# Mutual promotion between aerosol particle liquid water and particulate nitrate enhancement leads to severe nitrate-dominated particulate matter pollution and low visibility

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Abstract. As has been the case in North America and Western Europe, the  $SO_2$  emissions substantially reduced in North China Plain (NCP) in recent years. A dichotomy of reductions in SO<sub>2</sub> and NO<sub>x</sub> 40 concentrations result in the frequent occurrences of nitrate (pNO<sub>3</sub><sup>-</sup>)-dominated particulate matter pollution over NCP. In this study, we observed a polluted episode with the particulate nitrate mass fraction in non-refractory PM<sub>1</sub> (NR-PM<sub>1</sub>) up to 44% during wintertime in Beijing. Based on this typical  $pNO_3$ -dominated haze event, the linkage between aerosol water uptake and  $pNO_3$ -enhancement, further impacting on visibility degradation, have been investigated based on field observations and theoretical 45 calculations. During haze development, as ambient relative humidity (RH) increased from ~10% up to 70%, the aerosol particle liquid water increased from ~1  $\mu$ g/m<sup>3</sup> at the beginning to ~75  $\mu$ g/m<sup>3</sup> at the fully-developed haze period. The aerosol liquid water further increased the aerosol surface area and volume, enhancing the condensational loss of  $N_2O_5$  over particles. From the beginning to the fullydeveloped haze, the condensational loss of N<sub>2</sub>O<sub>5</sub> increased by a factor of 20 when only considering 50 aerosol surface area and volume of dry particles, while increasing by a factor of 25 considering extra surface area and volume due to water uptake. Furthermore, aerosol liquid water favored the thermodynamic equilibrium of HNO<sub>3</sub> into the particle phase under the supersaturated HNO<sub>3</sub> and NH<sub>3</sub> in the atmosphere. All above results demonstrated that the  $pNO_3^{-1}$  is enhanced by aerosol water uptake with elevated ambient RH during haze development, in turn, facilitating the aerosol taking up water due to 55 the hygroscopicity of particulate nitrate salt. Such mutual promotion between aerosol particle liquid water and particulate nitrate enhancement can rapidly degrade air quality and halve visibility within one day. Reduction of nitrogen-containing gaseous precursors, e.g., by control of traffic emissions, is essential in mitigating severe haze events in NCP.

## 60 1 Introduction

Aerosol particle hygroscopicity plays an important role in air quality deterioration and cloud formation (Yu, 2009;Fitzgerald, 1973;Kreidenweis and Asa-Awuku, 2014;Wang and Chen, 2019;McFiggans et al., 2006) and can also directly influence aerosol measurements (Chen et al., 2018a). In atmospheric environments influenced by anthropogenic activities, particulate secondary inorganic compounds are 65 often dominated by particulate sulfate and nitrate (Heintzenberg, 1989), which originate from the oxidation of sulfur dioxide (SO<sub>2</sub>) and nitrogen oxides (NO<sub>x</sub>) via multiple chemical pathways (Calvert et al., 1985; Cheng et al., 2016; Wang et al., 2016; Gen et al., 2019a, b). The abundance of secondary inorganic components is one of the most important factors determining particle hygroscopicity (Swietlicki et al., 2008), thereby governing the aerosol liquid water content under ambient moist conditions. Increased aerosol particle liquid water could accelerate secondary inorganic and organic 70 aerosol formation by decreasing the kinetic limitation of mass transfer of gaseous precursors and providing more medium for multiphase reactions (Mozurkewich and Calvert, 1988;Cheng et al., 2016; Wang et al., 2016; Ervens et al., 2011; Kolb et al., 2010).

Sulfuric acid (H<sub>2</sub>SO<sub>4</sub>) is formed from the oxidation of SO<sub>2</sub> via gaseous and multiphase reactions. H<sub>2</sub>SO<sub>4</sub>

75 is subsequently fully or partly neutralized by gaseous NH<sub>3</sub> taken up on particles, resulting in the formation of (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> and / or NH<sub>4</sub>HSO<sub>4</sub>. Any remaining NH<sub>3</sub> is available to neutralize HNO<sub>3</sub> to form particulate NH<sub>4</sub>NO<sub>3</sub> (Seinfeld. and Pandis., 2006) (and further excess NH<sub>3</sub> can neutralize any available HCl to form particulate NH<sub>4</sub>Cl). Over the past several decades, substantial efforts have reduced emissions of both SO<sub>2</sub> and NO<sub>x</sub> improving the local and regional air quality all over the world.

- For example, SO<sub>2</sub> and NO<sub>x</sub> emissions were reduced by 82% and 54% in the majority of European Environment Agency member countries between 1990 and 2016 (<u>https://www.eea.europa.eu/data-and-maps/indicators/main-anthropogenic-air-pollutant-emission\_s/assessment-4</u>). In consequence, an increasing trend of NO<sub>3</sub><sup>-/</sup>SO<sub>4</sub><sup>2-</sup> molar ratio was observed in long-term measurements at Leipzig, Germany (Spindler et al., 2004) and at some other European sites from the European Monitoring and Evaluation Programme (EMEP) (Putaud et al., 2004).
  - In the recent years, China has also managed to reduce SO<sub>2</sub> emissions by 75% during 2007~2015 (Li et al., 2017a) and declined by ~15.1% per year during 2013~2017 (Vu et al., 2019), whereas NO<sub>x</sub> emissions declined only by ~10% between 2011 and 2015 (de Foy et al., 2016) and by ~ 4.3% per year during 2013~2017 (Vu et al., 2019). The strict emission control reduced the PM<sub>2.5</sub> mass concentration and the corresponding chemical components in China significantly (Vu et al., 2019). The annual mean PM<sub>2.5</sub> mass loading decreased by 39.6% during 2013~2017 in Beijing-Tianjin-Hebei region, and the

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 $SO_4^{2-}$  and  $NO_3^{-}$  mass concentrations in the PM<sub>2.5</sub> declined by 40% and 34% respectively during 2015~2017 in Beijing (Vu et al., 2019). However, NH<sub>3</sub> emissions have been observed by satellites to increase by ~30% from 2008 to 2016 over the North China Plain (NCP) (Liu et al., 2018). The faster

95 reduction rate of SO<sub>2</sub> than NO<sub>x</sub> emissions in conjunction with elevated NH<sub>3</sub> level, made it reasonable of switching dominant inorganic component in fine aerosol particles from sulfate to nitrate in the recent years similar like European countries (Sun et al., 2015;Hu et al., 2017;Hu et al., 2016;Wu et al., 2018;Guo et al., 2014;Huang et al., 2014;Huang et al., 2010;Ge et al., 2017;Xu et al., 2019a;Xie et al., 2019;Li et al., 2018). Field measurements in Beijing show that annually averaged NO<sub>3</sub><sup>-</sup>/SO<sub>4</sub><sup>2-</sup> molar ratio of NR-PM<sub>1</sub> (non-refractory PM<sub>1</sub>) in 2012 (1.3~1.8) (Sun et al., 2015) has significantly increased

compared to that in 2008 (0.9~1.5) (Zhang et al., 2013). Comparably, the NO<sub>3</sub><sup>-</sup>/SO<sub>4</sub><sup>2-</sup> molar ratio of PM<sub>2.5</sub> in Beijing increased substantially, from 1.5 before 2013 to 3.33 in 2017 (Xu et al., 2019a).

Over the NCP region, heavy haze events are typically associated with enhanced ambient RH levels. This leads to an increased aerosol liquid water content (Wu et al., 2018), which will enhance the particulate nitrate formation by increasing the reactive uptake of precursors and the thermodynamic 105 equilibrium of ammonium nitrate (Cheng et al., 2016; Wang et al., 2016; Wang et al., 2017; Yun et al., 2018; Yue et al., 2019). To date, few studies reported aerosol liquid water content over NCP region (Wang et al., 2018;Bian et al., 2014;Cheng et al., 2016;Wu et al., 2018;Ge et al., 2019). However, the observational and theoretical analysis of the relationship between particulate nitrate enhancement and associated liquid water during haze events in China has been infrequently reported (Wu et al., 2018).

In this study, a self-amplification effect between particulate nitrate and liquid water is demonstrated by examining a nitrate-dominated fine particle Beijing pollution episode. The facilitation of particulate nitrate enhancement by abundant aerosol liquid water is subsequently theoretically explored through the impacts of liquid water on thermodynamic equilibrium and heterogeneous reactions. Finally, the

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corresponding impacts on light extinction coefficient, and visibility degradation are estimated. These 115 results improve our quantitative understanding of the development of haze events over the NCP and on formulating emission reduction strategies, as well as may also provide insights for other polluted regions.

## 2 Measurements and Methods

#### 120 **2.1 Location and instrumentation**

Measurements were conducted within the framework of the BEST-ONE (Beijing winter finE particle STudy- Oxidation, Nucleation, and light Extinctions) field campaign from January 1 to March 5, 2016, at the Huairou site (40.42°N, 116.69°E), located in a rural environment, north of Beijing, China. Detailed information about the sampling site was described in Tan et al. (2018). A weather station (Met

- 125 one Instrument Inc., USA) was performed to measure meteorological parameters (ambient RH, temperature, wind speed, wind direction) and detailed aerosol particle physical and chemical properties were recorded using a suite of state-of-the-science instrumentation. Hygroscopic growth factor (HGF) of sub-micrometer aerosol particles was measured using a Hygroscopicity-Tandem Differential Mobility Analyzer (H-TDMA, TROPOS, Germany) (Wu et al., 2011;Massling et al., 2011;Wang et al.,
- 130 2018;Wu et al., 2016;Liu et al., 1978) and data retrieval followed the TDMA<sub>inv</sub> method in Gysel et al. (2009). The hygroscopicity parameter (κ) was estimated using by the κ-Köhler approach (Petters and Kreidenweis, 2007;Köhler, 1936). Size-resolved NR-PM<sub>1</sub> was recorded by an Aerodyne High-Resolution Time-of-Flight Aerosol Mass Spectrometry (HR-ToF-AMS, Aerodyne Research, Inc., USA) (DeCarlo et al., 2006). Regular calibration procedures followed as reported in Jayne et al. (2000) and
- Jimenez et al. (2003) and composition dependent correction followed as in Middlebrook et al. (2012). Gaseous HNO<sub>3</sub> and NH<sub>3</sub> were measured using Gas-Aerosol Collector (GAC) coupled with Ion Chromatography (IC) (Dong et al., 2012). Mass concentration of equivalent black carbon in aerosol particles (Petzold et al., 2013) was recorded by Multi Angle Absorption Photometer (MAAP, Model

5012, Thermo Fisher Scientific, USA) with a laser wavelength of 670 nm (Petzold and Schönlinner,
2004). Furthermore, particle number size distribution (PNSD) in the size range of 3 nm~10 µm was measured using a Mobility Particle Size Spectrometer (MPSS, Model 3776+3085 3775+3081, TSI, USA), following the recommendations described in Wiedensohler et al. (2012) and an Aerodynamic Particle Size Spectrometer (APS, Model 3021, TSI, USA) (Wu et al., 2008;Pfeifer et al., 2016). Detailed description on H-TDMA, HR-ToF-AMS and GAC-IC can be found in the supporting information.

#### 2.2 Estimation of aerosol particle liquid water

Given the absence of direct liquid water measurement, size-resolved liquid water was calculated using the corresponding HGFs measured at RH=90% (50, 100, 150, 250, 350 nm in stokes diameter), PNSD data (3 nm~10 µm) and meteorological parameters (RH, T), following the method proposed in Bian et al. (2014), referred to below as H-TDMA-derived liquid water. Briefly, the measured PNSD with 57 size bins were fitted using a four-mode lognormal distribution. The classification of four modes and the fitting results are shown in Table S1 and Figure S4. Good agreement between measured values and fitted PNSD was achieved, which indicates the reliability of the four-mode lognormal fitting method. Based on four-mode lognormal fitting results, the particle number size distribution and number fractions

of each mode can be obtained. It has been assumed that particles from the same mode have constant particle hygroscopicity ( $\kappa$ ). Under the assumption of constant particle hygroscopicity in each mode (shown in Table S1), the  $\kappa$  values for each mode ( $\kappa_1$ ,  $\kappa_2$ ,  $\kappa_3$ ) can be calculated by Eq. [1] from the known number fraction of fitted four modes and the  $\kappa$  values of measured particle size from H-TDMA measurement.

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$$\kappa = \sum_{i=1}^{4} \kappa_i f_i$$
 [1]

Here,  $\kappa_i$  and  $f_i$  represent the  $\kappa$  value and the particle number fraction of the i mode. Then, the calculated  $\kappa$  values for each mode and the derived number fraction of each size bin were used to obtain the  $\kappa$  distribution for each size bin. Figure S5 shows the comparison of calculated sized-resolved  $\kappa$  distribution and the  $\kappa$  measured by H-TDMA, the good agreement showed the reliability of the method.

165 Then, based on  $\kappa$ -Kölher theory (Petters and Kreidenweis, 2007;Köhler, 1936), the size-resolved *HGFs* at ambient RH were calculated. Finally, liquid water of size-resolved particles can be derived by calculating the differentials between the dry and wet PNSD of aerosol particles in Eq. [2]:

Liquid water = 
$$\frac{\pi}{6} N_j D_{p,j}^3 \left( HGF(D_p, RH)^3 - 1 \right) * \rho_w$$
 [2]

where j represents the bin number of measured PNSD,  $N_j$  and  $D_{p,j}$  represent the number concentration and the diameter of dry particles of the j<sup>th</sup> bin, respectively, while, *HGF* and  $\rho_w$ , are the hygroscopic growth factor of aerosol particles and water density (1 g/cm<sup>3</sup>), respectively.

#### 2.3 Condensation rate of trace gases

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The condensation rate (k) of trace gases (dinitrogen pentoxide, N<sub>2</sub>O<sub>5</sub>, referred as k\_N<sub>2</sub>O<sub>5</sub>) was calculated by the method of Schwartz (1986), shown in Eq. [3]. In order to illustrate the influences of the dry and wet PNSD due to water uptake on condensation rate of gases, the PNSD of the dry and wet

particles (obtained by applying the HGF estimated from H-TDMA-derived liquid water method) were used.

$$k = \frac{4\pi}{3} \int_0^\infty \left(\frac{r^2}{3D_g} + \frac{4r}{3C_g\gamma}\right)^{-1} r^3 \frac{dN}{d\log r} \, d\log r \quad [3]$$
$$C_g = \sqrt{\frac{3RT}{M}} \quad [4]$$

- 180 Where, *r* represents radius of the particles,  $D_g$  represents the binary diffusion coefficient evaluated following Maitland (1981) (1.18\*e<sup>-5</sup> m<sup>2</sup>/s).  $C_g$  is the kinetic velocity of the gas molecules, calculated in Eq. [4]. Here, R and M are the ideal gas constant (8.314 kg.m<sup>2</sup>/mol/K/s<sup>2</sup>) and molar mass of the gas, respectively while T represents the ambient temperature. dN/dlogr is the number size distribution and  $\gamma$ is the uptake coefficient of the gas.
- The uptake coefficient of N<sub>2</sub>O<sub>5</sub> was estimated following the method proposed in Chen et al. (2018b) and Chang et al. (2016) and references therein. The influences of RH, temperature, multiple inorganic particle compositions, secondary organic aerosol (SOA) and primary organic aerosol (POA) are considered. The uptake suppression effect of N<sub>2</sub>O<sub>5</sub> due to the presence of SOA was considered following the method in Anttila et al. (2006). Based on our source apportionment using Positive matrix factorization (SoFi tool, ME2, Francesco Canonaco, PSI), two oxygenated organic aerosol factors (OOA), usually interpreted as SOA, and three POA factors were determined. The fraction of SOA in the total organic aerosol (OA) was 60%~90% during the observed period, which is quite consistent with the results of a previous study in Beijing (Huang et al., 2014). Hence, 75% was used as the ratio of SOA/OA in our model to estimate the suppression effect of SOA on the uptake of N<sub>2</sub>O<sub>5</sub> following the

work of Anttila et al. (2006). The reaction of chloride with N<sub>2</sub>O<sub>5</sub> was not considered in this study due to its limited mass concentration (on average 5% of the PM<sub>1</sub> mass concentration during the marked haze period), which could cause minor uncertainty in the k\_N<sub>2</sub>O<sub>5</sub> calculation. The detailed information regarding the estimation  $\gamma_{N2O5}$  is given in Chen et al. (2018b), and influences of different chemical components on  $\gamma_{N2O5}$  is summarized in the Table 1 of Chen et al. (2018b).

#### 200 2.4 Equilibrium of NH4NO3

The equilibrium dissociation constant of  $NH_4NO_3$  (*Kp*) under dry conditions was calculated as a function of ambient temperature (Seinfeld. and Pandis., 2006) in the following Eq. [5].

$$lnK_p = 84.6 - \frac{24220}{T} - 6.1ln\left(\frac{T}{298}\right)$$
 [5]

Taking into account the associated liquid water, the equilibrium vapor pressure of HNO<sub>3</sub> and NH<sub>3</sub> was calculated by employing the Extended-Aerosol Inorganic Model (E-AIM) Model II H<sup>+</sup> - NH<sub>4</sub><sup>+</sup> - SO<sub>4</sub><sup>2-</sup> -NO<sub>3</sub><sup>-</sup> - H<sub>2</sub>O (Clegg et al., 1998) using HR-ToF-AMS data, NH<sub>3</sub> from GAC-IC, and meteorological parameters (RH, T). In this calculation, a simplified ion pairing scheme was performed to ensure the ion balance of the input chemical composition following the method in Gysel et al. (2007).

#### 2.5 Light extinction coefficient and visibility calculation

210 Size-resolved chemical composition of the NR-PM<sub>1</sub> from HR-ToF-AMS, mass concentration of equivalent black carbon from MAAP, PNSD data and the H-TDMA-derived liquid water were used to calculate light extinction coefficient (including light absorption and scattering) and visibility degradation of size-resolved particles by the Mie scattering theory described in Barnard et al. (2010).

Here, size-resolved equivalent black carbon mass concentration was inferred by the particle mass size
distribution measurement from single particle soot photometer in PKUERS. The method of redistribution of liquid water and HR-ToF-AMS data has been described in the supporting information (Text S1, HR-ToF-AMS introduction section). Thus, with the re-distributed datasets as the input of the Mie scattering theory, the light extinction coefficient for atmospheric particles in the absence and presence of liquid water with a size range of 100~2500 nm in stokes diameter can be derived. Due to
lack of measurements on aerosol particle morphology and mixing state, we assume particles are spherical as described in Barnard et al. (2010). To perform Mie calculation, the complex reflective index of each component is given in Table 1 of Barnard et al. (2010) and references therein. This method shows good agreement with measurements in Mexico City and is consistent as the regional atmospheric chemistry model WRF-Chem. Here, Ext\_550nm\_wet and Ext\_550nm\_dry represent the

225 calculated light extinction coefficient for particles in the presence and absence of liquid water at an incident light wavelength of 550 nm. The corresponding visibility degradation (VIS) for dry/wet particles was calculated from the light extinction coefficient following the Koschmieder Eq. [6].

$$VIS = \frac{3.912}{Ext_{-}5 \ 5 \ nm}$$
[6]

# **3** Results and Discussion

#### 230 **3.1 Nitrate-dominated fine particulate matter pollution**

Figure 1 illustrates a summary of chemical composition of NR-PM<sub>1</sub>, ambient RH, size distribution and total aerosol particle liquid water, size distribution and total aerosol surface area concentration during

the period of February 29 to March 5, 2016 in the BEST-ONE campaign. During this period, polluted episodes occurred under stagnant meteorological conditions with low wind speed (Figure S6) and

- 235 elevated ambient RH (Figure 1a). As marked 'haze period' in Figure 1, an obvious increase of NR-PM<sub>1</sub> was observed. The secondary inorganic components (sulfate, nitrate and ammonium) were dominant components of the NR-PM<sub>1</sub>, accounting for up to 73% during the 'haze period'. Particularly, nitrate was the major contributor of the secondary inorganic components and accounted for up to ~44% of NR-PM<sub>1</sub> mass, while sulfate contributed for ~12% on average.
- In the recent decade, severe haze events with high aerosol mass loading occurred frequently in Beijing during wintertime (Hu et al., 2016;Hu et al., 2017;Sun et al., 2014;Sun et al., 2015). To mitigate the air pollution, the Beijing government implemented strict emission controls. The total mass loading of particulate matter has reduced substantially in the recent years (http://sthjj.beijing.gov.cn/). With decreasing in PM mass concentration, the mass fraction of particulate nitrate during these haze events in Beijing enhanced substantially. In 2014, the highest fraction of nitrate in PM<sub>1</sub> was reported as ~20% and increased to ~35% in 2016 (Xu et al., 2019b), which is comparable to the ratio (44%) in this study. The

particulate nitrate became more dominant in secondary inorganic compounds other than particulate sulfate with the air quality improvement over NCP.

As one of the main hydrophilic compounds in atmospheric aerosol particles, the ability of water uptake

is comparable between deliquescent (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> and NH<sub>4</sub>NO<sub>3</sub> particles with same sizes and ambient RH (Kreidenweis and Asa-Awuku, 2014;Wu et al., 2016), (http://umansysprop.seaes.manchester.ac.uk/). However, compared to (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, NH<sub>4</sub>NO<sub>3</sub> particles have a lower deliquescence RH (62%, 298 K) than (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> (80%, 298 K) (Kreidenweis and Asa-Awuku, 2014) and easily liquefy (Li et al.,

2017b). In addition, NH<sub>4</sub>NO<sub>3</sub> particles are semi-volatile, the co-condensation of semi-volatile 255 compounds and water (Topping et al., 2013;Hu et al., 2018) could be significant. Therefore, the switching from sulfate-dominated to nitrate-dominated aerosol chemistry may impact on aerosol water uptake. The interaction between aerosol particle liquid water and particulate nitrate formation and visibility degradation should be reconsidered.

#### 3.2 Mutual promotion between liquid water and particulate nitrate enhancement

Lu et al. (2019) conducted a box model to calculate the potential particulate nitrate formation during the same investigated period of the BEST-ONE project. They found out that HNO<sub>3</sub> from daytime photooxidation of NO<sub>2</sub> was the major source of the particulate nitrate (>75%), whereas the contribution of N<sub>2</sub>O<sub>5</sub> pathway was lower than 25% (Lu et al., 2019). In the following discussion, the enhancement of particulate nitrate during the 'haze period' is elucidated by theoretical calculations of condensational loss rate of N<sub>2</sub>O<sub>5</sub>, and the thermodynamic equilibrium of NH<sub>4</sub>NO<sub>3</sub> and HNO<sub>3</sub>. In particular, the role of aerosol water uptake in particulate nitrate formation is comprehensively investigated.

N<sub>2</sub>O<sub>5</sub> is an important gaseous precursor for particulate nitrate formation via its hydrolysis to form HNO<sub>3</sub> during nighttime (Brown et al., 2006). Liquid water can enhance aerosol surface areas and volumes, thereby increasing the available heterogeneous reacting medium. Across the development of 'haze period', the estimated liquid water increased from ~1 µg/m<sup>3</sup> at the beginning (March 2, 14:00~18:00 p.m.) to ~75 µg/m<sup>3</sup> when the haze was fully developed (March 4, 4:00~8:00 a.m.). The total surface area and volume concentrations of particles were increased by the liquid water by 2~3% at the beginning and by up to ~25 and ~40% in the fully-developed haze compared to the 'dry' values,

respectively (see Figure S7 and S8). Additionally, from the beginning to the fully-developed haze, the 275 uptake coefficient of N<sub>2</sub>O<sub>5</sub> was enhanced by a factor of 9 from 0.002 to 0.018, and the k\_N<sub>2</sub>O<sub>5</sub> increased by a factor of 20 (dry particles); while, considering the increased particle surface area and volume due to water uptake, the respective value of enhanced k\_N<sub>2</sub>O<sub>5</sub> was by a factor of 25 (Figure 2a). Apart from providing extra reacting medium, the abundant liquid water can liquefy the aerosol particles and may reduce any kinetic limitation of mass transfer for reactive gases (Koop et al., 2011;Shiraiwa et al., 2011) and impact thermodynamic equilibrium of semi-volatile compounds (Kulmala et al., 1993;Topping et al., 2013) to contribute to secondary aerosol formation. Our previous study provided the observational evidence that particles may have transitioned from the solid phase to the liquid phase

as RH increased from 20% to 60% during wintertime in Beijing (Liu et al., 2017). In this study, the ambient RH increased from ~10% up to 70% during the haze period, suggesting a likely transition of particles from the solid to liquid phase. Such phase transition may facilitate particulate nitrate formation by increasing diffusion coefficients of dissolved precursors.

To illustrate the facilitation of particulate nitrate enhancement from HNO<sub>3</sub> in the presence of liquid water, we performed the theoretical calculation of equilibrium between particulate NH<sub>4</sub>NO<sub>3</sub> and gaseous NH<sub>3</sub> and HNO<sub>3</sub> under dry and ambient conditions, respectively. The dissociation constant of NH<sub>4</sub>NO<sub>3</sub> (*Kp*) in dry condition was calculated using Eq. [5] without considering the influence of the liquid water. As shown in Figure 3, the equilibrium *Kp* in the dry condition ranged from 0.06 (275.3 K) to 4.61 (291.5 K) ppb<sup>2</sup> during the 'haze period'. Taking account of the aerosol liquid water, the equilibrium vapor pressure of HNO<sub>3</sub> and NH<sub>3</sub> over particles was calculated by E-AIM Model II (www.aim.env.uea.ac.uk). Note that this calculation assumes negligible interaction between dissolved

organic components and the activity of  $NO_3^-$ . In the presence of aerosol associated water, the product of 295 equilibrium vapor pressure of NH<sub>3</sub> and HNO<sub>3</sub> calculated from E-AIM was 10~60% lower than the equilibrium Kp in the dry condition during the marked 'haze period'. This means, the presence of aerosol liquid water changed the equilibrium and would favor the particulate nitrate enhancement. However, the aerosol particles didn't reach the equilibrium between particulate NH<sub>4</sub>NO<sub>3</sub> and the gases  $(NH_3 + HNO_3)$  during the investigated period, as the measured product of the NH<sub>3</sub> and HNO<sub>3</sub> partial 300 pressure  $(2.55 - 9.63 \text{ ppb}^2)$  was supersaturated compared to the equilibrium values in both dry and deliquescent particles. In this case, the partitioning of gaseous NH<sub>3</sub> and HNO<sub>3</sub> in the atmosphere into the particle phase could be accelerated and leaded particulate nitrate enhancement as increasing of ambient RH. Owing to the nature of highly hydrophilic, the increased ammonium nitrate mass fraction leads to further water uptake. Such a mutual promotion of particulate nitrate and aerosol liquid water 305 enhancement becomes more pronounced with the increasing pollution throughout the haze event owing to the simultaneously increasing ambient RH. Consistently, a significant co-increase of particulate nitrate and aerosol liquid water was observed during haze development as shown in Figure 4. At first, a steep increase of particulate nitrate in total nitrate mass ratio (from ~12% to ~98%) was observed as the aerosol liquid water enhanced up to  $\sim 20 \ \mu g/m^3$ . And then, the particulate nitrate mass kept increasing 310 with further increase of aerosol liquid water. We observed that,  $\sim 98\%$  of nitrate was present as particle phase when aerosol liquid water was higher than ~20  $\mu$ g/m<sup>3</sup>. The function between the particulate nitrate fraction in the total nitrate is given in Figure 4. It is worth noting that N<sub>2</sub>O<sub>5</sub> hydrolysis during nighttime can contribute extra HNO<sub>3</sub> in the wet denuding method within GAC-IC system. This effect explains the slightly underestimation of the particulate fraction during nighttime when aerosol liquid 315

water is less than 10  $\mu$ g/m<sup>3</sup> (Figure 4). However, the general consistency of this function between daytime and the nighttime (Figure 4) suggests a negligible influence of N<sub>2</sub>O<sub>5</sub> interference on our analysis during the investigated period.

- Except for aerosol liquid water, aerosol pH is also an important factor on the particulate nitrate 320 formation, higher pH is favorable for the equilibrium of HNO<sub>3</sub> into the particle phase (Nah et al., 2018). pH of the fine aerosol particles was calculated by ISORROPIA II (Fountoukis and Nenes, 2007) during the investigated period. The model was running in 'forward mode' with chemical composition of NR-PM<sub>1</sub> (NO<sub>3</sub><sup>-</sup>, SO<sub>4</sub><sup>2-</sup>, Cl<sup>-</sup>, NH<sub>4</sub><sup>+</sup>) and gas precursors (HNO<sub>3</sub>, HCl, NH<sub>3</sub>) by GAC-IC as inputs. And the model was running in 'metastable mode', assuming no solid existed in the system. Generally, the fine
- aerosol particles became more acidic with pH dropping from ~8 down to ~4 when NR-PM<sub>1</sub> mass concentration increased from ~12  $\mu$ g/m<sup>3</sup> up to >300  $\mu$ g/m<sup>3</sup> as shown in Figure 5 and Figure 6. This declining trend of pH is not favorable for the HNO<sub>3</sub> partitioning into the particle phase (Nah et al., 2018). However, a clear enhanced trend of molar ratio of particulate nitrate in the total nitrate as a function of NR-PM<sub>1</sub> mass concentration was observed correspondingly (as shown in Figure 5 and
- 330 Figure 6). Therefore, in this case the increase of aerosol liquid water is more likely to be the driving factor of particulate nitrate formation compared to the influence of pH.

It is worth noting that a similar co-condensation effect between water vapor and semi-volatile organic components (Topping and McFiggans, 2012;Topping et al., 2013;Hu et al., 2018) could promote the haze formation as well, for which there may be some evidence in the current case. Such a co-condensation effect will lead to the enhancement of semi-volatile organic and inorganic (e.g., nitrate) material with the increasing RH in a developing haze. The associated water will favor partitioning of

both HNO<sub>3</sub> and semi-volatile organic materials to the particle phase depending on the organic solubility, providing a linkage between the development of increasing organic and inorganic particle mass.

#### 3.3 The key role of liquid water on visibility degradation

- Aerosol particles grow up in size as ambient RH increases, further enhances their extinction coefficient and impacts visibility (Zhao et al., 2019;Kuang et al., 2016). In this section, size-resolved extinction coefficient of aerosol particles was estimated, and the influences of liquid water on the extinction coefficient and visibility were quantitatively evaluated. As shown in Figure 7a, the total light extinction coefficient of dry and wet aerosol particles enhanced by a factor of 4.3 and 5.4, respectively, from the
- 345 beginning to a fully-developed haze. Correspondingly, the calculated visibility without considering liquid water degraded significantly from ~10 km to less than 2 km within 48 hours during the marked 'haze period'. The contribution of aerosol associated water to visibility impairment was negligible in the beginning (2%), while it was significant (up to 24%) in the fully-developed haze (Figure 7b). This indicates that liquid water facilitated visibility degradation during haze development.
- The influences of liquid water on visibility degradation varied with aerosol particle size. The sizeresolved chemical composition data showed that the inorganic species, mainly particulate nitrate, were dominant components in the aerosol particles within the size range of 300~700 nm (Figure S3). Correspondingly, the particles in this size range contained most of the liquid water (50~80% of the total aerosol liquid water content of PM<sub>1</sub>). According to discussion in Sec. 3.2, the mutual promotion effect between liquid water and particulate nitrate can promote their mass loading enhancement. Aerosol
- 355 between liquid water and particulate nitrate can promote their mass loading enhancement. Aerosol particles in this size range experienced the most significant enhancement of light extinction due to water

uptake (Figure 8a and 8b) and contributed 70~88% of the total extinction coefficient of the total NR-PM<sub>1</sub> (Figure S9). In conclude, the rapid particulate nitrate enhancement enhanced the aerosol extinction coefficient during haze developing, while the aerosol water uptake further enhanced the visibility degradation by increasing extinction coefficient and promoting particulate nitrate enhancement.

It is worth noting that the enhanced dimming effect will further shallower the planetary boundary layer (PBL), which, in turn, depresses the dilution of water vapor and particulate matter in the atmosphere, hence leads to a higher RH and aerosol particle mass loading (Tie et al., 2017). Such effect is beyond the scope of this study.

## 365 4 Conclusions and implication

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In this study, we observed a particulate nitrate-dominated (up to 44% of non-refractory  $PM_1$  mass concentration) particulate matter pollution episode, which is typical during winter haze in Beijing, China. A clear co-increase of aerosol particle liquid water and particulate nitrate was observed, demonstrating the mutual promotion between them via observation-based theoretical calculations.

As shown in Figure 9, the water uptake by hygroscopic aerosols increased the aerosol surface area and volume, enhancing the condensational loss of N<sub>2</sub>O<sub>5</sub> over particles and favoring the thermodynamic equilibrium of HNO<sub>3</sub> into the particle phase under the supersaturated ambient HNO<sub>3</sub> and NH<sub>3</sub>. The enhanced particulate nitrate from the above pathways increased the mass fraction of particulate nitrate, which had a lower deliquescence RH than sulfate and resulted in more water uptake at lower ambient RH (Kreidenweis and Asa-Awuku, 2014). Hence, the increased aerosol particle surface area and

volume concentrations due to water uptake, in turn facilitates particulate nitrate enhancement. Hence, a feedback loop between liquid water and particulate nitrate enhancement is built up. Therefore the enhanced particulate nitrate components can accelerate the feedback compared with sulfate-rich pollution over the NCP region in the past (Hu et al., 2016). This self-amplification can rapidly degrade air quality and halve visibility within one day. Our results highlight the importance of reducing the particulate nitrate and its precursors (e.g.  $NO_x$ ) for mitigation of haze episodes in NCP region.

## Data availability

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The observational dataset of the BEST-ONE campaign can be accessed through the corresponding author Z. Wu (<u>zhijunwu@pku.edu.cn</u>).

385 The E-AIM model can be accessed via <u>http://www.aim.env.uea.ac.uk/aim/aim.php</u>.

# **Author contributions**

Z.W., Y.W. and Y.C. conceived the study. Y.Z., M.H., and A.K.S developed BEST-ONE field campaign program. Y.W., Z.W., D.S., Z.D., S.H.S., R.S., G.I.G., P.S., T.H., K.L., L.Z., C.Z., A.K.S., Y.Z., and M.H. participated in this campaign and collected the dataset. Y.W. conducted aerosol particle
liquid water calculation under guide of Y.B. and thermodynamic equilibrium of particulate ammonium nitrate under guidance of G.M. Y.C. calculated the uptake coefficient of N<sub>2</sub>O<sub>5</sub>, optical properties and visibility. Y.W. and Y.C. cowrite the manuscript with the inputs from all co-authors. Z.W., G.M., A.K.S., S.H.S., G.I.G., P.S., T.H., A.V., and A.W. proofread and help improve the manuscript. All authors discussed the results.

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Figure 1: The time series of (a) NR-PM<sub>1</sub> chemical composition measured by the HR-ToF-AMS and ambient RH (red solid line), (b) size-segregated aerosol particle liquid water and the total mass concentration of liquid water with smaller than 1 μm in aerodynamic diameter (red solid line), (c) size-segregated aerosol particle surface area and total aerosol particle surface area
645 without considering particle hygroscopic growth during February 29 to March 5, 2016.



Figure 2: The time series of condensational loss rate of N<sub>2</sub>O<sub>5</sub> (k\_N<sub>2</sub>O<sub>5</sub>) with the calculation of dry particle number size distribution (PNSD) and wet PNSD during February 29 to March 5, 2016.



Figure 3: The comparison of the calculated temperature-dependent dissociation constant of NH4NO3 (Kp) (Seinfeld. and Pandis., 2006) in the absence of liquid water, the product of equilibrium vapor pressure of gaseous NH3 and HNO3 from E-AIM (AIM\_pNH3pHNO3), and the
product of mixing ratios of gaseous NH3 and HNO3 measured by GAC-IC (M\_pNH3pHNO3). Here, Kp is colored by the ambient temperature ranging 265~293K during February 29 to March 5, 2016.



Figure 4: The relationship between aerosol particle liquid water and the molar ratio of particulate nitrate in the total nitrate, m<sub>NH4NO3</sub>/(m<sub>HNO3</sub> + m<sub>NH4NO3</sub>) (left axis) during the nighttime 18:00~07:00+1 (green solid triangle) and the daytime at 07:00 ~ 18:00 (red solid triangle), and mass concentration of particulate nitrate as a function of aerosol liquid water (right axis) during the period of during February 29 to March 5, 2016. Here, particulate nitrate was measured by 665 HR-ToF-AMS and the HNO3 in the gas phase was measured by GAC-IC. Aerosol liquid water was calculated by H-TDMA-derived method.



Figure 5. The time series of chemical composition measured by HR-ToF-AMS (left axis), calculated aerosol pH by ISORROPIA II (inner right axis) and molar ratio of particulate nitrate in the total nitrate (gas+particle phase) shown on outer right axis during February 29 to March 5, 2016.



Figure 6. The pH of the fine aerosol particles (left axis) and the molar ratio of particulate nitrate in the total nitrate (gas+particle phase) (right axis) as a function of NR-PM<sub>1</sub> mass concentrations.



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Figure 7: The time series of (a) calculated total extinction coefficient at wavelength of 550 nm with the consideration of dry and wet PNSD, referred as Extinction coefficient\_dry and Extinction coefficient\_wet, (b) calculated visibility with the consideration of dry and wet PNSD, referred as Visibility\_dry and Visibility\_wet, respectively during February 29 to March 5, 2016. Visibility degradation percentage is (Visibility\_wet-Visibility\_dry)/Visibility\_dry, representing the visibility degradation in the presence of liquid water.



Figure 8: (a) Size-segregated light extinction coefficient at wavelength of 550 nm for wet particles (Extinction coefficient\_wet), (b) size-segregated difference between Extinction coefficient\_wet and Extinction coefficient\_dry, representing light extinction coefficient difference with and without considering liquid water during February 29 to March 5, 2016.



Figure 9: The scheme of the mutual promotion effect between aerosol liquid water and particulate

690 nitrate