1	Unambiguous identification of N-containing oxygenated organic molecules
2	using CI-Orbitrap in an eastern Chinese megacity
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28 S1. Other ancillary measurements

29 The mass concentration of ambient particles was measured by particle monitor (TEOM 1405DF, Thermo, 30 USA). SO₂, O₃ and NO_x concentrations were measured using a SO₂ analyzer (Model 43i, Thermo, USA), a O₃ 31 analyzer (Model 49i, Thermo, USA) and a NO_x analyzer (Model 42i, Thermo, USA) with the detection limits of 32 0.1 ppby, 0.5 ppby and 0.4 ppby, respectively. The above instruments were pre-calibrated before the campaign. The 33 solar radiation was measured on the rooftop of the building. Atmospheric N₂O₅ concentrations were measured by 34 an iodide CI-API-TOF. The concentrations of NO₃ radicals were estimated under the assumption that NO₃, NO₂ and 35 N₂O₅ could reach an equilibrium quickly in tropospheric conditions (Brown and Stutz, 2012). VOC precursors were 36 measured by an online GC-MS (7890A-5975C, Agilent, USA).

37 S2. Overview of the campaign

38 An overview of the measurement data, illustrating the air quality as well as the meteorological 39 conditions (global radiation, temperature, wind direction, wind speed, and RH), concentrations of trace gases and pollutants (PM2.5, O3, NOx, N2O5, and TVOCs) during the campaign, is provided in this section 40 41 as shown in Figure S1 and Table S1. Firstly, the maximum intensities of global radiation on individual 42 days were in a range of 637-867 W m⁻², indicating strong photochemical activities during the daytime of 43 the campaign. The relative humidity (RH) exhibited a clear diurnal variation pattern with a range of 21-44 91%. The wind (0-7 m/s) from the north to northeast prevailed during the campaign and frequently resulted in increased PM_{2.5} concentrations due to the transport. The PM_{2.5} concentration were in a range 45 of 6-59 µg m⁻³ (5-95 % percentile). The 5-95 % percentile ranges of [O₃], [NO_x], and [TVOC] were 4.6-46 58.6 ppbv, 8.9-69.6 ppbv, and 15.2-77.5 ppbv, respectively. O₃ showed an obvious diurnal variation, 47 48 peaking at 13:00 - 15:00. Diurnal variations of NO_x and TVOC showed high concentrations over the rush hours. A PM episode with mean PM_{2.5} concentration of 56.4 µg m⁻³ occurred from November 6th to 49 November 8th, accompanied by a high concentration of both TVOC and NO_x, indicating the same origins 50 of air pollutants. While the N₂O₅ remained in low concentration levels in general, three peak 51 concentrations up to about 600 pptv appeared at nighttime during November 6th-8th. 52

53 S3. Positive matrix factorization (PMF)

54 Positive matrix factorization (PMF) allows for time-resolved mass spectra to be expressed as a linear 55 combination of a finite number of factors, assuming that the factor profiles are constant and unique. Since 56 this method does not require a priori information about the factors, it is an ideal technique for extracting 57 information from ambient measurements where the detailed chemistry, sources, and atmospheric 58 processes are complex. PMF has already been used in source apportionment analysis of OOMs in previous 59 studies (Yan et al., 2016; Zhang et al., 2019, 2022). In this study, PMF was performed using the Igor-based 60 interface Source Finder (SoFi, v6.3), run by the multilinear engine (ME-2) (Canonaco et al., 2013). The 61 data for the PMF model inputs were prepared according to the method described in previous studies (Zhang et al., 2022). Note that the orbitrap analyzer does not measure signal below a certain threshold 62 63 resulting in incomplete time series for species present at low concentration level. Therefore, the species 64 characterized by incomplete time series with more than 90% missing data and the spectra with more than 65 80% missing were removed (Zhang et al., 2022).

PMF analysis in this work was performed in 2-10 factors as shown in Figure S3. Five runs for each 66 67 solution show good consistencies in both Q/Qexp and explained variation, indicating the small model uncertainty. The change of Q/Qexp, which decreases stepwise from 2.61 (assuming two factors) to 0.65 68 69 (assuming nine factors). Since the absolute value of Q/Q_{exp} might be misleading, the trend of Q/Q_{exp} is useful to determine the minimum factor number (Ulbrich et al., 2009), a large decrease in Q/Qexp indicates 70 71 that the additional factor may explain a large fraction of unaccounted variability in the data. The third 72 factor significantly decreases the Q/Qexp value from 2.61 to 1.83, suggesting the importance of the third 73 factor. By adding the third factor, the model can explain 79.4 % of the data variation, in comparison to 75.4 % when only two factors are assumed. This improvement in model performance also implies the 74 75 addition of third factor is crucial. The second largest increase in the explained fraction (from 79.4 % to 76 81.3 %) happens when adding the fourth factor and the Q/Qexp value decreases from 1.83 to 1.51. When 77 model contains 5, 6, 7, 8, 9 and 10 factors, the Q/Qexp values are about 1.27, 1.07, 0.94 and 0.83, 0.73 78 and 0.66 respectively while the explained fraction by mode are in a range of 82.9-87.5 %.

Since the PMF analysis is a pure mathematical method without any prior physical or chemical 79 80 assumptions, choosing the best factor number is critical before describing the PMF results. In terms of 81 trends, more factors would get more freedom to follow subtle variations of the matrix, however, artificially choosing too many factors will over analyze the matrix, resulting in the split of physically meaningful 82 83 source apportionment into meaningless ones. The timeseries and diurnal variations of factors are shown 84 in Figure S4 and Figure S5. The two-factor solution leads to a distinct daytime factor and a night factor. 85 In the three-factor solution, the timeseries of first two factors are more or less the same as those in the 86 two-factor case, but the variation pattern of second factor has changed in the daytime, the new factor tracks 87 the PM_{2.5} concentration well in two PM episodes, and exhibits a ush-hour peak in the morning. The four-88 factor solution results in two daytime factors originated from the old daytime factor. When five factors are 89 assumed, an additional nighttime factor appears. When six factors are assumed, an afternoon rush-hour 90 factor appears. For seven factors, the derived new factor has no strong correlation with any independent 91 tracer. Herein, we concluded that the PMF solution with six factors is the optimal solutions and chose 92 to limit our further analysis to the six-factor solution because it is not possible to distinguish the 93 identification of "real" factors without significant correlations. The factor profiles in the six-factor solution 94 could be seen in Figure S6.

In the aspect of variation patterns, we classify the six factors into three types. The first two factors are related to the daytime photochemical activities and defined as daytime factor-1 and daytime factor-2. The third factor and fourth factor show clear nocturnal patterns and defined as nighttime factor-1 and nighttime factor-2. The fifth factor and the sixth factor are more related to the emission episode and thus defined as episode factor-1 and episode factor-2. Table S1 shows the peak times and fingerprint molecules of the factors.

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- 102





Figure S1. Timeseries of key measurements during the field campaign.



107 Figure S2. The relative contributions of OOMs with different carbons to the extremely low-volatility organic compounds (ELVOC, C* < 3

 $\times 10^{-5} \text{ µg m}^{-3}$) and low-volatility organic compounds (LVOC, $3 \times 10^{-5} \le C^* < 3 \times 10^{-1} \text{ µg m}^{-3}$).





Figure S3. Mathematical diagnostics of PMF solutions, including the overall changes of Q/Qexp and the explained variation from two-factor noise-factor solutions. For each number of factors, five seed runs were performed to test the consistency of the solution.



116 Figure S4. Timeseries of factors in 2-6 factor solutions of PMF. The panels from top to bottom are 2-factor solution, 3-factor solution, 4-







Figure S5. Diel variation patterns of factors in 2-6 factor solutions of PMF. The panels from top to bottom are 2-factor solution, 3-factor solution, 4-factor solution, 5-factor solution and 6-factor solution, respectively.



.22 Figure S6. The factor profiles in the six-factor solution, the black ones represent fingerprint molecules.





125 Figure S7. Timeseries of PM_{2.5} concentration (top panel), and the episode factor-1 (bottom panel).



Figure S8. Timeseries of N₂O₅ concentration (top panel), estimated NO₃ concentration (middle panel), and the nighttime factor-1 (bottom
panel).

Table S1 Summary of the Factors in six-factor solution

Factor	Factor	Peak time	Fingerprint molecules	
	Daytime factor-1	12:00-14:00	C _n H _{2n-4} O ₁₀ N ₂ (n=8-10)	
Daytime	Daytime factor-2	12:00-14:00	$C_nH_{2n}O_8N_2$ (n=4-5), $C_nH_{2n-2}O_8N_2$ (n=5-6)	
NT 1 //	Nighttime factor-1	19:00-23:00	$C_{10}H_{16}O_9N_2,C_{10}H_{16}O_8N_2,C_8H_{14}O_8N_2$	
Nighttime	Nighttime factor-2	20:00-06:00	C ₅ H ₈ O ₉ N ₂ , C _n H _{2n} O ₇ N ₂ (n=5-8)	
D · 1	Episode factor-1	PM episode	C _n H _{2n-2} O ₈ N ₂ (n=5-11)	
Episode	Episode factor-2	Afternoon rush-hour (16:00)	C _n H _{2n-2} O ₈ N ₂ (n=5-9)	

Table S2 Averaged nO_{eff} of 2N-OOMs in the four cases

Case	$\overline{[2N - OOM_{Aro}]}$	$\overline{[2N - OOM_{All}]}$	$\overline{[2N - OOM_{MT}]}$	$[2N - OOM_{Total}]$
CL _{day}	5.6	4.0	4.8	4.6
CL _{night}	4.8	3.9	4.6	4.2
PL _{day}	5.3	4.0	4.9	4.3
PL_{night}	4.8	3.9	4.5	4.1

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