

High Frequency of Urban New Particle Formation on the Tibetan Plateau: Quantifying Formation Rates, Growth Rates, and CCN Production

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Abstract

New particle formation (NPF) strongly influences aerosol number size distributions and cloud condensation nuclei (CCN), yet its characteristics and climatic relevance in high-altitude urban environments over the Tibetan Plateau (TP) remain poorly constrained. We conducted in situ observations at Tibet University (TU) in Lhasa during the ~~spring monsoon~~ and ~~autumn observation periods~~~~pre-monsoon seasons~~. NPF event frequencies were 86.2 % during the ~~autumn observation period~~~~monsoon season~~ and ~~55.462.1~~ % during the ~~spring observation period~~~~pre-monsoon period~~. ~~Observation days were classified as NPF, Undefined, or Non-Event days based on PNSD evolution, supported by size-segregated particle number concentrations and particle growth rates. Observation-period comparisons indicate clear differences in the factors associated with NPF: NPF during the autumn observation period may have been favored by greater sulfur-precursor availability, whereas no single dominant controlling factor could be identified during the spring observation period. Estimated CCN concentrations were substantially higher on NPF days, with a mean concentration of $2294.4 \pm 1512.7 \text{ cm}^{-3}$, compared with $873.9 \pm 367.0 \text{ cm}^{-3}$ on Undefined days and $1278.1 \pm 667.4 \text{ cm}^{-3}$ on Non-Event days.~~ ~~monsoon season NPF was mainly associated with sulfur precursor availability, whereas no single clear controlling factor was evident during the pre-monsoon period. NPF substantially enhanced CCN at TU, with mean concentrations of $3351 \pm 1919 \text{ cm}^{-3}$ on Class I days, $1540.2 \pm 872 \text{ cm}^{-3}$ on Class II days, $880.5 \pm 374 \text{ cm}^{-3}$ on Undefined days, and $1278.1 \pm 667 \text{ cm}^{-3}$ on Non-Event days.~~ The impact of NPF on CCN depends not only on nucleation, but also on the continued growth and survival of newly formed particles. This study provides new insights into urban NPF characteristics and ~~the contribution of NPF~~ its potential contribution to CCN on the TP, improving our understanding of aerosol-climate interactions in high-altitude urban environments.

34 1 Introduction

35 The Sixth Assessment Report of the Intergovernmental Panel on Climate Change (IPCC) highlights that the aerosol climatic
36 effect is a major source of uncertainty in climate change assessment, primarily attributed to the spatio-temporal variability of
37 aerosol concentration and size distribution (Arias et al., 2021). New Particle Formation (NPF) and subsequent particle growth
38 significantly alter aerosol concentration, size distribution, and Cloud Condensation Nuclei (CCN) concentration (Hong et al.,
39 2023). Endowed with unique geographical and atmospheric conditions, the Tibetan Plateau (TP) hosts aerosols that are prone
40 to long-distance transport. This capability allows TP aerosols to modify atmospheric concentration and cloud properties along
41 their path more readily than those in low-altitude regions, thereby impacting both regional atmospheric environment and global
42 climate (Hu et al., 2024). Thus, in-depth exploration of NPF patterns over the TP is crucial for accurately evaluating TP aerosol
43 impact on the global atmosphere and climate.

44 NPF has also been frequently observed at high-altitude mountain and free-tropospheric sites outside the TP, where mountain-
45 valley circulation, boundary-layer transport, and low-temperature conditions can strongly influence precursor supply and
46 particle formation. At the Nepal Climate Observatory–Pyramid in the Himalayas, frequent NPF was previously observed under
47 the influence of valley winds (Venzac et al., 2008). More recent molecular-level observations at the same site showed that up-
48 valley transport supplied gaseous precursors that were oxidized into low-volatility organic compounds, producing new
49 biogenic particles that could subsequently enter the free troposphere (Bianchi et al., 2021). At the Chacaltaya high-altitude
50 station in Bolivia, NPF and subsequent particle growth produced substantial enhancements in CCN-relevant particle
51 concentrations under both boundary-layer-influenced and free-tropospheric conditions (Rose et al., 2017). These studies
52 demonstrate that high-altitude NPF can be strongly affected by vertical transport and organic precursor chemistry. In contrast,
53 Lhasa represents a high-altitude urban environment influenced by traffic, combustion, construction activities, and elevated
54 anthropogenic gaseous precursors. It therefore provides an opportunity to investigate how urban emissions modify mountain
55 NPF processes that have previously been studied mainly at remote background sites.

56 Research on NPF in the TP background atmosphere is extensive, leading to a comprehensive understanding of its
57 environmental and climatic effects. For instance, a study at the National Atmospheric Background Monitoring Sub-station on
58 Moshi Daban Mountain (3295 m a.s.l.) in Menyuan, Qinghai Province (Gao et al., 2025), found that NPF events occurred on
59 approximately 80 % of the observation days. These events typically started in the morning, with aerosol particle sizes rapidly
60 growing from approximately 5 nm to 150 nm, exhibiting the characteristic "banana-shape" curve. Intensive measurements at
61 Nam Co station (4730 m a.s.l.) in the central TP (Tang et al., 2023) explored NPF frequency and mechanisms, revealing
62 significant seasonal differences: approximately 15 % in the pre-monsoon season but up to 80 % during the summer monsoon.
63 Comprehensive analysis of the CS, gaseous precursors, and meteorological factors, combined with SO₂ and Volatile Organic
64 Compounds (VOCs) simulations, concluded that organics-involved condensation is the dominant NPF mechanism, with CS
65 and gaseous sulfuric acid having negligible influence. Furthermore, the high NPF frequency during the summer monsoon
66 season may be closely linked to frequent southerly and south-westerly airflow. Focusing on the south-eastern TP (Lai et al.,

2024), research investigated NPF jointly driven by anthropogenic and biogenic factors. Using field measurements and a chemical transport model, they found frequent NPF events on clear-sky days during the pre-monsoon season, contributing significantly to CCN concentration. Observations confirmed that Highly Oxygenated Organic Molecules (HOMs) from monoterpene oxidation participate in the nucleation process in this region. Liu et al. (2024) further showed that the frequent NPF at high-altitude sites in the south-eastern TP is mainly driven by HOMs, including products from monoterpene, and unexpectedly, sesquiterpenes and diterpenes oxidation. Moreover, anthropogenic NO_x emissions were found to play a crucial regulatory role in particle nucleation. Overall, NPF frequency is high in the TP background atmosphere, and its formation is closely related to temperature, relative humidity, and chemical precursor concentrations.

Aerosols generated by NPF in the TP background atmosphere can contribute up to half of the total atmospheric aerosol number concentration and significantly enhance CCN concentration. A study at the world's highest station, Chacaltaya (Rose et al., 2017)(~~Rose et al., 2017~~), analysed 61 % of NPF events, finding they led to increased CCN number concentration, with the probability reaching 79 % during the wet season due to faster particle growth. The NPF impact on CCN concentration varies seasonally, causing CCN to increase by a factor of 1.54 and 1.36 in spring and winter, respectively. During the summer monsoon season, at a supersaturation of 1.2 %, aerosol number concentration and CCN concentration surged by a factor of 2 and 0.6, respectively, compared to the pre-monsoon season. Furthermore, considering that small particles formed during NPF may continue to grow to CCN activation size over several days, the potential CCN number during the monsoon season may far exceed instantaneous local measurements (Tang et al., 2023). Comparative observations by Shang et al. (2022) indicated that the NPF contribution to total aerosol and CCN concentrations is more significant in the clean atmosphere than that of the polluted atmosphere.

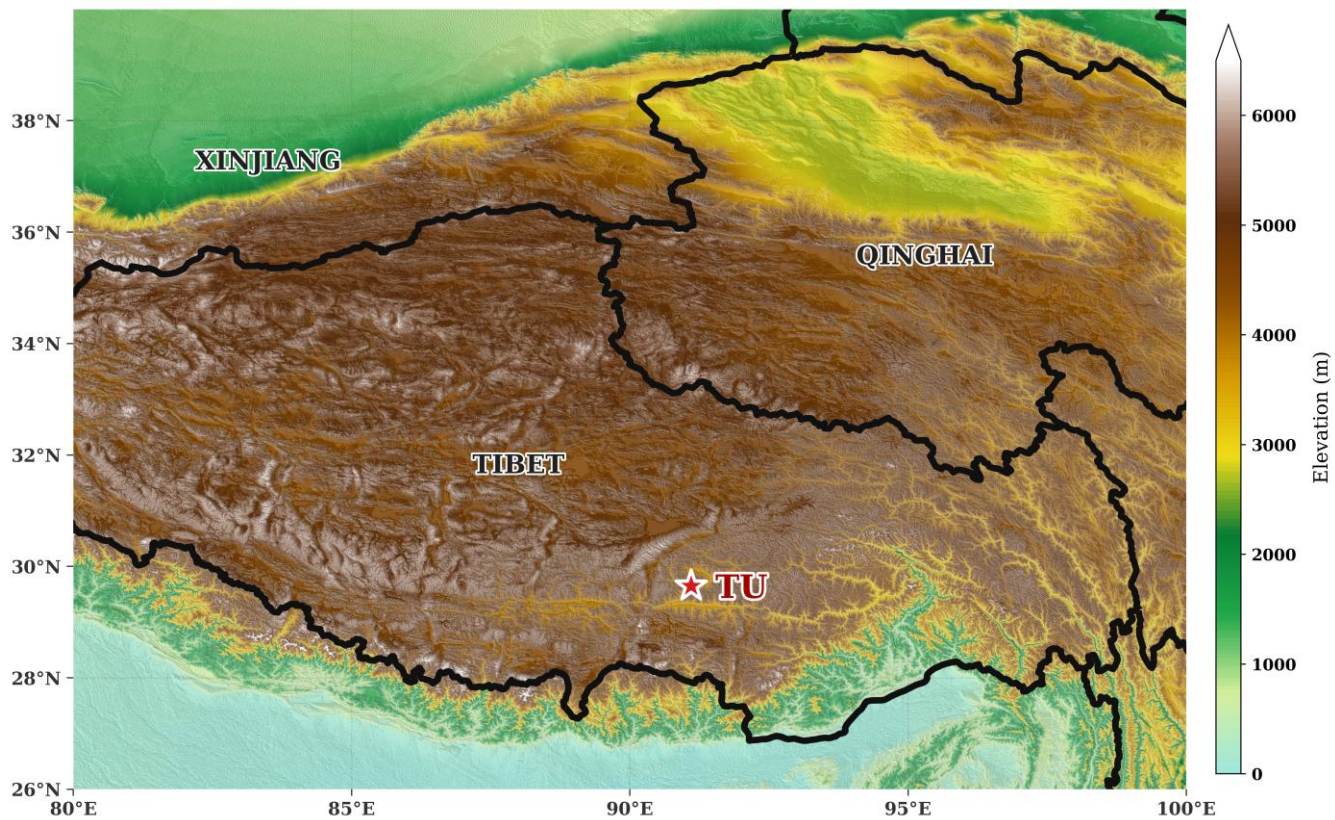
Despite the extensive work summarized above, our understanding of new-particle formation over the TP remains strongly biased toward remote background sites. The urban TP atmosphere—characterised by markedly stronger primary emissions and elevated concentrations of both SO₂ and NO_x—has, so far, received no systematic exploration of its NPF characteristics, governing mechanisms, or quantitative contribution to CCN. This knowledge gap hampers the accurate representation of aerosol sources in regional climate models and precludes a full assessment of the role of urban NPF over the TP in continental CCN budgets. To fill this gap, we carried out seasonal-scale particle number size distribution (PNSD) measurements in Lhasa (3650 m a.s.l.), a high-altitude urban environment on the Tibetan Plateau, during the ~~springpre-monsoon~~ and ~~autumn observation periodsmonsoon seasons~~. The specific aims are (i) to establish an event classification and to quantify the frequency and intensity of urban NPF, (ii) to elucidate the seasonal differences in precursor–meteorology interactions driving NPF, and (iii) to evaluate the instantaneous and seasonal enhancement of CCN concentrations attributable to newly formed particles at the TP urban environment. The observational methodology is described in Section 2, results and discussions are presented in Section 3, and conclusions together with implications for aerosol–climate interactions are summarized in the last Section.

98 2 Methods

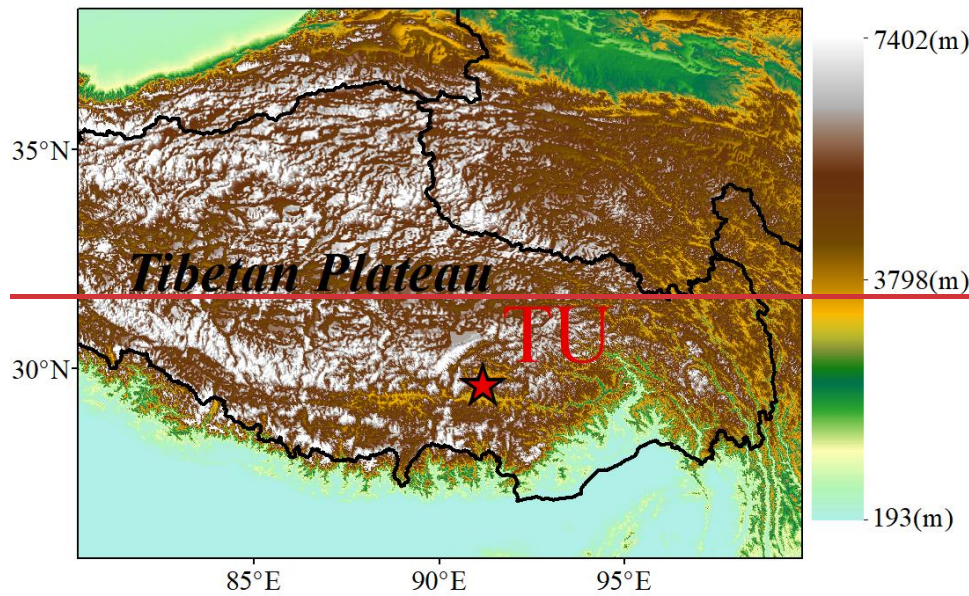
99 2.1 Sampling sites

100 As shown in Fig. 1, the study site is located at Tibet University (TU, 29 °38 ' N, 91 °10 ' E; 3650 m a.s.l.) in Lhasa, which lies
101 in the south-central TP. Lhasa serves as the capital of the Tibet Autonomous Region and its political, economic, and cultural
102 hub. As one of the largest urban centers on the TP, it is a typical high-altitude city where aerosols are predominantly driven by
103 local sources, including traffic emissions, residential coal/biomass combustion, construction activities, and dust resuspension
104 (Zhao et al., 2022). Traffic-related particle concentrations can exhibit pronounced spatial and temporal heterogeneity within
105 complex urban environments, which may not be fully captured by fixed-site measurements (Yeganeh et al., 2026). This spatial
106 heterogeneity should therefore be considered when interpreting the influence of local urban emissions at the TU site. The site
107 is situated in a broad river valley surrounded by mountains with elevations up to 5500 m a.s.l. Its climate is strongly influenced
108 by the Asian Summer Monsoon and the East Asian Winter Monsoon, with spring and autumn functioning as transitional
109 seasons between these two climate systems (Li et al., 2025). In situ measurements were conducted during two periods: the
110 autumn observation~~monsoon~~ period (9 October–10 November 2024) and the spring observation~~pre-monsoon~~ period (25
111 March–22 April 2025).

112



113



114 **Figure 1: Regional topography and location of the Tibet University (TU) observation site in Lhasa, Tibet Autonomous Region, China.**
 115 **Color shading denotes elevation above sea level (m), black lines indicate provincial-level administrative boundaries, and the red star**
 116 **marks the TU observation site.**

117 2.2 Instrumentation

118 In this study, a comprehensive Scanning Mobility Particle Sizer (SMPS) system was employed to measure the PNSD from 3
119 to 800 nm with a 5-minute time resolution. This system consisted of two separate instruments from TSI Inc. A nanoparticle
120 SMPS (TSI, Model 3087) was used to measure particles in the size range of 3 to 40 nm, while a standard SMPS (TSI, Model
121 3080) measured particles from 20 to 800 nm. The two datasets were then merged to create a complete and continuous PNSD.
122 To ensure consistent and accurate measurements, the instruments were housed in a temperature-controlled container
123 maintained at 25 °C. This setup was located on the rooftop of a campus building, with the aerosol inlet positioned
124 approximately 20 meters above ground level. This elevated sampling height was chosen to minimize the influence of localized
125 ground-level emission sources like traffic exhaust and suspended road dust. Furthermore, a silicon diffusion dryer was placed
126 upstream of the SMPSs to reduce the relative humidity of the sampled air to below 40 %, preventing particle growth from
127 water vapour condensation. Ambient aerosols were sampled through a short, conductive tube with minimal bends to reduce
128 size-dependent diffusion losses. All SMPS data underwent rigorous corrections for these diffusion losses and for multiple
129 charging effects to ensure the reliability of the measured particle number concentrations. We also utilized meteorological data
130 from an on-site automatic weather station, which provided essential parameters including temperature (T), relative humidity
131 (RH), wind speed (WS), wind direction (WD), Sulfur Dioxide (SO₂), Nitrogen Dioxide (NO₂), and Ozone (O₃) for a
132 comprehensive analysis of the atmospheric conditions. [Because gaseous H₂SO₄ was not directly measured, a relative H₂SO₄-
133 related photochemical proxy was additionally calculated for periods with available spectral measurements during the autumn
134 observation period; the calculation procedure and associated limitations are described in Sect. S1.](#)

135 2.3 Calculation of variables characterizing new particle formation

136 The Condensation Sink (CS, unit: s⁻¹) is utilized in this study as a core parameter to quantify the regulating effect of pre-
137 existing aerosols on condensable vapors (Kerminen et al., 2018; Wu et al., 2021). Its calculation is based on the kinetic theory
138 of dilute gases and assumes sulfuric acid (H₂SO₄) as the characteristic condensable substance (Dal Maso, 2005). The
139 calculation process utilizes measured PNSD data and incorporates the influence of ambient temperature (T) and pressure (P).
140 First, the diffusion coefficient of sulfuric acid vapor in the air (D_{x, air}) is calculated using an empirical formula based on T and
141 P. Subsequently, the mean free path of the condensable vapor (λ) is calculated using the molar mass of sulfuric acid (M_x =
142 98.08 g/mol). Based on λ and the size-resolved particle radius (R_i), the Knudsen number (Kn_i = λ/R_i) is determined. To correct
143 the condensation rate from the continuous regime to the transition and free molecular regimes, the transition regime correction
144 factor (β) is introduced (Dinoi et al., 2021):

$$145 \quad \beta = \frac{Kn + 1}{0.377Kn + 1 + \frac{4}{3\alpha}(Kn^2 + Kn)} \quad (14)$$

Here, the sticking coefficient (α) is set to 1.0. Finally, the size-resolved condensation sink (CSi) is determined by $4\pi \cdot D_{x,air} \cdot R_i \cdot \beta_i \cdot n_i$, and the total CS value is quantitatively calculated by numerically integrating CSi over all particle size ranges.

$$CS = 2\pi D \int_0^\infty d_p \beta_m(d_p) \times n(d_p) dd_p = 2\pi D \sum_i \beta_i d_{pi} N_i \quad (22)$$

The Coagulation Sink (CoagS, unit: s^{-1}) is another key parameter in atmospheric aerosol dynamics, quantifying the efficiency of pre-existing particles (scavenger particles) in removing specific size-range target particles from the system via the coagulation mechanism (Kulmala et al., 2001). Similar to CS, CoagS is calculated through PNSD integration.

$$CoagS_{d_p} = \int K(d_p, d_{p'}) n(d_{p'}) dd_{p'} \cong \sum_{d_{p'}=d_p}^{d_{p'}=\max} K(d_p, d_{p'}) N_{d_{p'}} \quad (33)$$

GR, defined as the rate at which the geometric mean diameter of nucleation mode particles increases from dp_1 to dp_2 , was calculated using the procedure outlined by (Kulmala et al., 2012).

$$GR = \frac{\Delta d_p}{\Delta t} = \frac{d_{p2} - d_{p1}}{t_2 - t_1} \quad (44)$$

To quantify the intensity of NPF events, we calculated the particle formation rate J (i.e., the rate of aerosol particle formation at a specific size) following the method described by (Baalbaki et al., 2021), which is derived from rearranging the equation governing the time evolution of particle number concentration (Kulmala et al., 2012). Specifically, J was computed for two size bins (dp): 3 nm (J_3), and 7 nm (J_7), with the upper size limits of the corresponding bins set to 7 nm and 20 nm, respectively. The growth rate term in the equation (Eq. 5) was defined as the average of GR measurements for total particles; this GR value was assumed constant during NPF events and set to zero outside the event duration or in non-event periods. The formation rate J_{dp} is expressed as:

$$J_{d_p} = \frac{dN_{d_p}}{dt} + CoagS_{d_p} \cdot N_{d_p} + \frac{GR}{\Delta d_p} \cdot N_{d_p} \quad (55)$$

CCN concentration is quantitatively calculated based on κ -Köhler theory (Petters and Kreidenweis, 2007). This method aims to determine the number of particles that can be activated at a specific supersaturation (S).

$$A = \frac{4\sigma_{s/a} M_w}{RT\rho_w} \quad (66)$$

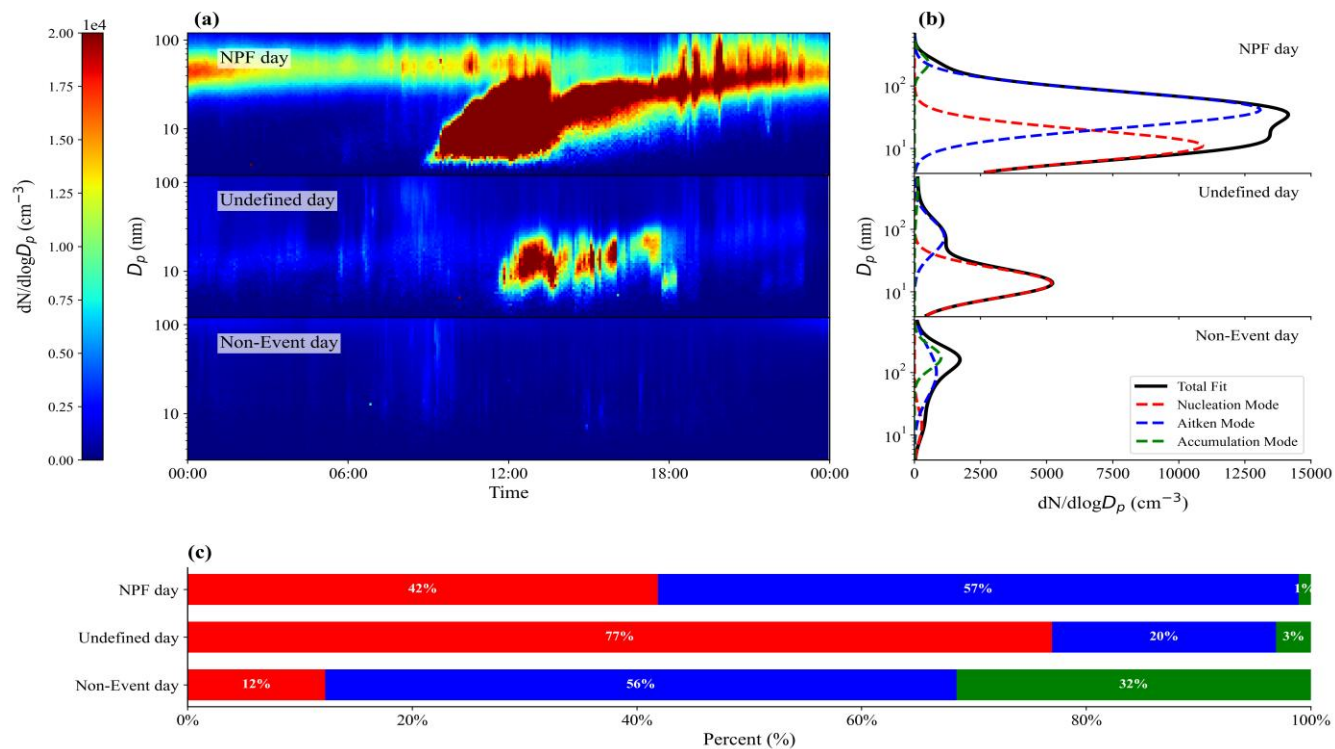
$$\kappa = \frac{4A^3}{27D_c^3 \ln^2 S_c} \quad (77)$$

In Eqs. (6) and (7), D_c is the critical dry particle diameter for CCN activation, S_c is the critical saturation ratio, and A is the Kelvin term. The parameter κ represents the influence of particle chemical composition on hygroscopicity and solution water activity. $\sigma_{s/a}$ is the surface tension of water against air. M_w and ρ_w are the molar mass and density of water, respectively. R is the universal gas constant, and T is the absolute temperature (K). In this study, κ and S were set to 0.12 and 1.2 %, respectively.

173 2.4 NPF Event Classification

174 In this study, atmospheric particles were divided into three diameter modes: the nNucleation Mmode (< 25 nm), the Aitken
175 Mode (25–100 nm), and the Aaccumulation Mmode (> 100 nm) (Wang et al., 2022; Hussein et al., 2020; Kalkavouras et al.,
176 2021; Shen et al., 2022). The classification followed the methodology proposed by Dal Maso et al. (2005), which has been
177 widely applied in previous NPF studies (Gao et al., 2025; Cramer et al., 2026; Lampilahti et al., 2025; Casans et al., 2025).
178 Based on visual inspection of the daily PNSD evolution, the observation days were classified as NPF days, Undefined days,
179 or Non-Event days. The classification was further supported by size-segregated particle number concentrations and
180 candidate particle growth rates.
181 NPF days were characterized by the appearance of a distinct new particle mode at small diameters followed by identifiable
182 growth toward larger sizes. Continuous growth throughout the entire day was not required because temporary interruptions
183 may result from changes in air masses, boundary-layer development, precursor availability, or missing measurements. All
184 remaining days were treated as non-NPF days and further divided into Undefined and Non-Event days. Undefined days showed
185 increases in small-particle concentrations or partial modal evolution, but the formation and growth patterns were too weak,
186 discontinuous, or ambiguous to be confidently classified as NPF. Non-Event days showed neither the distinct appearance of a
187 new particle mode nor its subsequent systematic growth. To improve the transparency of the classification, the mean N_{3-25} and
188 N_{25-100} concentrations during 09:00-18:00 local time and candidate GR_{7-20} values derived from continuous growth trajectories
189 within 07:00-18:00 were included as supporting metrics. These metrics were used to support the day-by-day classification
190 rather than as fixed classification thresholds. The revised classification, event timing, particle number characteristics, and
191 candidate GR_{7-20} values for individual observation days are provided in Table S1. Daily PNSD plots for NPF, Undefined, and
192 Non-Event days are presented in Figs. S1-S3. The former Class I and Class II categories were merged into the NPF-day
193 category for all main statistical analyses, while the original classification is retained in Table S1 only for comparison and
194 traceability.
195 Figure 2 presents representative dates for the NPF, Undefined, and Non-Event categories. Figure 2a shows their diurnal PNSD
196 evolution. On the NPF day, the number concentration of small particles began to increase at approximately 09:00 local time,
197 followed by an identifiable shift of the particle mode from the nucleation mode toward the Aitken mode, forming the
198 characteristic “banana-shaped” pattern. On the Undefined day, the modal evolution was relatively weak, discontinuous, or
199 ambiguous. On the Non-Event day, no distinct formation of a new particle mode followed by systematic growth was observed.
200 Figure 2b shows the corresponding mean PNSD and modal fitting results for the nucleation mode, Aitken mode, and
201 accumulation mode. The NPF day was characterized by substantial contributions from both the nucleation mode and Aitken
202 mode, consistent with the formation and subsequent growth of newly formed particles. The Undefined day was dominated by
203 particles in the nucleation mMode but showed relatively limited development toward larger sizes, whereas the Non-Event day
204 exhibited greater relative contributions from pre-existing particles in the Aitken mode and accumulation mode. Figure 2c
205 shows the relative contributions of the three particle modes. On the NPF day, particles in the nucleation mode, Aitken mode,

206 and accumulation mode contributed 42 %, 57 %, and 1 % of the total particle number concentration, respectively, consistent
207 with the growth of newly formed particles toward the Aitken mode. On the Undefined day, the corresponding contributions
208 were 77 %, 20 %, and 3 %, respectively, indicating that particles in the nucleation mode dominated but their subsequent growth
209 remained limited or ambiguous. On the Non-Event day, the corresponding contributions were 12 %, 56 %, and 32 %,
210 respectively, reflecting greater relative contributions from pre-existing particles in the Aitken mode and accumulation mode.
211 Fig. 2a illustrates typical examples of the diurnal evolution of the NPF event categories. On Class I days, the 3-nm particle
212 number concentration began to increase strongly around 09:00, and the mode diameter steadily grew from the Nucleation
213 Mode to the Aitken Mode, persisting until 0:00, displaying the classic “banana shape”. On Class II days, the number
214 concentration also began to increase around 09:00, but the growth of the mode diameter only lasted for a few hours, peaking
215 around 13:00. On Undefined days, although an increase in the number concentration of the Nucleation Mode was observed
216 in the morning, the mode diameter remained in the Nucleation Mode between 09:00 and 13:00 without significant growth.
217 On Non-Event days, neither an increase in Nucleation Mode number concentration nor growth into the Aitken Mode was
218 observed. Figure 2c presents a stacked percentage bar chart that quantifies the relative number contributions of nucleation,
219 Aitken, and accumulation modes to the total particle number concentration across these four classified NPF event types. On
220 Class I days, nucleation mode particles account for 42 % of the total number concentration, while Aitken mode particles
221 dominate with a 57 % contribution, and accumulation mode particles contribute a mere 1 %, indicating a scenario where
222 freshly formed nucleation mode particles undergo rapid growth into the Aitken mode. For Class II days, the nucleation mode
223 overwhelmingly dominates, representing 87 % of the total particle number, with Aitken and accumulation modes
224 contributing only 11 % and 2 %, respectively, which reflects relatively inefficient particle growth after initial formation.
225 Undefined days exhibit a similar modal contribution pattern to Class II days, with nucleation mode accounting for 78 %,
226 Aitken mode for 19 %, and accumulation mode for 3 %, suggesting some uncertainty in the particle growth process. In
227 contrast, Non-Event days show a more balanced contribution from the three modes, with nucleation mode at 12 %, Aitken
228 mode at 56 %, and accumulation mode at 32 %, reflecting the steady-state particle size distribution of the background
229 atmosphere in the absence of active NPF events. Collectively, this chart highlights the distinct modal contribution
230 characteristics of different NPF event types, providing critical statistical insights into the atmospheric physicochemical
231 processes governing new particle formation and growth.



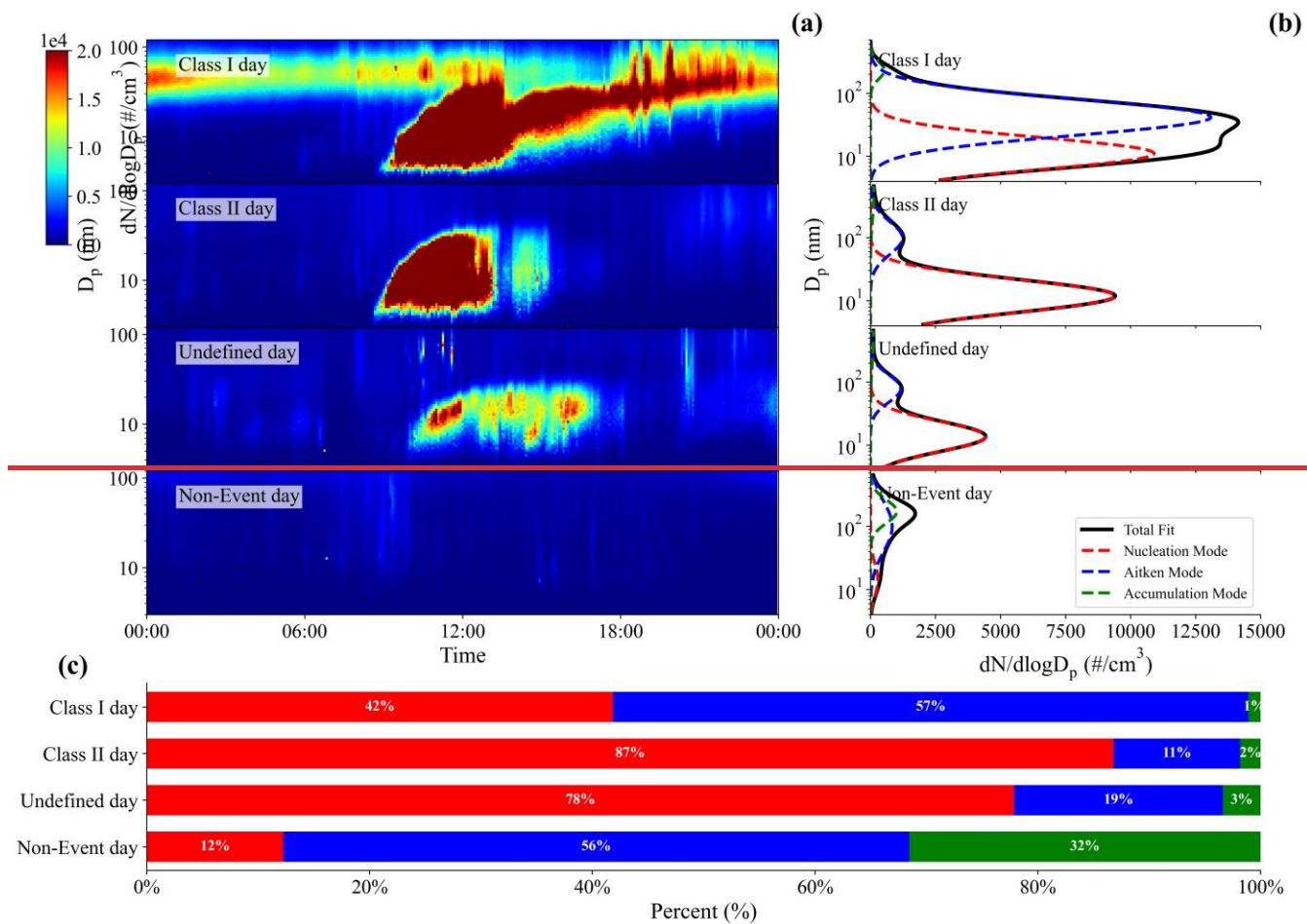


Figure 2: (a) Particle number size distribution evolution, (b) averaged modal fitting, and (c) relative number contributions for four classified day categories.

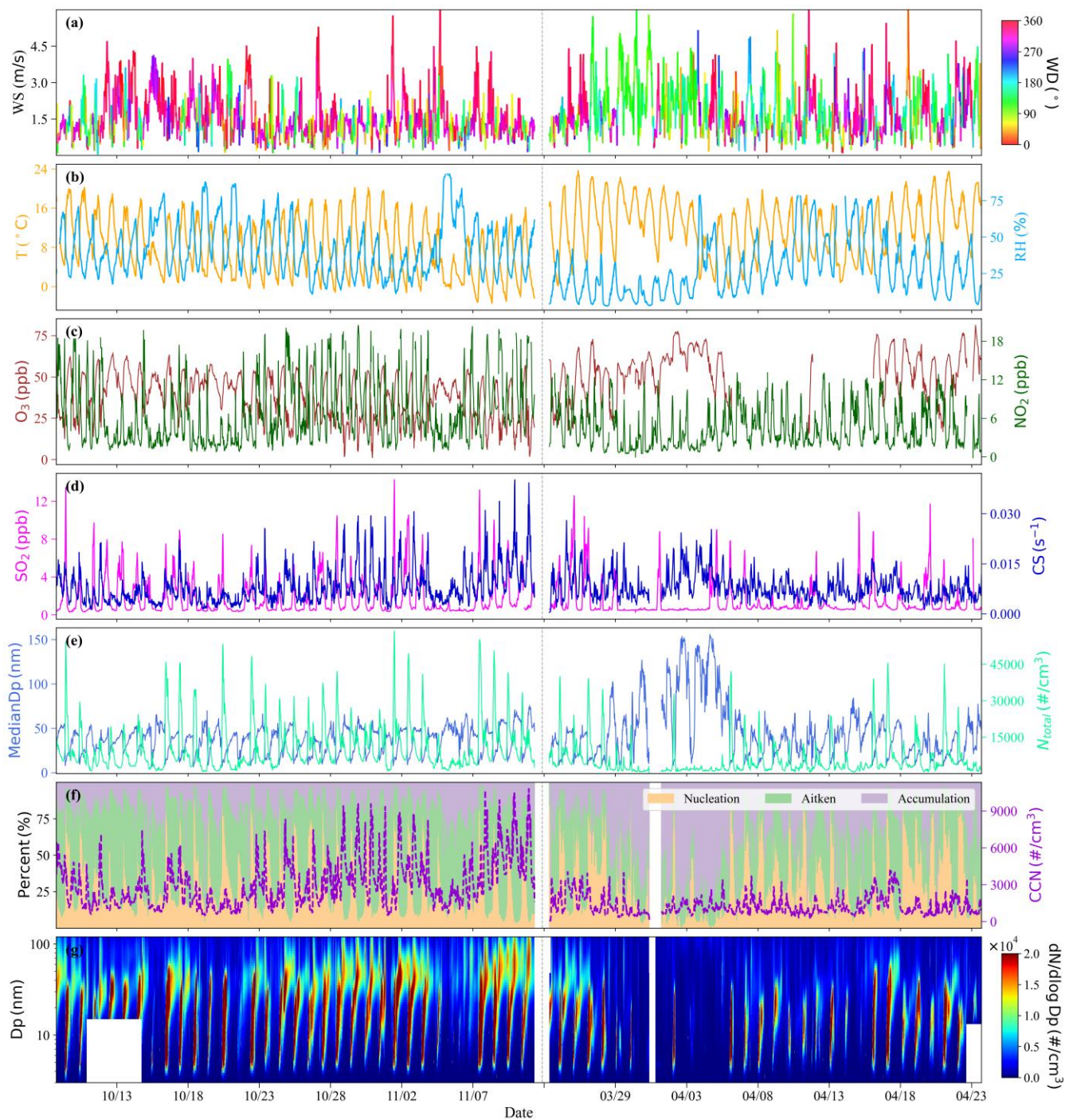
237 3 Results and discussion

238 3.1 Overview of the field observations

239 Meteorological conditions and gaseous pollutants exhibited strong short-term fluctuations and clear diurnal variability during
240 both measurement periods. As shown in Fig. 3a–b, ~~wind speed/wind direction~~ WS, WD and RH varied substantially on synoptic
241 and diurnal timescales. The ~~autumn observation period~~ monsoon period was characterized by higher RH (41.60 ± 17.78 %) ~~than the~~
242 ~~spring observation~~ pre-monsoon period (27.46 ± 18.23 %). Wind speeds were comparable between the two periods
243 (1.57 ± 0.92 m s⁻¹ in the ~~autumn observation~~ monsoon period and 1.87 ± 1.07 m s⁻¹ in the ~~spring observation~~ pre-monsoon
244 period), indicating predominantly low-to-moderate ventilation conditions that may facilitate episodic accumulation of
245 precursors and aerosols. Figure 3c–d presents the temporal evolution of SO₂, CS, O₃, and NO₂. O₃ was generally higher during
246 the ~~spring observation~~ pre-monsoon period (52.47 ± 14.11 ppb) than during the ~~autumn observation~~ monsoon period ($37.87 \pm$
247 12.75 ppb), suggesting a more oxidizing background in the ~~spring observation~~ pre-monsoon period. In contrast, SO₂ and NO₂
248 showed intermittent spikes in both ~~observation period~~ seasons, implying episodic impacts from transport and/or local
249 emissions. Notably, the CS remained similar across ~~observation period~~ seasons (0.0079 ± 0.0053 s⁻¹ in the ~~autumn~~
250 ~~observation~~ monsoon period and 0.0081 ± 0.0038 s⁻¹ in the ~~spring observation~~ pre-monsoon period), indicating that the
251 ~~observation-period difference~~ -contrast in NPF occurrence is unlikely driven solely by differences in the pre-existing particle
252 sink.

253 NPF events were frequently observed at the TU site and were associated with pronounced differences between the two
254 observation periods in PNSD and estimated CCN concentrations. In total, 413 NPF events, 6 Undefined days, and 9 Non-
255 Event days were identified during the 58 measurement days. The autumn observation period included 25 NPF days, 1
256 Undefined day, and 3 Non-Event days, whereas the spring observation period included 18 NPF days, 5 Undefined days, and 6
257 Non-Event days. Based on Fig. 3g, NPF occurrence displayed a marked ~~observation-period~~ contrast, with NPF observed on
258 86.2 % of the measurement days during the ~~autumn observation~~ monsoon period but only on ~~55.162.1~~ 55.162.1 % during the ~~spring~~
259 ~~observation~~ pre-monsoon period. Consistent with this contrast, Fig. 3e shows that the ~~autumn observation~~ monsoon period
260 exhibited a substantially higher total particle number concentration ($N_{3-700} = 10541 \pm 9771$ cm⁻³) than the ~~spring~~
261 ~~observation~~ pre-monsoon period (5335 ± 6886 cm⁻³), while the median particle diameter (median D_p) was smaller in the
262 ~~autumn observation~~ monsoon period (36.55 ± 13.99 nm) than in the ~~spring observation~~ pre-monsoon period (45.38 ± 33.60
263 nm), indicating a stronger contribution from smaller particles during the ~~autumn observation~~ monsoon period. Figure 3f further
264 presents the estimated CCN concentrations and the fractional contributions from different modes. CCN levels were higher
265 during the ~~autumn observation~~ monsoon period (2812.99 ± 1665.10 cm⁻³), accompanied by more frequent and more persistent
266 enhancements in the nucleation- and Aitken-mode fractions and episodic reductions in the accumulation-mode fraction,
267 suggesting a shift of the particle population toward the small-to-intermediate size range. In contrast, the ~~spring observation~~ pre-
268 monsoon period showed lower CCN (1152.05 ± 608.91 cm⁻³), weaker and less persistent increases in the nucleation- and
269 Aitken-mode fractions, and a more dominant background contribution from the accumulation mode. Together, these

270 observations highlight strong [observation-period differences](#) of particle sources and CCN-relevant size structure at the TU site
271 and provide context for the process-based analysis in the following sections.



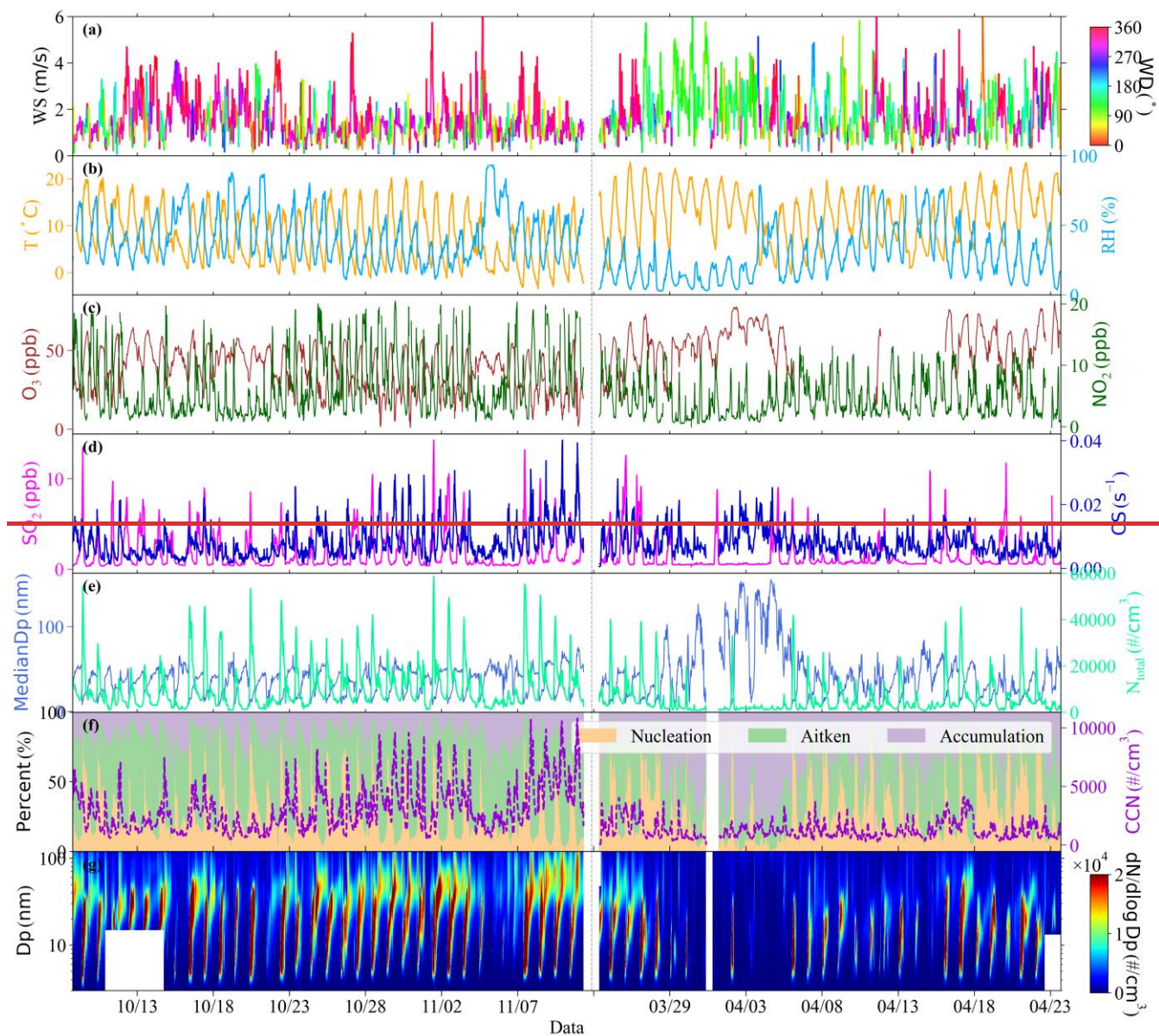


Figure 3: Time series of (a) wind speed and wind direction, (b) ambient temperature and relative humidity, (c) O₃ concentration and NO₂ concentration, (d) SO₂ concentration and Condensation Sink, (e) median particle diameter and particle number concentration in the size range of 3–700 nm, (f) CCN concentration and percentage contributions of the nucleation mode, Aitken mode, and accumulation mode, (g) PNSD during the spring-pre-monsoon and autumn observationmonsoon periods.

278 3.2 PNSD and number concentration responses

279 3.2.1 ~~M~~Seasonal mean PNSD and modal contributions

280 ~~The mean PNSD and modal contributions reveal clear differences in the campaign-mean particle size structure and surface-~~
281 ~~area/volume partitioning between the autumn and spring observation periods. Seasonal mean PNSD and modal contributions~~
282 ~~reveal a distinct background size structure and surface/volume partitioning between the monsoon and pre-monsoon periods.~~
283 ~~Figure 4a shows that the mean PNSD during the two observation periods at the TU site was typically multimodal, reflecting~~
284 ~~the combined influences of secondary particle production, particle growth, and the pre-existing aerosol population. Figure 4a~~
285 ~~shows that the seasonal mean PNSD at the TU site is typically multimodal, reflecting the combined influences of secondary~~
286 ~~particle production, growth, and the pre-existing aerosol population. The total number concentration differed markedly~~
287 ~~between the two observation periods. A pronounced seasonal contrast is evident in the total number concentration:~~ the mean
288 N_{3-700} during the ~~autumn observation~~monsoon period ($10541 \pm 9771 \text{ \# cm}^{-3}$) was 1.98 times higher than that during the ~~spring~~
289 ~~observation~~pre-monsoon period ($5335 \pm 6886 \text{ \# cm}^{-3}$) (Fig. 4a). In terms of modal partitioning, Fig. 4b indicates that while the
290 Aitken mode dominates number concentration in both periods, the surface area and volume are strongly controlled by the
291 accumulation mode in the ~~spring observation~~pre-monsoon period, accounting for 85 % (surface area) and 96 % (volume). In
292 contrast, during the ~~autumn observation~~monsoon period the Aitken mode contributes substantially more to surface area (42 %)
293 and volume (25 %) than in the~~spring observation period~~ (14 % and 4 %, respectively), consistent with a particle population
294 shifted toward smaller-to-intermediate sizes. These ~~differences between the two observation periods~~seasonal differences
295 provide a baseline for interpreting event-day diurnal changes and their implications for CCN-relevant size fractions.

296

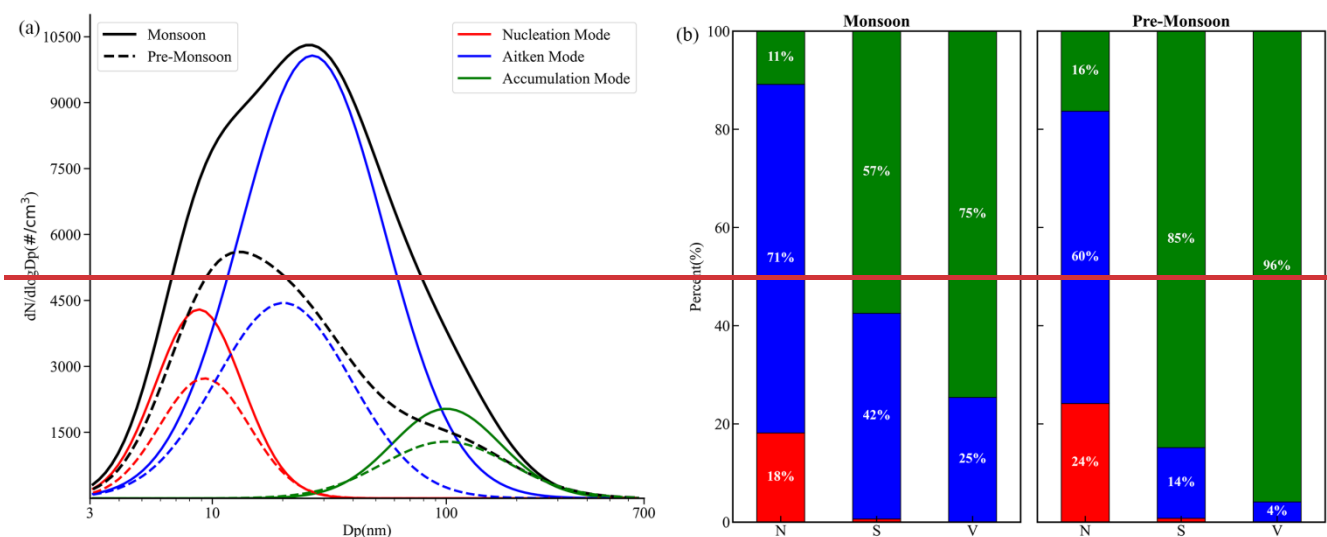
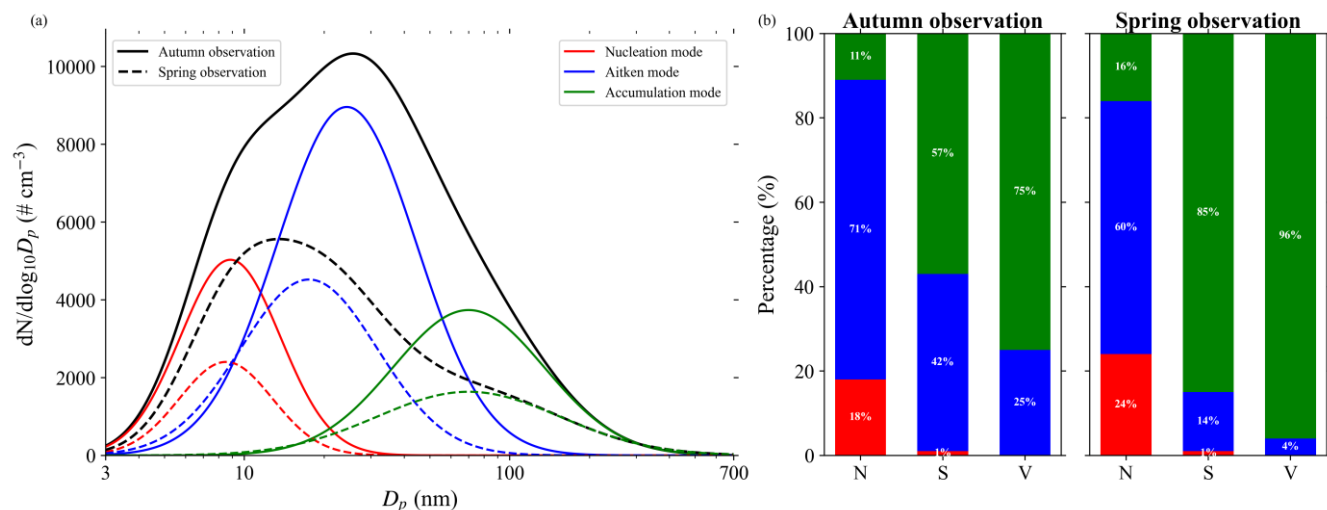
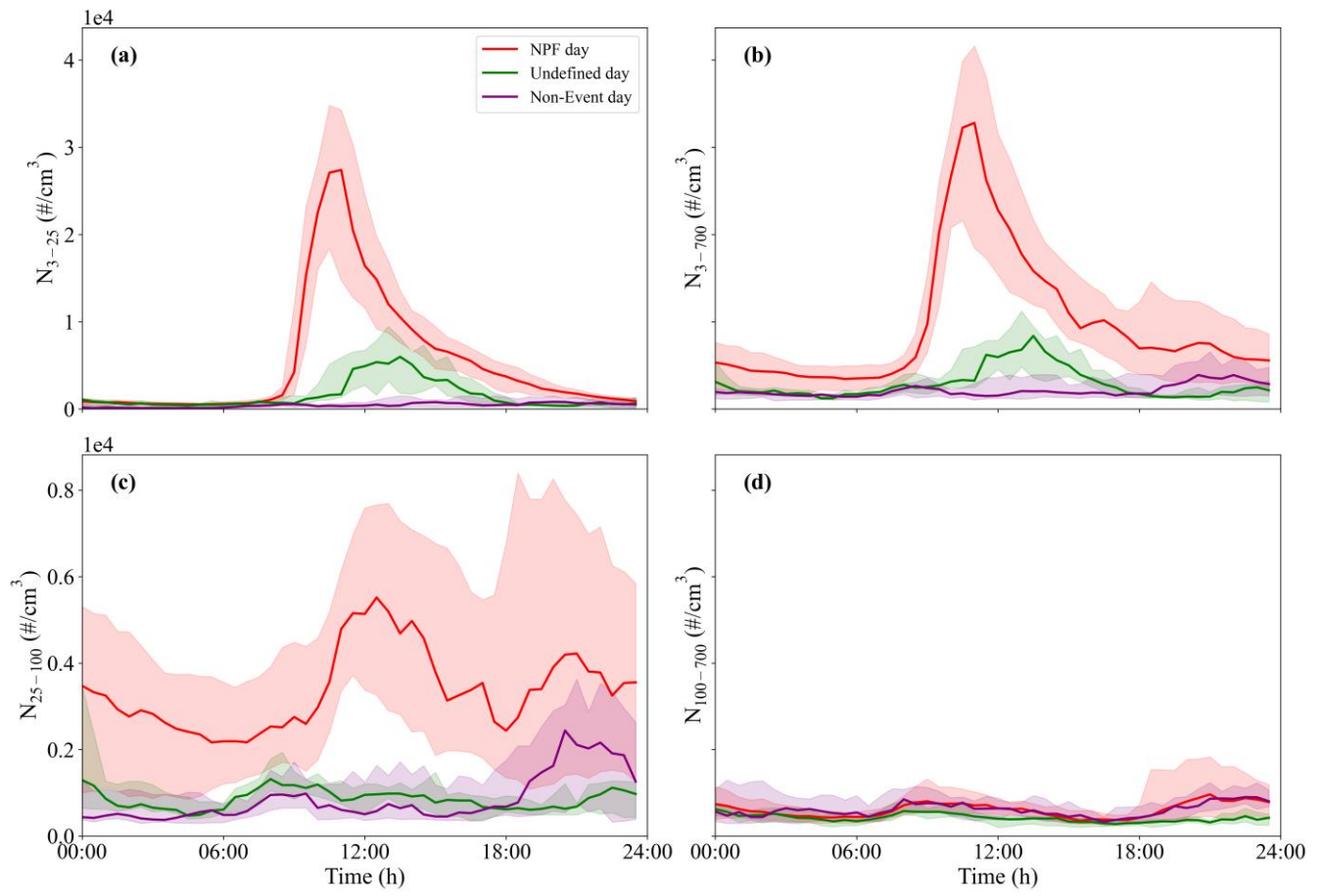


Figure 4: (a) Particle number size distribution characteristics and (b) modal contributions to number (N), surface area (S), and volume (V) distributions during the spring and autumn observation monsoon and pre-monsoon periods.

3.2.2 Diurnal variations of size-segregated number concentrations

The diurnal profiles of size-segregated particle number concentrations differed clearly among NPF, Undefined, and Non-Event days. Event day diurnal profiles show a clear sequence from daytime ultrafine particle production to enhanced Aitken mode concentrations, whereas non-event days exhibit weaker and less structured variability. Figure 5 summarizes diurnal profiles of particle number concentrations for the nucleation mode (N₃₋₂₅), Aitken mode (N₂₅₋₁₀₀), accumulation mode (N₁₀₀₋₇₀₀), and total measured size range (N₃₋₇₀₀) across the three day categories. On NPF days, N₃₋₂₅ increased rapidly during the morning, reached a pronounced maximum at approximately 11:00–12:00 local time, and subsequently declined (Fig. 5a). Undefined days

308 exhibited a weaker and broader increase from late morning to afternoon, whereas Non-Event days maintained substantially
309 lower N_{3-25} concentrations without a preceding N_{3-25} peak. N_{3-700} showed a similar pattern, with the highest and most
310 pronounced daytime maximum occurring on NPF days (Fig. 5b). N_{25-100} on NPF days exhibited a broad daytime enhancement
311 from approximately 11:00 to 15:00, followed by a secondary increase in the evening (Fig. 5c). The daytime enhancement is
312 consistent with the growth of newly formed particles into the Aitken mode, whereas the evening increase may also be
313 influenced by local urban emissions. Undefined days showed relatively weak variations in N_{25-100} , while Non-Event days
314 exhibited a more evident evening increase without a preceding increase in the nucleation mode. Differences in $N_{100-700}$ among
315 the three categories were relatively small, although modest morning and evening enhancements were observed (Fig. 5d). These
316 diurnal statistics indicate that the impact of NPF on the size distribution depends not only on daytime particle production, but
317 also on the persistence of growth into the Aitken mode.



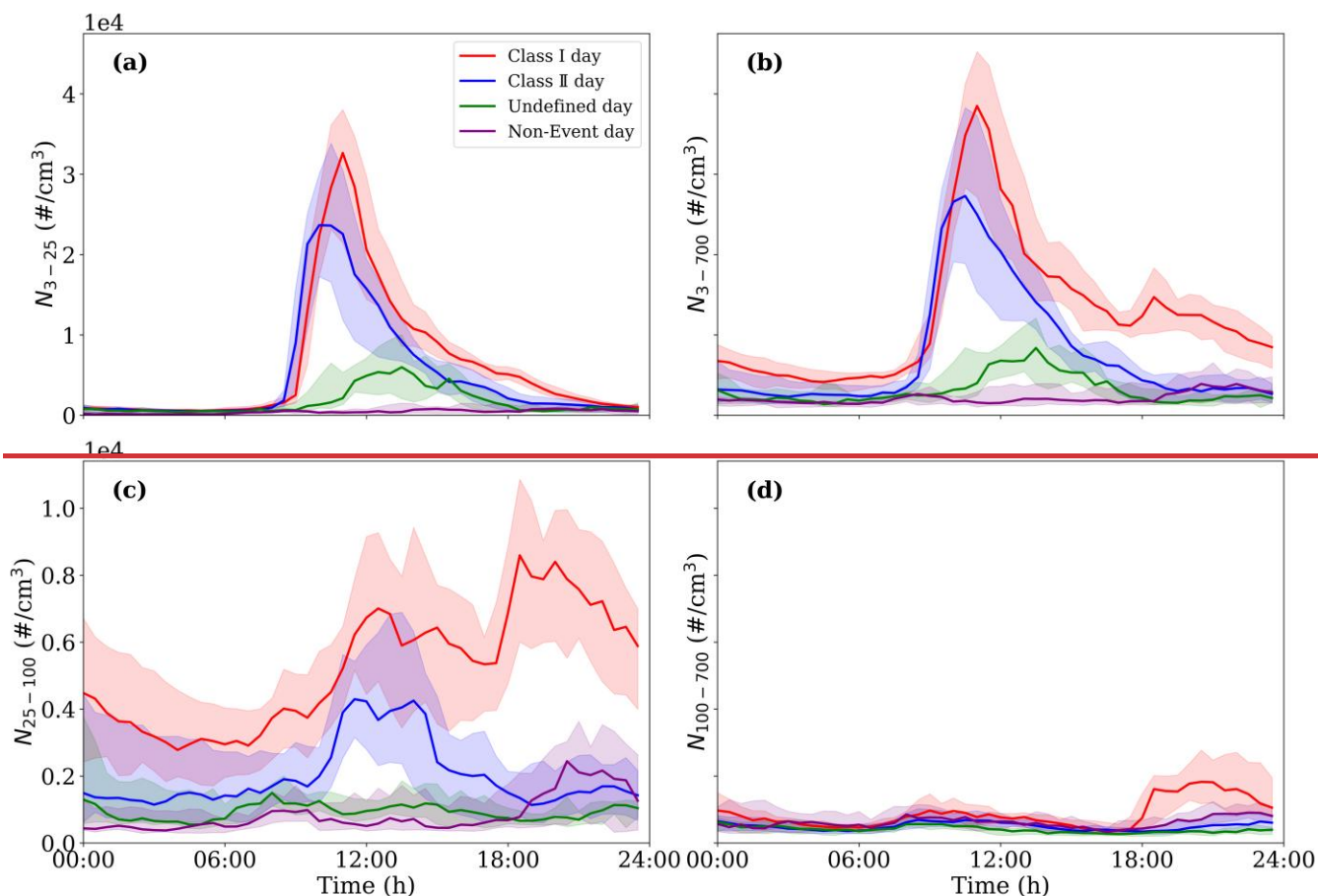


Figure 5: Diurnal variation of aerosol number concentrations ($\# \text{ cm}^{-3}$) at the size ranges of (a) 3–25 nm, (b) 3–700 nm, (c) 25–100 nm, and (d) 100–700 nm, during on NPF, Undefined, and Non-Event days. Shaded areas represent the interquartile range (25th to 75th percentiles).

3.3 Factors controlling the formation and development of NPF events

3.3.1 Formation and growth characteristics

Figure 6 summarizes the diurnal variations in particle formation rates at 3 nm (J_3) and 7 nm (J_7) for the three day categories. On NPF days, both formation rates increased rapidly during the morning and reached pronounced maxima at approximately 09:00–11:00, followed by a gradual decline. Undefined days exhibited weaker and less persistent daytime enhancements, whereas Non-Event days generally showed the lowest formation-rate signals and lacked a distinct morning peak. These contrasting diurnal patterns support the occurrence of active daytime particle formation on NPF days.

During 08:00–16:59, the mean J_3 and J_7 on NPF days were $1.19 \pm 2.00 \text{ cm}^{-3} \text{ s}^{-1}$ and $2.17 \pm 3.26 \text{ cm}^{-3} \text{ s}^{-1}$, respectively. The observed J_3 was of the same order of magnitude as the formation rates reported at Nam Co ($J_3 = 1.15 \pm 0.58 \text{ cm}^{-3} \text{ s}^{-1}$). The formation and growth characteristics of NPF were quantified using particle formation rates (J) and growth rates (GR), and the

333 diurnal evolution of formation rates in the 3–7 nm and 7–20 nm size ranges is summarized in Fig. 6. On NPF days, both Class
 334 I and Class II events exhibit pronounced daytime increases in (J_{3-7}), consistent with active photochemically driven particle
 335 production during the morning to midday period, whereas Non-Event days show substantially weaker formation. Averaged
 336 over the daytime period (08:00–16:59), the mean formation rates in the initial nucleation stage (3–7 nm) were $1.13 \pm 1.60 \text{ cm}^{-3} \text{ s}^{-1}$
 337 for Class I events and $1.15 \pm 1.69 \text{ cm}^{-3} \text{ s}^{-1}$ for Class II events, indicating strong daytime particle production for both event
 338 classes. These values are close to results from the Nam Co station ($J_4: 1.15 \pm 0.58 \text{ cm}^{-3} \text{ s}^{-1}$) (Tang et al., 2023) and the Mt.
 339 Yulong station ($J_3: 1.18 \text{ cm}^{-3} \text{ s}^{-1}$) (Shang et al., 2018). ~~As particles evolved into the 7–20 nm range, the formation rate (J_{7-20})~~
 340 ~~further intensified to approximately $3.6 \text{ cm}^{-3} \text{ s}^{-1}$, remaining within the reasonable range established by high-altitude~~
 341 ~~environmental observations. The mean GR_{3-7} was $3.73 \pm 1.12 \text{ nm h}^{-1}$, whereas the mean GR_{7-20} was $2.03 \pm 1.72 \text{ nm h}^{-1}$. These~~
 342 growth rates were broadly comparable to values at ~~Regarding growth kinetics, the observed GR_{3-7} ($2.31\text{--}3.16 \text{ nm h}^{-1}$) and~~
 343 GR_{7-20} ($2.97\text{--}3.51 \text{ nm h}^{-1}$) were comparable to values at Nam Co ($\text{GR}_{4-25}: 4.0 \pm 1.2 \text{ nm h}^{-1}$) and Mt. Yulong (3.2 nm h^{-1}), but
 344 higher than the value reported ~~significantly exceeding those~~ at the NCO-P station ($1.18 \pm 0.7 \text{ nm h}^{-1}$) (Venzac et al.,
 345 2008) ~~(Venzac et al., 2008)~~ and Mt. Daban (2.0 nm h^{-1}) ~~(Du et al., 2015)~~. In summary, the J and GR values at TU are well
 346 within the ranges reported for global high-altitude environments, indicating that the site experiences characteristic high-altitude
 347 NPF with strong daytime formation and sufficiently rapid growth to promote downstream enhancement of Aitken-mode
 348 number concentrations.

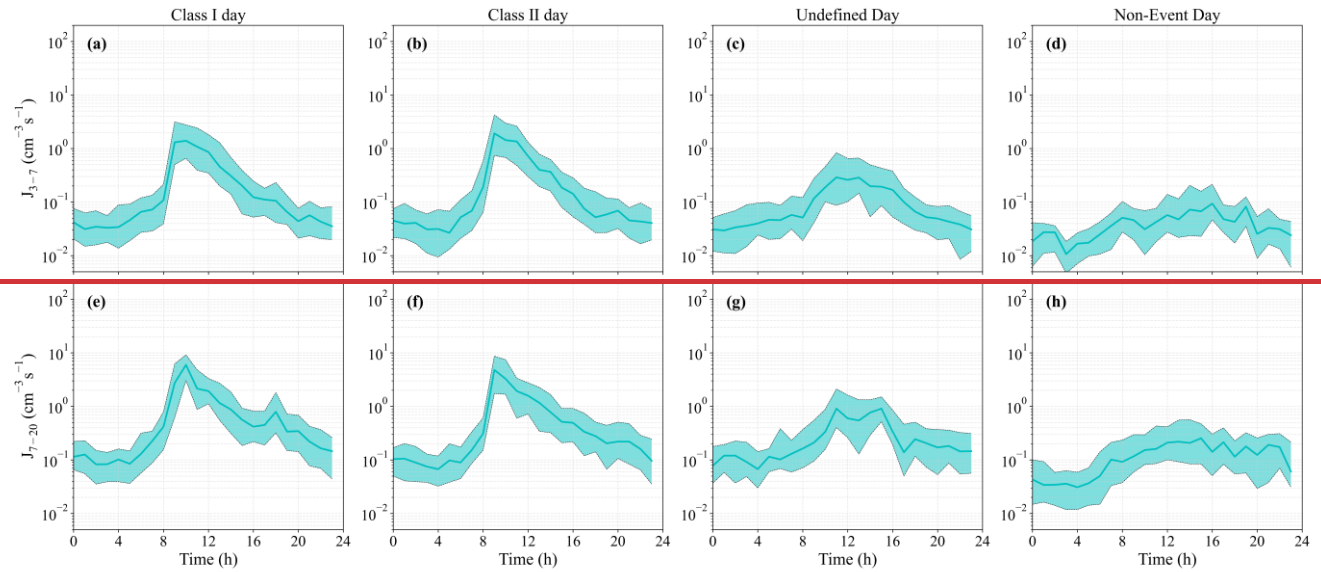
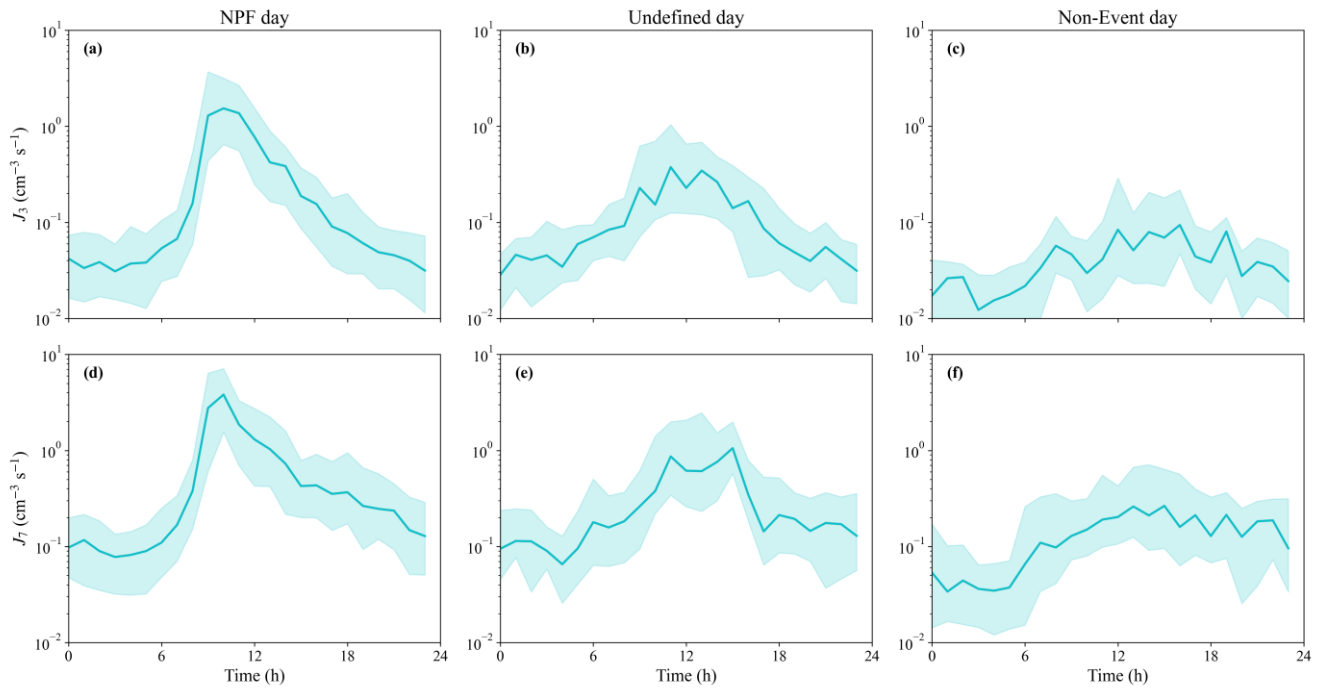


Figure 6: Diurnal variations of the formation rates J_3 during (a) **Class I NPF** day, (b) **Class II day**, (c) Undefined day, and (d) Non-Event day periods. The corresponding diurnal variations of the formation rate J_7 were shown in panels (e), (f), (g), and (h). The shaded areas represent the 25th and 75th percentile boundaries, and the solid lines denote the median values.

3.3.2 Condensation sink and its role

CS represents the ability of the pre-existing aerosol population to remove condensable vapors via condensation, while CoagS describes the loss of freshly formed clusters and ultrafine particles through collision and coalescence with larger background particles (Victor et al., 2024). Figure 7 shows the diurnal variations of CS and CoagS for different day types and observation periods. Both sink terms exhibit clear diurnal structure and substantial variability, indicating that the removal capacity for vapors and nascent particles can change markedly over the course of a day and between different regimes.

Consistent with many studies reporting that NPF tends to be favored under relatively low CS (Cai et al., 2017; Saha et al., 2018; Kalivitis et al., 2019; Kalkavouras et al., 2020; Aktypis et al., 2023), the TU site nevertheless shows that NPF can occur even when CS is not minimal. The mean CS levels ($\pm 1\sigma$) ~~are were~~ $0.00769 \pm 0.00445 \text{ s}^{-1}$ on NPF days, $0.00531 \pm 0.00453 \text{ s}^{-1}$ ~~on for~~ Undefined days, and $0.0064 \pm 0.0044 \text{ s}^{-1}$ ~~on for~~ Non-Event days (Fig. 7 a–d). Thus, CS on NPF days ~~is was~~ slightly higher than that on Non-Event days, and the NPF-day mean CS at TU is higher than values reported for remote plateau background sites in China (Tang et al., 2023), yet lower than those reported for the major urban environment of Beijing (Du et al., 2017), suggesting an intermediate sink environment characteristic of a high-altitude urban setting. Moreover, the mean CS values were broadly similar between the autumn and spring observation periods ~~monsoon~~ ($0.0069 \pm 0.0041 \text{ s}^{-1}$) and ~~spring pre-monsoon~~ ($0.0062 \pm 0.0047 \text{ s}^{-1}$) observation periods (Fig. 7 e–f), ~~indicating that differences in the pre-existing particle sink are unlikely to be the primary driver of the seasonal contrast in NPF occurrence. Therefore, differences in the pre-existing particle sink were unlikely to be the primary explanation for the contrast in NPF frequency between the two observation periods.~~

In contrast, CoagS shows a stronger separation among day types and observation periods. The mean CoagS values were $(4.25 \pm 2.58) \times 10^{-5}$ ~~(4.25 ± 2.58) 5.27 × 10⁻⁵~~ $\pm 2.62 \times 10^{-11} \text{ s}^{-1}$ on NPF days, $(2.18 \pm 1.23) \times 10^{-5}$ ~~(2.18 ± 1.23) 3.66 × 10⁻⁵~~ $\pm 2.39 \times 10^{-11} \text{ s}^{-1}$ ~~for~~ Undefined days, and $(2.54 \pm 1.43) \times 10^{-5}$ ~~(2.54 ± 1.43) 2.57 × 10⁻⁵~~ $\pm 1.53 \times 10^{-11} \text{ s}^{-1}$ ~~for~~ Non-Event days (Fig. 7 g–h). The mean CoagS was also higher during the autumn observation period $(4.49 \pm 2.57) \times 10^{-5}$ ~~(4.49 ± 2.57) × 10⁻⁵~~ s^{-1} than during the spring observation period $(2.88 \pm 1.97) \times 10^{-5}$ ~~(2.88 ± 1.97) × 10⁻⁵~~ s^{-1} (Fig. 7 i–j). Seasonally, CoagS is also markedly higher during the autumn ~~monsoon~~ $(4.85 \times 10^{-11} \pm 2.57 \times 10^{-11} \text{ s}^{-1})$ than during the spring observation ~~pre-monsoon period~~ $(2.88 \times 10^{-11} \pm 1.93 \times 10^{-11} \text{ s}^{-1})$. Because CoagS reflects the scavenging loss of newly formed clusters and ultrafine particles, its elevated value during the autumn observation period indicates a stronger constraint on particle survival. However, NPF occurred more frequently during this period, suggesting that particle-production processes were sufficiently strong to offset the enhanced coagulation loss.

Taken together, these results suggest that the occurrence and development of NPF at TU are better interpreted as the outcome of competition between particle-production and loss processes rather than as being controlled by the sink alone. This interpretation is consistent with previous observations that NPF can be observed under moderate sinks when vapor production is sufficiently strong (Zhang et al., 2021; Sellegri et al., 2019; Wang et al., 2022; Baalbaki et al., 2021). In this framework, CoagS provides an additional constraint on whether newly formed clusters can survive long enough to grow into larger sizes,

387 while CS limits the availability of low-volatility vapors required for sustained growth; hence, event intensity and downstream
 388 impacts are expected to depend on the joint evolution of formation/growth rates and sinks over the diurnal cycle.

389

390

391

392 **Figure 7: Diurnal variations of condensation sink and coagulation sink for different event classifications and observation**
 393 **periodsseasons.**

394 **3.3.3 Meteorology and precursor modulation, with transport context**

395 Meteorological conditions during the study period exhibited clear diurnal variability, consistent with the dependence of NPF
 396 on atmospheric state and precursor availability reported in previous studies (Hakala et al., 2022; Kalkavouras et al., 2020;

397 [Kalkavouras et al., 2019; Victor et al., 2024; Singh et al., 2025](#)). The site was predominantly under clear to partly cloudy
398 conditions with sufficient solar radiation, occasionally interrupted by precipitation. Notably, an extended rainfall episode
399 occurred from 4 to 6 November during the autumn observation period, during which no NPF events were identified. The
400 absence of NPF during this episode may have been associated with wet scavenging and reduced photochemical production of
401 condensable vapors under rainy and cloudy conditions.

~~To quantify how meteorology and key gaseous precursors are associated with NPF, we compared SO₂, O₃, and NO₂ together
402 with temperature, RH, and wind speed across day types and between the two observation periods. To examine the associations
403 of meteorological conditions and key gaseous precursors with NPF, we compared SO₂, O₃, and NO₂, together with temperature,
404 RH, and wind speed, across day types and between the two observation periods. The mean temperature were observed: mean
405 temperature was higher in the spring observation period (11.38 ± 5.77 °C) than in the autumn observation period ($7.77 \pm$
406 5.60 °C), whereas RH was substantially higher during the autumn observation period (42.02 ± 18.11 %) than during the
407 spring observation period (28.57 ± 20.42 %). Wind speed was slightly higher during the spring observation period ($2.46 \pm$
408 1.78 m s⁻¹) than during the autumn observation period (1.97 ± 1.32 m s⁻¹), indicating comparatively weaker ventilation
409 during the autumn observation period, which may have favored the episodic accumulation of gaseous precursors and particles.
410 During the autumn observation period, NPF days exhibited substantially higher SO₂ and NO₂ than non-NPF days: SO₂ averaged
411 1.92 ± 2.08 ppb on NPF days, clearly exceeding 0.73 ± 0.75 ppb on non-NPF days, while NO₂ averaged 7.74 ± 5.69 ppb on
412 NPF days compared with 4.38 ± 2.99 ppb on non-NPF days. The sector plots further showed that elevated SO₂ and NO₂
413 occurred more frequently under southerly winds on NPF days, whereas concentrations in the corresponding sectors were
414 generally lower on non-NPF days (Fig. 8). Because SO₂ oxidation by OH[·] produces H₂SO₄, and H₂SO₄ is widely recognized
415 as a key nucleation precursor and critical for early particle growth, the higher SO₂ concentrations observed on NPF days
416 suggest that sulfur-containing precursors may have played an important role in NPF during the autumn observation period
417 (Olin et al., 2020; Yao et al., 2018; Kerminen et al., 2018). Conditions during the study period show clear diurnal variability in
418 temperature and relative humidity, consistent with the strong dependence of NPF on atmospheric state and precursor
419 availability reported in previous studies (Hakala et al., 2022; Kalkavouras et al., 2020; Kalkavouras et al., 2019; Victor et al.,
420 2024; Singh et al., 2025). The site was predominantly under clear to partly cloudy conditions with sufficient solar radiation,
421 occasionally interrupted by precipitation. Notably, an extended rainfall episode occurred from 4 to 6 November during the
422 autumn observation monsoon period, during which no NPF events were identified. To quantify how meteorology and key
423 gaseous precursors are associated with NPF, we compared SO₂, O₃, and NO₂ together with temperature, RH, and wind speed
424 across day types and . were observed: mean temperature was higher in the spring observation pre monsoon period ($11.38 \pm$
425 5.77 °C) than in the autumn observation monsoon period (7.77 ± 5.60 °C), whereas RH was substantially higher in the autumn
426 observation period monsoon (42.02 ± 18.11 %) than in the spring observation pre monsoon period (28.57 ± 20.42 %). Wind
427 speed was slightly higher in the spring observation pre monsoon period (2.46 ± 1.78 m s⁻¹) than in the autumn
428 observation monsoon period (1.97 ± 1.32 m s⁻¹),~~

During the ~~autumn observation period~~ monsoon season, NPF days exhibited substantially higher SO₂ and NO₂ than non-NPF days: SO₂ averaged 1.92 ± 2.08 ppb on NPF days, clearly exceeding 0.73 ± 0.75 ppb on non-NPF days, while NO₂ averaged 7.74 ± 5.69 ppb on NPF days compared with 4.38 ± 2.99 ppb on non-NPF days. Because SO₂ oxidation by OH[•] produces H₂SO₄, and H₂SO₄ is widely recognized as a key nucleation precursor and critical for early particle growth, (Olin et al., 2020; Yao et al., 2018; Kerminen et al., 2018). Although elevated NO₂ can alter atmospheric radical chemistry and potentially reduce OH availability under some conditions, the clearly elevated SO₂ observed on NPF days suggests that plentiful SO₂ increases the potential for H₂SO₄ formation. This interpretation is further supported by Fig. S54, which shows that during the particle-formation period SO₂ concentrations on NPF days are clearly higher than those on non-NPF days. Moreover, for periods with available spectral measurements, the relative H₂SO₄-related proxy was also higher on NPF days during the particle-formation period (Fig. S5), providing additional evidence for an association between photochemical H₂SO₄ production and autumn NPF. These results suggest that the increased availability of sulfur-containing precursors may have partly compensated for potentially unfavorable effects associated with elevated NO_x and supported NPF occurrence (Gao et al., 2025). Although the proxy is not a direct measurement of H₂SO₄, these results support an important contribution from sulfur-containing precursors to autumn NPF rather than demonstrating their exclusive control.

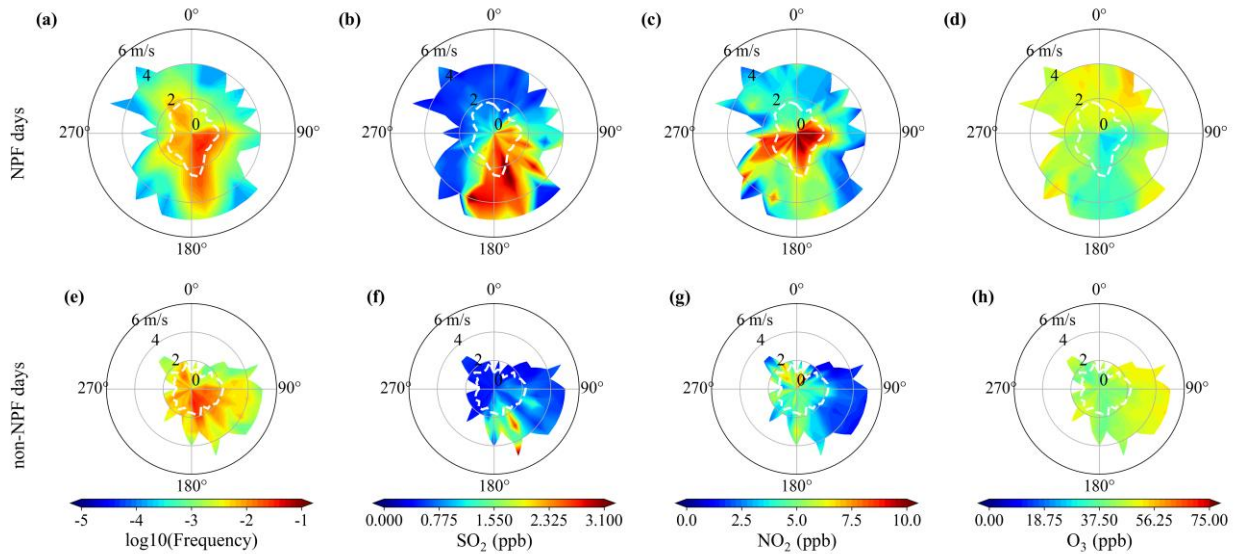


Figure 8: Wind-sector analysis comparing the distributions of (a) wind speed and wind direction frequency together with sector-averaged concentrations of (b) SO₂, (c) NO₂, and (d) O₃, for NPF days during the ~~autumn observation monsoon~~ period; panels (e), (f), (g), and (h) show the corresponding wind-sector frequency and gas concentrations for non-NPF days during the ~~autumn observation monsoon~~ period.

During the ~~spring observation pre monsoon~~ period, no single, well-defined factor controlling NPF occurrence was evident. As shown in As shown in Figs. 9 and S6, SO₂, NO₂, and O₃ exhibited differences between NPF and non-NPF days during certain hours and under specific wind conditions. However, no consistent separation between the two categories was observed

across all gaseous species or throughout the particle-formation period. Wind speed and wind direction likewise did not show a distinctive pattern that consistently characterized NPF days. These results suggest that spring NPF was more likely associated with the combined effects of multiple processes, including precursor availability, atmospheric oxidation, and boundary-layer evolution, rather than with any single dominant factor (Figs. 9 and S6, although SO₂, NO₂, and O₃ differ to some extent between NPF and non-NPF days, the overall separation remains limited, while wind speed and wind direction also fail to show a consistent or distinctive signal. These results separation between the two categories was observed across all gaseous species or throughout the particle formation period. Wind speed and wind direction likewise did not show a distinctive pattern that consistently characterized NPF. These results suggest that spring NPF was more likely associated with the combined effects of several processes, including precursor availability, the spring observation period pre-monsoon NPF is more likely governed by the combined effects of multiple factors, including precursor supply, atmospheric oxidation, and boundary layer evolution, rather than by any single dominant factor (Wu et al., 2024). Given the relatively weak and inconsistent contrasts in the measured inorganic gaseous precursors, a contribution from VOC-related oxidation processes cannot be excluded, particularly for the subsequent growth of newly formed particles the possible contribution of VOC related oxidation processes to the initial growth of newly formed particles deserves attention (Zhang et al., 2026; Yang et al., 2026). However, because direct observations of VOCs and their oxidation products are not available in this study, their specific role cannot be further constrained and still requires additional investigation.

Overall, the comparison between the two observation periods indicates differences in the factors associated with NPF, although the relatively short measurement periods preclude establishing a general seasonal pattern. During the autumn observation period monsoon season, NPF was associated with higher SO₂ concentrations and an enhanced relative H₂SO₄-related proxy during the particle-formation period, suggesting an important contribution from sulfur-containing precursors rather than demonstrating a uniquely sulfur-precursor-driven regime. Elevated NO₂ may have modified atmospheric radical chemistry, but its net effect on NPF could not be determined from the available measurements. In contrast, during the spring observation period pre-monsoon season, no single dominant factor was identified. Instead, precursor availability, atmospheric oxidation, boundary-layer evolution, and potentially VOC-related oxidation may have jointly affected particle formation and growth. Because H₂SO₄, ammonia, amines, VOCs, and their oxidation products were not directly measured, their respective contributions could not be quantified, and the detailed molecular mechanisms during both observation periods remain uncertain.

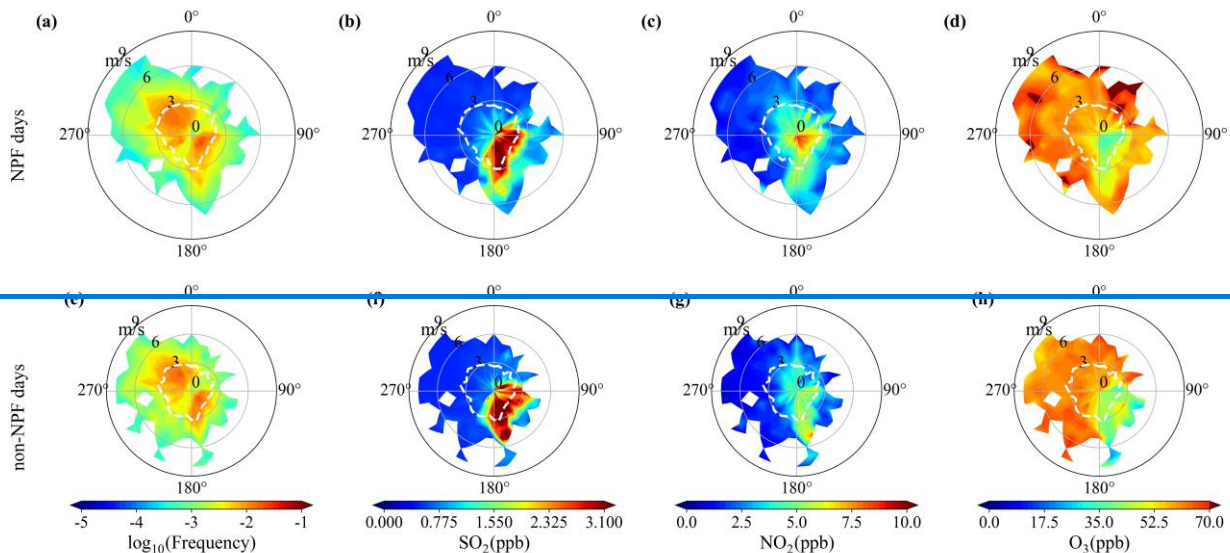
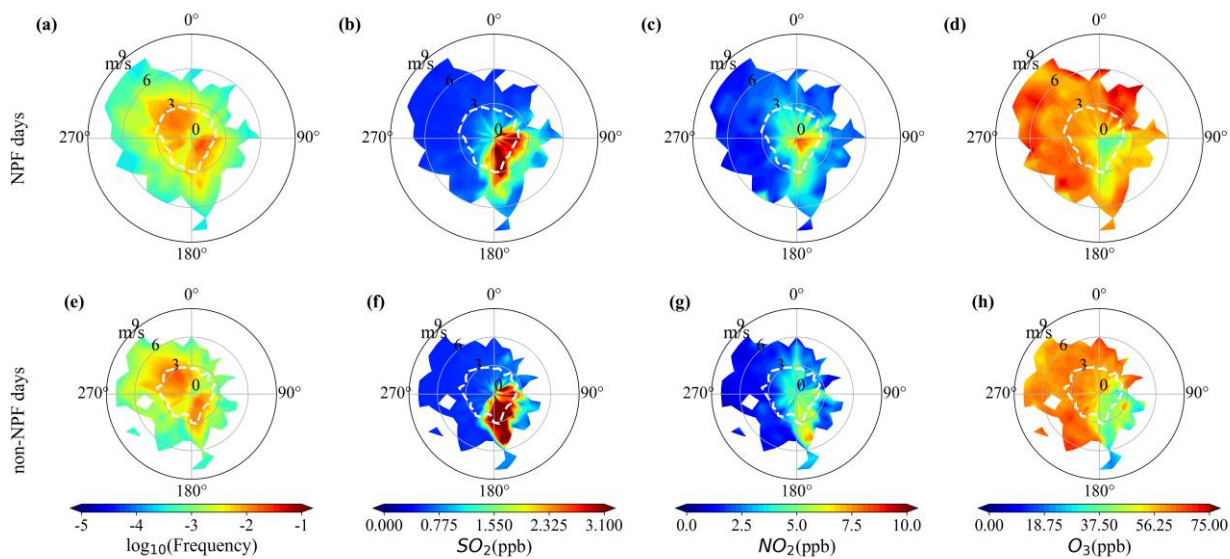


Figure 9: Wind-sector analysis comparing the distributions of (a) wind speed and wind direction frequency together with sector-averaged concentrations of (b) SO₂, (c) NO₂, and (d) O₃, for NPF days during the spring observationpre-monsoon period; panels (e), (f), (g), and (h) show the corresponding wind-sector frequency and gas concentrations for non-NPF days during the spring observationpre-monsoon period.

3.4 Modulation of CCN by NPF in urban air

To quantify the influence of NPF on CCN in the urban atmosphere of the Tibetan Plateau, we examined both the diurnal evolution and the distribution of estimated CCN for different event categories as well as for the two seasonal-periods

(~~monsoon~~spring and ~~pre-monsoon~~autumn). Because direct measurements of CCN concentration were unavailable in this study, CCN was estimated from the measured PNSD using the κ -Köhler framework described in Sect. 2.3, assuming a constant particle hygroscopicity parameter κ of 0.12 and a supersaturation S of 1.2 %. Because the estimated CCN concentrations depend on these assumptions, a sensitivity analysis was conducted using κ values of 0.08, 0.12, and 0.20 and supersaturations of 0.6 %, 0.8 %, and 1.2 % (Fig. S7). Although the absolute CCN concentrations varied with κ and S , estimated CCN remained consistently higher during the autumn observation period than during the spring observation period across the tested parameter combinations.
The estimated CCN concentrations were higher than $N_{100-700}$, because the assumed κ and supersaturation corresponded to a critical activation diameter below 100 nm, allowing part of the Aitken-mode particle population to contribute to the estimated CCN.

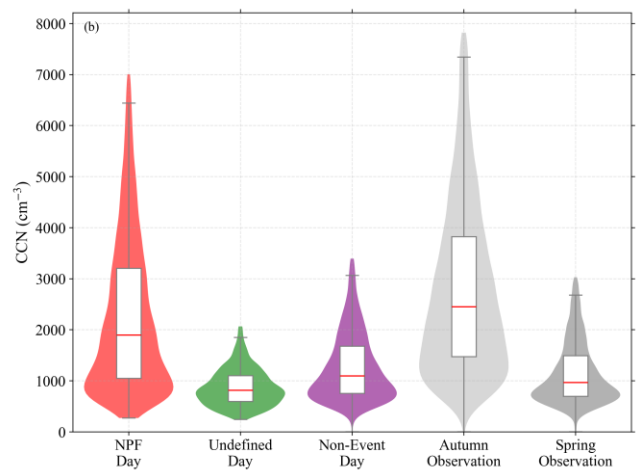
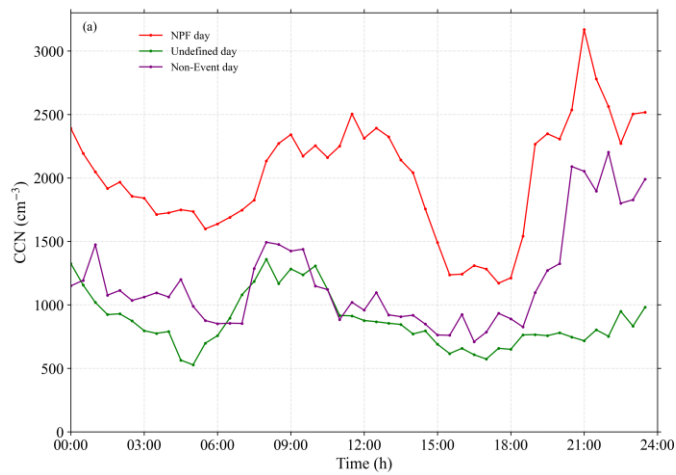
Figure 10a shows clear differences in the diurnal variation of estimated CCN concentrations among NPF, Undefined, and Non-Event days. NPF days generally exhibited the highest CCN concentrations, with pronounced daytime variations and a marked enhancement during the late afternoon and evening. This late-day enhancement is consistent with the continued growth and survival of newly formed particles into CCN-relevant sizes (Shen et al., 2019). Undefined days showed the lowest CCN concentrations and relatively weak diurnal variability. Non-Event days generally exhibited intermediate CCN concentrations, including an evening increase that may reflect the influence of pre-existing particles and local urban emissions rather than daytime NPF. The distributions in Fig. 10b likewise show that estimated CCN concentrations were generally higher and more broadly distributed on NPF days than on Undefined and Non-Event days. To further examine the role of particle growth, NPF days were divided into high- and low- N_{25-100} groups using the observation-period-specific median of daily mean N_{25-100} during 09:00–18:00. The high- N_{25-100} group exhibited higher estimated CCN concentrations, particularly in the afternoon and evening (Fig. S8), supporting the importance of continued particle growth and survival for potential CCN production.

The distributions during the two observation periods show substantially higher estimated CCN concentrations during the autumn observation period than during the spring observation period (Fig. 10b), consistent with the higher NPF frequency reported in Sect. 3.1. The ~~autumn observation period~~monsoon median CCN ~~is was~~ 2448.9 (1471.5–3820.6) cm^{-3} with a mean of $2813.0 \pm 1665.2 \text{ cm}^{-3}$, whereas the ~~spring observation period~~pre-monsoon median ~~is was~~ 964.7 (696.4–1489.6) cm^{-3} with a mean of $1152.1 \pm 608.9 \text{ cm}^{-3}$. These observation-period differences suggest that the more frequent occurrence of NPF, together with subsequent particle growth during the autumn measurements, may have contributed to the higher estimated CCN-relevant particle concentrations at the TU site.

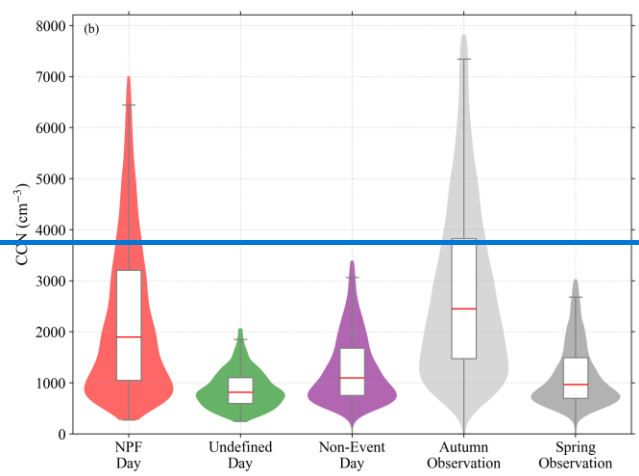
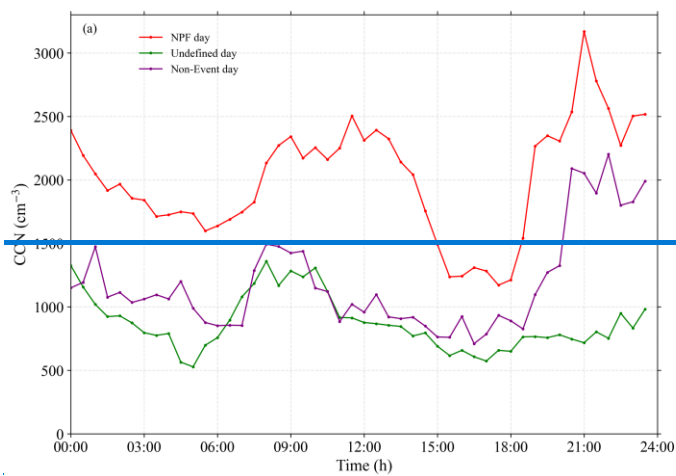
Overall, our results demonstrate that NPF is an effective source of CCN in the TP urban atmosphere, but its impact is strongly modulated by event type and observation periodseason. The pronounced late-day CCN enhancement and the broad, high-CCN distribution on Class I days highlight the importance of sustained post-formation growth and particle survival in determining the strength of the CCN response. At the ~~observation period~~seasonal scale, the markedly elevated CCN levels during the ~~autumn observation period~~monsoon indicate that frequent NPF events, together with favorable growth conditions, jointly sustain a high CCN background. This implies that secondary aerosol formation may play a more important role in aerosol–cloud interactions over high-altitude urban regions. Overall, our results indicate that NPF days were associated with higher

521 estimated CCN concentrations in the TP urban atmosphere, although the magnitude of the estimates depended on the assumed
522 κ and supersaturation. The pronounced late-day enhancement on NPF days, particularly in the high- N_{25-100} group, highlights
523 the importance of sustained post-formation growth and particle survival in determining the potential CCN response. At the
524 observation-period scale, the higher estimated CCN concentrations during the autumn observation period may reflect the
525 combined influence of frequent NPF and favorable particle-growth conditions. Because CCN concentrations were estimated
526 rather than directly measured, these results indicate potential CCN production rather than direct confirmation of CCN
527 activation.

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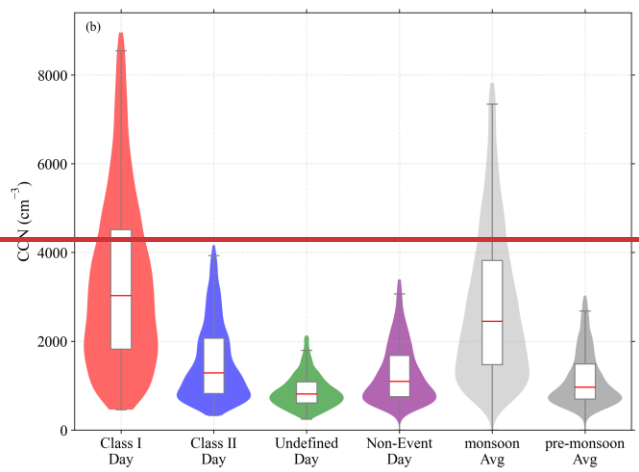
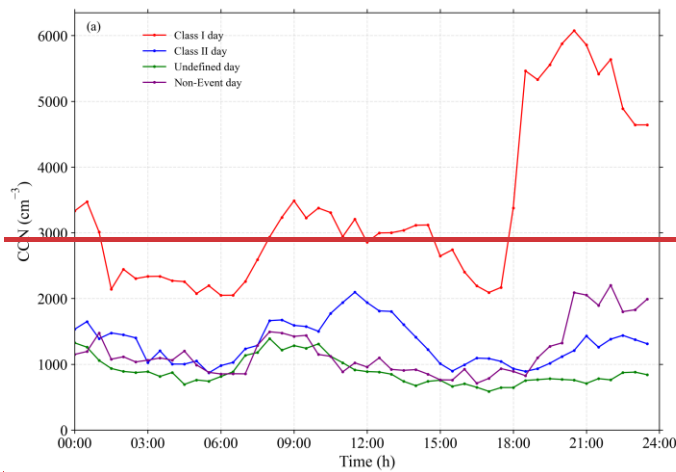


Figure 10: Diurnal variation (a) and statistical distribution (b) of CCN concentration during different days, classified by NPF events. (a) Diurnal variations and (b) statistical distributions of estimated CCN concentrations for NPF, Undefined, and Non-Event days and for the autumn and spring observation periods. CCN concentrations were estimated using $\kappa = 0.12$ and $S = 1.2\%$.

4 Conclusion

This study provides a comprehensive characterization of NPF in Lhasa, a high-altitude urban center on the Tibetan Plateau, with particular emphasis on the controlling factors of NPF and its implications for CCN during the ~~pre-monsoon~~spring and autumn observation periods~~monsoon seasons~~. Frequent NPF was observed throughout the campaigns. The ~~contrast between the two observation periods~~ in NPF occurrence was pronounced, with event frequencies of 86.2 % ~~during in~~ the autumn observation period~~monsoon~~ and 55.4~~62.1~~ % ~~during the spring observation period~~pre-monsoon, indicating that the NPF regime in Lhasa is highly sensitive to ~~variations in meteorology and atmospheric chemistry between the observation periods~~seasonally ~~varying meteorology and atmospheric chemistry~~. The results indicate that differences in the pre-existing particle sink alone cannot explain the contrast in NPF frequency between the two observation periods, and that precursor availability and atmospheric oxidation likely also played important roles. The CS remained comparable between the two observation periods~~seasons~~, suggesting that the ~~observed contrast change~~ in NPF frequency cannot be explained by sink suppression alone. Instead, the chemical–meteorological context differs substantially across ~~observation periods~~seasons. During the autumn observation period, NPF days were associated with elevated SO_2 and a higher relative H_2SO_4 -related proxy during the particle-formation period, suggesting that sulfur-containing precursors likely contributed to particle formation and early growth. ~~During change the autumn observation periodmonsoon, NPF days were associated with elevated SO_2 , consistent with intensified sulfur related photochemical production of condensable vapors under abundant radiation, pointing to a regime where sulfuric acid driven nucleation and early growth are favored.~~ In contrast, the spring observation period~~pre-monsoon~~ NPF did not show a single, well-defined controlling factor, but was more likely influenced by the combined effects of precursor supply, atmospheric oxidation, and boundary-layer evolution. Under such conditions, VOC-related oxidation may contribute to the early growth of newly formed particles, although its specific role remains to be further constrained. A key outcome of this work is that NPF days were associated with higher estimated CCN concentrations in Lhasa, and that the potential CCN response depended critically on whether newly formed particles continued to grow and survive after formation. Estimated CCN concentrations were generally higher on NPF days than on Undefined and Non-Event days. Moreover, NPF days with higher N_{25-100} exhibited greater afternoon and evening CCN enhancements than those with lower N_{25-100} , supporting the importance of continued particle growth and survival in determining whether newly formed particles reach CCN-relevant sizes. The mean estimated CCN concentration during the autumn observation period was more than twice that during the spring observation period, suggesting that frequent NPF together with favorable particle-growth conditions may have contributed to higher CCN-relevant particle concentrations during the autumn measurements. These findings

improve our understanding of the potential contribution of urban NPF to CCN-relevant particle populations in high-altitude environments over the Tibetan Plateau.

A key outcome of this work is that NPF substantially enhances CCN in Lhasa, and the magnitude of the CCN response is governed by event strength and, critically, by whether newly formed particles can sustain growth after nucleation to reach CCN activation diameters. CCN concentrations followed a clear hierarchy of Class I > Class II > non-event > undefined, indicating that strong NPF events can maintain a markedly elevated CCN population relative to background conditions. The diurnal evolution further shows that the CCN enhancement is most pronounced from late afternoon to evening on Class I days, consistent with a persistent “banana-shaped” growth pattern and efficient survival of freshly formed particles into the size range where they can be activated as CCN. During the two observation periods, On a seasonal basis, mean CCN during the autumn observation periodmonsoon exceeded that during the spring observation periodpre-monsoon by more than a factor of two, implying that frequent NPF combined with favorable growth conditions can raise the CCN background level in this high-altitude urban region and potentially influence aerosol–cloud interactions over the TP. Deepening our understanding of NPF over the Tibetan Plateau is essential for more accurately assessing the impacts of aerosols in the TP on the global atmospheric environment and climate, and it can provide a solid scientific basis for the formulation of relevant policies.

Despite the insights gained in this study, several limitations should be acknowledged. First, the day classification was based primarily on the daily PNSD evolution and modal development, supported by particle number concentrations in the 3 – 25 and 25 – 100 nm ranges and particle growth rates. Although these supporting metrics reduced reliance on visual inspection alone, some classification uncertainty remained for days with weak, discontinuous, or ambiguous particle evolution; these days were therefore retained as Undefined days and included in the non-NPF group for comparisons between NPF and non-NPF conditions. Second, CCN concentrations were not directly measured but estimated from the measured PNSD using the κ -Köhler framework, which may introduce uncertainties associated with the assumed hygroscopicity parameter and supersaturation. Therefore, the estimated CCN values are more suitable for interpreting relative differences and response patterns than for over-interpreting their absolute magnitudes. In addition, due to the lack of direct observations of VOCs and their oxidation products, the mechanisms of NPF, especially during the spring observationpre-monsoon period, could not be further constrained. Future studies should combine longer-term continuous observations with direct CCN measurements and additional observations of VOCs, H₂SO₄, and related oxidation products, in order to better identify the controlling factors of NPF and its contribution to CCN in high-altitude urban environments over the Tibetan Plateau.

Code and data availability

The elevation data used in this study are openly available from the Geospatial Data Cloud at <https://www.gscloud.cn>.

The raw observational datasets used in this study have been deposited in Zenodo and can be freely downloaded from <https://doi.org/10.5281/zenodo.19876491>.

594 The codes used for data processing and figure generation have been deposited in a separate Zenodo repository and can be
595 freely downloaded from <https://doi.org/10.5281/zenodo.19879177>.

596 **Author contributions**

597 QM and LW wrote the manuscript. QM, LW, GZ, CY, JS and TZ discussed the original idea and results. JS, FS, TL, YG, and
598 YC conducted the field measurement. QM, LW and JS processed and analysed the data.

599 **Competing interests**

600 The authors declare that they have no conflict of interest.

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