Characterization of nighttime formation of particulate organic nitrates based on high-resolution aerosol mass spectrometry in an urban atmosphere in China

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10 Abstract. Organic nitrates are important atmospheric species that significantly affect the cycling of NOx and ozone production. 11 However, characterization of particulate organic nitrates and their sources in polluted atmosphere is a big challenge and has 12 not been comprehensively studied in Asia. In this study, an Aerodyne high-resolution time-of-flight aerosol mass spectrometer 13 (HR-ToF-AMS) was deployed at an urban site in China from 2015 to 2016 to characterize particulate organic nitrates in total 14 nitrates with high time resolution. Based on the cross validation of two different data processing methods, organic nitrates 15 were effectively quantified to contribute a notable fraction of organic aerosol (OA): 9-21% in spring, 11-25% in summer, 9-16 20% in autumn, while very small fraction in winter. The good correlation between organic nitrates and fresh secondary organic 17 aerosol (SOA) at night as well as the diurnal trend of size distribution of organic nitrates indicated a key role of nighttime 18 secondary formation in Shenzhen, which is consistent with that reported in the US and Europe. The size distribution of organic 19 nitrates also implied that organic nitrates were mainly a local product and could experience substantial loss during air mass 20 transport. Furthermore, theoretical calculations of nighttime SOA production of NO₃ reactions with volatile organic 21 compounds (VOCs) measured during the spring campaign were performed, resulting in three biogenic VOCs (α -pinene, 22 limonene, and camphene) and one anthropogenic VOC (styrene) identified as the possible key VOC precursors for particulate 23 organic nitrates. The comparison with similar studies in the literature implied that nighttime particulate organic nitrates 24 formation is highly relevant with NOx levels. This study proposes that unlike the documented cases in the United States and 25 Europe, modeling nighttime particulate organic nitrate formation in China should incorporate not only biogenic VOCs but also 26 anthropogenic VOCs with high SOA yield for urban air pollution, which needs the support of relevant smog chamber studies 27 in the future.

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31 **1. Introduction**

Organic nitrates (ON) in aerosols have an important impact on the fate of NO_X and ozone production (Lelieveld et al., 2016), which can be formed in a minor channel of the reaction between peroxy radicals and NO (R1 and R2) (usually, an increased fraction of this reaction leads to the formation of alkoxy radicals and NO₂ (R3)) or via the NO₃-induced oxidation of unsaturated hydrocarbons (R4). Even though some recent studies have suggested that the formation of organic nitrates from peroxy radicals and NO may play a larger role than previously recognized (Teng et al., 2015, 2017), yields of organic nitrates via NO₃ reacting with alkenes are generally much higher (Fry et al., 2009; Ayres et al., 2015; Boyd et al., 2015, 2017).

38 $\operatorname{RH} + \operatorname{OH} + O_2 \rightarrow RO_2 + H_2O$ (R1)

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	2	2	2	
$RO_2 + NO \rightarrow$	RONO	2		(R2)
$RO_2 + NO \rightarrow$	RO + L	NO_2		(R3)
$\mathbf{R} = R' + \mathbf{N}O$	$\rightarrow R($	ONO_{2}) <i>R'</i>	(R4)

Several methods have been developed to directly measure total organic nitrates (gas + particle) in the real atmosphere. For 42 43 example, Rollins et al.(2012) used a thermal-dissociation laser-induced fluorescence technique (TD-LIF) to observe organic 44 nitrates in the United States; Sobanski et al. (2017) measured organic nitrates in Germany using the thermal dissociation cavity 45 ring-down spectroscopy (TD-CRDS). Field and laboratory studies around the world indicated that particulate organic nitrates 46 could contribute a large portion of secondary organic aerosol (SOA) (Rollins et al., 2012; Xu et al., 2015a; Fry et al., 2013; 47 Avres et al., 2015; Boyd et al., 2015; Lee et al., 2016). Recently, researchers have proposed some estimation methods for 48 particle-phase organic nitrates based on aerosol mass spectrometry (AMS) with high time resolution (Farmer et al., 2010; Hao 49 et al., 2014: Xu et al., 2015a, 2015b). Ng et al. (2017) reviewed the nitrate radical chemistry and the abundance of particulate 50 organic nitrates in the United States and Europe, and further concluded that particulate organic nitrates are formed substantially 51 via NO₃+BVOC chemistry, which plays an important role in SOA formation. Unfortunately, relevant Chinese datasets are 52 scarce yet and not included in this review. This was because (1) the contributions of organic nitrates in SOA and total nitrates 53 in Chinese atmosphere remain poorly understood; (2) the anthropogenic and biogenic precursor emissions in China are 54 significantly different from those in the United States and Europe, and thus cannot be easily estimated. To our best knowledge, 55 few studies have investigated the concentrations and formation pathways of particulate organic nitrates in China. Xu et al. 56 (2017) estimated the mass concentration of organic nitrogen in Beijing using AMS, but in this study they ignored the 57 contribution of NO_{X^+} family, which are the major fragments of organic nitrates.

58 Shenzhen is a megacity of China in a subtropical region, where NOx involved photochemical reactions are very active, given 59 considerable biogenic and anthropogenic VOC emissions (Zhang et al., 2008). To assess the evolution of particulate organic 60 nitrates in a polluted urban atmosphere, we deployed an Aerodyne high-resolution time-of-flight aerosol mass spectrometer 61 (HR-ToF-AMS) and other instruments in Shenzhen from 2015 to 2016 in this study. Organic nitrates and their contributions 62 to OA in different seasons were estimated by different methods using the HR-ToF-AMS datasets, based on which, the 63 secondary formation pathway of particulate organic nitrates in Shenzhen was further explored.

64 **2. Experiment methods**

65 2.1 Sampling site and period

- 66 The sampling site (22.6°N, 113.9°E) was on the roof (20 m above ground) of an academic building on the campus of Peking
- 67 University Shenzhen Graduate School (PKUSZ), which is located in the western urban area in Shenzhen (Figure 1). This site
- is mostly surrounded by subtropical plants without significant anthropogenic emission sources nearby, except a local road that
- 69 is ~100 m from the site. In this study, we used the statistical data from the Meteorological Bureau of Shenzhen Municipality
- 70 (http://www.szmb.gov.cn/site/szmb/Esztq/index.html) as the reference data to determine the sampling periods for four
- 71 different seasons during 2015-2016, as shown in Table 1.



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Figure 1. The location of the sampling site.

Table 1.Meteorological conditions, PM₁ species concentrations and relevant parameters for different sampling periods in
 Shenzhen

75 Shenzhen.

	Sampling period	4.1-4.30, 2016	8.1-8.31, 2015	11.4-11.30,2015	1.21-2.3, 2016
		Spring	Summer	Autumn	Winter
Meteorology	T (°C)	24.5±2.5	29.0±3.0	23.6±3.7	10.7±4.7
	RH (%)	78.0±12.7	71.2±17.5	68.2±15.8	75.4±18.7
	WS (m s^{-1})	1.4±0.8	1.0±0.7	1.2±0.7	1.5±0.8
	Org	4.3±3.2	10.0±6.9	7.8±5.9	5.1±3.5
Species	SO_4^{2-}	3.2±1.8	5.8±3.3	2.3±1.5	1.9±1.2
Species	Total NO ₃	0.96±1.4	0.91±0.90	1.3±1.4	1.6±1.0

_3	NH4 ⁺	1.4±0.8	2.0±1.1	1.1±0.8	1.2±0.6
(µg m ⁻³)		0.14+0.10	0.03+0.05	0.22+0.36	0.64+0.85
	Cl	0.14±0.19	0.05±0.05	0.22±0.30	0.04±0.85
	BC	1.9±2.1	2.4±1.6	3.5±2.6	2.4±1.5
	Total	12.0±8.9	15.1±13.8	11.8±9.5	12.2±7.2
	RNULANOZ	2.80	3.20	3.32	3.48
ON relevant	\n4N03	2.74	C 1 4	4.20	2.55
	R _{obs}	3.74	6.14	4.30	3.33
parameters	Fraction of positive numbers of	99%	99%	84%	47%
	Robs- RNH4NO3				

76 2.2 Instrumentation

77 2.2.1 High Resolution Time-of-Flight Aerosol Mass Spectrometer

78 During the sampling periods, chemical composition of non-refractory PM₁ was measured by an Aerodyne HR-ToF-AMS, and 79 detailed descriptions of this instrument are given in the literature (DeCarlo et al., 2006; Canagaratna et al., 2007). The setup 80 and operation of the HR-ToF-AMS can be found in our previous publications (Huang et al., 2010, 2012; Zhu et al., 2016). To 81 remove coarse particles, a PM_{2.5} cyclone inlet was installed before the sampling copper tube with a flow rate of 10 l min⁻¹. 82 Before entering the AMS, the sampled air was dried by a nafion dryer (MD-070-12S-4, Perma Pure Inc.) to eliminate the 83 potential influence of relative humidity on particle collection (Matthew et al., 2008). The ionization efficiency (IE) calibrations 84 were performed using pure ammonium nitrate every two weeks. The relative ionization efficiencies (RIEs) used in this study 85 were 1.2 for sulfate, 1.1 for nitrate, 1.3 for chloride, 1.4 for organics and 4.0 for ammonium, respectively (Jimenez et al., 2003). 86 Composition-dependent collection efficiencies (CEs) were applied to the data according to the method in Middlebrook et al. (2012). The instrument was operated at two ion optical modes with a cycle of 4 min, including 2 min for the mass -sensitive 87 88 V-mode and 2 min for the high mass resolution W-mode. The HR-ToF-AMS data analysis was performed using the software 89 SOUIRREL (version 1.57) and PIKA (version 1.16) written in Igor Pro 6.37 (Wave Metrics Inc.) (http://cires1.colorado.edu/jimenezgroup/ToFAMSResources/ToFSoftware / index.html). 90

91 **2.2.2 Other co-located instruments**

In addition to the HR-ToF-AMS, other relevant instruments were deployed at the same sampling site. An aethalometer (AE-31, Magee) was used for measurement of refractory black carbon (BC) with a resolution of 5 min. An SMPS system (3775 CPC and 3080 DMA, TSI Inc.) was used to obtain the particle number size distribution in 15–615 nm (mobility diameter) with a time resolution of 5 min. Ozone and NO_X were measured by a 49i ozone analyzer and a 42i nitrogen oxide analyzer (Thermo Scientific), respectively. In the spring campaign, ambient VOC concentrations were also measured using an on-line VOC monitoring system (TH-300B, Tianhong Corp.), including an ultralow-temperature preconcentration cold trap and an

- 98 automated in-situ gas chromatograph (Agilent 7820A) equipped with a mass spectrometer (Agilent 5977E). The system had 99 both a flame ionization detector (FID) gas channel for C2-C5 hydrocarbons and a mass spectrometer (MS) gas channel for 100 C5–C12 hydrocarbons, halohydrocarbons and oxygenated VOCs. A complete working cycle of the system was one hour and 101 included five steps: sample collection, freeze-trapping, thermal desorption, GC-FID/MS analysis, heating, and anti-blowing 102 purification. The sample collection time was 5 min. The sampling flow speed was 60 ml min⁻¹. The anti-blowing flow speed 103 was 200 ml min⁻¹. The calibration of over 100 VOCs was performed using mixed standard gas before and after the campaign. 104 Detection limits for most compounds were near 5 pptv. More description of this instrument can be found in Wang et al. (2014). 105 2.3 Organic nitrates estimation methods
- In this study, we used two independent methods to estimate particulate organic nitrates based on the AMS data, following the approaches in Xu et al. (2015b). The first method is based on the NO⁺/NO₂⁺ ratio (NO_X⁺ ratio) in the HR-AMS spectrum. Due to the very different NO_X⁺ ratios of organic nitrates and inorganic nitrate (i.e., R_{ON} and R_{NH4NO3}, respectively) (Farmer et al., 2010; Boyd et al., 2015; Fry et al., 2008; Bruns et al., 2010), the NO₂⁺ and NO⁺ concentrations of organic nitrates (*NO*_{2,ON} and *NO*_{ON}) can be quantified with the HR-AMS data via Eqs. (1) and (2), respectively (Farmer et al., 2010):

111
$$NO_{2}^{+}{}_{,ON} = \frac{NO_{2}^{+}{}_{,Obs} \times (R_{obs} - R_{NH4NO3})}{R_{ON} - R_{NH4NO3}}$$
(1)

$$NO^{+}_{ON} = R_{ON} \times NO_{2,ON}$$
(2)

- 113 where R_{obs} is the NOx⁺ ratio from the observation. The value of R_{ON} is difficult to determine because it varies between 114 instruments and precursor VOCs. However, R_{NH4NO3} was determined by IE calibration using pure NH₄NO₃ every two weeks 115 for each campaign and the results showed stable values: In spring, the average R_{NH4NO3} was 2.66 for the first IE calibration and 116 2.94 for the second one; in summer, the average $R_{\rm NH4NO}$ was 3.05 and 3.34 for the first and second IE calibrations, respectively; 117 in autumn, the average R_{NH4NO3} was 3.33 and 3.31 for the first and second IE calibrations, respectively; in winter, the average 118 R_{NH4NO3} was 3.45 and 3.51 for the first and second IE calibrations, respectively. We adopted the R_{ON}/R_{NH4NO3} estimation range 119 (from 2.08 to 3.99) for variation of precursor VOCs in the literature to determine R_{ON} (Farmer et al., 2010; Boyd et al., 2015; 120 Bruns et al., 2010; Sato et al., 2010; Xu et al., 2015b), and thus two R_{ON} values were calculated for each season to provide the 121 upper bound (NO_{3 org ratio 1}) and lower bound (NO_{3 org ratio 2}) of NO_{3,org} mass concentration.
- 122 The second method is based on the traditional positive matrix factorization (PMF) analysis of HR organic mass spectra, which 123 resolves different organic factors (Zhang et al., 2011; Ng et al., 2010; Huang et al., 2013). Combined with NO⁺ and NO₂⁺ ions, 124 the same analysis of HR organic mass spectra was performed to separate NO+ and NO2+ ions into different organic factors 125 and an inorganic nitrate factor (Hao et al., 2014; Xu et al., 2015b). The PMF analysis procedures in this study can be found in 126 our previous publications (Huang et al., 2010; Zhu et al., 2016; He et al., 2011), resulting in three organic factors and one 127 inorganic factor in spring, summer and autumn: a hydrocarbon-like OA (HOA) characterized by $C_nH_{2n+1}^+$ and $C_nH_{2n-1}^+$ and O/C 128 of 0.11 to 0.18, a less-oxidized oxygenated OA (LO-OOA) characterized by $C_xH_yO_z^+$ especially $C_2H_3O^+$ and O/C of 0.28 to 129 0.70, a more-oxidized oxygenated OA (MO-OOA) also characterized by $C_xH_yO_z^+$ especially CO_2^+ and O/C of 0.78 to 1.24,

- 130 and a nitrate inorganic aerosol (NIA) characterized by overwhelming NO⁺ and NO⁺, as indicated in Fig S6. According to the 131 diagnostic plots of the PMF analysis shown in Figure S2 to S4, the same organic factors as those in the traditional PMF analysis 132 of only organic mass spectra were obtained. The NO⁺ and NO₂⁺ ions were distributed among different OA factors and the NIA factor; thus the concentrations of nitrate functionality $(NO_{org}^+ \text{ and } NO_{2,org}^+)$ in organic nitrates (NO_{3, org}) are equal to the sum of 133 134 NO_2^+ and NO^+ via Eqs. (3) and (4), respectively (Xu et al., 2015b):
- 135
 - $NO_{2.ora}^{+} = \sum ([OA \ factor]_{i} \times f_{NO2.i})$ (3)
- $NO_{org}^{+} = \sum ([OA \ factor]_{i} \times f_{NO,i})$ 136

where [OA factor]_i represents the mass concentration of OA factor i, and f_{NO2} i and f_{NO2} i represent the mass fractions of NO₂⁺ 137 138 and NO⁺, respectively.

(4)

139 It should be noted that the 4-factor solution seemed to have a "mixed factor" problem to some extent (Zhu et al., 2018). For 140 example, HOA mixed with COA (clear $C_3H_3O^+$ in m/z 55 for spring, summer and autumn) (Mohr et al., 2012), and BBOA 141 mixed with LO-OOA (clear m/z 60 and 73 signals in LO-OOA in autumn) (Cubison et al., 2011). However, running PMF with 142 more factors would produce unexplained factors but less influence on the apportionment of NO⁺ and NO₂⁺ ions between organic nitrates and inorganic nitrate (Table S1). In addition, the standard deviations of NO⁺ and NO₂⁺ ions in the OA factors 143 144 across different FPEAK values (from -1.0 to 1.0) were very small (Table S2). Therefore, the 4-factor solution was used for 145 quantifying organic nitrates in spring, summer and autumn.

146 3. Results and discussion

147 3.1 Organic nitrates estimation

148 Table 2 shows the concentrations of nitrate functionality in organic nitrates (i.e., $NO_{3, org}$), estimated by both the NO^+/NO_2^+ 149 ratio method and PMF method, as well as their contributions to the total measured nitrate. It should be noted that the small 150 difference between the average R_{obs} and R_{NH4NO3} in winter leads to a large portion of negative data using the NO⁺/NO₂⁺ ratio 151 method (Table 1). The result from the PMF method shows that the contribution of organic nitrates in total nitrates is only 4.2% 152 in winter (Figure S6), suggesting a negligible contribution of organic nitrates. Thus, we will only discuss organic nitrate 153 estimation results in spring, summer and autumn. The analytical outcomes by the NO⁺/NO₂⁺ ratio method and by the PMF 154 method consistently suggest that organic nitrates had the highest ambient concentration (0.34-0.53 $\mu g m^3$) and proportion in 155 total nitrates (41-64%) in summer among the different seasons. This finding agrees with the finding in (Ng et al., 2017) and it 156 implies a seasonal trend in comparison with that of total nitrates in Table 1. Assuming the average molecular weight of organic 157 nitrates of 200 to 300 g mol⁻¹ (Rollins et al., 2012), we found that organic nitrates contributed 9-21% to OA in spring, 11-25% 158 in summer, and 9-20% in autumn.

159 In the PMF method, the mass fractions of organic nitrates in HOA, LO-OOA and MO-OOA were 31%, 49% and 20%, 160 respectively, in spring; 28%, 52% and 20%, respectively, in summer; 30%, 46% and 24%, respectively, in autumn. The major

161	fraction of organic nitrates occurring in LO-OOA for the three seasons implied that organic nitrates were mostly related to
162	fresher secondary OA formation. The NIA factors in all seasons were dominated by but are not limited to NO^+ and NO_2^+ .
163	Some organic fragments, such as CO_{2^+} and $C_2H_3O^+$, are also part of these factors, which agreed with the findings in the
164	literature (Hao et al., 2014; Xu et al., 2015b; Sun et al., 2012). This indicated the potential interference of organics in the NIA
165	factor. It is also worth to be noted that the NO^+/NO_2^+ ratios in NIA (2.93 for spring, 3.53 for summer and 3.54 for autumn)
166	were higher than that for pure NH_4NO_3 (Table 1), indicating an underestimation of $NO_{3.org}$ concentration by the PMF method.
167	This finding may also explain the reason that the concentration of NO3.org estimated using the PMF method was always close
168	to the lower estimation bound of $NO_{3.org}$ concentration estimated using the NO^+/NO_2^+ ratio method in each season (Table 2).
169	Table 2. Summary of organic nitrates estimations using the NO^+/NO_2^+ ratio method and the PMF method

Sampling period		NO ⁺ /NO ₂ ⁺ rat	2 ⁺ ratio method I			PMF method	
	NO _{3,org}	(µg m ⁻³) ^a	NO _{3,01}	rg/NO ₃	<i>NO</i> _{3,org} (µg m ⁻³) ^b	NO _{3,org} /NO ₃	
	lower	upper	lower	upper			
Spring	0.12	0.19	13%	21%	0.12	12%	
Summer	0.34	0.53	41%	64%	0.39	43%	
Autumn	0.21	0.33	16%	25%	0.21	16%	
Winter	/	/	/	/	0.07	4.2%	

^a NO_{3, org} for upper bound is denoted as NO_{3_org_ratio_1}, and NO_{3, org} for lower bound is denoted as NO_{3_org_ratio_2}.

^b NO_{3, org} estimated using the PMF method is denoted as NO_{3_org_PMF}.

172 To further verify the reliability of the estimated results of organic nitrates, the NO_{3, org} concentration time series calculated by 173 the two methods in each season are shown in Figure 2a. The computed correlation coefficients (R) are good (0.82 for spring, 174 0.82 for summer and 0.77 for autumn), indicating that similar results were achieved. The inorganic nitrate (NO_{3 inorg}*) obtained 175 by subtracting NO_{3 org ratio 1} from total measured nitrates also correlated well with the inorganic nitrate estimated using the PMF method (R=0.92 for spring, 0.87 for summer and 0.86 for autumn). , While they were distinctive from those of inorganic 176 177 nitrate (Figure 2b), which indicates that organic nitrates had been well separated from inorganic nitrate in this study, the diurnal 178 trends of organic nitrates obtained by the two methods were similar in each season, with lower concentrations in the daytime 179 and higher concentrations at night.



181 **Figure 2.** (a) Time series of NO_{3, org} concentration estimated by the NO⁺/NO₂⁺ ratio method and PMF method for each

182 season; (b) correlations between $NO_{3_{org_ratio_1}}$ and $NO_{3_{org_PMF}}$ (left panel); diurnal trends of organic nitrates and $NO_{3_{org}}$

183 estimated by the different methods (right panel).

184 **3.2 Correlation between organic nitrates and OA factors**

185 As indicated by the results in the PMF method, the majority of organic nitrates were associated with LO-OOA in spring, 186 summer and autumn in the urban atmosphere in Shenzhen, implying a dominant secondary origin of organic nitrates. To further 187 confirm this relationship, we made the correlation analysis between organic nitrates estimated by the NO⁺/NO₂⁺ ratio method 188 and the three factors resolved by the PMF analysis with only organic mass spectra in the three seasons. Generally, organic 189 nitrates were found better-correlated with LO-OOA (R=0.69-0.77 in Figure 3) than with HOA and MO-OOA (R=0.03-0.69 in 190 Figures S6-S8), which is consistent with the fact that the majority of organic nitrates were associated with LO-OOA in the 191 PMF method. However, the moderate correlation between organic nitrates and HOA implied possibility of direct emissions of 192 organic nitrates. Furthermore, we found a noticeably improved correlation between LO-OOA and organic nitrates at night 193 (19:00-6:00) and a reduced correlation during the daytime (7:00-18:00) in Figure 3, especially in summer, implying that 194 organic nitrates formation might be more closely related to secondary formation at night.



196 **Figure 3.**Correlation of NO_{3_org_ratio_1} and LO-OOA in each season for the whole day (a), at night (b) and in the daytime (c).

197 **3.3 Size distribution characteristics of organic nitrates**

198 In this section, we used the NO^+/NO_2^+ ratio as an indicator to investigate the size distribution of organic nitrates. The size 199 distributions of the NO⁺ and NO₂⁺ signals for the different seasons have been shown in Figure S11. Due to the lack of HR-200 PToF data, our analyses used the UMR-PToF data (m/z 30 and 46), which may bring in the interferences of $CH_2O_X^+$. However, 201 for all four seasons, the contributions of $CH_2O_X^+$ in m/z 30 and 46 in the HR data of PM₁ were less than 10% (Figure S10). 202 which indicates that the interferences were negligible in this study. The average size distributions of different aerosol species and NO^+/NO_2^+ ratio in four seasons are shown in Figure 4a. It is clearly found that the NO^+/NO_2^+ ratio exhibited a decreasing 203 trend in spring, summer and autumn, while it kept constant in winter, similar to the value of R_{NH4NO3} (red dotted line in Figure 204 205 4). It should also be noted that in spring, summer and autumn, the lowest values of NO⁺/NO₂⁺ ratio occurring at >1 μ m 206 approximated to the corresponding seasonal values of R_{NH4NO3}. These characteristics clearly indicated that organic nitrates 207 existed mostly in fresh particles with smaller sizes. Different from the bulk OA and inorganic spices, very limited amount of 208 organic nitrates exist in larger aged particles, implying that they could be easily removed by deposition and/or chemical 209 degradation during air mass transport. In addition, the diurnal trends of size distribution of NO⁺/NO₂⁺ ratio in spring, summer 210 and autumn in Figure 4b show apparent higher values at small sizes at night, suggesting an important nighttime local origin of 211 organic nitrates. Combining with the analysis in section 3.2, local nighttime secondary formation of organic nitrates in warmer 212 seasons in the urban polluted atmosphere in Shenzhen is highlighted. This is consistent with the previous findings in the US 213 and Europe that the nighttime NO₃+VOCs reactions serve as an important source for particulate organic nitrates (Rollins et al., 214 2012; Xu et al., 2015a, 2015b; Fry et al., 2013; Lee et al., 2016). We will then explore the nighttime NO₃+VOCs reactions in 215 Shenzhen in the following section.



Figure 4.(a) Average size distributions of aerosol species and NO⁺/NO₂⁺ ratio (red dotted line represents $R_{\rm NH4NO3}$); (b) diurnal trends of size distribution of NO⁺/NO₂⁺ ratio.

219 **3.4 Nighttime particulate organic nitrates formation via NO₃+VOCs**

Since on-line VOCs measurement was only performed during the spring campaign (described in section 2.2), the following theoretical analysis of NO₃+VOCs reactions applies only to the spring case. NO₃+VOCs reactions would yield a large mass of gas- and particle-phase organic nitrates (Rollins et al., 2012; Nah et al., 2016; Boyd et al., 2015, 2017; Xu et al., 2015a, 2015b; Lee et al., 2016). We used Eq. (9) to roughly judge the production potential (PP) of organic nitrates from a NO₃+VOC reaction: [Production Potential]_{NO3+VOCi} = $K_i \cdot [VOC_i] \cdot [NO_3]$ (9)

Where K_i represents the reaction rate coefficient for NO₃ radical and a VOC; $[VOC_i]$ is the concentration of the specific VOC 225 226 $[NO_3]$ is the concentration of NO₃ radical. It should be noted that no organic nitrates yield parameter was introduced in Eq. 227 (9), because only a few organic nitrate yields for BVOCs were available in the literature (Fry et al., 2014; Ng et al., 2017). However, given the fact that the values of $K_i \cdot [VOC_i] \cdot [NO_3]$ for different VOC species can differ by orders of magnitude, not 228 229 multiplying the organic nitrate yields (ranging from 0 to 1) would not significantly affect the PP ranking of VOCs. In the spring 230 campaign, the diurnal variations of NO₂, O₃ and estimated NO₃ radical concentrations are shown in Figure S12. It was found 231 that as a comparison to the nighttime NO₃ radical concentrations reported in literature in the United States (Rollins et al., 2012; 232 Xu et al., 2015a), high concentrations of NO₂ (19.93 \pm 2.31 ppb) at night leaded to high yield of NO₃ radical (1.24 \pm 0.76 ppt) 233 in Shenzhen, as calculated in Text S1.

The reaction rate coefficients of typical measured nighttime VOC concentrations with NO₃ radical and the production potentials are listed in Table S3 and shown in Figure 5. These VOCs were considered based on their higher ambient concentrations and availability for reaction kinetics with NO₃ radical. According to the distribution of production potential, five biogenic VOCs (BVOCs) (i.e., α -pinene, limonene, camphene, β -pinene and isoprene) and one anthropogenic VOC (styrene) were identified as notable VOC precursors with high production potential, while the sum of production potential from the other VOCs was negligible as shown in Figure 5b.



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Figure 5. (a) Mean concentrations of VOCs and the corresponding calculated production potential of NO₃+VOC at night
 during the spring campaign; (b) production potential ranking of VOCs at night during the spring campaign.

Based on the production potential evaluation above, we further explore SOA yield of NO₃+the six notable VOC precursors according to the analysis method of particulate organic nitrate formation in Xu et al. (2015a). Briefly, NO₃ and ozone are two main oxidants for SOA formation from VOCs at night. Based on the concentrations of oxidants and the reaction rate constants for VOCs with NO₃ and ozone, the branching ratio of each VOC that reacts with NO₃ can be estimated as in Eq. (10). By combining the estimated branching ratios and SOA yields from chamber studies (Table 3), potential SOA production from these VOCs can be calculated as in Eq. (11) (Xu et al., 2015a):

250
$$branching \ ratio_{species \ i+NO3} = \frac{k_{[species \ i+NO3]} \times [NO_3]}{k_{[species \ i+NO3]} \times [NO_3] + k_{[species \ i+O3]} \times [O_3]}$$
(10)

251
$$[SOA]_{species,oxidant} = [species] \times branching \ ratio_{species,oxidant} \times yield_{species,oxidant} \ (11)$$

The results in Table 3 show that all six notable VOC species were prone to react with NO₃ radical instead of O₃ at night, and

253 the estimated potential SOA production from NO₃+VOCs reactions using SOA mass yields in the literature was 0-0.33 μ g m⁻³ 254 ³ for α -pinene, 0.09-1.28 μ g m⁻³ for limonene, 0.24 μ g m⁻³ for styrene, 0.004-0.06 μ g m⁻³ for β -pinene and 0.002-0.02 μ g m⁻³ 255 for isoprene. The SOA yield from camphene is currently unknown in the literature. It is seen that the average observed 256 nighttime concentration of particulate organic nitrates during the spring campaign (0.39-0.83µg m⁻³, converting NO_{3.org_ratio_1},

257 NO_{3.org_PMF} in Figure 6 into organic nitrates assuming the average molecular weight of organic nitrates of 200 to 300 g mol⁻¹)

258 was well within the estimated SOA concentration ranges produced by α -pinene, limonene and styrene in Table 3, indicating

259 that these three VOCs were the key VOC precursors in urban atmosphere in Shenzhen. Considering both the production 260 potentials and SOA yields, the contributions of β -pinene and isoprene to nighttime formation of particulate organic nitrates

could be negligible.

262 **Table 3.** Average concentrations, reaction branching and SOA production of α -pinene, limonene, styrene, camphene, β -

263 pinene and isoprene with respect to different oxidants at night in the spring campaign.

Species	Concentration	Rate coe	efficient ^a	Branchi	ng ratio	SOA yield from	SOA from
	(ppbv)					the literature	$VOCs + NO_3$
	-	NO ₃	O ₃	NO ₃	O ₃	(with NO ₃)	(µg m ⁻³)
α-pinene	0.39	6.64E ⁻ 12	7.2E ⁻ 17	0.962	0.038	0-0.16 ^b	0-0.33
Limonene	0.14	1.22E-11	1.54E-16	0.957	0.043	0.12-1.74°	0.09-1.28
Styrene	0.19	1.50E-12	1.70E-17	0.941	0.059	0.23 ^d	0.24
Camphene	0.28	6.20E-13	9.0E ⁻ 19	0.992	0.008	/	/
β-pinene	0.01	2.51E-12	1.50E-17	0.968	0.032	0.07-1.04 ^e	0.004-0.06
Isoprene	0.032	6.96E-13	1.27E-17	0.908	0.091	0.02-0.24 f	0.002-0.02

^a Rate coefficients for all species except camphene are from the Master Chemical Mechanism model

265 (http://mcm.leeds.ac.uk/MCM/; under 25 °C), rate coefficients for camphene are from Martínez et al. (1999) and Atkinson et

266 al. (1990).

^b Hallquist et al. (1999); Spittler et al. (2006); Perraud et al. (2010); Fry et al. (2014); Nah et al. (2016).

^c Fry et al. (2011, 2014); Spittler et al. (2006); Boyd et al. (2017).

^d Cabrera-Perez et al. (2017).

^e Griffin et al. (1999); Fry et al. (2009); Fry et al. (2014); Boyd et al. (2015).

^f Rollins al. (2009); Ng et al. (2008).

The estimation of potential SOA production above suggests significant contributions of α -pinene, limonene, and styrene to SOA, and the significant contribution of camphene is also possible. Thus, we further explored the diurnal variations of the PPs of these four VOCs. Figure 6 shows the diurnal trends of BC, LO-OOA, NO_{3.org_ratio_1}, NO_{3.org_PMF}, and the PPs of the four VOCs during the spring campaign. There were two apparent nighttime growth periods (i.e., I: 19:00–22:00 and II: 2:00–6:00) for both NO_{3.org_ratio_1} and NO_{3.org_PMF}. During Period I, BC maintained a relatively higher level, suggesting stable anthropogenic emissions. In contrast, the increases of all the PPs during Period I indicated that these precursors contributed to the organic

278 nitrate growth. After 22:00, while the PPs still showed a rapid growth, BC and organic nitrates began to decrease, implying

possible existence of other important anthropogenic VOC precursors, which were not identified by the GC-FID/MS analysis but would dominate the formation of organic nitrates at this stage. During Period II, the anthropogenic emissions remained at a stable lower level, as indicated by BC, while all the PPs increased with organic nitrates again, indicating that these four precursors also contributed to, or could dominate, this organic nitrate growth. As shown in Figure S13, organic nitrates correlated better with the PPs (R=0.63–0.74) than with LO-OOA (R=0.19–0.31) or BC (R=0.02–0.05) during Period II at the spring campaign, suggesting the significant contributions of the NO₃ reactions with these precursors.



- **Figure 6.** Diurnal trends of BC, LO-OOA, $NO_{3.org_PMF}$ and production potential (PP) of α -pinene, limonene, styrene, and camphene during the spring campaign.
- 288 It should be noted that, all previous studies on nighttime organic nitrate formation in the US and Europe focused on mechanisms 289 of NO₃ reactions with BVOCs (Hallquist et al., 1999; Spittler et al., 2006; Perraud et al., 2010; Fry et al., 2014; Nah et al., 290 2016; Boyd et al., 2015, 2017). In this study, however, we found that anthropogenic VOCs could also play significant roles in 291 particulate organic nitrate formation at night. Besides styrene, one of major aromatics (Cabrera-Perez et al., 2016), there were 292 also other important anthropogenic VOC precursors that we did not identify in the spring campaign. In China, styrene has been 293 actually identified as an important VOC of non-methane hydrocarbons (NMHCs) in urban areas, and has a notable contribution 294 to ozone formation and SOA production (An et al., 2009; Yuan et al., 2013; Zhu et al., 2019). This study highlights the possible 295 key roles of anthropogenic VOC precursors in nighttime particulate organic nitrate formation in urban atmosphere in China, 296 and relevant smog chamber studies for anthropogenic VOCs+NO₃ reactions are needed to support parameterization in 297 modeling.

298 **3.5** Comparison with other similar studies and implications

299 Table 4 shows the average ambient temperatures, average concentrations of NO, NO₂, monoterpenes, NO_{3,org}, the ratio of 300 NO_{3.org} to NO_{3.total} and the ratio of organic nitrates to total organics in several similar field campaigns available in the literature, 301 which implies the key role of NO₃+VOCs reactions for nighttime particulate organic nitrate formation. In general, the variation 302 of the particulate organic nitrates concentration is within an order of magnitude (0.06-0.98 μ g/m³) among the different sites. 303 Higher concentrations of particulate organic nitrates generally is associate with higher NOx concentrations rather than BVOC 304 concentrations. On the other hand, although the BVOC concentrations in Bakersfield were far less than that in other campaigns, 305 the concentration of particulate organic nitrates there showed an intermediate level among all the campaigns. Therefore, it is 306 suggested that the formation of particulate organic nitrates may be more relevant with NOx than BVOCs, which is consistent 307 with the finding that the organic nitrate production was dominated by NOx in the southeastern US (Edwards et al., 2017). In 308 the spring campaign of this study, we examined the correlation between organic nitrates and NO₂ or VOCs (by the sum of α -309 pinene, limonene, styrene and camphene) at night (Figure S14) and found a significant correlation of organic nitrates with NO_2 310 (R=0.40-0.47) rather than with VOCs (R=0.22-0.23), which further suggests that the organic nitrates formation was driven by 311 the NOx-involved NO₃ chemistry.

Table 4. Average ambient temperatures, average concentrations of monoterpenes, NO_{3,total}, NO_{3,org}, NO_{3,org}/ NO_{3,total} and the ratio of organic nitrates to total organics (ON/Org) for different

314	field campaigns around the world	The ON results at the Euror	ean and US sites are from	Kiendler-Scharr et al (201	6) and No et al (2017)
511	nota campaigns around the world.	The or results at the Laron		included belluit of ul. (20)	(0) (1) (1) (2) (1) (2)
	10	1			

Sampling site	Site	Sampling	Temperature	NO	NO ₂	Monoterpenes	NO _{3,org}	NO _{3,org} /	ON/Org	Reference/Note
	type	period	(°C)	(ppbv)	(ppbv)	(ppbv)	(µg m ⁻³)	NO _{3,total}		
Bakersfield, US	rural	May-June, 2010	23.0		8.2	0.045 (α-pinene)	0.16	0.28	0.23	Rollins et al. (2012)/
						0.004 (β-pinene)				NO3,org measured by TD-LIF
						0.034 (limonene)				
Woodland Park,	high	July-August,	15.0		1.2	0.25 (monoterpene)	0.06	0.86	0.09	Fry et al. (2013)/
US	attitude	2011								Use AMS data to
										estimate NO3,org
Centreville, US	rural	June-July, 2013	24.7	0.1	1.1	0.350 (α-pinene)*	0.08	1.00	0.10	Xu et al. (2015a)
						0.312 (β-pinene)*				Xu et al. (2015b)/
						0.050 (limonana)*				Use AMS data to
						0.050 (innonene).				estimate NO3,org
Barcelona,	urban	March, 2009	13.3	11.0	23.6	0.423	0.48	0.13	0.13	Mohr et al. (2012)
Spain						(monoterpene)				Pandolfi et al. (2014) /
										Use AMS data to
										estimate NO3,org
Shenzhen,	urban	April, 2016	24.5	8.0	19.4	0.391 (α-pinene)*	0.16	0.17	0.11	This study/
China						0.013 (B-pinene)*				Use AMS data to
						0.127 (P pinene)				estimate NO3,org
						0.137 (limonene)*				

315 *BVOC concentration at night.

316 4. Conclusions

317 An Aerodyne HR-ToF-AMS was deployed in urban Shenzhen for about one month per season during 2015–2016 to 318 characterize particulate organic nitrates with high time resolution. We discovered high mass fractions of organic nitrates in 319 total organics during warmer seasons, including spring (9-21%), summer (11-25%) and autumn (9-20%), while particulate 320 organic nitrates were negligible in winter. The correlation analysis between organic nitrates and each OA factor showed high 321 correlation (R=0.77 in spring, 0.91 in summer and 0.72 in autumn) between organic nitrates and LO-OOA at night. The diurnal 322 trend analysis of size distribution of NO^+/NO_2^+ ratio further suggested that organic nitrates formation mainly occurred at night. 323 It also suggested that organic nitrates concentrated on smaller sizes, indicating that they were mostly local product. The 324 calculated theoretical nighttime production potential of NO₃ reactions with VOCs measured in spring showed that six VOC 325 species (i.e., α -pinene, limonene, styrene, camphene, β -pinene and isoprene) were prominent precursors. The SOA yield 326 analysis and the nighttime variation of production potential further indicated that α -pinene, limonene, camphene and styrene 327 could contribute significantly to nighttime formation of particulate organic nitrates in spring in Shenzhen, highlighting the 328 unique contribution of anthropogenic VOCs as a comparison to that documented in previous studies in the US and Europe. 329 Finally, the comparison of the results in this study with other similar studies implied that nighttime formation of particulate 330 organic nitrates is more relevant with NOx levels.

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