The Important Contribution of Secondary Formation and Biomass Burning to Oxidized Organic Nitrogen (OON) in a Polluted Urban Area: Insights from In Situ FIGAERO-CIMS Measurements

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Abstract. To investigate the sources and formation mechanism of oxidized organic nitrogen (OON), field measurements of OON

- were conducted using an iodide-adduct chemical ionization mass spectrometer equipped with a Filter Inlet for Gases and AEROsols (FIGAERO-CIMS) during fall of 2018 in the megacity of Guangzhou, China. Using levoglucosan as tracer of biomass burning emissions, the results show that biomass burning ($49 \pm 23\%$) and secondary formation ($51 \pm 23\%$) accounted for comparable fractions to the total particle-phase OON (pOON), while $24 \pm 25\%$ and $76 \pm 25\%$ to the gas-phase OON (gOON), respectively, signifying the important contribution of biomass burning to pOON and secondary formation to gOON in this urban area.
- 30 Calculations of production rates of gOON indicated that hydroxyl radical (42%) and nitrate radical (NO₃) (49%) oxidation pathways potentially dominated the secondary formation of gOON. High concentration of NO₃ radical during the afternoon was observed, demonstrating that the daytime NO₃ oxidation might be more important than the previous recognition. Monoterpenes, found to be major precursors of secondary gOON, were mainly from anthropogenic emissions in this urban area. The ratio of secondary pOON to O_x ([O_x] = [O₃] + [NO₂]) increased as a function of relative humidity and aerosol surface area, indicating that heterogeneous
- 35 reaction might be an important formation pathway for secondary pOON. Finally, the highly oxidized gOON and pOON with 6 to 11 oxygen atoms were observed, highlighting the complex secondary reaction processes of OON in the ambient air. Overall, our results improve the understanding of the sources and dynamic variation of OON in the urban atmosphere.

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1 Introduction

Oxidized organic nitrogen (OON, including organic nitrates (ONs) and nitroaromatics), acting as an important reservoir of

40 atmospheric nitrogen oxides ($NO_x = NO + NO_2$) (Fisher et al., 2016; Romer Present et al., 2020; Romer et al., 2016; Ditto et al., 2022), substantially influence the NO_x cycling, formation of ozone (O_3) (Farmer et al., 2011; Perring et al., 2013), and secondary organic aerosol (SOA) (Lee et al., 2016; Rollins et al., 2012), thus affect air quality, climate, and ecosystem nutrient cycling (Kiendler-Scharr et al., 2016; Pye et al., 2015). A comprehensive and in-depth understanding of dynamic variations of *in situ* OON (including in gas phase (gOON) and particle phase (pOON)) and their sources is crucial for accurately assessing their environmental

45 impacts.

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With the rapid development of measurement techniques, high-time resolution measurement of OONs has become more available. Currently, online measurement of OONs can be conducted by the following routes: (I) by thermodenuder dissociation to NO₂ and then detection by laser-induced fluorescence (TD-LIF) (Day et al., 2002; Rollins et al., 2010) or cavity-related spectroscopy (Keehan et al., 2020; Sadanaga et al., 2016); (II) by using aerosol mass spectrometer (AMS) (Decarlo et al., 2006) based on 50 NO_2^+/NO^+ apportionment (Farmer et al., 2010; Fry et al., 2013; Hao et al., 2014; Day et al., 2022; Xu et al., 2015) and/or thermodenuder (Xu et al., 2021b); and (III) by using chemical ionization mass spectrometer (CIMS) with different ionization sources, typically with iodide-adduct chemistry (Huang et al., 2019; Lee et al., 2014; Lee et al., 2016) or extractive electrospray ionization (Bell et al., 2021; Lopez-Hilfiker et al., 2019; Pospisilova et al., 2020). Although the first two methods can quantify nitrate functional group (-ONO₂ or -NO₂) in bulk; CIMS, by taking advantage of soft ionization, can provide information on molecular compositions 55 and better comprehend particle-phase ONs (pON) and nitroaromatics at the molecular level (Lee et al., 2014; Pospisilova et al., 2020: Wang et al., 2020b; Salvador et al., 2021). In general, nitroaromatics have also been included in the quantification of ONs by CIMS under negative ionization mode, due to the difficulty encountered in distinguishing the nitro functional group $(-NO_2)$ from – ONO₂ and -NO₂ groups based solely on chemical formulas of ions (Huang et al., 2019). So far, gOON and pOON (containing 4-12 oxygen atoms) formed from multiple oxidation process of volatile organic compounds (VOCs) have been quantified by a high-60 resolution time-of-flight CIMS installed with a Filter Inlet for Gases and AEROsols (FIGAERO-CIMS) in the forests (Lee et al., 2018; Lee et al., 2016) and at rural sites (Huang et al., 2019; Chen et al., 2020). However, limited measurement results were reported in the polluted urban areas (Le Breton et al., 2019).

Both primary emission and secondary formation can contribute to mass concentrations of ambient OON. Biomass burning and/or fossil fuel combustion have been suggested to be important primary emission sources of gOON (Liu et al., 2017; Palm et al., 2020; Peng et al., 2021) and pOON (Gaston et al., 2016; Mohr et al., 2013; Wang et al., 2019; Zhang et al., 2016). Furthermore,

secondary formation of OON in biomass burning plumes has also been observed. For example, Juncosa Calahorrano et al. (2021)

observed the existence of gOON in aged plumes of wildfires. Kodros et al. (2020) showed that pOON could not only be directly emitted from laboratory-generated biomass burning emissions but also formed quickly through nitrate radical (NO₃) oxidation within biomass burning plumes. Based on aircraft measurements, Palm et al. (2020) found that the VOCs and vapors evaporated from

- 70 primary biomass burning could be quickly subjected to radical-driven oxidation, thus contributing to the formation of SOA including nitroaromatics. For the secondary formation pathway in ambient air, gOON is formed mainly through the oxidation of VOCs by hydroxyl radical (OH), NO₃, and ozone in the presence of NO_x (Ng et al., 2017; Perring et al., 2013). Functionalization of gOON in ambient air reduces their volatility, leading to condensation of gOON on particles to form secondary pOON (Capouet and Müller, 2006).
- 75 Previous studies indicated that the oxidation of biogenic VOCs by NO₃ dominated gOON formation at a forest-urban site in Germany (56% of average gOON production rate) (Sobanski et al., 2017), as well as at a boreal forest site in Finland (70% of total gOON production rate) (Liebmann et al., 2019) and the southeast US (84% of monoterpene organic nitrate mass) (Ayres et al., 2015; Pye et al., 2015; Xu et al., 2015). For urban areas, the contributions of the above-mentioned three secondary formation pathways to total gOON remain poorly understood (Yu et al., 2019). Initiation of oxidation by OH under high NO_x condition is 80 traditionally regarded as the main formation pathway for urban OONs during the day (Perring et al., 2013). However, Hamilton et al. (2021) recently found that a large fraction of isoprene-derived OONs was formed through unexpected NO₃ oxidation pathway in the afternoon in the Beijing urban area. Thus, a better understanding of the OON sources and formation mechanism in urban areas
 - is still needed.

In this study, quantitative measurements of gOON and pOON were carried out using a high-resolution time-of-flight 85 FIGAERO-CIMS and an AMS in a Chinese megacity. The contributions of biomass burning and secondary formation to ambient total gOON and pOON measured by CIMS were quantified, and the secondary oxidation pathways were systematically explored based on the production rates of gOON. Finally, the molecular compositions of ambient OON measured by CIMS were comprehensively investigated.

2 Experimental Methods

90 2.1 Sampling site.

Measurements were conducted on the campus of Guangzhou Institute of Geochemistry, Chinese Academy of Sciences (23.14°N, 113.36°E) in the urban area of megacity Guangzhou, during the coordinated campaign "Particles, Radicals, and Intermediates from oxidation of primary Emissions over the Great Bay Area" (PRIDE-GBA) (Wu et al., 2020). The observation site is located 25 m above the ground on the ninth floor of the highest building on the campus. The campus was surrounded with 95 industrialized and urbanized downtown areas in a typically subtropical climate, thus strongly influenced by both anthropogenic and biogenic emissions, as shown in Fig. S1. The average ambient temperature and relative humidity (RH) during the campaign were 23.7 ± 2.9 °C and $71.9 \pm 17.4\%$, respectively. The site was mostly affected by northerly winds with an average speed of 4.5 ± 2.18 m s⁻¹.

2.2 Measurement and analysis.

100 2.2.1 Operation of FIGAERO-CIMS

- During the campaign, a CIMS installed with a long time-of-flight detector (10000 < m/∆m < 11000) with an iodide source (Aerodyne Research Inc, USA) was deployed (Lee et al., 2014; Wang et al., 2020d). The FIGAERO inlet was installed with the CIMS to measure speciated gOON and pOON (Lopez-Hilfiker et al., 2014; Bannan et al., 2019; Schobesberger et al., 2018; Thornton et al., 2020). The detailed performance and calibration information of the CIMS can be found in a recent paper about this campaign (Ye et al., 2021). A brief description is introduced here. The sampling flow rate is 3.8 L min⁻¹ for gas sampling line and 5 L min⁻¹ for particle line. A PM_{2.5} cyclone inlet and a nafion dryer (Perma Pure, model PD-07018T-12MSS) were set ahead of the particle sampling inlet of the FIGAERO to keep the filter for aerosol sampling not getting wet due to the high ambient RH (72 ± 17%) in this campaign. Recent studies show that aerosol in equilibrium with semi-/intermediate-volatility organic compounds (S/IVOCs) will be perturbed by the removal of the gases by the nafion (Liu et al. 2019). However, in this study, the retention time for particles
- 110 through the nafion dryer was ~0.12 s, which might lead to a very small change of S/IVOCs signal on such a timescale (< a few percents) based on the partitioning delay model (Pagonis et al., 2017). In addition, an accurate correction for S/IVOCs loss in nafion dryer is also not available in current (Liu et al. 2019). Thus, no S/IVOCs correction on aerosol phase was performed in this study.</p>

In general, to measure the gOON and pOON, the FIGEARO was operated alternately at two main stages during the measurement: (i) For the first 24 min in one hour cycle, ambient air was continuously sampled into two inlets, i.e., gas and particle inlets. The gas inlet was connected to an ion-molecule reaction region (IMR) of the CIMS. An X-ray source was used in this campaign, which has lower ionization efficiency compared to the polonium-210 radioactive source used in the previous studies (Faxon et al., 2018; Lee et al., 2021; Palm et al., 2019). Therefore, a higher pressure (370–390 mbar) in the IMR than in previous studies (e.g., 93–200 mbar) (Faxon et al., 2018; Lee et al., 2021; Palm et al., 2021; Palm et al., 2019) was used to achieve similar strength of reagent ions in the CIMS system. The sampled VOCs were first ionized in the IMR (VOC•I⁻), in which the primary ions were generated by flowing 2 mL/min 1000 ppm methyl iodide in 2.4 L/min N₂ through the X-ray source, and then moved into the mass spectrometer for measurement at a resolution of 1 s. Moreover, the ambient air was also introduced in the particle sampling inlet where a sliding Teflon tray held a polytetrafluoroethylene membrane filter (Zefluor®, Pall Inc., USA) for aerosol collection for 24 min. (ii) Next, after 24 min when the gas-phase measurement was completed, a linear actuator was used to move the Teflon tray on which the filter

was placed in front of the IMR, while the gas phase inlet was blocked. Then, ultrahigh-purity N₂ gas at a flow rate of 2 L min⁻¹ was
passed through a stainless steel "heating tube" to thermally desorb the collected particles on the filter into gas phase, and then into the IMR and mass spectrometer for measurement. The temperature of N₂ flow was ramped from room temperature to 175 °C in 12 min and then kept at 175 °C for another 20 min. The temperature of the IMR was kept almost constant by setting the temperature constant (80 °C) of the heater strip in the IMR. Meanwhile, the room temperature, which was maintained by an air-conditioner, was relatively stable (23.7 ± 2.9 °C). The gas sampling line inside the room was covered by heat insulation associated with a heating
cable to hold the temperature of sampling gas steady. These protocols reduce the effect of the temperature dependence of IMR, as indicated by Robinson et al. (2022) that I⁻ CIMS sensitivity may be influenced by the temperature of IMR.

The background signal of the gas-phase measurement was determined by pure N₂ signal at the last 3 min within the 24min sampling time (Palm et al., 2019). The background signal of the particle measurement was determined by the measured signals from every sixth 1-h running cycle, in which particle-free air was obtained with ambient air passing through a High-Efficiency Particulate Air (HEPA) filter set ahead of the FIGAERO filter (Ye et al., 2021). The TofWare software (version 3.0.3) was used to perform high-resolution peak fitting of the CIMS mass spectra.

2.2.2 Oxidized Organic nitrogen quantification based on CIMS measurement

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Based on the CIMS measurement, speciated OON (nitrogen-containing oxygenated hydrocarbons, 339 closed-shell compounds with oxygen versus carbon atom ratio no less than 3, $C_{\geq 1}H_{\geq 1}O_{\geq 3}N_{1-2}$) in both gas and particle phases were quantified. These OON compounds can be fitted well in the HR analysis after the background signals have been removed. In this study, nitroaromatics were also a subset of OON due to (I) their similar chemical and optical properties as the bulk OON compounds (He et al., 2021; Lin et al., 2017) and (II) interferences between ONs and nitroaromatics in the CIMS measurement due to desorption and fragmentation (Ye et al., 2021). The composition of OON was mainly contributed by the CHON (one nitrogen atom containing species), and the CHON₂ (two nitrogen atom containing species) only contributed 6.8% of the gOON and 8.3% of the pOON.

- 145 For quantification, 39 species in total including levoglucosan (C₆H₁₀O₅), 4-nitrophenol (C₆H₅NO₃), 2,4-dinitrophenol (C₆H₄N₂O₅), and 4-nitrocatechol (C₆H₄NO₄) were calibrated with standard compounds, where the effect of humidity on the sensitivities was also accounted for (Ye et al., 2021). Their calibration factors were shown in the excel file of the supplement zip package of Ye et al. (2021). For other uncalibrated species, a voltage scanning procedure was carried out every few days throughout the campaign to determine their sensitivities (including ON species) (Bi et al., 2021a, b; Lopez-Hilfiker et al., 2016). Lopez-Hilfiker et al. (2016) and Iyer et al. (2016) have verified the connections among the binding energy of the iodide-adduct bond, the voltage
- dissociating iodide adducts and the sensitivity of corresponding species. The relationship between the voltage difference (dV) and

signal fraction remaining of an iodide-molecule adduct is established by scanning the dV between the skimmer of the first quadrupole and the entrance to the second quadrupole ion guide of the mass spectrometer. This relationship curve of an individual iodide adduct can be fitted by a sigmoid function and yields two parameters: S₀, the relative signal at the weakest dV compared to the signal under

of all the uncalibrated OON species was determined, after taking into account the relative transmission efficient of all the ions. The

- 155 operational dV; dV_{50} , the voltage at which half of the maximum signal is removed (i.e., half the adducts that could be formed are declustered). A sigmoidal fit was then applied to the results of all the iodide adducts. An empirical relationship between relative sensitivity (1/S₀) and dV_{50} of each ion (includes levoglucosan) based on average values of the entire campaign was obtained. By linking the relative sensitivity of levoglucosan with its absolute sensitivity based on the authentic standard, the absolute sensitivity
- detailed data of these response factors can be found in the supporting information of Ye et al. (2021). Three OON species which are 4-nitrophenol (C₆H₅NO₃), 2,4-dinitrophenol (C₆H₄N₂O₅), and 4-nitrocatechol (C₆H₅NO₄) were calibrated in both authentic standards and voltage scanning methods. By comparing their sensitivity (Fig. S3), the uncertainty of the voltage scanning method can be roughly estimated. Detailed description of the calibration curves and the application of the calibration curve to estimate the sensitivity can be refferred to the supporting information text of Ye et al. (2021). In general, the voltage scanning method underestimates (32–56%) the sensitivity of OON in this study compared to the values using the standard compounds as real. This uncertainty was comparable with 30% uncertainty of all analytes in Bi et al. (2021b) and 60% uncertainty of total carbon in Isaacman-Vanwertz et al. (2018) measured by the Iodide-CIMS. Finally, an avergae underestimation of 47% on sensitivity was taken as the uncertainty of the whole OON mass loading in this study.
- In summary, nitroaromatics, i.e., nitrophenol (C₆H₅NO₃), methyl nitrophenol (C₇H₇NO₃), dinitrophenol (C₆H₄N₂O₅), nitrocatechol (C₆H₅NO₄), methyl nitrocatechol (C₇H₇NO₄), and nitrosalicylic acid (C₇H₅NO₅), which were assumed to be identified with the ions containing the same molecule compositions detected by the CIMS (Wang et al., 2018; Wang and Li, 2021; Chen et al., 2021b), accounted for 18% and 5% of total gOON and pOON mass concentrations, respectively. Some nitroaromatic signal may be detected as elemental formulas other than those listed above, and some of the signal at the elemental formulas identified here as nitroaromatics may have contribution from ON species. While the uncertainty exists, it is likely that ONs dominated the OON observed during this campaign. For the total OON, the Iodide-CIMS may underestimate or poorly detect some types of OON, e.g., simple alkyl or keto nitrates (Lee et al., 2016). Moreover, the thermal fragmentation reactions that result from heating on the FIGAERO filter may also lead to underestimation of OON due to the loss of the nitrogen-containing groups, such as peroxy nitrates, which have the propensity to thermally dissociate into NO₂ and other non-nitrogen-containing species (Lee et al., 2016).

In the ambient air, the $C_6H_{10}O_5$ measured in the particle phase using the CIMS was probably composed by levoglucosan and its isomers (mannosan and galactosan) (Ye et al., 2021). The isomer measurement of $C_6H_{10}O_5$ in this campaign have revealed that the levoglucosan contributed $90 \pm 2\%$ mass loading of the three isomers of C₆H₁₀O₅ (Jiang et al., 2023), thus C₆H₁₀O₅ signal in this study probably can be used as a tracer for biomass burning emission (Bhattarai et al., 2019). The good correlation (R=0.78) between C₆H₁₀O₅ and another biomass burning tracer potassium (K⁺) (Wang et al., 2017c; Andreae, 1983), also supports this statement (Fig. S11a). Multiple studies show that levoglucosan might be degraded due to photochemistry (Lai et al., 2014; Bai et

al., 2013; Hennigan et al., 2010). We calculated the ambient photochemical age based on the ratios of two hydrocarbons (m+p-xylene and ethylbenzene) that react at different rates with OH radicals (Yuan et al., 2013; Wu et al., 2020; De Gouw et al., 2005). A daily average OH concentration of 1.5×10⁶ molecule cm⁻³ was assumed here (Mao et al., 2009; Wang et al., 2020d; Chen et al., 2021a). The estimated results show that the average diurnal photochemical age varied from 0.2 days during the night to maximum 0.5 days in the daytime in this campaign (Chen et al., 2021a), which was lower than the lifetime of levoglucosan (>1 day -26 days)
determined in laboratory and field studies (Hennigan et al., 2010; Hoffmann et al., 2010; Lai et al., 2014; Bai et al., 2013; Bhattarai et al., 2019). It suggests that the levoglucosan observed in this study shall be stable for being the tracer of biomass burning emissions.

2.2.3 Other instruments

Besides the CIMS, a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS, Aerodyne Research Inc., hereinafter referred to as "AMS") was used to provide online quantitative measurement of submicron non-refractory aerosols (PM₁) 195 at a time resolution of 4 min (Canagaratna et al., 2007; Decarlo et al., 2006). In addition to the total organic aerosol (OA), the mass concentration of -ONO₂ group from pON (pOrgNO_{3, AMS}) was also estimated by NO₂^{+/} NO⁺ ratio method (Farmer et al., 2010; Fry et al., 2013; Day et al., 2022; Xu et al., 2015), positive matrix factorization (PMF) method (Hao et al., 2014), and thermodenuder (TD) method (Xu et al., 2021b) based on the AMS data. The NO_{2}^{+}/NO^{+} ratio method was based on the different ratios of NO_{2}^{+} to NO⁺ fragmented from pOrgNO_{3, AMS} and inorganic nitrate. The PMF method was performed by including the NO⁺ and NO₂⁺ ions 200 into the PMF analysis combined with spectral matrix of organic ions. The TD method was conducted based on the difference of volatility between pOrgNO3, AMS and inorganic nitrates in particles. Detailed description and intercomparison of these three methods can be found in the supporting information (Text S1), where some insights into the pros and cons of the AMS-based methods are also presented. The comparative analysis among the CIMS and these three methods by the AMS aided in evaluating the measurement accuracy of OON in this study. Herein, pON estimated by NO_2^+/NO^+ ratio method was selected as representative data for the 205 following discussion due to the better performance achieved by this method. More detailed information on calibrations and operations of the AMS during this campaign can be found elsewhere (Chen et al., 2021a).

VOCs were measured by online gas chromatography-mass spectrometry and using a flame ionization detector (GC-MS/FID) (Wuhan Tianhong Instrument Co., Ltd.) at a time resolution of 1 h and by proton transfer reaction time-of-flight mass spectrometry (PTR-ToF-MS, IONICON Analytik) at a time resolution of 10 s (Wu et al., 2020; Yuan et al., 2017). A 56-component

VOC gas standard was used for daily calibration of the GC-MS/FID (Wang et al., 2020c). For the PTR-ToF-MS, a 16-component VOC gas standard was used for daily calibration under both dry (RH < 1%) and ambient humidity during the whole campaign, and an additional 23-component VOC gas standard was used during the last period of the campaign (Wang et al., 2020a; Wu et al., 2020). The uncertainties for the VOC measurements by both instruments were below 20%. Trace gases, i.e., O₃ (TL43i), NO/NO₂ (TL42i), and CO (TL48i), were measured using Thermo Fisher Scientific instruments at a time resolution of 1 min (Wang et al., 2020d). Meteorological parameters were measured on a Vantage Pro2 weather station (Davis Instruments) at a time resolution of

10 s.

2.3 Calculation of the production rates of gas-phase oxidized organic nitrogen.

The production rates of gOON from VOCs oxidized by OH, NO₃, and O₃ were calculated using the reactant concentrations and reaction rate coefficients combined with formation branching ratios and yields (Liebmann et al., 2019), of which the detailed calculation process can be found in Text S2. In this calculation, the production rates mainly from organic nitrates were shown. The parameters for secondary nitroaromatics are not available, thus, was not included here. The concentrations of VOCs and O₃ were obtained from direct measurements, while those of OH radicals were derived from a box model simulation with the Master Chemical Mechanism v3.3.1 (MCM v3.3.1) (Wang et al., 2020c; Wolfe et al., 2016). The NO₃ radical was calculated based on the measured N₂O₅ by the CIMS (Ye et al., 2021) based on temperature equilibrium between these two species (Brown and Stutz, 2012; Chen et al., 2022). The remaining parameters were obtained from the previous studies (Liebmann et al., 2019; Perring et al., 2013). The different VOC species and corresponding parameters are listed in Table 1. An overall uncertainty of 56% was estimated by the Monte Carlo method through 10,000 calculations in this method. The detailed uncertainties of different parameters can be found in Text S3.

Table 1. The VOC species and their average mass concentrations with standard deviations; The reaction rate coefficients, branch ratios for OH and O₃ pathway and yields for NO₃ pathway used for the calculations of gOON production rates.

OH-initiated pathway VOC species	Average concentration (ppb)	k_{OH} at 298K (cm ³ molecules ⁻¹ s ⁻¹)	α^{RO_2}
Isoprene	0.15 ± 0.17	$1.00 imes10^{-10}$	0.070
<i>d</i> -Limonene ^a	0.07 ± 0.07	$1.70 imes10^{-10}$	0.230
α -pinene ^a	0.07 ± 0.07	$5.30 imes 10^{-11}$	0.180
Propane	6.23 ± 4.92	$1.09 imes 10^{-12}$	0.036
iso-Butane	1.56 ± 1.27	2.12×10^{-12}	0.096
n-Butane	2.80 ± 2.35	$2.36 imes 10^{-12}$	0.077
Cyclopentane	0.09 ± 0.05	$4.97 imes 10^{-12}$	0.045
iso-Pentane	1.17 ± 1.01	$3.60 imes10^{-12}$	0.070
n-Pentane	0.65 ± 0.65	$3.80 imes 10^{-11}$	0.105
2,2-Dimethylbutane	0.03 ± 0.02	$2.23 imes10^{-12}$	0.152
2,3-Dimethylbutane	0.05 ± 0.05	$2.23 imes10^{-12}$	0.152
2-Methylpentane	0.26 ± 0.27	$5.20 imes 10^{-12}$	0.097
3-Methylpentane	0.25 ± 0.25	$5.20 imes10^{-12}$	0.109

		12		
n-Hexane	0.50 ± 0.75	$5.20 imes 10^{-12}$	0.141	
2,4-Dimethylpentane	0.03 ± 0.03	3.34×10^{-12}	0.140	
Methylcyclopentane	0.09 ± 0.09	$5.60 imes 10^{-12}$	0.140	
Cyclohexane	0.05 ± 0.05	$6.97 imes 10^{-12}$	0.160	
n-Heptane	0.09 ± 0.15	$6.76 imes 10^{-12}$	0.178	
Methylcyclohexane	0.07 ± 0.09	$9.64 imes 10^{-12}$	0.170	
n-Octane	0.04 ± 0.05	$8.11 imes 10^{-12}$	0.226	
Nonane	0.03 ± 0.03	$9.70 imes 10^{-12}$	0.393	
n-Decane	0.02 ± 0.02	$1.10 imes 10^{-11}$	0.417	
Benzene	0.43 ± 0.16	$1.22 imes 10^{-12}$	0.034	
Toluene	1.75 ± 1.86	$5.96 imes 10^{-12}$	0.029	
Ethylbenzene	0.28 ± 0.30	$7.00 imes10^{-12}$	0.072	
m-p-Xylene	0.79 ± 0.82	$2.30 imes 10^{-11}$	0.074	
o-Xylene	0.29 ± 0.31	$1.36 imes 10^{-11}$	0.081	
Isopropylbenzene	0.01 ± 0.01	$6.30 imes 10^{-12}$	0.110	
m-Ethyltoluene	0.03 ± 0.03	$1.86 imes10^{-11}$	0.094	
p-Ethyltoluene	0.02 ± 0.02	$1.18 imes 10^{-11}$	0.137	
1,3,5-Trimethylbenzene	0.02 ± 0.02	$5.76 imes 10^{-11}$	0.031	
o-Ethyltoluene	0.02 ± 0.02	$1.19 imes 10^{-11}$	0.106	
1,2,4-Trimethylbenzene	0.06 ± 0.06	$3.25 imes 10^{-11}$	0.105	
1,2,3-Trimethylbenzene	0.02 ± 0.01	$3.25 imes 10^{-11}$	0.119	
Propene	0.37 ± 0.37	$2.63 imes 10^{-11}$	0.015	
trans-2-Butene	0.03 ± 0.03	$6.40 imes 10^{-11}$	0.034	
1-Butene	0.07 ± 0.05	$3.14 imes 10^{-11}$	0.025	
cis-2-Butene	0.02 ± 0.02	$5.64 imes 10^{-11}$	0.034	
1-Pentene	0.03 ± 0.02	$3.14 imes 10^{-11}$	0.059	
trans-2-Pentene	0.01 ± 0.02	$6.70 imes 10^{-11}$	0.064	
cis-2-Pentene	0.01 ± 0.01	$6.50 imes 10^{-11}$	0.064	
1-Hexene	0.02 ± 0.01	$3.70 imes 10^{-11}$	0.055	
	A	k at 208K		
NO ₃ -initiated pathway VOC species	Average	k_{NO_3} at 298K	$lpha_i$	
	concentration (ppb)	$(cm^3 molecules^{-1} s^{-1})$		
Isoprene	0.15 ± 0.17	$6.95 imes 10^{-13}$	0.700	
<i>d</i> -Limonene	0.07 ± 0.07	1.22×10^{-11}	0.670	
α-pinene	0.07 ± 0.07	6.21×10^{-12}	0.150	
Phenol	0.04 ± 0.03	3.92×10^{-12}	0.251	
Cresol	0.03 ± 0.03	1.37×10^{-11}	0.128	
Styrene	0.17 ± 0.26	1.50×10^{-12}	0.251 ^b	
	Average	k_{0_3} at 298K		
O ₃ -initiated pathway VOC species	concentration (ppb)		α^{O3}	
.		$\frac{(\text{cm}^3 \text{ molecules}^{-1} \text{ s}^{-1})}{1.20 \text{ molecules}^{17}}$	1.000	
Isoprene	0.15 ± 0.17	1.28×10^{-17}	1.000	
d-Limonene	0.07 ± 0.07	2.20×10^{-16}	0.750	
α -pinene	0.07 ± 0.07	$9.40 imes10^{-17}$	0.800	

Note that all the reaction rate coefficients and the formation branching ratios or yields are from MCM v3.3.1 and previous studies (Perring et al., 2013; Liebmann et al., 2019; Atkinson and Arey, 2003). ^a We assumed total monoterpenes measured from PTR-MS are composed of *d*-limonene and α -pinene with a ratio of 1:1 based on the anthropogenic origins of monoterpene, as discussed in text S2. ^b The parameters were assumed to be equal to that of phenol.

235 **3 Results and Discussion**

3.1 Quantification and chemical composition of oxidized organic nitrogen

Fig. 1a shows a subset of the time series of pOON measured using the CIMS (i.e., pOON_{CIMS}) and pOrgNO_{3, AMS} (-ONO₂

+ NO₂ group) measured with the AMS. Total OA measured with the AMS is also shown in Fig. 1a. During the entire campaign, a

moderate correlation (Pearson correlation coefficient, R = 0.42, Fig. S6) was observed between pOON_{CIMS} and pOrgNO_{3, AMS}, which
is probably due to the high uncertainty of pOrgNO_{3, AMS} estimation from the AMS when organic nitrate fraction in total nitrate signal is low (<10%, the details can be found in Text S1), as well as is probably due to the fact that only -ONO₂/NO₂ groups not entire ON molecule was measured with AMS. The description of the correlation coefficient (R) within this study is defined based on the interpretation by Dancey and Reidy (2007), as shown in detail in Text S1. When the mass concentration of total nitrates from the AMS is below 5 µg m⁻³ (corresponding to pOrgNO_{3, AMS} signal fraction in total nitrate signal >60%), an improved correlation between pOON_{CIMS} and pOrgNO_{3, AMS} is found (R = 0.86, Fig. 1b), validating the robustness of pOON_{CIMS} measured here. A similar moderate correlation between pOrgNO_{3, AMS} and pOON_{CIMS} was also observed at a rural site in the southwest Germany (R = 0.52) (Huang et al., 2019), and a much better agreement (R = 0.82) was obtained in the southeast US when the pOrgNO_{3, AMS} fraction in total nitrate is above 70% (Lee et al., 2016; Xu et al., 2015).

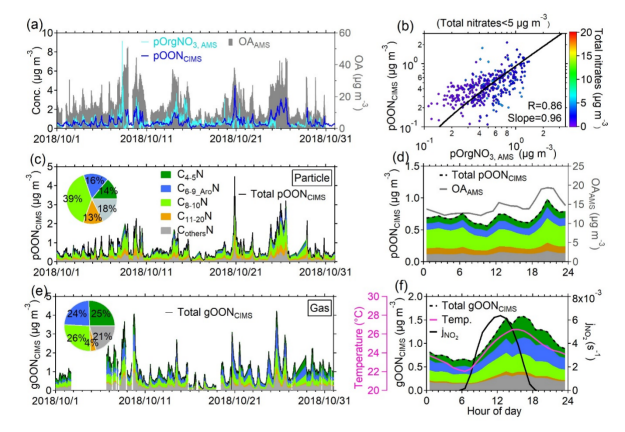


Figure 1. Time series and variations of OON during the PRIDE-GBA campaign. (a) Time series of pOON_{CIMS} and pOrgNO₃, AMS. Time series of total OA detected by the AMS is shown on the right axis. (b) Scatterplot of pOON_{CIMS} versus pOrgNO₃, AMS during the campaign. The term "total nitrates <5 μg m⁻³" indicates the data used in this scatterplot is under the condition that the mass concentration of total nitrates (including organic nitrate and inorganic nitrate) measured by the AMS is lower than 5 μg m⁻³. The points are color-coded using the total nitrate signals measured by the AMS. The scatterplot from all AMS and CIMS measurement can be found in Fig. S6. The logarithm was applied to both of the axes. Time series of (c) pOON_{CIMS} and (e) gOON_{CIMS}, as well as the time series of their C_xN groups from the CIMS measurement. The insets show their average

mass contributions to total gOON_{CIMS} and pOON_{CIMS} during the campaign, respectively. The average diurnal variations of (d) pOONCIMS and its CxN groups, as well as OA; (f) Average diurnal variations of total gOONCIMS and its CxN groups, photolysis rate of NO₂ (j_{NO2}), and temperature during the entire campaign. All the diurnal variations calculated throughout the manuscript are based on the average values. All the linear fitting are based on the orthogonal distance regression (ODR)

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algorithm in this study. All the acronyms can be found in appendix A.

The average concentrations of the gOON_{CIMS} and pOON_{CIMS} measured using the CIMS were 1.00 ± 0.67 and 0.66 ± 0.53 μ g m⁻³, respectively. Moreover, an average concentration of 0.60 ± 0.46 μ g m⁻³ for pOrgNO_{3 AMS} during the campaign was obtained. Notably, different size cuts between the AMS (PM₁) and the CIMS (PM_{2.5}) should only play a minor role in the quantification of 265 pOON, as the measured total aerosol mass concentrations of PM_{2.5} and PM₁ during the campaign are very similar (a regression slope of 0.96) (Chen et al., 2021a). The average mass-weighted chemical compositions for gOON_{CIMS} and pOON_{CIMS} observed in this campaign were determined to be $C_{6.6}H_{9.4}N_{1.1}O_{5.3}$ and $C_{8.5}H_{12.2}N_{1.1}O_{6.5}$, respectively, corresponding to molecular weight (MW) of 189 ± 7.8 and 234 ± 7.9 g mol⁻¹. The molecular weight of pOON_{CIMS} in this study (234 ± 7.9 g mol⁻¹) is comparable to the reported values in forest (256 g mol⁻¹) and rural (220 and 296 g mol⁻¹) sites (Lee et al., 2018; Huang et al., 2019; Chen et al., 2020). If such 270 average MW of pOON_{CIMS} was applied, the pOON_{AMS} (in addition to -ONO₂/-NO₂ groups, the organics part was also accounted for) would be 2.3 μ g m⁻³ based on multiplying a factor of 3.8 to pOrgNO_{3. AMS}, which is well within the range of 0.06–2.94 μ g m⁻³ of pOONAMS as reported in the previous studies around the world, as shown in Fig. S7. On the other side, if only -ONO2/-NO2 groups are considered to calculate pON_{CIMS} to be pOrgNO_{3, CIMS}, the calculated pOrgNO_{3, CIMS} can explain $28 \pm 18\%$ of pOrgNO₃. AMS, which is consistent with the fraction (23%) of total functionalized OA detected using the CIMS versus total OA measured using the AMS (Ye et al., 2021). The detailed analysis process on comparison uncertainty between AMS and CIMS can be found in Text

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S3 of the supporting information.

For this study, an average mass fraction of $15 \pm 12\%$ of pON_{AMS} in total OA (14.7 ± 8.20 µg m⁻³ on average) measured by the AMS was observed (calculation method referred to Takeuchi and Ng (2019)). In spite of the absolute mass concentrations of pON varied largely in different studied urban environments, the pON/OA ratios are very similar ($15 \pm 3\%$ on average, Fig. S7), suggesting a potential similar pON formation process/fate in the urban areas (Day et al., 2010; Rollins et al., 2012; Xu et al., 2015).

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To further illustrate the contributions of different components of speciated gOON and pOON measured by the CIMS, five C_xN groups, namely (1) $C_{4-5}N$, (2) $C_{6-9 \text{ Aro}}N$, (3) $C_{8-10}N$, (4) $C_{11-20}N$, and (5) $C_{others}N$ (Figs. 1c and 1e) were categorized only based on the number of carbon atoms in the molecules. The C_{6-9 Aro}N group was recognized based on the number of 6–9 carbon atoms and positive aromaticity index (0–1) (Text S4) (Koch and Dittmar, 2016; Wang et al., 2019; Koch and Dittmar, 2006). C_{11–20}N group contains species with large carbon backbones from 10 to 20 and/or oligomers, CothersN includes the remaining nitrogen-containing short-chain ions which are not possible to fit into the previous four categories, such as the ions with carbon atoms less than 4 (e.g., C₃H₇NO₄ and C₃H₇NO₅) and the ions with 6–9 carbon atoms excluded in C_{6-9 Aro}N group (e.g., C₆H₉NO₃ and C₇H₁₁NO₅). In general, it is observed that the contributions of C₄₋₅N (25%) and C_{6-9 Aro}N (24%) to total gOON are higher than those to total pOON (14% and 16%, respectively), while C₈₋₁₀N (39%) and C₁₁₋₂₀N (13%) contributed more to total pOON than to total gOON (26% and 4%, respectively). These results are consistent with the lower volatility for C₈₋₁₀N and C₁₁₋₂₀N than those for C₄₋₅N and C_{6-9 Aro}N due to the longer backbones of compounds in the former groups (Kroll and Seinfeld, 2008; Odum et al., 1996). In general, the time series of all groups for the gOON shows similar variability as the pOON (correlation coefficient R > 0.6), except for C₄₋₅N groups that show a positive correlation of R = 0.31 (Fig. S8). The slightly poor correlation of C₄₋₅N groups between gas and aerosol phase was probably caused by less partitioning of substantial formed isoprene-oxidized gOON in the daytime to the pOON compared to other
long-chain compounds. The regression slope between gOON and pOON of each category (3.61 to 1.80) decreases with the increase of the carbon number, as shown in Fig. S8, which is reasonable considering their gas/particle partitioning balances (Odum et al., 1996).

The overall average diurnal variations of gOON and pOON and their C_xN groups are shown in Figs. 1d and 1f. Despite the boundary layer expansion in the daytime (Fig. S9a), the gOON peaks in the afternoon and drops slowly with tail toward the night. The enhancement of the gOON increases with the j_{NO2} (Fig. 1f), indicating that daytime secondary formation is an important source of gOON (Sobanski et al., 2017). The primary biomass burning (e.g., levoglucosan as a tracer) and vehicle emissions (e.g., NO/NO_x as tracers), which are also potential sources for gOON, usually show enhancement during nighttime (levoglucosan and isomers in Fig. S11f and NO/NO_x in Fig. S9b) and morning rush hour time (NO/NO_x only), respectively, in their diurnal variations, indicating that both primary sources are unlikely to contribute to the gOON enhancement during daytime. In contrast to the gOON, the pOON and nechanisms for the pOON compared to the gOON, e.g., biomass burning contribution during nighttime (Rollins et al., 2012). A more detailed analysis about the sources of the gOON and pOON is presented in the next section.

In general, the average diurnal concentration of each gOON (pOON) group shows a similar trend (Figs. S9d and S9g). The fraction of C₄₋₅N group in gOON enhances slightly during daytime (Fig. S9e), which might be due to strong photochemical formation of isoprene-nitrates (Fisher et al., 2016; Reeves et al., 2020; Mayhew et al., 2022; Hamilton et al., 2021). The C₈₋₁₀N fraction in total pOON slightly enhances during nighttime (Fig. S9h), which was probably due to the primary emissions and the formation of monoterpene-nitrates by NO₃ oxidation chemistry (Peng et al., 2021; Fisher et al., 2016).

3.2 Source apportionment of oxidized organic nitrogen.

- To elucidate the sources of OON measured using the CIMS, the correlations between OON and a wide range of trace 315 species representing different sources were explored to filter the best tracer for source apportionment. The scatterplots of gOON and pOON (hereinafter represent the $gOON_{CIMS}$ and $pOON_{CIMS}$) with selected species, including particle-phase $C_6H_{10}O_5$ (levoglucosan and its isomers) measured using the CIMS; m/z 60 measured using the AMS, benzene, NO_x, and CO are shown in Fig. 2 and Fig. S10. It was observed herein that the scatterplots of gOON (and pOON) versus $C_6H_{10}O_5$, the tracer for biomass burning emissions (Li et al., 2021b; Simoneit, 2002; Simoneit et al., 1999), exhibit two different regression slopes during daytime and 320 nighttime (Figs. 2a and 2c). However, different regression slopes were not observed in the scatterplots of OON versus other tracers, e.g., CO, benzene. It suggests the biomass burning which usually peaks during the night might be an important source for OON. The time series of OON, in particular pOON, indeed peak consistently with $C_6H_{10}O_5$ during high concentration episodes (Figs. S12a and S13a), indicating biomass burning emissions contributed substantially to OON during this campaign. For other biomass burning tracer, e.g., m/z 60, did not show separated regression slope with OON, which is probably due to elevated background of m/z 60 325 contributed by non-biomass burning emissions (Cubison et al., 2011; Mohr et al., 2009). Another two potential biomass burning tracers, i.e., $C_7H_8O_2$ (methoxyphenol and its isomers) and $C_8H_8O_4$ (vanillic acid and its isomers), the former of which exhibits relatively low concentration and the latter shows larger background than $C_6H_{10}O_5$ (levoglucosan and its isomer) (Fig. S11), are both not the ideal biomass burning tracers in this study.
- In contrast to that $C_6H_{10}O_5$ and OON peak consistent with each other, the time series of OON did not peak during the episodes with strong influences of vehicle emissions (as indicative of high NO and NO_x concentrations >50 ppb) (Harrison et al., 2003; Wormhoudt et al., 2015), as shown in Figs. S14a and S15a. In addition, the anti-correlations between OON and NO/NO_x during some of these episodes were even found, indicating that vehicle emission is not a significant source of primary OON. Furthermore, it was found that the diurnal variation of pOON peaks one-hour earlier than NO_x in the morning time, thus, supports their different origins (Fig. S15g). The coincidence of the peaking time between pOON and NO_x during nighttime is probably more influenced by biomass burning. Another piece of evidence is that multiple laboratory studies found negligible emission of gOON from emission tests of vehicle exhaust based on Iodide-CIMS direct measurement (Le Breton et al., 2019; Li et al., 2021a). Thus, the biomass burning emissions and secondary formation should be the main sources of OON observed in this campaign.

To quantify the contributions of biomass burning (OON_{bb}) and secondary formation (OON_{sec}) to OON in both gas and particle phases, the following Eqs. (1) and (2) were proposed to allocate the OON sources:

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$$OON_{bb} = ([OON_{measured}]/[levo.])_{bb} \times [levo.]$$

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(1)

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(2)

Where [levo.] is the concentration of particulate $C_6H_{10}O_5$ measured using the CIMS, ([$OON_{measured}$]/[levo.])_{bb} is the averaged ambient concentration ratio determined from slopes between ambient OON and particulate $C_6H_{10}O_5$ during selected episodes with strong influences of biomass burning (Figs. S12 and S13). This approach relies on a concept similar to the widely-used "elemental carbon tracer method" for source apportionment of primary and secondary organic carbon (Turpin and Huntzicker, 1995). A similar method was used by Salvador et al. (2021) to quantify the sources of nitro-aromatic compounds. The episodes were selected based on the following three criteria: (1) the peak concentration of $C_6H_{10}O_5$ should be above 0.2 µg m⁻³ for selecting periods strongly influenced by biomass burning plumes; (2) the regression coefficient R between gOON (pOON) and $C_6H_{10}O_5$ during each episode should be >0.7; and (3) the number of the fitting points during each episode should be above 4, due to the hourly data of particlephase $C_6H_{10}O_5$ were used. The averaged ambient concentration ratios were determined to be $3.95 \pm 1.67 \ \mu g \ m^{-3}/\mu g \ m^{-3}$ for gOON and $5.05 \pm 1.01 \ \mu g \ m^{-3}/\mu g \ m^{-3}$ for pOON, as presented in Table S1. The variability of [OON_{measured}]/[levo.] ratios from multiple biomass burning episodes for pOON is 20%. The slightly larger ratio uncertainty for gOON (42%) is mainly due to active gas-phase reaction and low contributions of biomass burning emissions to total gOON, as discussed below.

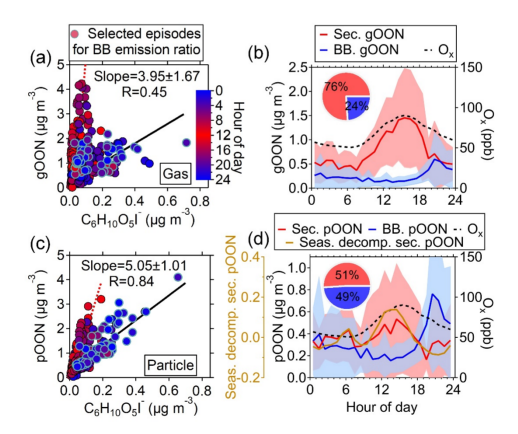


Figure 2. Scatterplots of (a) gOON and (c) pOON versus C₆H₁₀O₅ measured by CIMS with the data points color-coded using hour of day. The blue circles indicate the data points from multiple strongly influenced episodes by biomass burning emission, of which the ratios between gOON or pOON and C₆H₁₀O₅ were used to determine the average ratio (black regression line).

The red dotted line means the line regression during the daytime. The diurnal variations of (b) gOON and (d) pOON from biomass burning (BB.) and secondary formation (sec.). The shaded areas mean the standard deviations. The seasonal decomposed secondary pOON (corresponding method is referred to Text S4) and O_x are also shown. The inset pies are the contributions from biomass burning and secondary formation to total gOON and pOON, respectively.

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The ratios, i.e., ([OON_{measured}]/[levo.])_{bb}, were obtained based on ambient measurement; therefore, the ratios might be influenced by secondary formation within biomass burning plumes, the OON_{bb} herein is referred to as the total primary and rapidly formed secondary OON from biomass burning emissions. OON_{sec} is defined as secondary OON from non-biomass burning sources, e.g., biogenic and non-biomass burning anthropogenic sources and possible OON slowly formed from biomass burning sources (i.e., next day), which shall be minor. By using this approach, the estimated diurnal variation and time series of OON_{bb} and OON_{sec} are shown in Figs. 2 and 3. Compared with the measured total OON, the OON_{sec} exhibits better agreement with the total gOON production rate in terms of both time series and correlation coefficients (R increases from 0.61 to 0.65 for gOON, and 0.19 to 0.45 for pOON, Fig. S16). In particular, better agreement was found during 24–26 October 2018 when the contribution of biomass burning to OON was high. The precursors, e.g., alkanes, alkenes, aromatics (phenol and cresol), and terpenes (isoprene and monoterpenes), considered in the calculation were also contributed by biomass burning (Liu et al., 2017; Gilman et al., 2015), especially during the strong biomass burning emission period. The particle-phase OON_{sec} also showed consistent variation (R=0.70) to semi-volatile oxygenated OA (SV-OOA), which was treated as freshly formed SOA during the day (Fig. S17) (Chen et al., 2021a), supporting the secondary origins of pOON_{sec}. These results validated the source apportionment of OON applied herein.

375 On average, biomass burning emissions accounted for $49 \pm 23\%$ of total pOON measured by the CIMS, while the contribution was much lower $(24 \pm 25\%)$ for gOON (Figs. 2b and 2d), indicating that biomass burning is one of the major sources for pOON measured by the CIMS, and gOON is predominately from secondary formation ($76 \pm 25\%$) (Huang et al., 2019; Lee et al., 2016). The uncertainty of these ratios representing the error of this source apportionment method, the detailed calculation process of which can be found in Text S3 of the supporting information. The high contribution of biomass burning to total pOON is also 380 consistent with the results of relevant previous studies (Mohr et al., 2013; Wang and Li, 2021; Wang et al., 2017c; Wang et al., 2019), in which substantial OON compounds were observed in biomass burning plumes. In this study, the nighttime enhancement of $C_6H_{10}O_5$ indicates that the influence of biomass burning at this site was mainly contributed by the plumes of the agricultural residue combustion transported from the vicinity of Guangzhou city, as shown in the MODIS wildfire point plot in Fig. S18 (Wang et al., 2017b; Yuan et al., 2010). Moreover, the ambient ratio of $C_6H_{10}O_5$ to water-soluble potassium (K⁺) (0.20 ± 0.04) observed in 385 this study also supports that the biomass burning at this site was contributed by the combustion of crop residuals (0.1-0.2) rather than wood combustion (5.8-24.0) (Cheng et al., 2013). Fig. S19 presents the Van Krevelen diagram of all OON compounds in this study. The appearance of OON compounds observed herein is linked with both fresh and aged biomass burning emissions (Wang et al., 2019). In general, most of the biomass burning OON were found in particle phases, indicating that the OON formed in biomass burning plumes have generally lower volatility than OON formed via O₃, NO₃ oxidation, or OON oxidized from non-biomass burning related precursors.

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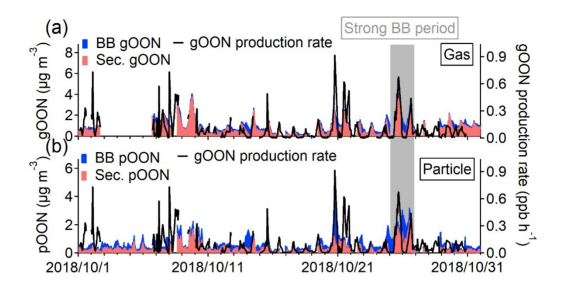


Figure 3. Stacked time series of secondary and biomass burning (a) gOON, (b) pOON. The gOON production rate is presented on the right axis in both of the figures. The grey period represents a strong biomass burning emission period during 24–26 October 2018, which was selected based on the high mass concentrations of C₆H₁₀O₅ and other biomass burning tracers in their time series in Fig. S11.

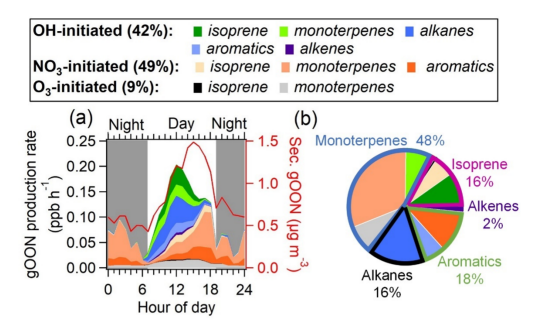
Based on the diurnal pattern, secondary gOON peaks during the afternoon $(1.49 \pm 0.49 \ \mu g m^{-3})$, which is consistent with the peaking time of O_x ($[O_x] = [O_3] + [NO_2]$), and then reduces rapidly to $0.43-0.83 \ \mu g m^{-3}$ at night (Fig. 2b). The similar averaged diurnal variations of gOON and O_x indicate that the daytime chemistry corresponds to the major formation pathway of gOON. Secondary pOON also shows a slight peak ($0.39 \pm 0.07 \ \mu g m^{-3}$) corresponding to O_x during daytime, however, exhibits much larger uncertainty than the secondary gOON. To elucidate this large uncertainty, a seasonal decomposition method (Hilas et al., 2006), which was performed by locally weighted linear regression to decompose the time series into three components, i.e., trend component, seasonal component and remainder, was applied (detailed process can be found in Text S4). By replacing seasonal variation with hour variation, the method can down weight the impact of daily peak intensity variation. Clear diurnal variation of secondary pOON after seasonal decomposition is displayed in Fig. 2d, which supports the daytime peak of secondary pOON. During nighttime, the concentration of pOON ($0.32 \pm 0.07 \ \mu g m^{-3}$) remains at a high level with a peak at 6 o'clock (Fig. 2d), indicating that there is a different formation pathway or formation yield for secondary pOON compared to secondary gOON at night. It is speculated that the enhanced secondary pOON formation at night is probably associated with higher yield of pOON from NO₃ chemistry, as well as heterogeneous reactions of NO₃ and N₂O₅ at particle surface, which is discussed in detail in the following section. Following Eqs. (1) and (2), the averaged concentration ratios for each category of particle-phase C_xN (pC_xN) were also calculated separately,

thus the contributions of biomass burning and secondary formation for each pC_xN group were estimated (Fig. S20). The contribution from biomass burning to each pC_xN group only shows small difference, ranging from 52% for the $C_{6-9 \text{ Aro}}N$ group to 40% for the $C_{4-5}N$ group, indicating biomass burning is an important source for OON at wide carbon numbers.

3.3 Secondary formation pathways of oxidized organic nitrogen.

To further elucidate the secondary formation mechanism of gOON, the diurnal patterns of gOON production rates from 415 the three pathways following the procedure mentioned in section 2.3 are calculated and shown in Fig. 4a and Fig. S21. Although the production rate did not consider the loss of OON, the calculation of production rate still serves as a useful tool to assess the formation pathway and precursor contribution to OON (Liebmann et al., 2019; Sobanski et al., 2017; Hamilton et al., 2021). As expected, the gOON production rates from OH- and O₃-initiated oxidation peaking (0.14 ppb h⁻¹ and 0.01 ppb h⁻¹, respectively) at noon (11:00–13:00) are due to the high concentrations of these two oxidants during this period (3.9×10^6 and 1.4×10^8 molecule cm⁻ 420 ³, respectively) (Wang et al., 2020c). Interestingly, the gOON production rate from NO₃-initiated oxidation peaks at around $0.10 \pm$ 0.18 ppb h⁻¹ from late afternoon to evening (16:00–19:00, Fig. 4a). This is mainly due to the high NO₃ concentration (2.08 \pm 1.32 ppt, Fig. S4d) when the low NO (1.37 ± 0.34 ppb) and moderate O₃ (53 ± 34 ppb) and NO₂ (31 ± 13 ppb) (Fig. S9b) concentrations appear during that period of the day. The precursors, i.e., cresol, phenol, isoprene, and monoterpenes, do not show rising concentrations during the period (16:00–19:00) except for some aromatics (Figs. S9c, S9f, and S9i), indicating that not the precursor 425 VOCs but the high NO₃ concentration is the main contributor for this enhancement of NO₃-initiated gOON production rate during the day. The inconsistent peaks for the secondary gOON production rate and secondary gOON mass loading may be due to the (i) counteraction of photochemical formation and degradation, and or (ii) effect of survivor bias from measured VOCs (Wang et al., 2022b; Perring et al., 2013).

In general, OH- and NO₃-initiated oxidation pathways dominated the secondary gOON formation in this study and 430 accounted for 42% and 49% of total gOON production rate, respectively, while the remaining was attributed to the O₃-initiated oxidation pathway (9%). Fig. S21 shows that the contribution to total gOON production rate from 8:00 to 14:00 mainly came from OH chemistry (73%), then quickly changed to NO₃ chemistry during the late afternoon (mean 55%) and onward (mean 86% at night). These results emphasize the importance of NO₃ chemistry for the gOON production rate during later daytime and the entire night in this urban area. The importance of NO₃ chemistry toward ON formation was also found in other locations. For example, 435 high contributions of NO₃ chemistry to secondary gOON production rate in Finnish boreal forest (41% during the day and almost 100% at night) (Liebmann et al., 2019) and isoprene-derived gOON production rate in Beijing urban area (32% in the afternoon and 86% at nighttime) (Hamilton et al., 2021) were observed. The low contribution of O_3 pathway (9%) at this site is also consistent with the estimated fraction (12%) at a forest area (Liebmann et al., 2019).



440 Figure 4. (a) Average diurnal variations of categorized gOON production rates and the concentration of secondary gOON for the whole campaign. (b) Contributions of various VOC precursors to total gOON production rate.

Fig. 4b displays the contributions to the gOON production rate from different VOC precursors, among which monoterpenes account for 48% of total gOON production rate. The remaining contribution is attributed to biogenic VOC, i.e., isoprene (16%), and other anthropogenic VOCs including aromatics (18%), alkanes (16%), and alkenes (2%). These results indicate that monoterpenes 445 are the largest contributor to the secondary formation of multifunctional gOON in this urban region. Relatively high concentrations of monoterpenes were observed during the campaign (mean: 0.14 ± 0.14 ppb; range: 0.003-1.5 ppb). Strong correlations of the daily-averaged values from nighttime monoterpenes with two tracers of volatile chemical product (VCP) sources (C8 aromatics, R = 0.85 and ethanol, R = 0.78), and moderate correlation with CO (R = 0.62) as well as biomass burning tracer $C_6H_{10}O_5$ (R = 0.50) (Fig. 5) were also found. These strong and moderate correlations with monoterpene are in contrast with the poor correlations of 450 these anthropogenic tracers with biogenic-derived isoprene during daytime (R = -0.09 to 0.20, Fig. S22). The ambient ratio of monoterpenes to CO (1.82 ppb ppm⁻¹) determined in Fig. 5c is significantly higher than their emission ratios from vehicle exhausts $(0.001-0.35 \text{ ppb ppm}^{-1})$ (Wang et al., 2022a). Combining these results, it is concluded that the monoterpenes observed during this campaign should be mainly of anthropogenic origin (Hellén et al., 2012), with VCPs as the potentially most important source. Recent studies in the US also demonstrated that monoterpenes are strongly emitted by the VCP sources in highly populated areas (Coggon 455 et al., 2021; Gkatzelis et al., 2021). In addition, a field study conducted in the tower located in the Guangzhou urban area found the

ambient monoterpenes at an altitude of 450 m were predominantly come from VCP sources (Li et al., 2022), which is consistent

with the findings here. In summary, considering monoterpenes of anthropogenic origin, anthropogenic VOCs accounted for 80% of total gOON production rate. Note that certain contributions of biogenic-derived monoterpenes might offset the anthropogenic-origin OON; however, other anthropogenic VOCs, such as long-chain or cyclic alkanes/alkenes (Wang et al., 2020a; Zhao et al., 2016), which are important precursors for OONs in urban areas (Lee et al., 2015; Lim and Ziemann, 2009; Matsunaga and Ziemann, 2010), were not considered herein due to the omission of data. Thus, anthropogenic contribution to the gOON estimated via secondary

gOON production rates might still be biased low.

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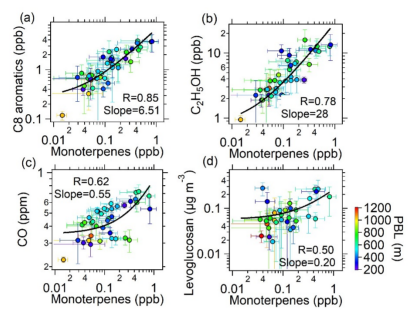
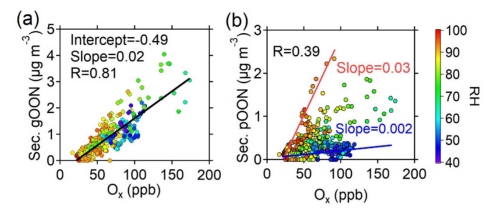


Figure 5. Scatter plots between the averaged concentrations of (a) C8 aromatics, (b) ethanol, (c) CO, and (d) C₆H₁₀O₅ versus
 monoterpenes at night (19:00–6:00 next day) during the entire campaign. The color represents the planetary boundary layer height (PBL). The error bars were the standard deviations of average values during nighttime. The logarithm was applied to both of the axes.

Fig. 6a shows a strong correlation between secondary gOON and O_x (R = 0.83, slope = 0.02 µg m⁻³/ppb), which is within expectation as the major channel of gOON formation between peroxy radicals (RO₂) and NO can lead to the formation of OONs and O₃ by continual radical propagation and photolysis of NO₂ (Perring et al., 2013; Xu et al., 2021a). Fig. 6b demonstrates that secondary pOON shows moderate correlation (R = 0.39) with O_x and secondary pOON/O_x ratio increases as a function of RH (from ~0.002 to ~0.03 µg m⁻³/ppb), aerosol liquid water content (ALWC) (Fig. S23b) and wet aerosol surface area (Fig. S23c). The elevated RH and ALWC may lead to an increase in the aerosol surface area to facilitate more highly functionalized and watersoluble gOON partitioning into particle phase and/or to promote NO₂, NO₃, and/or N₂O₅ uptake onto aerosol phase (George et al., 2015; Ceases and Abbett 2010). Meanour exceeds become more liquified at higher BU, thus leading to the increase in melanulur

475 2015; George and Abbatt, 2010). Moreover, aerosols become more liquified at higher RH, thus leading to the increase in molecular diffusion in aerosols to promote heterogeneous reactions (George et al., 2015) of NO₃ with unsaturated species (Xiao and Bertram, 2011; Zhao et al., 2011), NO₂ with aromatic species, and N₂O₅ with alcohols to form pOON (Gross et al., 2009; Lee et al., 2015).

Indeed, lower gas/particle partitioning coefficient (saturation mass concentration, C^*) of OON at RH >70% than low RH (<70%) was found (Fig. S24a), which supports the favored pOON formation from heterogeneous reactions. A study in an anthropogenicemission-dominated region also showed the heterogeneous reactions through N₂O₅ uptake can explain around half of the formation of particle-phase alkyl nitrates (Lee et al., 2015), thus signifying the important contribution from heterogeneous reactions to pOON. The secondary pOON/O_x ratio shows a much worse correlation with temperature (Fig. S23d), indicating that lower temperatureinduced higher partition to particles contributes little to higher pOON during nighttime.



485 Figure 6. Scatterplots of (a) secondary gOON and (b) secondary pOON versus O_x color-coded using RH during the campaign. The red and blue lines are plotted for guiding eyes.

Furthermore, the lifetime of gOON in this study can be approximately estimated by a steady-state approach (gOON mixing ratio versus total production rate) as shown in Liebmann et al. (2019). A scatterplot of the secondary gOON versus the secondary gOON production rate at 1 hour time resolution is shown in Fig. S24b. The roughly estimated lifetime of secondary gOON is around 0.54–0.78 h during daytime and nighttime, respectively, which is shorter than the lifetime of alkyl nitrates (~2 ± 3 h) found in the boreal forest (Liebmann et al., 2019). This might be associated with the stronger photolysis, oxidative degradation, and different chemical composition of gOON in urban area compared to boreal forest (Perring et al., 2013). The incomplete measurement of gOON by the CIMS was one of the potential reasons as well. For the lifetime of pOON, a modeling study including explicit formation mechanism as conducted by Lee et al. (2016) is required for systematic explorations in the future.

495 **3.4** Molecular chemical compositions of oxidized organic nitrogen.

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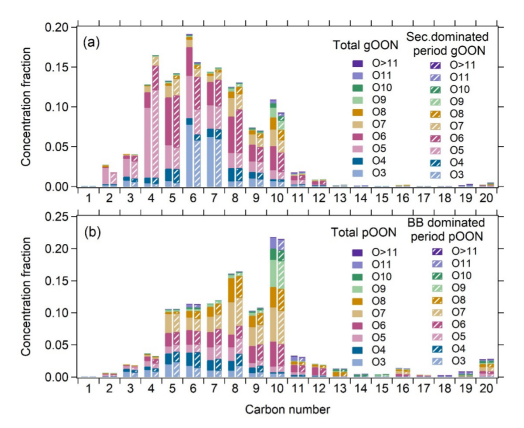
In this section, the molecular components of the gOON and pOON measured by the CIMS categorized with different oxygen and carbon atom numbers are briefly discussed. Fig. 7 shows the overview of the distribution of molecular OON during the entire campaign, as well as secondary-dominated period for gOON and biomass burning-dominated period for pOON. In general, the gOON abundance is dominated by C_5 - C_8 and C_{10} compounds (65–70%), while the pON shows peaks around C_8 and C_{10} compounds. The highly oxidized molecules (containing at least 6 oxygen atoms) contributed 44% and 71% to gOON and pOON,

respectively. By comparing the ion distribution of total gOON in the whole campaign versus in the period dominated by secondary sources (Fig. 7a), we found that the mass concentration fraction of C_4 compounds in total gOON is much enhanced, while C_6 and C_{10} compounds are significantly decreased. The enhanced C_4 compounds (e.g., $C_4H_7NO_5$ as the most abundant ion in gOON, Fig. S19) are probably contributed by the isoprene oxidation during the day (Wennberg et al., 2018; Brownwood et al., 2021). The decreases of C_6 and C_{10} compounds in gOON during secondary-dominated period indicate a large fraction of these compounds might come from biomass burning sources. For example, $C_6H_5NO_3$ (nitrophenol and its isomers) in C_6 is the second most (7.4%)

abundant compound in gOON. Multiple ambient studies had shown that biomass burning emission can contribute substantially to $C_6H_5NO_3$ (Mohr et al., 2013; Wang et al., 2018), e.g., 58% in Beijing (Song et al., 2021), consistent with the finding shown here.

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The almost identical distributions of pOON during the whole campaign and biomass burning-dominated period mainly 510 result from approximate contributions from biomass burning (49% and ~70%, respectively). The C₁₀ compounds contribute both high (~22%) in pOON during the whole campaign and biomass burning-dominated period (biomass burning contribution of pOON > 60%, Fig. S25b), confirming the important contribution of biomass burning to this type of compounds (Fig. 7b). In pOON, the abundance of C₁₀H_xNO_y ($y \ge 6$) in pOON (19%) is higher than that in gOON (9%), which is reasonable due to their low volatility with multiple functional groups (Odum et al., 1996). The extremely oxidized C₁₀ compounds (containing at least 8 oxygen atoms) 515 might come from multiple oxidation steps and autoxidation mechanism (Iyer et al., 2021; Zhao et al., 2018; Pye et al., 2019; Shen et al., 2021; Mayorga et al., 2022), demonstrating that the complex secondary formation processes indeed happened within the biomass burning plumes. There are high mass peaks at the positions corresponding to C₁₆ and C₂₀ species in pOON (Fig. 7b), which are oligomers. These ions might come from direct emission and/or oxidation process in biomass burning plumes, e.g., dimerization of C₈ and C₁₀ species (Wu et al., 2021; Lee et al., 2018).



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Figure 7. Concentration fractions of (a) gOON categorized based on carbon numbers and oxygen numbers during the whole campaign and the secondary (sec.) formation-dominated period (13:00–15:00) (Fig. S25a). Similar plot is displayed for (b) pOON during the whole campaign and the biomass burning (BB) dominated period (19:00–21:00) (Fig. S25b).

4 Conclusions

525 The mass concentrations, sources, and formation mechanism of gas- and particle-phase OON were systematically investigated in a megacity in southern China. The good comparison of pOON measured by AMS and FIGAERO-I-CIMS indicates that the CIMS can measure a fraction of $28 \pm 18\%$ of total pOON in this study. Compared to AMS, the missed pOON mass measured by CIMS is probably due to the lack of detection of less-polar OON (keto/alkyl ON) and/or non-nitrogen-containing pOON resulting from the loss of $-NO_2$ group by thermal desorption in CIMS measurement.

530 Using C₆H₁₀O₅ (~90% of levoglucosan) as the biomass burning tracer for source apportionment, almost half of the pOON measured by the CIMS is attributed to biomass burning in this study, underscoring the important contribution of biomass burning to pOON in this urban area. Biomass burning is a very common source across the world. The proposed estimation method in this study might help to clarify the exact biomass burning contribution to OON and their potential atmospheric implication. Note that the sources of the undetected pOON from CIMS are still unknown, which shall be further investigated. The gOON measured by CIMS was mainly produced by secondary formation processes (76%), initiated by OH (42%) and NO₃ (49%) chemistry. The significant contribution of NO₃ chemistry to gOON and potentially to other secondary products, e.g., SOA, was not only observed

in Guangzhou but also in the megacity of Beijing (Hamilton et al., 2021), highlighting the important daytime NO₃ chemistry in urban areas. This indicates that the importance of daytime NO₃ chemistry might not be unique and should be considered in other locations that are impacted by strong anthropogenic emissions. By ranking the precursors of OONs in the ambient atmosphere, monoterpenes are determined to be the most important VOC precursors for secondary gOON formation. Multiple evidences suggest

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that the monoterpene observed in this study is anthropogenic-origins from VCP source. The monoterpene isomer measurement, which has seldomly been carried out in current Chinese urban areas, is highly recommended for future studies to evaluate the impact of biogenic and anthropogenic emissions (e.g., VCPs) on ozone and other secondary products in the ambient atmosphere.

Furthermore, in this study, it is also found that heterogeneous reactions might contribute substantially to the secondary formation of pOON in urban areas; however, the detailed mechanism and its quantified contribution to pOON in ambient air are still unclear, which warrants further investigation. A thorough comparison between modelled and measured pOON might shed light on this question. Highly functionalized gOON and pOON, as well as oligomers can be formed through multigenerational oxidation and autoxidation during biomass burning plumes, highlighting the complexity of sources and chemical processes in the urban environment. The results of this study provide valuable data and insights to understand the chemistry of reactive organic nitrogen 550 in urban areas.

Data Availability

The 2022) used to data sets (Cai, evaluate the conclusions in the study available at are https://data.mendeley.com/datasets/s8s6wk32fy/1. Figures were made with Igor Pro version 6.37 and Igor Pro version 8.04, available under the Igor Pro license at https://www.wavemetrics.com. The more detailed data can be provided by contacting the corresponding authors.

Competing Interests

The authors declare no conflicts of interest relevant to this study.

Author contributions

YC and CY contributed equally to this work. YC and WH: writing, visualization, and validation; YC, CY, WC, WH, YP,
560 BW, XH, LH, SG, and BY: data curation, methodology; CY, WC, WH, WS, YP, SH, JQ, SW, CW, CW, ZW, BY: experiment, investigation, and formal analysis; WH, BY, MS, and XW: conceptualization, supervision, project administration and funding acquisition.

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570 Appendices A: Summary of the acronym

Table A1. The summary of the acronym and corresponding full name in this study.

Acronym	Full name	Acronym	Full name
ALWC	aerosol liquid water content	ONs	organic nitrates
C ₁₁₋₂₀ N	oxidized organic nitrogen molecules with 11–20 carbon atoms	OON	oxidized organic nitrogen
C ₄₋₅ N	oxidized organic nitrogen molecules with 4–5 carbon atoms	OON _{bb}	oxidized organic nitrogen from biomass burning
$C_{6-9 Aro}N$	oxidized organic nitrogen molecules with 6–9 carbon atoms and benzene ring	OON _{sec}	oxidized organic nitrogen from secondary formation
$C_{8-10}N$	oxidized organic nitrogen molecules with 8–10 carbon atoms	O _x	odd oxygen, sum of O_3 and NO_2
CHON	oxidized organic nitrogen with only one nitrogen atom	pC_xN	particle-phase C _x N
CHON ₂	oxidized organic nitrogen with two nitrogen atoms	PMF	positive matrix factorization
CIMS	chemical ionization mass spectrometer	pON	particle-phase organic nitrates
CothersN	oxidized organic nitrogen molecules not in other four group	pOON	particle-phase oxidized organic nitrogen
C _x N	oxidized organic nitrogen molecules with x carbon atoms	pOON _{AMS}	particle-phase oxidized organic nitrogen derived from aerosol mass spectrometer measurement.
dV	voltage difference	pOON _{CIMS}	particle-phase oxidized organic nitrogen measured by chemical ionization mass spectrometer
dV_{50}	the voltage at which half the signal is removed (i.e., half of iodide adducts dissociate)	pOrgNO _{3, AMS}	nitrate functional group from particle- phase oxidized organic nitrogen measured by aerosol mass spectrometer
FIGAERO-I- CIMS	an iodide-adduct chemical ionization mass spectrometer equipped with a Filter Inlet for Gases and AEROsols	pOrgNO _{3,CIMS}	nitrate functional group in particle-phase oxidized organic nitrogen based on the data by chemical ionization mass spectrometer
GC-MS/FID	gas chromatography coupled with mass spectrometry and flame ionization detector	PRIDE-GBA	Particles, Radicals, and Intermediates from oxidation of primary Emissions over the Great Bay Area
gON	gas-phase organic nitrates	PTR-ToF-MS	proton transfer reaction time-of-flight mass spectrometry
gOON	gas-phase oxidized organic nitrogen	RH	relative humidity

gOON _{CIMS}	gas-phase oxidized organic nitrogen measured by the chemical ionization mass spectrometer	\mathbf{S}_0	the relative signal at weakest dV compared to the signal under operational dV
HR-ToF-AMS	high-resolution time-of-flight aerosol mass spectrometer	TD	thermodenuder
IMR	ion-molecule reaction region	TD-LIF	thermal dissociation laser induced fluorescence
MW	molecular weight	VCP	volatile chemical product
NO _x	Sum of NO and NO ₂	VOCs	volatile organic compounds

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