.70	BDL	37.32	2.37	3.47 ± 3.34	3.41 ± 3.68	$\textbf{3.48} \pm \textbf{2.64}$	
NO _v (ppbv)	BDL	86.74	17.47	20.36 ± 17.07	22.55 ± 17.90	18.47 ± 16.35	BDL	69.58	8.45	10.14 ± 7.73	11.22 ± 8.74	$\textbf{8.45} \pm \textbf{5.42}$	
Temp. (°C)	-6.4	18.6	4.9	5.3 ± 4.9	$\textbf{8.3} \pm \textbf{4.8}$	3.0 ± 3.7	14.9	36.8	26.1	25.9 ± 4.5	27.3 ± 4.5	23.9 ± 3.8	
RH (%)	19	102	72	68 ± 18	55 ± 18	79 ± 11	25	99	82	76 ± 16	71 ± 17	85 ± 10	

Note: The statistical analysis was based on data with time resolution of 1 min for trace species and 5 min for meteorological parameters. According to the time of sunset and sunrise throughout the experiment, the daytime was defined as 08:00-17:00 (05:00-19:00) local time for winter-spring (summer), and the nighttime was defined as 19:00-06:00 (20:00-04:00) local time for winter-spring (summer). BDL means below detection limit.

1 μ g·m⁻³. As a secondary photochemical product, O₃ exhibited typical high levels in summer with several peaks over 150 ppbv as a result of enhanced photochemistry.

Fig. 3 shows the average diurnal patterns of HONO and related species and parameters during the two-phased field study. Most of the pollutants and parameters presented similar diurnal variations in the two campaigns except for HONO/NOx. HONO concentrations displayed a typical diurnal pattern in both campaigns, rising during the nighttime with a peak before sunrise and decreasing during the daytime with a valley around the late afternoon. The accumulation of HONO during nighttime indicated steady HONO sources at night. Given that the lifetime of HONO during midday was less than 30 min, the considerable HONO concentrations during daytime suggested the existence of significant HONO sources. Besides, despite the greater HONO photolysis and several times lower concentrations of related pollutants (NOx, NOy, and PM_{2.5}) in summer than in winter-spring, the HONO concentrations in summer were just a little lower than those in winter-spring, indicating enhanced HONO sources in summer. The NO concentrations began to rise in the late night and peaked in the early morning during both campaigns, with a much sharper and wider peak in winter-spring than in summer. The diurnal patterns of NO_2 , NO_y , and $PM_{2.5}$ were similar,

peaking in the morning and then decreasing to the valley in the afternoon. O3 showed a typical diurnal pattern in both campaigns, increasing since sunrise and reaching the maximum at around 15:00 LT, and then decreasing until the next day before sunrise and reaching the minimum. The ratio of HONO/NO_x (or HONO/NO₂) has been used to characterize the heterogeneous formation of HONO from NO₂ (e.g., Xu et al., 2015). As the $HONO/NO_2$ ratio may be subject to interferences from direct HONO emissions, the HONO/NOx ratio should be a more reliable parameter to indicate the efficiency of the heterogeneous NO2-to-HONO conversion. The diurnal pattern of HONO/NOx displayed noticeable seasonal variations. In winter-spring, it decreased since late night and reached the minimum in the late afternoon, then increased rapidly before midnight and stayed stable for a few hours until the late night. In summer, an extra peak of HONO/NOx showed up in the midday, which was comparable to the nighttime peak. This peculiar HONO/NO_x peak indicated some enhanced HONO sources at noon in summer. A detailed discussion about the potential HONO sources in the daytime will be given in section 3.3.



Fig. 3. Diurnal variations of (a) NO, (b) NO₂, (c) NO_y, (d) HONO, (e) j(HONO), (f) O₃, (g) PM_{2.5}, and (h) HONO/NO_x during the two-phased field experiment in winter-spring and summer. The error bars represent one standard deviation.

3.2. Nocturnal HONO formation

Heterogeneous conversion of NO₂ to HONO is the commonly accepted HONO source. The HONO conversion rate divided by NO₂ (C_{HONO}, h⁻¹) is a widely used scaling approach for the calculation of nocturnal NO₂-to-HONO conversion frequency (Su et al., 2008a; Yu et al., 2009). Equation (E2), which is suitable for the C_{HONO} calculation at rural sites with little or negligible effects of vehicle emissions, is used in this study (Su et al., 2008a). In E2, P_{HONO} is the HONO production rate with the unit of ppbv·h⁻¹, [NO₂] is the mean NO₂ concentration during time t₁ to t₂, [HONO]_{t₁/2} is the HONO concentration at time t₁ or t₂. To

calculate the C_{HONO} , several criteria were adopted for the case selection. First, only the nocturnal data were considered to avoid the complicated daytime sources and sinks of HONO. Note that the heterogeneous conversion of NO₂ to HONO is the most commonly accepted HONO source at nighttime, when both anthropogenic emissions (in particular at such a rural site in this study) and the atmospheric photochemistry are in their minimum intensity. Second, both HONO concentration and HONO/NO₂ ratio presented increasing trends for a relatively long period (e.g., more than 2 h). Third, the concentrations of other pollutants (such as NO, NO₂, and CO, etc.) were stable to ensure that the air mass was consistent. Last, the weather conditions were stable with steady wind direction and small wind speeds (e.g., $<3 \text{ m} \cdot \text{s}^{-1}$). According to the above criteria, 10 and 7 cases were chosen for the winter-spring and summer phases, respectively, as listed in Table S1.

$$C_{HONO} = \frac{P_{HONO}}{[\overline{NO_2}]} = \frac{[HONO]_{t_2} - [HONO]_{t_1}}{(t_2 - t_1)[\overline{NO_2}]}$$
(E2)

The calculated C_{HONO} showed a significant variation between winterspring and summer. While C_{HONO} ranged from 0.48 to 1.54% h^{-1} with an average of 0.96 \pm 0.33% h^{-1} in winterspring, it varied from 0.90 to 4.06% h^{-1} with a mean of 2.47 \pm 0.98% h^{-1} in summer. C_{HONO} presented a strong correlation with ambient temperature (r = 0.76) and a moderate relationship with RH (r = 0.57), which was consistent with the study of Wen et al. (2019) in a marine boundary layer. The average C_{HONO} at our site was within the range of 0.43–3.36% h^{-1} that determined from other studies around the world (Liu et al., 2019; Xu et al., 2015). Seasonal variation of C_{HONO} has also been found by Liu et al. (2019) at a suburban site in Nanjing in the subtropical zone with four distinct seasons.

3.3. Daytime HONO formation

To investigate the strength of the daytime HONO sources and their seasonal variations, we performed a budget analysis based on the description by equation (E3) (Liu et al., 2019; Su et al., 2008b). The HONO sources include direct emission from anthropogenic combustions (Pemis), the well-accepted homogeneous formation from reaction R3 (P_{OH+NO}) , and the remaining other sources (P_{other}) . The sinks of gaseous HONO contain the homogeneous reaction of R2 (L_{OH+HONO}), photolysis (L_{phot}), and dry deposition (L_{dep}). In addition, vertical and horizontal transport (F_v and F_h) could affect the HONO budget as well. Assuming that no HONO was emitted from anthropogenic activity at this coastal site (considering little anthropogenic emission sources nearby) and the influence of transport was negligible (given the short lifetime of HONO during the daytime), equation (E4) could be derived from E3 for the calculation of $P_{\text{other}}.$ In E4, $k_{\text{OH}+\text{HONO}}$ denotes the reaction rate constant of reaction R2 with a value of 2.5 \times 10⁻¹² \times exp(260/T) $\mbox{cm}^3\mbox{-molecules}^{-1}\mbox{-s}^{-1}$ (Atkinson et al., 2004); k_{OH+NO} denotes the reaction rate constant of reaction R3 with a value of 7.4 \times 10^{-31} \times $(T/300)^{-2.4} \times [N_2] \text{ cm}^3 \cdot \text{molecules}^{-1} \cdot \text{s}^{-1}$ (Atkinson et al., 2004); ν_{HONO} is the dry deposition velocity of HONO in the daytime using an empirical value of $2 \text{ cm} \cdot \text{s}^{-1}$ (Harrison et al., 1996); H is the mixing height of HONO with an assumed value of 200 m (Alicke et al., 2002).

$$\frac{\Delta[\text{HONO}]}{\Delta t} = (P_{\text{emis}} + P_{\text{OH+NO}} + P_{\text{other}}) - (L_{\text{OH+HONO}} + L_{\text{phot}} + L_{\text{dep}}) + F_{\text{v}} + F_{\text{h}}$$
(E3)

$$\begin{split} P_{other} = j(HONO)[HONO] + k_{OH+HONO}[OH][HONO] + \frac{\Delta[HONO]}{\Delta t} \\ + \frac{\nu_{HONO}}{H}[HONO] - k_{OH+NO}[OH][NO] \end{split} \tag{E4}$$

Fig. 4 shows the hourly average HONO budgets during the daytime in the winter-spring and summer campaigns. Seasonal consistencies of HONO sinks but significant seasonal differences of HONO sources were found. During both campaigns, HONO photolysis (Lphot) was the major loss pathway, followed by dry deposition (Ldep) and the negligible homogeneous reaction of R2 ($L_{OH+HONO}$). During noontime (11:00-14:00 LT) in winter-spring, the average loss rate of L_{phot}, L_{dep}, and L_{OH+HONO} was 0.57, 0.07, and 0.01 ppbv·h⁻¹, respectively. During noontime in summer, in comparison, the average loss rate of L_{phot}, L_{dep}, and $L_{OH+HONO}$ was 0.53, 0.04, and 0.01 ppbv·h⁻¹, respectively. Despite the similar HONO sinks, the HONO sources presented significant differences between the two campaigns. In winter-spring, the homogeneous reaction of R3 (P_{OH+NO}) was the dominant HONO source with a mean production rate of 0.43 $ppbv \cdot h^{-1}$ during noontime, while P_{other} was 0.14 ppbv·h⁻¹, accounting for 25% of the HONO production. During noontime in summer, however, the average production rate of P_{OH+NO} was only 0.08 ppbv h^{-1} , whilst P_{other} was up to 0.50 ppbv h^{-1} and accounted for 87% of the HONO production. The strength of Pother during noontime at our site was at the lowest level compared with other studies, whose results ranged from 0.22 to 3.05 $ppbv \cdot h^{-1}$ (see Liu et al. (2019) and references therein). The seasonal variation of Pother in this study was consistent with that at an urban site in Beijing, where the value in summer (3.05 ppbv· h^{-1}) was higher than that in spring (2.63 ppbv· h^{-1}) and winter (1.30 $ppbv\cdot h^{-1}$) (Wang et al., 2017). In comparison, it differed from that at a suburban site in Nanjing where higher values occurred in spring (1.73 ppbv \cdot h⁻¹) and winter (1.15 ppbv \cdot h⁻¹) than in summer (1.00 ppbv \cdot h⁻¹) (Liu et al., 2019). The varying seasonal differences of the HONO budget demonstrate that multi-season observations are required at diverse sites for HONO study.

The elevated strength and proportion of P_{other} in summer suggest the existence of some strongly enhanced HONO source(s) in the summer season. To investigate the potential daytime HONO sources, we summarized the correlations between the daytime (08:00-17:00 LT) P_{other} with relevant trace species and meteorological parameters (see Table 2).



Fig. 4. Hourly average HONO production and loss rates during the daytime in (a) winter-spring and (b) summer.

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Table 2

Correlations of the diurnal Pother against related pollutants and parameters.

Parameter	Total		Winter-Spring		Summer	Summer		BB case		NBB case	
	r	N	r	Ν	r	Ν	r	Ν	r	Ν	
RH	-0.04	989	0.06	455	-0.24	534	0.88	29	0.08	19	
j(NO ₂)	0.24	1002	-0.10	455	0.29	547	-0.11	29	0.89	19	
CO	0.23	1001	0.25	454	0.39	547	0.08	29	0.73	19	
SO ₂	0.22	1000	0.28	453	0.59	547	0.47	29	0.35	19	
NO	0.01	1002	0.19	455	0.09	547	0.74	29	0.63	19	
NO ₂	0.05	1002	0.18	455	0.47	547	0.81	29	0.51	19	
NOz	0.17	1002	-0.02	455	0.57	547	0.51	29	0.36	19	
PM _{2.5}	0.15	968	0.20	439	0.36	529	0.74	29	0.20	19	
PM _{2.5} *NO ₂	0.08	968	0.22	439	0.53	529	0.83	29	0.52	19	
PM _{2.5} *NO _z	0.11	968	0.10	439	0.54	529	0.64	29	0.30	19	
j(NO ₂)*NO ₂	0.09	1002	0.02	455	0.56	547	0.80	29	0.78	19	
j(NO ₂)*NO _z	0.30	1002	-0.14	455	0.61	547	0.38	29	0.67	19	
j(NO ₂)*PM _{2.5}	0.24	968	0.07	439	0.40	529	0.54	29	0.65	19	
j(NO ₂)*NO ₂ *PM _{2.5}	0.08	968	0.08	439	0.57	529	0.82	29	0.75	19	
j(NO ₂)*NO _z *PM _{2.5}	0.18	968	0.00	439	0.56	529	0.58	29	0.51	19	

Note: The daytime period was set as 08:00-17:00 here following section 3.3. The time resolution of all the data used for the correlation calculation is 30 min.



Fig. 5. Diurnal variations of HONO, NO, NO_x, NO_y, O₃, PM_{2.5}, j(HONO), and meteorological parameters during (a) the none biomass burning (NBB) case on June 5th and (b) the biomass burning (BB) case from June 16th to 17th 2017.

The correlation analysis revealed different seasonal characteristics of the potential daytime HONO sources. In winter-spring, P_{other} showed no linear correlations with $j(NO_2)$ -related parameters ($r \le 0.1$), suggesting that the other HONO source in winter-spring had no relationship with the photo-related HONO formation mechanisms. And the weak positive correlations (r < 0.3) between P_{other} with other pollutants give no significant indications of any potential HONO source. In summer, direct emission sources were ruled out first because P_{other} showed no correlation with NO (r = 0.09), which is the tracer of fresh plume emitted by fuel combustion. Some studies have proposed that photo-enhanced heterogeneous reactions of NO_2 and photolysis of nitrate and nitric acid may be important daytime HONO sources (Finlayson-Pitts et al., 2003; Zhou et al., 2003; Ye et al., 2016a). Here we examined the correlations of P_{other} with NO_2 and NO_z , taking NO_z as a proxy of nitrate and nitric acid (highly time resolved measurements of nitrate and nitric acid

were unavailable in this study). P_{other} showed overall moderate correlations with NO₂ (r = 0.47) and NO_z (r = 0.57). When taking solar radiation (j(NO₂) as a proxy) into consideration, the correlations became stronger, with j(NO₂)*NO₂ (r = 0.56) and j(NO₂)*NO_z (r = 0.61). On one hand, the increased correlation indicates the photo-induced heterogeneous formation mechanism of HONO. On another hand, according to the correlation strength, it can be argued that the photolysis of nitrate and nitric acid as well as photo-enhanced heterogeneous conversion of NO₂ should be important HONO sources in the YelRD region. These results agree with a recent study that demonstrated aqueous phase photolysis of particulate nitrate as an important source of HONO (Gen et al., 2019).

3.4. Effects of biomass burning

As June is the harvest month of winter wheat in North China, deteriorated air quality could happen during this period due to the wheat straw burning. As shown in Fig. 2, highly elevated concentrations of HONO, K⁺ in fine particles, PM_{2.5}, and NO_y were observed on June 16th and 17th during which the daily fire maps (see Fig. S1 (https://firms.modaps.eosdis.nasa.gov)) displayed intensive fire points around North China. On June 5th, by contrast, almost no fire point was detected around our study site. Therefore, June 16th-17th was considered as the typical biomass burning (BB) case, and June 5th was considered as the none-biomass burning (NBB) case.

Significant differences in the pollution characteristics were found between the BB case and the NBB case. Fig. 5 presents the diurnal variations of HONO and several relevant pollutants and parameters during the two cases. The average HONO concentration was about three times higher in the BB case (0.41 \pm 0.42 ppbv) than in the NBB case (0.14 \pm 0.07 ppby). In the NBB case, the maximum HONO concentration occurred in the late night and was about 0.45 ppby. In the BB case, the HONO concentration increased unusually in the early morning, and the peak went across the whole morning with a substantially elevated maximum of 1.60 ppbv on June 16th and 2.00 ppbv on 17th, which was also the maximum value observed throughout the summer campaign. While the average concentrations of NO_x were the same in both cases (both of 5.4 \pm 3.1 ppbv), the average concentration of NOv was over two times higher in the BB case (18.6 \pm 8.2 ppbv) than in the NBB case (8.0 \pm 3.3 ppbv), indicating that the BB plume was much chemically aged than the NBB plume. The average concentration of PM2.5 in the BB case (84.9 \pm 37.3 $\mu g {\cdot} m^{-3}$) was about six times higher than that in the NBB case (14.2 \pm 7.3 µg·m⁻³), implying much heavier particulate pollution, and as detailed below, aerosol played a more important role in the heterogeneous formation of HONO during the BB case. The O3 concentration doubled in the BB case (96 \pm 42 ppbv), indicating more serious photochemistry pollution.

The strength of the other HONO sources (P_{other}), as shown in Fig. S2, increased substantially in the BB case. To explore the potential HONO source and the influence of BB on HONO chemistry during this episode, Table 2 lists the correlations of P_{other} against various pollutants and parameters in the BB and NBB cases. In both cases, differing from the

whole summer campaign, Pother presented moderate correlations with NO, indicating a considerable contribution of direct HONO emissions. In the NBB case, Pother was highly correlated with j(NO2), and all the correlations with NO2, NO2, and PM2.5 increased significantly after considering the solar radiation effect. In the BB case, however, no correlation of Pother with j(NO2) was found, and all the relationships with NO₂, NO₂, and PM_{2.5} weakened more or less when considering the solar radiation. This phenomenon implies that the role of the photo-related heterogeneous formation mechanism of HONO was suppressed during the BB period, which should be ascribed to the dimming effect of high aerosol loading from BB emissions. Compared with the NBB case, the correlation of Pother with PM2.5 drastically enhanced in the BB case, implying significantly increased effects of particles. As the indicator of particle loading, PM_{2.5} could contribute to HONO chemistry through either producing HONO via the photolysis of particulate nitrate or providing the aerosol surface for HONO heterogeneous formation. Because of the weaker relationship of Pother with j(NO2)*PM2.5 compared with PM_{2.5}, and the stronger correlation of Pother with PM_{2.5}*NO₂ (or PM_{2.5}*NO_z) compared with NO₂ (or NO_z), we argue that particles may play a major role in HONO formation by providing heterogeneous reaction surface during the BB episode. It should be noted that such correlation analysis may be inherently subject to some uncertainties, and more detailed process studies are needed to better understand the HONO formation mechanisms in the BB plumes.

3.5. Contribution to the OH source

Photolysis of HONO can make a considerable contribution to the OH radical during the daytime. To evaluate the net OH production rate from HONO photolysis, equation (E5) was used for the calculation considering the homogeneous reactions of R1 to R3. For comparison, the OH production rate from the photolysis of O_3 , another significant precursor of OH, was calculated using equation (E6) based on the reactions of R14 to R16. $P_{OH}(HONO)$ is the net OH production rate from HONO photolysis, and $P_{OH}(O_3)$ is the OH production rate from O_3 photolysis. The reaction rate constants were taken from Atkinson et al. (2004).

$$O_3 + h\nu \rightarrow O(^1D) + O_2 \quad (\lambda < 340 \text{ nm}) \tag{R14}$$



Fig. 6. Production rates of OH from the photolysis of HONO and O₃ in (a) winter-spring, (b) summer, (c) the non-biomass burning (NBB) case in summer, and (d) the biomass burning (BB) case in summer.

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$$O(^{1}D) + H_{2}O \rightarrow 2OH$$
(R15)

$$O(^{1}D) + M \rightarrow O(^{3}P) + M \quad (M = N_{2} \text{ or } O_{2})$$
(R16)

 $P_{OH}(HONO) = j(HONO)[HONO] - k_{OH+HONO}[HONO][OH]$

$$-k_{OH+NO}[NO][OH]$$
(E5)

$$P_{OH}(O_3) = j(O^1D)[O_3] \cdot \frac{2k_{O(^1D)+H_2O}[H_2O]}{k_{O(^1D)+H_2O}[H_2O] + k_{O(^1D)+N_2}[N_2] + k_{O(^1D)+O_2}[O_2]}$$
(F6)

Fig. 6(a) and (b) present the average diurnal OH production rates from photolysis of HONO and O₃ in winter-spring and summer, respectively. In winter-spring, HONO photolysis made a much greater contribution to OH than O₃ throughout the daytime, with the peak P_{OH} (HONO) of 0.61 ppbv h⁻¹ in the late morning and the peak P_{OH} (O₃) of 0.15 ppbv $\cdot h^{-1}$ around the noon. In summer, while HONO contributed more OH than O_3 did before the late morning with the peak $P_{OH}(HONO)$ of 0.73 ppbv·h⁻¹, O₃ photolysis became the more significant OH source during the rest of the daytime with the peak $P_{OH}(O_3)$ of 2.22 ppbv·h⁻¹ around the noon. On average, during the daytime of 08:00-17:00 LT, P_{OH} (HONO) accounted for 78% of P_{OH} (HONO + O₃) in winter-spring and 26% in summer. The seasonal difference in the relative contribution of HONO and O₃ to OH was the combined effect of the seasonal changes in the concentrations of HONO and O3 and in the photochemistry. Specifically, HONO showed higher concentrations in winter-spring than in summer, whilst O₃ showed much higher levels in summer than in winter-spring, owing to the seasonal variation of sunlight and hence photochemical processes. To sum up, HONO plays a significant role in OH formation and hence affects the atmospheric oxidation capacity in both seasons.

Fig. 6(c) and (d) display the diurnal OH production rates from HONO and O₃ photolysis in the NBB case and BB case, respectively. Compared with the NBB case, P_{OH}(HONO) in the BB case increased greatly, especially in the morning. P_{OH}(HONO) in the BB case was about five times larger than that in the NBB case with a peak of 5.0 ppbv \cdot h⁻¹ in the late morning, and $P_{OH}(O_3)$ in the BB case reached the peak of 3.7 ppbv·h⁻¹ around noon with the average value of over two times higher than that in the NBB case. The differences of POH(HONO) and POH(O3) are consistent with the concentration differences of HONO and O3 between the two cases. In terms of relative contribution, while POH(HONO) was larger than P_{OH}(O₃) only before 08:00 in the NBB case, P_{OH}(HONO) in the BB case was larger than POH(O3) before 10:00 and was over three times higher than POH(O3) during 06:00-10:00. These results indicate the significant impacts of BB on HONO and atmospheric oxidation capacity, and BB should be taken into consideration in atmospheric modelling study and pollution control policy-making.

4. Summary and conclusions

The paper presents the HONO measured during the winter-spring and summer in 2017 at a rural coastal site in North China and demonstrates the significant seasonal variations in HONO pollution and chemistry, including the pollution characteristics, nocturnal C_{HONO}, daytime budgets, and the impacts on atmospheric oxidation capacity. More severe air pollution was observed in winter-spring than in summer, with the mean HONO mixing ratio of 0.26 ± 0.28 ppbv in winter-spring and 0.17 ± 0.19 ppbv in summer. The HONO/NO_x ratio in summer was about twice of that in winter-spring, and it reached a peak at noon in summer, which indicated the photo-enhanced heterogeneous formation of HONO. The mean nocturnal C_{HONO} in summer was over twice of that in winter-spring. Budget analysis indicated a much stronger daytime P_{other} in summer (0.14-0.56 ppbv·h⁻¹) than in winter-spring (0.02-0.23 ppbv·h⁻¹), and the enhanced photo-induced HONO formation mechanisms were suggested through correlation analysis. Photolysis of nitrate

and nitric acid may be the most significant HONO sources, followed by the comparable photo-enhanced heterogeneous conversion of NO₂. HONO photolysis was the major contributor to OH in winter-spring and also a significant OH source in summer. Biomass burning was found to make a significant contribution to HONO through direct HONO emission and aerosol release, which provided HONO precursors and heterogeneous reaction surface. Biomass burning affected HONO chemistry by enhancing the effects of aerosol and weakening the role of photolysis on HONO formation. Besides, HONO photolysis could be enhanced to be the major OH contributor in the morning in the BB case. Therefore, biomass burning should be taken into consideration in HONO study and in making regional pollution control policy, especially during the harvest seasons.

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Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

CRediT authorship contribution statement

Rongrong Gu: Formal analysis, Writing - original draft. Penggang Zheng: Investigation, Data curation. Tianshu Chen: Investigation, Data curation. Can Dong: Writing - review & editing. Ya'nan Wang: Investigation, Methodology. Yiming Liu: Investigation, Methodology. Yuhong Liu: Investigation. Yuanyuan Luo: Investigation, Data curation. Guangxuan Han: Resources. Xinfeng Wang: Investigation. Xuehua Zhou: Investigation. Tao Wang: Supervision, Writing - review & editing. Wenxing Wang: Supervision, Resources. Likun Xue: Conceptualization, Funding acquisition, Project administration, Writing - review & editing.

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Appendix A. Supplementary data

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