

1 **Dominant Influence of Biomass Combustion and Cross-Border Transport on Nitrogen-Containing Organic**
2 **Compound Levels in the Southeastern Tibetan Plateau**

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18 **Abstract**

19 The Tibetan Plateau (TP) is highly susceptible to climate change and the nitrogen-containing organic compounds
20 (NOCs) in fine particulate matter (PM_{2.5}) represent one of the large uncertainties in affecting the climate in high-
21 altitude areas. Previous studies have shown that NOCs play a vital role in the nitrogen budget of PM_{2.5}. However,
22 our understanding of the composition and sources of NOCs in PM_{2.5}, particularly in TP, is limited. Here, we aim
23 to enhance our understanding of NOCs in the TP region by examining their identification, concentration levels,
24 sources, and origins. We conducted field sampling at a regional background sampling site in Gaomeigu, in the
25 southeastern margin of TP from March 11th to May 13th in 2017, followed by laboratory analysis of the NOCs
26 collected on the filters. The daily mass concentrations of NOCs ranged from 714.4 to 3887.1 ng m⁻³, with an
27 average of (2119.4 ± 875.0 ng m⁻³) during the campaign. This average concentration was approximately 40%
28 higher than that reported at a typical regional site in the North China Plain (NCP), highlighting a more significant
29 presence of NOCs in the Tibetan area. Biomass burning and secondary sources were identified as the major
30 contributors to total NOCs. This was further substantiated by a regional air quality model, which indicated that
31 over 80% of the aerosol in the southeast of TP originated from neighboring countries. This study enhances our
32 understanding of NOCs' contribution to PM_{2.5} in TP and their potential impacts on the climate stability in high-
33 altitude areas.

34

35 **Keywords:** Southeastern Tibetan Plateau, Nitrogen-containing organic compounds, Source apportionment,
36 Receptor model,

37

38 **1 Introduction**

39 The Tibetan Plateau (TP), located near densely populated and industrialized regions, is particularly susceptible to
40 climate change (Meng et al., 2013; Duo et al., 2015; Li et al., 2015; Yuan et al., 2016; Zhao et al., 2022). The
41 dry season features prevalent natural forest fires and anthropogenic burning activities, such as the combustion of
42 agricultural residues, leading to substantial emissions of atmospheric pollutants (Zhao et al., 2015; Ran et al.,
43 2022; Arun et al., 2021). Consequently, aerosol concentrations in the TP, especially during the premonsoon period,
44 have risen markedly (Han et al., 2020). Previous studies in the Tibetan region have mainly focused on
45 carbonaceous organic aerosols (OA), with nitrogen-containing organic compounds (NOCs) garnering less focus
46 (Zhang et al., 2020; Zhang et al., 2019; Chen et al., 2014). NOCs play an important role in modulating climate,
47 primarily through their light absorption abilities which influence aerosol radiative effects (Li et al., 2023). These
48 compounds actively contribute to the formation of new particulate matter and secondary organic aerosols (SOAs),
49 affecting cloud properties and the Earth's energy balance (Lin et al., 2021; Yu et al., 2024). The anthropogenic
50 augmentation of nitrogen emissions has notably disrupted the global nitrogen cycle, with NOC deposition
51 emerging as a significant source of reactive nitrogen (Li et al., 2023). The increased input of reactive nitrogen
52 from human activities, such as fertilizer production, adversely affects terrestrial and aquatic ecosystems and
53 human health by impacting air, soil, and water quality (De Vries, 2021). These effects have profound implications
54 for atmospheric chemistry and climate, necessitating a deeper understanding of NOC sources and atmospheric
55 processes in the climate-sensitive region of TP.

56 The pre-monsoon period features meteorological conditions that facilitate the long-range transport of NOC-
57 containing aerosols onto the TP, with prevailing atmospheric circulations transporting pollutants from neighboring
58 countries in southwest China (Wang et al., 2019a). Anthropogenic biomass burning is more intensive during the
59 pre-monsoon period and the incoming NOCs associated with biomass burning may have the potential to alter the
60 chemical composition of the atmosphere, influence cloud microphysics, and affect the regional radiative balance
61 during a critical time of hydrological accumulation and ecological transition (Tan et al., 2021). Given the TP's
62 significance in the Asian water cycle and its role as a global climate regulator, the poorly characterized
63 atmospheric behavior of NOCs during the pre-monsoon season represents a significant knowledge gap (Li et al.,
64 2023).

65 Over the past decade, studies on NOCs have primarily focused on identifying their sources and concentrations
66 (Song et al., 2017; Boreson et al., 2004; Barbaro et al., 2015; Lin et al., 2021). More than 200 NOCs have been
67 detected in the atmosphere, originating from a variety of natural sources such as animals, vegetation, ocean, and
68 husbandry, as well as anthropogenic sources including sewage treatment, combustion processes, vehicle emissions,
69 and industrial activities (Zhu et al., 2020; Zhang and Anastasio, 2003b; Shi et al., 2010; Ho et al., 2019; Wang et
70 al., 2022). Determining the sources of NOCs in the atmosphere remains challenging. For example, studies have
71 identified the sources of specific NOCs like amines, amino acids, amides, nitriles, urea, and nitrophenol (Ge et
72 al., 2011). Notably, Amines are prevalent in both urban and rural areas in America, mainly derived from industrial
73 and animal husbandry (Sorooshian et al., 2008). Biomass burning and animal farming are known emission
74 pathways for amino acids (Zhang and Anastasio, 2003a). Furthermore, investigations have shown that a
75 significant portion of water-soluble organic nitrogen (WSON) may form secondarily, as indicated by its
76 correlations with water-soluble ionic species like nitrate (NO_3^-), sulfate (SO_4^{2-}), and ammonia (NH_4^+) (Ho et al.,
77 2015). Amides can react with atmospheric acidic particles, forming secondary aerosols (Priestley et al., 2018).
78 Although previous studies have focused on identifying sources of prevalent NOCs (e.g., amino acids and amines)
79 via tracer correlations, uncertainties about specific NOC concentrations and their sources persist. Recent studies
80 have employed receptor models for source apportionment (Yu et al., 2024), yet a comprehensive understanding
81 of NOCs is still lacking.

82 In this study, we collected fine particulate matter (PM_{2.5}) samples during the pre-monsoon season at a high-
83 altitude, remote location near the Sino-Burmese border along the southeastern edge of the TP. The collected
84 samples were analyzed to determine their NOCs as well as carbonaceous components, water-soluble ions, and
85 elements. The objectives of the study were to investigate the general attributes and chemical composition of NOCs,
86 ascertain the contribution of various sources to these compounds, and identify the source regions influencing
87 PM_{2.5} and specific chemical constituents in the area.

88 **2 Experimental**

89 **2.1 Sampling**

90 Aerosol sampling was conducted at the Lijiang Astronomical Station, the Chinese Academy of Sciences
91 (26.70°N, 100.03°E, 3260 m above the sea level, Fig. S1) in Gaomeigu from March 11th to May 13th 2018. The
92 location is approximately 2 km away from the Gaomeigu village and 30 km from Lijiang City, located on the
93 southeastern edge of the TP (Zhao et al., 2019; Wang et al., 2019a). The surrounding area comprises farmland
94 and forests, with no obvious industrial proximity. Two highways are situated about 6 km from the sampling site.
95 Daily PM_{2.5} samples were collected using a high-volume sampler (model TE-6070 Tisch Inc., Village of Cleves,
96 OH, USA) at a flow rate of 1.13 m³ min⁻¹. The aerosol samples were collected on quartz fiber filters (20.3 cm ×
97 25.4 cm, Whatman QM/A, Clifton, NJ, USA) that had been pre-heated to 780 °C for 3 h for removing
98 carbonaceous materials. The sampling equipment was positioned approximately 10 m above the ground level on
99 a building's rooftop. All sampled filters were enveloped in clean aluminum foils and stored at -20 °C in a freezer
100 until subsequent analysis in the laboratory. To account for background levels, field blank filters were processed
101 and analyzed as the same method as the PM samples. All data presented was subtracted by field blank values.

102 **2.2 NOCs analysis**

103 A total of 64 PM_{2.5} samples were analyzed to determine the target NOCs in this study. Amines and amino acids
104 were quantified with the derivatization and analytical procedures by the Waters' AccQ-Tag method (Cohen and
105 Michaud, 1993; Ho et al., 2015; Ho et al., 2019). For sample extraction, a 4.3 cm² filter was cut into pieces and
106 subjected to ultrasonic extraction with 5 mL of Milli-Q water (18 MΩ cm) twice in a water bath at 25 °C. Each
107 extract was then filtered through a 0.45 μm filter and concentrated to 0.5 mL using a rotary evaporator under
108 vacuum. The resulting extracts were reacted with 6-aminoquinoly1-N-hydroxysuccinimidyl carbamate (AQC) to
109 produce fluorescent derivatives. The AccQ-Fluor reagent kit (WAT052880, Waters Corporation, Milford, MA,
110 USA) consists of AQC and AccQ. Tag borate buffer, and AccQ. Tag Eluent A was used for the derivatization
111 process. The derivatized sample extracts were reconstituted and stored in a desiccator at room temperature before
112 analysis. In the HPLC analysis, the derivates and calibration standards were injected into the high-performance
113 liquid chromatography (HPLC, 1200 Series, Agilent Technology, Santa Clara, CA, USA) equipped with a
114 fluorescence detector. The sample vials were heated at 55 °C for 10 minutes using the oven within the system.
115 The mixture was separated using a column (3.9 × 150 mm AccQ.Tag Amino Acid Analysis Silica base) bonded
116 with a 4-μm C-18 reversed-phase column at 37 °C and detected at an absorption wavelength of 395 nm. The
117 linearity of the calibrations was assessed by the correlation coefficient ($R^2 > 0.999$), and the minimum detection
118 limits (MDLs) for the target organic nitrogen species ranged from 0.036 to 0.086 nmol m⁻³. To ensure the
119 reliability of the analysis, one replicate analysis of the ambient sample was conducted for every 10 samples.
120 Additionally, ambient samples spiked with known amounts of internal and external standards were analyzed to
121 assess potential interference from the sample matrix.

122 For alkyl amides, alkyl nitriles, isocyanates, and cyclic NOCs, the extraction procedures were the same as those
123 used for the free amino acids (FAAs). After extraction, combination, filtration, and concentration, the extracts
124 were mixed with 50 μL of borate buffer to adjust the pH to 9.1. The solutions were then diluted with a
125 water/acetone mixture (3:1, v/v) to a final volume of 150 mL. To this mixture, 40 mL of dansyl chloride in acetone
126 and 10 mL of an internal standard were added. The resulting mixture underwent a derivatization reaction, which
127 involved vortex agitation for 1 minute and subsequent ultrasound irradiation at 35 $^{\circ}\text{C}$ for 15 minutes, following
128 the method described by Ruiz-Jiménez et al. (2012). The reaction vials were kept in the dark until the analysis.
129 The derivatized products were introduced into the HPLC system, which was equipped with a 2.1×150 mm C18
130 column (3.5- μm particle size, Waters Sunfire), and coupled with an ion-trap mass spectrometer (Esquire 3000,
131 Bruker Daltonics). The linearity of the calibrations for these compounds was evaluated using the correlation
132 coefficient ($R^2 > 0.999$). The MDL for the target organic nitrogen species ranged from 0.005 to 0.019 nmol m^{-3} .

133 Urea was identified and quantified using a direct injection method on an HPLC system coupled with a
134 photodiode array detector (DAD) (1200 Series, Agilent Technology). The separation of urea was achieved using
135 a 4.6×150 mm C18 column (4- μm particle size, Cogent Bidentate), and its detection was performed at an
136 absorption wavelength of 210 nm (Ho et al., 2019). The calibration of the method exhibited a high correlation
137 coefficient ($R^2 > 0.999$), indicating a strong linear relationship between the concentration of urea and the detector
138 response. The MDL for urea was determined to be 0.05 ng mL^{-1} , denoting the lowest concentration of urea that
139 could be reliably detected using the analytical method. By employing this direct injection approach, along with
140 the specific column and detection parameters, accurate identification and quantification of urea in the samples
141 were achieved. The high linearity of the calibration and low MDL underscore the sensitivity and reliability of the
142 method for analyzing urea content in the study.

143 2.3 Auxiliary measurements

144 Organic carbon (OC), elemental carbon (EC), organic markers including polycyclic aromatic hydrocarbons (PAHs)
145 and levoglucosan, and elemental components of $\text{PM}_{2.5}$ including Ca, Ti, V, Mn, Fe, Cu, As, Br, Pb, and Zn were
146 also determined (Table S1). Further details regarding the chemical analyses, including processes, accuracies,
147 precisions, and quality assurance/quality control (QA/QC) procedures of auxiliary data, can be found in Text S1
148 in Supplement Information.

149 2.4 Estimation of secondary organic carbon (SOC)

150 In this study, an approach called the minimum R^2 (MRS) method was utilized to estimate [SOC] concentration
151 (Wu and Yu, 2016) which is deduced using the following equations:

$$152 \quad [\text{SOC}] = [\text{OC}] - [\text{POC}] \quad (1)$$

$$153 \quad [\text{POC}] = [\text{EC}] \times (\text{OC/EC})_{\text{primary}} \quad (2)$$

154 where [OC] and [EC] represent the measured concentrations, [POC] represents the primary organic carbon
155 concentrations, and $(\text{OC/EC})_{\text{primary}}$ denotes an estimate of the primary OC/EC ratio. We calculated a series of
156 $(\text{OC/EC})_{\text{primary}}$ values to achieve the lowest coefficient of determination (R^2) between [SOC] and [EC], as shown
157 in Fig. S2. This minimization of R^2 allows the accurate deduction of SOC levels, considering the relationship
158 between [SOC] and [EC].

159

160 2.5 Source apportionment

161 Source apportionment using Positive Matrix Factorization (PMF) with the multilinear engine (ME-2) was

162 performed by employing the source finder tool SoFi v6.7 (Canonaco et al., 2013). The analysis involved aligning
163 daily measurements of seven nitrogen organic classes with concurrent measurements of three carbonaceous
164 materials (EC, POC, and SOC), one water-soluble inorganic ion (K^+), and 10 elements (Ca, Ti, V, Mn, Fe, Cu,
165 As, Br, Pb, and Zn) in the $PM_{2.5}$ fraction. The characteristics of the input species and the correlation matrix of
166 each species can be found in Table S2 and Fig. S3, respectively, providing statistical information for the analysis.
167 Details of the PMF and ME-2 analysis can be found in the supplementary (Text S1). Briefly, we first performed
168 unconstrained PMF with a factor number of 2-12 and examined the factor profile and time series (Fig. S4-7). 7-
169 factor factors were determined as the optimum solution (Fig. S8 and S9). To reduce the mixing between the factors,
170 a constrained PMF analysis using the “a value” approach of the ME-2 solver was applied (Canonaco et al., 2013).
171 The 7-factor solution with the constrained matrix is shown in Table S2. The constrained run was performed by
172 adding constraints in the base run resolved factor profiles so that the tracers are only present in the corresponding
173 sources (Wang et al., 2019b).

174

175 **2.6 Potential source contribution function (PSCF)**

176 The potential source contribution function (PSCF) was used to identify the likely pollution regions that influenced
177 PMF factors based on back trajectories. PSCF analysis was performed using Zefir (Petit et al., 2017). Each
178 trajectory includes a range of latitude–longitude coordinates every 1-hr backward in a whole day. The studying
179 field is from 20 to 30 °N, and 90 to 105 °E, which includes more than 95% of the area covered by all the paths.
180 The set of trajectory data for each arriving elevation level contained two trajectories per day. More details of the
181 PSCF analysis can be found in Text S1.

182

183 **2.7 Community Multiscale Air Quality**

184 The Community Multiscale Air Quality (CMAQ) model (Version 5.4) was applied to assess the transport of
185 aerosols from neighboring countries in southwest China. The CMAQ model was configured with the aero7 aerosol
186 module and cb6r5 gas-phase mechanism (Murphy et al., 2021). The model adopted a horizontal grid resolution
187 of 27 km, consisting of 34 vertical layers.

188 To generate the necessary meteorological fields for the CMAQ simulations, the Weather Research and
189 Forecasting (WRF) model (version 4.4) was utilized. The initial and boundary conditions for WRF were obtained
190 from the National Centers for Environmental Prediction (NCEP) Final (FNL) dataset, which is a reanalysis dataset.
191 For the domestic emission inventory, the Multiresolution Emission Inventory for China (MEIC) was employed.
192 Additionally, the MIX inventory was used to account for emissions from other countries (Li et al., 2017).

193 Two simulation cases were conducted: one considering only domestic emissions (i.e., MEIC), and the other
194 considering emissions from both domestic and other countries (i.e., MEIC + MIX). By employing the zero-out
195 method, the differences between these two cases represented the contribution of emissions from other countries
196 to the $PM_{2.5}$ levels in the study area. The CMAQ simulations were performed from March 9th to March 27th, 2018,
197 with the first 3 days considered a spin-up period for the model. The simulation period covered the first two weeks
198 of the campaign, encompassing the period before and during the initial pollution event from March 22nd to March
199 26th. CMAQ reproduced the measured $PM_{2.5}$ at GMG reasonably well when considering both MEIC and MIX in
200 the emission inventory, with a correlation coefficient of $r > 0.9$ between the modeled and measured $PM_{2.5}$ and a
201 slope of 0.61 (Fig. S11).

202 3 Results and discussion

203 3.1 Overview of NOC Concentration

204 Figure 1 illustrates the concentration variations of NOCs, carbonaceous aerosols, and meteorological parameters
205 in Gaomeigu during the campaign. The daily mass concentrations of NOCs range from 714.4 to 3887.1 ng m⁻³,
206 with an average of 2119.4 ± 875.0 ng m⁻³. This average is approximately 40% greater than the NOCs concentration
207 observed at a regional site in Xianghe, China (1270 ng m⁻³) (Wang et al., 2022). The NOCs are classified into
208 major (> 10% contribution) and minor (< 10% contribution) compounds, as detailed in Table 1, with the major
209 classes including FAAs, amines, and urea. The average concentrations of these major NOCs are 1922.6 ± 790.5
210 ng m⁻³, dominated by FAAs (58.9%), followed by amines (28.0%), and urea (13.7%). Minor NOC species such
211 as alkyl amides, alkyl nitriles, isocyanates, and cyclic NOCs have average concentrations of 45.1 ± 18.6 ng m⁻³,
212 4.68 ± 1.75 ng m⁻³, 10.9 ± 4.73 ng m⁻³, and 136.2 ± 61.6 ng m⁻³, respectively.

213 As shown in Fig. 1, the campaign is segmented into five periods (EP1-EP5) based on NOC concentration
214 variations. The clean period featured a temperature consistently above 9 °C and an average OC concentration of
215 2137.3 ± 296.7 ng m⁻³. Elevated wind speeds during this period (4.4 ± 1.3 m s⁻¹) enhanced atmospheric dispersion
216 relative to other polluted periods. Notably, average NOC concentration increased during pollution periods,
217 reaching 1482.6 ± 346.4 ng m⁻³, which is more than triple the level observed during the clean period (451.8 ± 65.2
218 ng m⁻³). Delving into the high NOC concentration periods individually, EP1 shows the highest aggregate
219 concentration of major NOCs, which is 4.3 to 5.0 times greater than during the clean period. The NOCs/POC
220 ratios were 0.773 (EP1), 0.774 (EP2), 0.674 (EP3), and 0.638 (EP4), presenting a stark contrast to the clean
221 period's ratio of 0.503. However, the NOCs/SOC ratio remains relatively stable across the phases. These trends
222 underscore the significant influence of primary sources during elevated NOC concentration periods. Conversely,
223 during the clean period, the source of NOCs appears more complex, suggesting a nuanced interplay of primary
224 and secondary sources. A more in-depth discussions on source apportionment are provided in Section 3.4.

225 3.2 Major NOC Classes

226 3.2.1 Free Amino acids (FAAs)

227 During the sampling campaign, the average FFA concentration is 1092.9 ± 443.37 ng m⁻³, in a range of 370.2 and
228 2033.2 ng m⁻³ (Table 1). This level is comparable with FAAs observed in regions such as rural Guangzhou, China
229 (Song et al., 2017), Arizona, U.S. (Boreson et al., 2004), and Antarctica's MZ Station, U.S. (Barbaro et al.,
230 2015) but is higher than in urban/suburban and marine regions like Nanchan, China (Zhu et al., 2020), California,
231 U.S. (Zhang and Anastasio, 2003b), Qingdao, China (Shi et al., 2010), Hong Kong, China (Ho et al., 2019).
232 Notably, the average FAAs concentration in this study is approximately four times higher than that reported in
233 Xianghe, China (Wang et al., 2022).

234 FAAs are classified into protein-type and non-protein-type categories. Table S3 provides an overview of
235 protein-type and non-protein-type FAAs, with mean concentrations of 989.5 ± 403.54 ng m⁻³ and 103.3 ± 41.76
236 ng m⁻³, respectively. Protein-type FAAs, including Asp, Ser, Glu, Gly, His, Thr, Ala, Pro, Cys, Tyr, Val, Met,
237 Lys, Ile, Leu, and Phe, accounts for 90.5% of total FAAs, with Glycine (Gly) being the most prevalent. These
238 findings are consistent with previous studies that identified Gly as the predominant FAA in Nanchang (Zhu et al.,
239 2020), Hong Kong (Ho et al., 2019), and Venice (Barbaro et al., 2011). Non-protein-type FAAs such as β-
240 alanine (β-Ala), γ-aminobutyric acid (γ-Ala), and ornithine (Orn) also contributed, with β-Ala representing 9.5%
241 of these FAAs.

242 Figure 2 illustrates a positive correlation between FAAs and O_x (NO₂ + O₃), indicating an association with

243 secondary formation processes post-precursor emissions. The average FAA concentration is 900 ng m⁻³ at O_x
244 levels below 70 ppb but rises above 1200 ng m⁻³ when O_x exceeds 85 ppb. Moreover, FAAs correlate strongly
245 with both POC (r = 0.95) and SOC (r = 0.90), indicating that secondary processes likely influence the FAA
246 formation, despite no obvious direct local emission near the sampling site. Secondary formation of FAAs can occur
247 through several mechanism, including direct photolysis, photochemical hydrolysis, and enzyme-based hydrolysis
248 (Mopper and Zika, 1987; Milne and Zika, 1993; Song et al., 2017). Given that the sampling site is subject to long-
249 range transport (discussed in Sect. 3.5), it is likely that free amino acids were secondarily produced by the
250 breakdown of proteins during the transport.

251 Moreover, Gly comprises 31% of total FAAs and shows a similar positive relationship with O_x. The Gly
252 concentration increases from 250 ng m⁻³ when the O_x is below 70 ppb to 400 ng m⁻³ when the O_x is above 85 ppb.
253 Its correlations with POC (r = 0.94) and SOC (r = 0.89) reinforce the impact of secondary formation processes,
254 similar to patterns observed in the North China Plain (NCP) region, China (Wang et al., 2022).

255 3.2.2 Amines and urea

256 The average concentration of amines during the sampling period is 563 ng m⁻³. Aliphatic amines dominate,
257 contributing 90% of the total amine, while aromatic amines constitute less than 1% (Fig. 3). The remaining 9%
258 includes other amine compounds including ethanolamine, galactosamine, 2-amino-1-butanol, and N-
259 methylformamide. During the pollution episodes, aliphatic amine concentrations exceed 600 ng m⁻³, with a
260 maximum of 1000 ng m⁻³. In contrast, during clean periods, these levels declined to ~200 ng m⁻³. The proportions
261 of aliphatic amines during pollution episodes are 90-91%, which decreases to 84% during clean periods, with an
262 increase in other concentrations.

263 Methylamine (MA) emerges as the predominant aliphatic amine, constituting 62% of the total aliphatic amines.
264 Ethylamine (EA) follows, contributing 28% to the total aliphatic amines. Dimethylamine (DMA), trimethylamine
265 (TMA), and other amine species together account for the remaining 10%. Both MA and EA exhibit negative
266 correlations with ambient temperature (Fig. 3), indicating the potential influence of temperature on gas-to-particle
267 partitioning. Below 12°C, the average MA concentration is around 400 ng m⁻³, which halves to 200 ng m⁻³ as
268 temperature increases above 18°C. Similarly, EA concentration is higher at lower ambient temperatures, around
269 195 ng m⁻³ below 12°C, decreasing to 100 ng m⁻³ above 18°C. Given their low molecular weight, MA and EA are
270 more prevalent in the gas phase at elevated ambient temperatures, where they also exhibit enhanced atmospheric
271 reactivity with acids, transforming into other compounds.

272 Both MA and EA show negative correlations with RH, with elevated concentrations at lower RHs (Fig. 3d).
273 This inverse relationship might be counterintuitive, given that higher RH typically promotes the partitioning of
274 low molecular weight amine into the particle phase. However, MA and EA, being atmospheric reactive amines,
275 are involved in in-particle reactions. Under high RH conditions, increased condensation of acids and/or reactive
276 organic compounds occur, which subsequently react with MA and EA, consuming them and thus establishing a
277 negative correlation with RH.

278 Urea is identified as the third major NOC species, with an average concentration of 266 ng m⁻³ during the
279 campaign. This value is approximately half that reported at a regional site in the NCP (Wang et al., 2022), though
280 the direct comparison is limited due to spatial and temporal differences. The urea level at this elevated site
281 highlights the notable role of agricultural fertilizers as a potential source. Urea can be released into the atmosphere
282 through agricultural activities and biomass burning (Wang et al., 2022), and it can also be formed secondarily in
283 the atmosphere through chemical reactions (Leung et al., 2024).

284 3.3 Minor NOC Classes

285 3.3.1 Alkyl amides and nitriles

286 In this study, the distributions and sources of alkyl amides in a range of C₆-C₂₀ were determined in Gaomeigu.
287 Figure S10 illustrates the distribution patterns of these species during the campaign, where the concentrations of
288 n-alkyl amides vary from 1.11 to 7.57 ng m⁻³, reflecting diverse emission sources. These amides can originate
289 from anthropogenic activities such as coal combustion and vehicular traffic, as well as biogenic processes. To
290 distinguish between these sources, we use the carbon preference index (CPI) and the oleamide to stearamide ratio
291 (Cheng et al., 2006). The CPI, calculated as the ratio of the sum of odd-numbered C₇-C₁₉ alkyl amides to even-
292 numbered C₆-C₂₀ alkyl amides, helps identify the dominant source: a CPI ≤ 1 indicates anthropogenic dominance,
293 whereas >1 suggests biogenic predominance (Abas and Simoneit, 1996). The results show that the CPI of alkyl
294 amides ranges from 0.46 to 0.75, with an average of 0.61 ± 0.05, emphasizing the anthropogenic impact on their
295 concentrations. Notably, the CPI values do not vary between the periods having low and high NOC concentrations,
296 suggesting consistent alkyl amide sources throughout the campaign, potentially influenced by long-range transport
297 and stable meteorological conditions.

298 Beyond the CPI, the R₁₈, which is a ratio of oleamide (C_{18:1}) and stearamide (C_{18:0}), serves as an indicator for
299 alkyl amide aging (Wang et al., 2022). This ratio provides insights into the precursor composition, oxidation
300 degradation, and transport processes influencing unsaturated amide concentrations (Nielsen et al., 2012). An R₁₈
301 < 1 implies the aging of alkyl amides due to long-range transport, whereas R₁₈ > 1 indicates local biomass-burning
302 emissions (Cheng et al., 2006). According to the results of this study, R₁₈ values range from 0.73 to 2.27, with
303 an average of 1.36 ± 0.35, suggesting the alternation between local and long-range transport (Cheng et al., 2006).

304 During the sampling period, the average concentration of alkyl nitriles is 4.69 ± 1.75 ng m⁻³ in Gaomeigu. As
305 shown in Table S3, hexadecanenitrile (C₁₆) is the most prevalent (0.49 ng m⁻³), followed by tetradecanenitrile (C₁₄)
306 (0.45 ng m⁻³). The concentrations of the other analyzed alkyl nitriles are below 0.4 ng m⁻³. The results are
307 consistent with the higher concentrations observed at the Xianghe site (Wang et al., 2022). Moreover, the CPI
308 values for alkyl nitriles were between 0.605 to 0.848, with an average of 0.702 ± 0.05, which points out the
309 anthropogenic influence on their levels. During high NOC concentration phases, the CPI values remain constant
310 (i.e., EP1: 0.72, EP2: 0.71, EP3: 0.71, and EP4: 0.72), compared to 0.75 during clean periods. This consistency
311 implies that anthropogenic sources predominantly influence alkyl nitrile concentrations regardless of the pollution
312 levels (Wang et al., 2022).

313 Furthermore, it is important to note that alkyl amides and nitriles might form as secondary products during
314 biomass burning through reactions between ammonia (NH₃) and FAAs (Simoneit et al., 2003). The link between
315 biomass burning and the generation of these compounds is reinforced by robust correlations with levoglucosan
316 and K⁺ in Fig. S3 (r > 0.88, p < 0.01), both recognized as markers for biomass burning (Wang et al., 2018; Liu
317 et al., 2021b). These evidences confirm that biomass burning is a key contributor to the occurrence of alkyl
318 amides and nitriles in the region.

319 3.3.2 Cyclic NOCs and isocyanates

320 The average mass concentration of cyclic NOCs is 136.2 ng m⁻³. This study identified five cyclic NOCs (Table
321 S3), with caprolactam being the most prevalent at 54.2 ng m⁻³ (39.8% of the total cyclic NOC), which is commonly
322 used in commercial manufacturing processes and lysine synthesis (Cheng et al., 2006). Other cyclic NOCs include
323 isoindole-1,3-dione (50.7 ng m⁻³, 37.2%), N-butyl-benzen-sulfonamide (NBBS) (22.1 ng m⁻³, 16.2%), N,N-
324 diethyl-m-toluamide (DEET) (5.79 ng m⁻³, 4.3%), and benzothiazolone (3.36 ng m⁻³, 2.5%). These compounds

325 are known to pose health risks (Cheng et al., 2006; Balducci et al., 2012), and they primarily originate from
326 industrial and agricultural activities (Wang et al., 2022; Richardson and Ternes, 2018; Trapp and Eggen, 2013). In
327 comparison with the findings of the Xianghe site (Wang et al., 2022), the concentrations of cyclic NOCs in this
328 study are lower, indicating the lower contributions of industrial sources. During the four high NOC emission
329 periods, the concentrations of cyclic NOCs are 2-4 times higher than those during the clean period, suggesting the
330 influence of pollution levels.

331 Isocyanates, commonly used in polyurethane resin production, are associated with several health threats,
332 including asthma, allergies, and skin reactions (Lesage et al., 2001). The average total mass concentration of
333 eight isocyanates is $10.89 \pm 4.73 \text{ ng m}^{-3}$ (Table 1) while the individual concentration of each isocyanate is given
334 in Table S3, including methyl isocyanate (MIC), toluene-2,4-diisocyanate (2,4-TDI), toluene-2,6-diisocyanate
335 (2,6-TDI), isophorone diisocyanate (IPDI), 1,6-hexamethylene diisocyanate (1,6-HDI), ethyl isocyanate (EIC),
336 phenyl isocyanate (PHI), and propyl isocyanate (PIC). Among these, TDI and HDI are predominantly used in
337 industry (Hejna et al., 2024). TDI is commonly utilized in various foam products (Akindoyo et al., 2016), while
338 HDI is essential in polyurethane paints and coatings (Golling et al., 2019). The presence of these isocyanates in
339 numerous products is linked to heightened health hazards, such as skin allergies, atopic dermatitis, and various
340 respiratory diseases (Nawrot et al., 2008).

341 3.4 Sources apportionment of NOCs

342 In this study, a constrained PMF analysis was applied to identify the sources of NOCs, which include biomass
343 burning, coal combustion, industry-related sources, crustal sources, traffic emissions, agriculture activities, and
344 secondary sources (Fig. 4).

345 Factor 1, attributed to biomass burning, was characterized by high loadings of K^+ (84.3%) and levoglucosan
346 (100%), recognized tracers for biomass-burning activities (Liu et al., 2021a; Lin et al., 2018). This factor also has
347 a notable Zn content (38.7%), indicative of wood burning (Salam et al., 2013). Biomass burning contributes 26.3%
348 to the total NOCs, emerging as the second-largest emission category. Factor 2, associated with coal combustion,
349 exhibits substantial loadings of As and also contains Cu, Pb, and EC. As and Pb are typical tracers of coal
350 combustion (Qin et al., 2019), and Cu is also associated with coal combustion (Hsu et al., 2016). Factor 3 is
351 recognized as industry-related emissions which is characterized by high loading of cyclic NOCs and isocyanates,
352 which are synthetic compounds (Wang et al., 2022). It also exhibits a significant characteristic value of Pb, which
353 can be released during industrial processes (Wang et al., 2015). This factor accounts for 7.6% of NOCs. Factor
354 4, characterized by crustal sources, had high loadings of Ti and moderate loadings of Mn, Fe, Ca, and arabitol.
355 These elements are acknowledged as crustal constituents (Gosselin et al., 2016), and arabitol is typically released
356 from soil fungal spores (Wang et al., 2018), contributing to 6.1% of the total NOCs. Factor 5, linked to traffic
357 emissions, showed high loadings of V, Br, Zn, and Cu. V acts as an indicator for heavy oil combustion in marine
358 vessels (Bian et al., 2018), and Br is a tracer of motor vehicle emissions (Guo et al., 2009). Emissions of Zn and
359 Cu are associated with brake, tire, and road wear (Salameh et al., 2018; Liu et al., 2021a). Factor 6, named
360 agriculture activities, exhibited relatively high loading of urea and moderate loadings of K^+ , Ca, and Mn in NOCs.
361 These elements are commonly used in agriculture (Ge et al., 2011), with K^+ being crucial for plant growth and
362 metabolic functions (Meena et al., 2014), and Mn playing roles in plant oxidation-reduction (Gonçalves et al.,
363 2022). This factor accounted for approximately 13% of NOCs. Factor 7, ascribed to secondary sources,
364 demonstrated considerable influence on the SOC (Secondary Organic Carbon) variation. It was responsible for
365 30.2% of the NOCs, emerging as the predominant emission source, highlighting the role of secondary production
366 in both local and regional pollutant formation.

367 Figure 5 illustrates the average contributions of the seven identified sources to each NOC species and the total

368 NOC. The analysis demonstrated that secondary sources and biomass burning were predominant, together
369 constituting over 50% of total NOCs (Figure 5a). Specifically, for FAAs (Figure 5b), secondary sources (39.6%)
370 and biomass burning (37.3%) are the two major contributors, while other sources accounted for less than 10%
371 including agriculture activities, crustal sources, industry-related, coal combustion, and traffic emissions. The
372 notable influence of secondary sources and biomass burning on FAAs could be attributed to increased
373 transportation and biomass/wildfire heating in the region, consistent with findings in a previous study (Zhang et
374 al., 2018).

375 In the context of amines, agriculture activities make a notable contribution (18.8%), twice as high as its
376 contribution to FAAs (9.3%). For alkyl amides and nitriles, secondary sources and biomass burning were the
377 primary contributors, each surpassing 30%. This contrasts with findings from another study in a different Chinese
378 region where biomass burning is predominant in these NOC categories (Wang et al., 2022). These significant
379 contributions from biomass burning and secondary sources underscore the impact of regional transportation on
380 NOC sourcing within this area.

381 **3.5 Influence from long-range transport and biomass burning in Gaomeigu**

382 Figure 6 presents the spatial distribution of PM_{2.5} concentrations during the high NOC events, analyzing
383 two scenarios: one with only domestic emissions (MEIC-China) and another incorporating both domestic
384 and foreign emissions (MEIC-China + MIX). With solely domestic emissions considered, PM_{2.5} levels at the
385 GMG and across the broader Tibet region, as well as western Sichuan and Yunnan, were relatively low, not
386 exceeding 5 $\mu\text{g m}^{-3}$ (Fig. 6a). However, incorporating international transport into the analysis revealed a
387 significant increase in PM_{2.5} levels at GMG, where daily concentrations exceeded 20 $\mu\text{g m}^{-3}$ (Fig. 6c).
388 Similarly, elevated PM_{2.5} concentrations, reaching above 40 $\mu\text{g m}^{-3}$, were observed in southeast Tibet and
389 western Sichuan and Yunnan. Figure 6b presents the relative contributions of domestic and international
390 emissions at GMG. The contribution from international transport varied from 25% to 92%, overshadowing
391 domestic sources, which did not exceed 25% for most of the time. Notably, during the high NOC events,
392 such as in EP1, where biomass burning and secondary sources contributed over half of the total NOCs (Fig.
393 S12), the contribution from international transport increased to over 80% for the study area (Fig. 6d).

394 The emission inventory used in this study did not include data on NOCs; hence, NOCs were not explicitly
395 simulated in the CMAQ model. However, the marked influence of international transport indicates that
396 PM_{2.5}-bound NOC species likely originated from international sources, corroborated by PSCF analysis
397 linking NOCs to specific PMF factors (Fig. S13), and by the observed correlation between bulk PM_{2.5} and
398 total NOCs (Fig. S14). The contribution hotspots in India and Myanmar indicate that the long-range transport
399 of biomass-burning emissions to the study area is facilitated by prevailing winds. Conversely, secondary
400 NOC sources were predominantly linked to air masses from Myanmar, implying proximate secondary
401 formation through atmospheric reactions of precursor gases and pollutants. The complex atmospheric
402 chemistry leading to secondary NOCs includes the oxidation of precursor compounds such as volatile
403 organic compounds (VOCs) and nitrogen oxides (NO_x). Other NOCs that were not measured in this study,
404 such as nitro-aromatics, were likely contributing to the NOCs and will be the focus of future research.

405 Similar spatial patterns were observed for factors related to coal combustion, industry-related sources,
406 crustal sources, traffic emissions, and agricultural activities. This implies that their contributions were
407 associated with the proximity of the sampling site to their respective source origins. For instance, NOCs
408 related to coal combustion were potentially transported from the nearby mining or industrial areas, while
409 industry-related sources could have originated from regional transmission or industrial activities in the

410 vicinity. Crustal sources, which involve the resuspension of dust particles, could be influenced by local soil
411 conditions and wind patterns.

412 **4 Conclusions**

413 In conclusion, this study provides valuable insights into the composition, sources, and transport of NOCs in the
414 study area. The average daily mass concentrations of NOCs during the campaign ranged from 714.4 to 3887.1 ng
415 m⁻³, with an average of 2119.4 ± 875.0 ng m⁻³. The major NOC species include free amino acids (FAAs), amines,
416 and urea, accounting for 58.9%, 28.0%, and 13.7% of the major NOCs, respectively. Minor NOC species such as
417 alkyl amides, alkyl nitriles, isocyanates, and cyclic NOCs were also identified. The PMF analysis revealed seven
418 distinct sources of PM_{2.5}, with biomass burning and secondary sources as the primary contributors to total NOCs.
419 Biomass burning sources exhibited hotspots of contribution from India and Myanmar, indicating long-range
420 transport. Secondary sources, predominantly originating from Myanmar, suggested the formation of NOCs during
421 the transport. This is confirmed by the CMAQ modeling. The study also revealed the possible aging of NOCs
422 from biomass-burning sources as they approached the measurement site, highlighting the impact of atmospheric
423 transformation processes. Contributions from industry-related sources, crustal sources, and agricultural activities
424 were influenced by both regional transmission and local emissions in the vicinity of the sampling site. Overall,
425 this research highlights the complex nature of NOCs and their sources, emphasizing the interplay between long-
426 range transport, regional emissions, atmospheric chemistry, and local influences. These findings contribute to our
427 understanding of air pollution dynamics and provide a basis for developing targeted mitigation strategies and
428 policies to reduce NOC emissions and their impacts on air quality and human health in the study area and similar
429 regions. For future research, we suggest further investigation into the specific chemical pathways involved in the
430 formation of NOCs during atmospheric transport, which could involve controlled laboratory experiments and field
431 studies. Additionally, more detailed source apportionment studies in different regions, including urban, rural, and
432 remote areas, would provide a comprehensive understanding of the sources and contributions of NOCs. By
433 addressing these areas, future research can further enhance our understanding of NOCs and inform effective policy
434 measures to mitigate their adverse effects.

435

436 **Declaration of competing interest**

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

439 **Credit authorship contribution statement**

440 Meng Wang: Conceptualization, Methodology, Validation, Formal Analysis, Writing - Original Draft.

441 Qiyuan Wang: Conceptualization, Writing - Review and Editing, Funding Acquisition.

442 Steven Sai Hang Ho: Formal analysis, Writing - Review, and Editing.

443 Jie Tian: Investigation.

444 Yong Zhang: Investigation, Formal analysis.

445 Shun-cheng Lee: Resources.

446 Junji Cao: Conceptualization, Writing - Review and Editing, Funding Acquisition, Supervision.

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453

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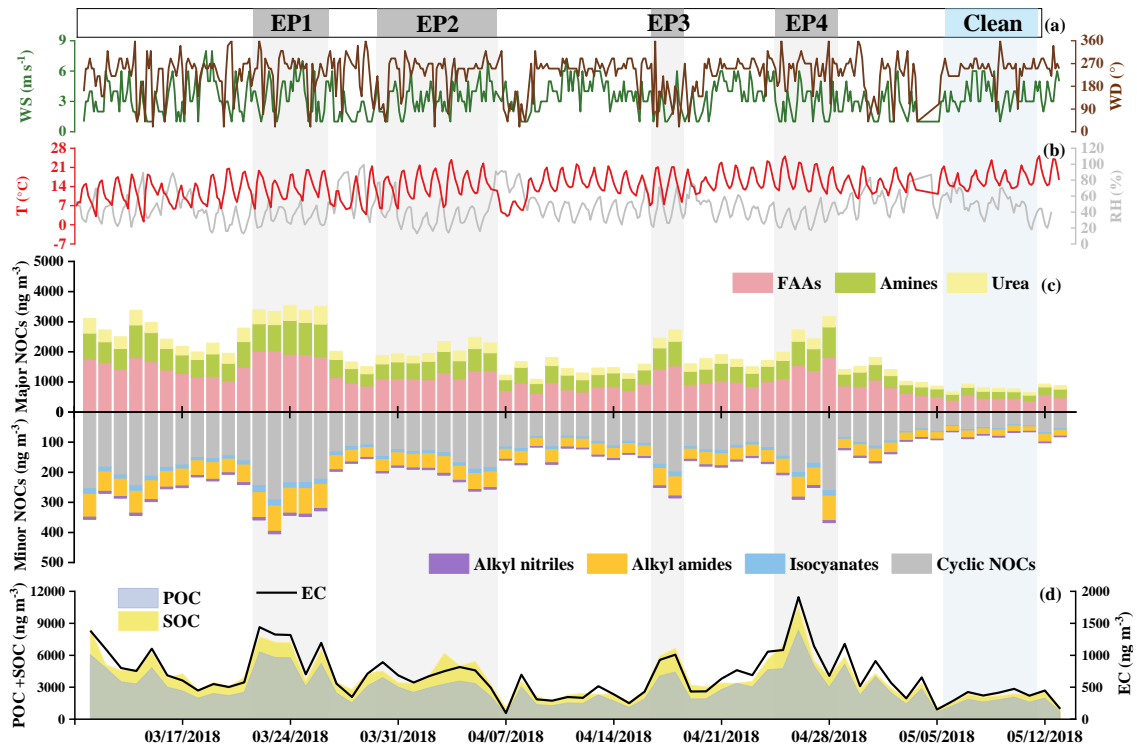
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662 **Table 1** Concentration levels of chemical compounds and groups in Gaomeigu, China. (ng m⁻³).

Species	Mean	SD ^a	Min ^b	Max ^c
NOCs (ng m⁻³)				
Major Compound Classes				
FAAs				
Protein FAAs	989.5	403.5	337.8	1857.5
Non-protein FAAs	103.3	41.8	32.5	206.8
Total FAAs	1092.9	443.4	370.2	2033.2
Amines				
Aliphatic Amines	508.9	225.9	158.5	1032.2
Aromatic Amines	2.6	1.0	1.0	5.3
Other Amines	51.8	16.8	29.2	100.7
Total Amines	563.3	240.2	190.2	1113.5
Urea	266.4	119.0	79.4	588.8
Total Major Compound	1922.6	790.5	649.0	3543.7
Minor Compound Classes				
Amides				
Alkyl amides (Odd)	13.1	5.8	4.1	26.6
Alkyl amides (Even)	21.4	8.9	6.6	41.2
Total Alkyl amides	45.1	18.6	14.9	84.6
Nitriles				
Alkyl nitriles (Odd)	1.9	0.7	0.8	3.5
Alkyl nitriles (Even)	2.7	1.0	1.0	4.8
Total Alkyl nitriles	4.7	1.7	1.8	8.2
Cyclic NOCs	136.2	61.6	42.1	291.9
Isocyanates	10.9	4.7	3.3	23.2
Total Minor Compound	196.8	86.1	65.4	404.4
Total NOCs	2119.4	875.0	714.4	3887.1

663 ^aSD represents standard deviation. ^bMin and ^cMax donate “minimum and maximum, respectively.

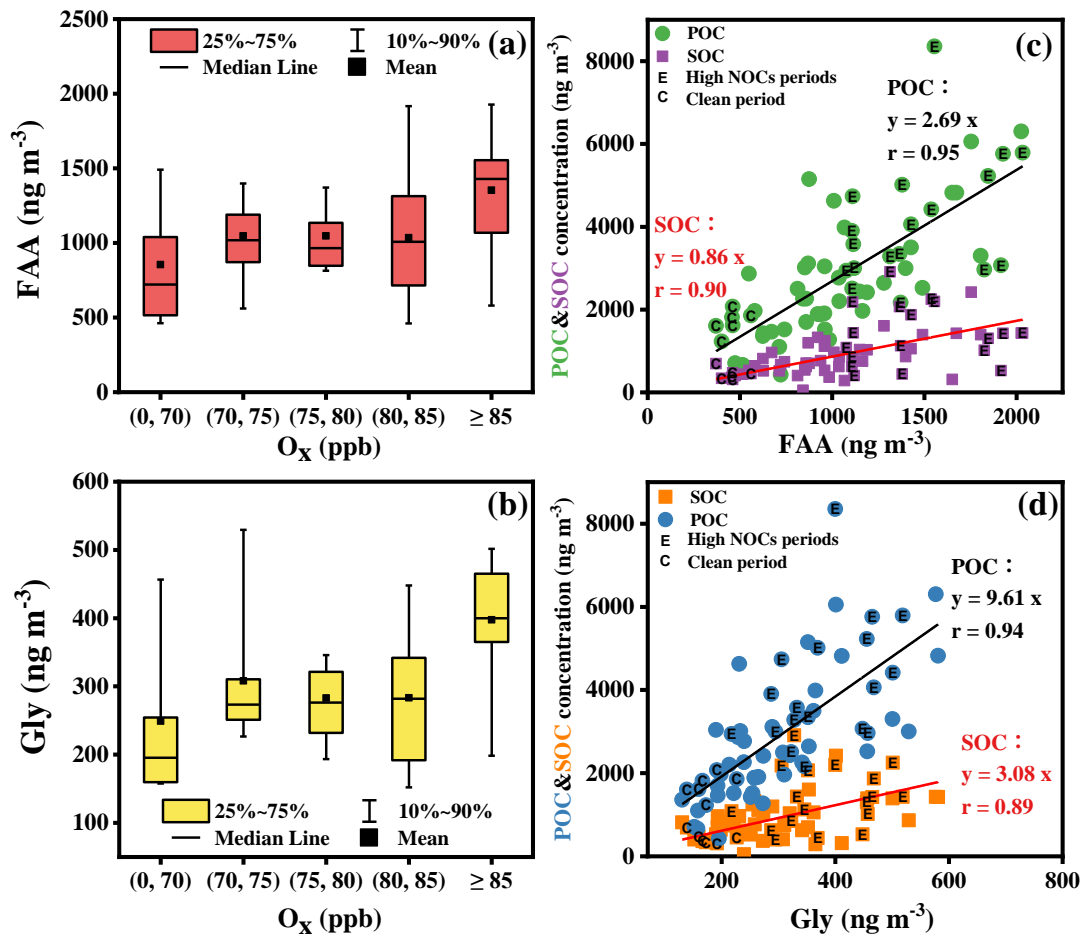
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666 **Figure 1** Hourly variations in meteorological parameters and daily chemical compositions of NOCs during
 667 different events in Gaomeigu in 2018 (EP1: 3/22 to 3/26; EP2: 3/30 to 4/6; EP3: 4/17 to 4/18; EP4: 4/25
 668 to 4/28; Clean period: 5/6 to 5/11).

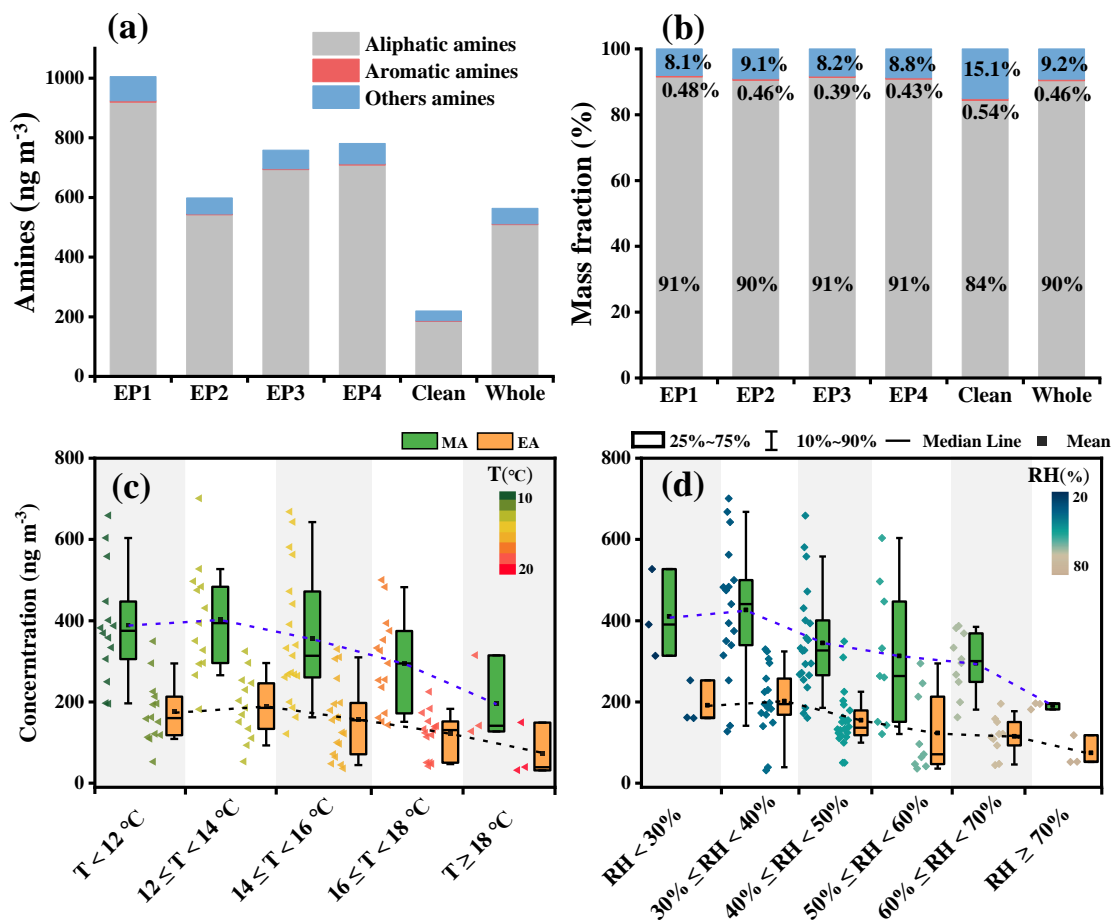
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671 **Figure 2** (a) FAA dependence of O_3 ; (b) Gly dependence of O_3 ; Correlation plots of POC&SOC
 672 concentration versus (c) FAA, and (d) Gly. The box represents the 25th (bottom) and 75th percentiles (top),
 673 and the box-whisker data represent the range from 10th to 90th percentiles.

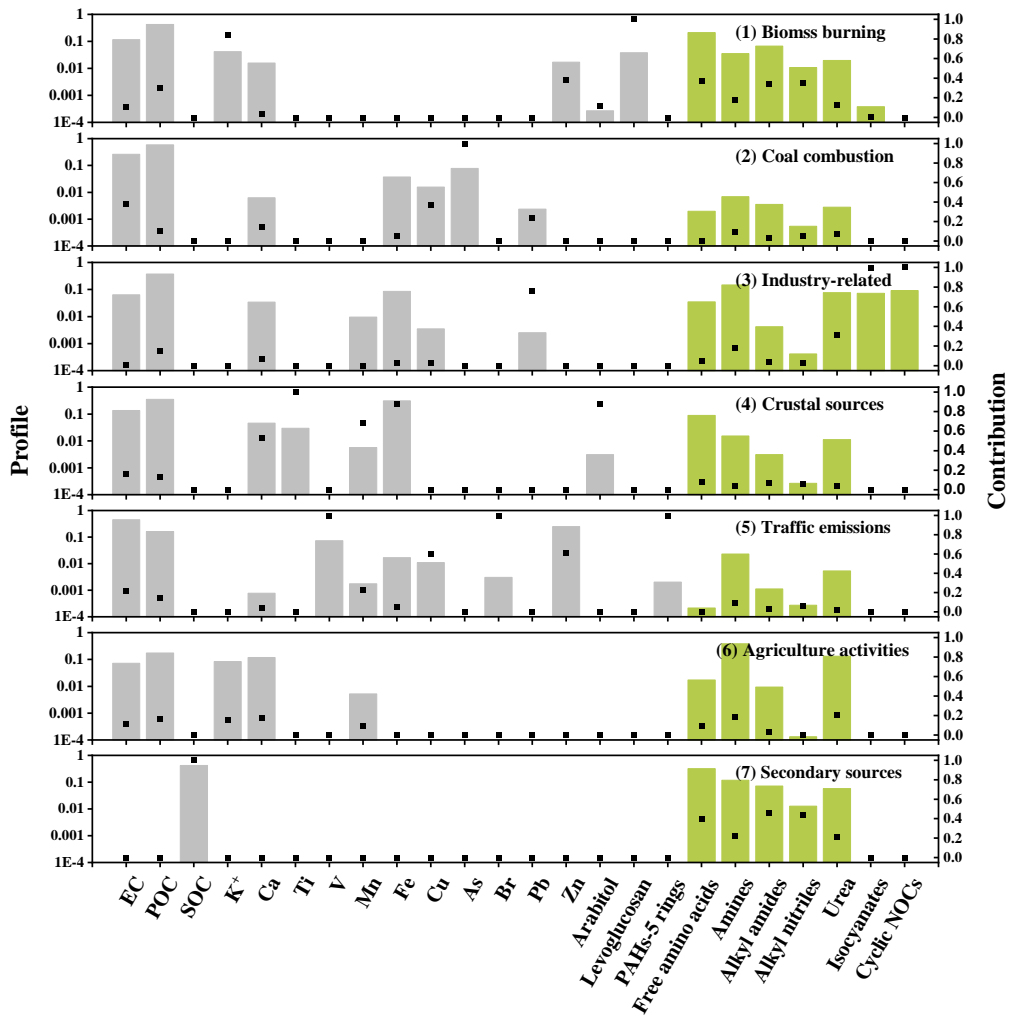
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677 **Figure 3** (a) Concentration and (b) composition of amines. (c) Temperature dependence of EA and MA,
678 and (d) RH dependence of EA and MA. The box represents the 25th (bottom) and 75th percentiles (top),
679 and the box-whisker data represents the 10th to 90th percentiles.

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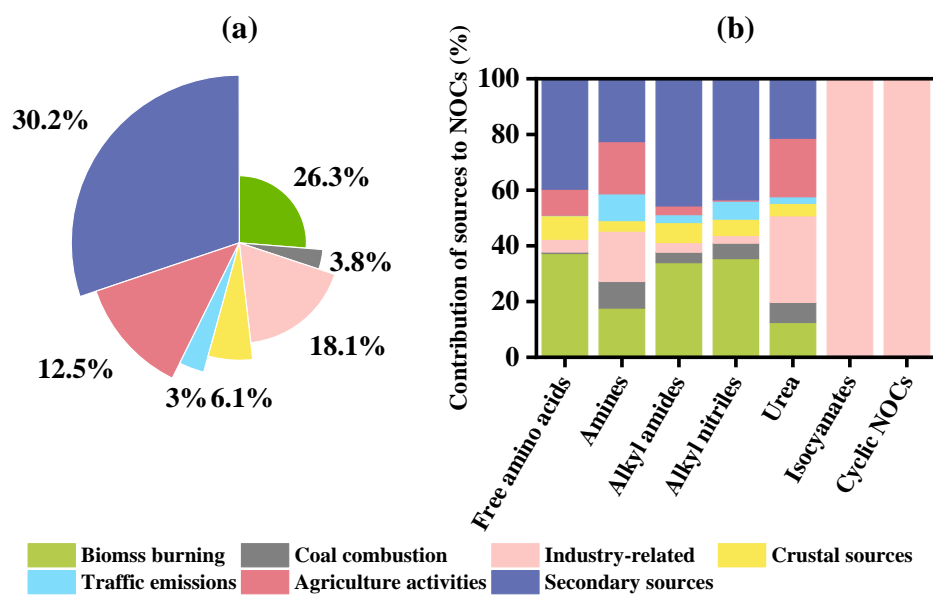


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682 **Figure 4** The factor profiles and explained variations in the ME-2 modeling.

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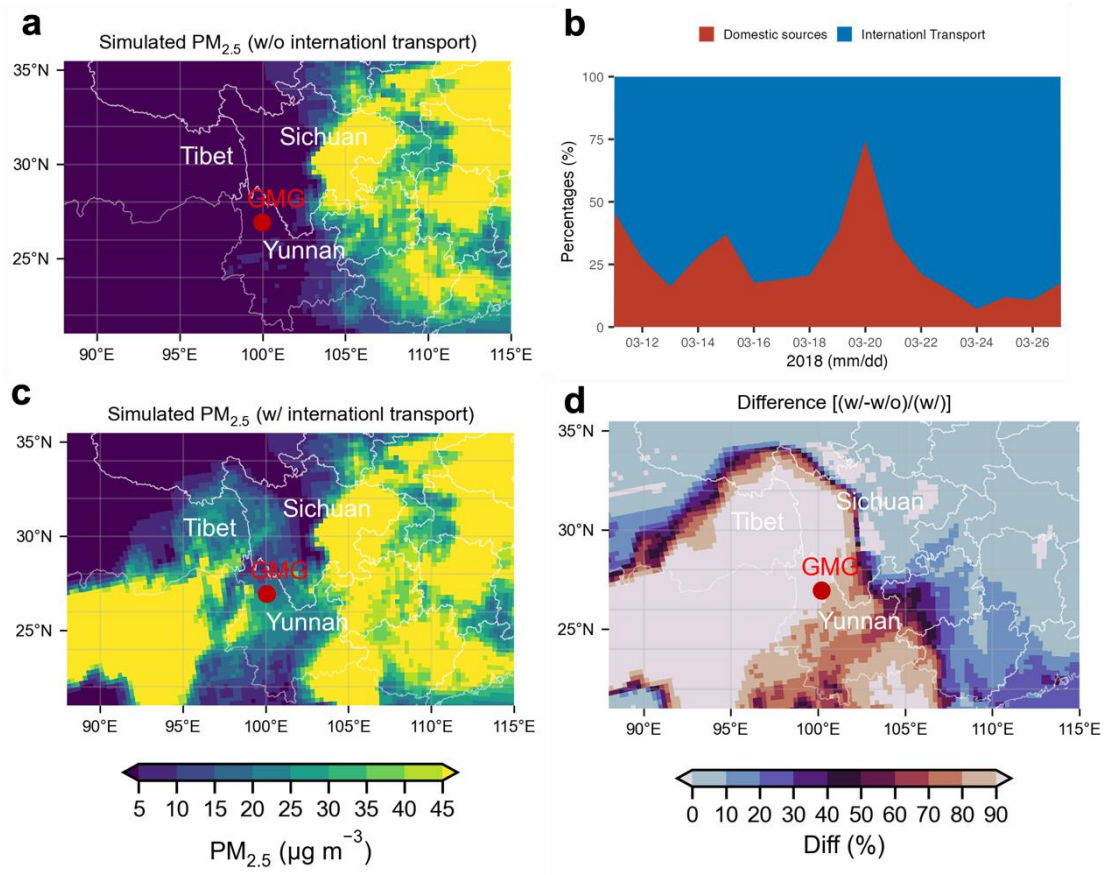
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686 **Figure 5** Contributions of each source to (a) total NOCs; and (b) seven classes NOC species.

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Figure 6. (a) Distribution of PM_{2.5} concentrations resulting solely from China's domestic emissions (MEIC-China only); (b) Proportionate contributions of domestic versus international PM_{2.5} transport during the simulation window of March 11th -27th 2018; (c) Distribution of PM_{2.5} incorporating both domestic and international transport influences (MEIC-China+MIX); (d) Difference of contribution of international transport to PM_{2.5} concentrations, derived from the differential analysis [(c)-(a)]/(c).