



# Seasonal variation in aerosol chemistry drives new particle formation and CCN activity in a coastal city, China: insights from year-long online measurements in Fuzhou

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**Abstract.** New particle formation (NPF) is an important source of cloud condensation nuclei (CCN), which affects the global climate. Continuous observations in the coastal city of Fuzhou, conducted from June 2021 to May 2022, aimed to study NPF events and their impact on CCN. A total of 46 NPF events were identified, with a frequency of 12.7 %. The average formation rate (FR) and growth rate (GR) of particles were  $3.94 \pm 8.26 \text{ cm}^{-3} \text{ s}^{-1}$  and  $5.20 \pm 1.78 \text{ nm h}^{-1}$ . The NPF events showed evident seasonal variation: spring (27.17 %), fall (9.89 %), winter (8.89 %), and summer (4.35 %). Spring NPF events were characterized by high FR ( $5.56 \text{ cm}^{-3} \text{ s}^{-1}$ ) and suppressed growth processes, while summer, under the dominance of marine winds, exhibited the lowest FR yet the highest GR among all seasons (peak value  $11.68 \text{ nm h}^{-1}$ ). The influence of NPF on the chemical composition of  $\text{PM}_{2.5}$  and CCN also showed seasonal differences. In summer, NPF generated substantial amounts of sulfate and nitrate, resulting in stronger particle hygroscopicity ( $> 0.6$ ). In fall and winter, higher concentrations of black carbon (BC) and primary organic carbon (POC) led to weaker  $\kappa_{\text{inorg}}$  ( $\leq 0.55$ ). XGBoost–SHAP attribution further quantified that FR is dominated by physical processes (nucleation mode 76.2 %, CS 13.8 %), with a sharp  $\text{NH}_3$  threshold at  $4 \mu\text{g m}^{-3}$  and a narrow temperature range (20–25 °C). For particle growth, temperature shows a positive linear effect above 20 °C, and  $\text{RH} > 60 \%$  consistently suppresses particle concentrations. The enhancement effect of NPF on CCN was most significant in summer ( $E_{\text{NCCN}} = 1.64$ ), accompanied by CCN growth. In spring, the high condensation sink (CS) suppressed growth, leading to an insignificant CCN enhancement effect. In fall and winter, NPF-induced CCN enhancement mainly occurred 3–5 h after the event, with increases ranging from 13 % to 65 %, particularly notable at high supersaturation levels (0.8 %–1.0 % SS).

## 1 Introduction

New particle formation (NPF) is a complex process in which gaseous precursors in the atmosphere nucleate and condense to form new particles, which subsequently grow through condensation, coagulation, and other processes. NPF contributes over 50 % of the global cloud condensation nuclei (CCN), significantly influencing cloud albedo, structure, lifetime, and solar radiation reaching the Earth's surface (Tröstl et al., 2016; Yao et al., 2018). Additionally, efficient nucleation and explosive growth of particles are important sources of haze formation in urban atmospheres, impacting air quality and public health (Kulmala et al., 2021).

Although its frequency may vary with season and location, NPF events fundamentally represent competition between aerosol particle sources and sinks. Current research indicates that secondary particle formation is driven by the photochemical oxidation of atmospheric gases. Sulfuric acid and highly oxidized molecules can act as nucleation precursors (Fan et al., 2018; Zaveri et al., 2022). Furthermore, ions may also play a role in particle nucleation, though their significance remains debated (Hirsikko et al., 2011; Kirkby et al., 2016). Pre-existing aerosol particles act as a sink for these precursors, small clusters, and newly formed particles, thereby suppressing NPF occurrence (McMurry and Friedlander, 1979). However, frequent NPF events also occur in heavily polluted cities (Sun et al., 2015; Yao et al., 2018). Therefore, the mechanisms governing NPF generation and growth under different atmospheric conditions are still under investigation.

NPF events can be described by the formation rate (FR) of nucleation-mode particles and the growth rate (GR) of newly formed particles (Kulmala et al., 2012). The formation rate of 3 nm particles in the boundary layer typically ranges from 0.01 to 10 cm<sup>-3</sup> s<sup>-1</sup>, while the typical growth rate in mid-latitudes ranges from 1 to 20 nm h<sup>-1</sup> (Kulmala et al., 2004). Yli-Juuti et al. (2011) reported typical growth rates of 1.8–10.7 nm h<sup>-1</sup> for 1.5–20 nm particles. Previous studies have shown that NPF is enhanced in the presence of sulfuric acid, alkaline substances, organic acids, and ions (Wang et al., 2011) but suppressed in the presence of nitrogen oxides (NO<sub>x</sub>) (Wildt et al., 2014), indicating significant synergistic effects in chemically complex mixtures (Guo et al., 2014). The availability of precursor vapors and the atmospheric chemical environment play decisive roles. H<sub>2</sub>SO<sub>4</sub> is a key nucleating species, and its stabilizing co-components, such as ammonia (NH<sub>3</sub>) and amines, can dramatically enhance FR (Dunne et al., 2016; Yao et al., 2018; Kirkby et al., 2016). For particle growth, condensation of low-volatility compounds is the dominant process. Sipilä et al. (2010) showed experimentally that early-stage growth is primarily driven by H<sub>2</sub>SO<sub>4</sub> condensation.

Additionally, NPF occurrence is constrained by atmospheric temperature and humidity (Yu et al., 2017; Yue and Hamill, 1979). Low temperatures promote nucleation,

whereas high temperatures suppress it (Sipilä et al., 2010; Dunne et al., 2016; Yu et al., 2017). Dunne et al. (2016) further showed that at low temperatures, the ion-enhancement effect is weak due to suppressed evaporation of neutral clusters, while at ambient temperatures, ions can increase the nucleation rate by about a factor of 15. Consequently, neglecting temperature dependence leads to a marked overestimation of NPF and CCN concentrations in summer (Yu et al., 2017). Hamed et al. (2011), based on observations at multiple continental sites, found that NPF events predominantly occur at relative humidity below 60 % and are rare above 80 %. The reason is that high relative humidity reduces ultraviolet radiation, lowering the production of OH and H<sub>2</sub>SO<sub>4</sub>; meanwhile, hygroscopic growth enhances the condensation sink, thereby suppressing new particle formation.

Aerosol chemical composition also influences aerosol hygroscopicity, altering its critical diameter and thereby affecting CCN activation and cloud formation (Petters and Kreidenweis, 2007; Williamson et al., 2019; Xu et al., 2020). Petters and Kreidenweis (2007) proposed hygroscopicity parameter ( $\kappa$ ) is widely used to quantify the water uptake ability and CCN activity of aerosols. Extensive experimental studies have well characterized the  $\kappa$  values of inorganic salts such as ammonium sulfate, ammonium nitrate, and ammonium bisulfate (Liu et al., 2014; Kuang et al., 2020; Cai et al., 2018; Wu et al., 2016). In the accumulation mode (150–1  $\mu$ m), inorganic species contribute about 90 % of the total  $\kappa$  (Liu et al., 2014).

Observations across different environments (e.g., mountain, urban) have shown that NPF events typically lead to a significant increase in  $N_{\text{CCN}}$  (Kuwata et al., 2008; Yue et al., 2011; Fan et al., 2018). Kuwata et al. (2008) observed a clear increase in  $N_{\text{CCN}}$  at different supersaturation levels after NPF events on Jeju Island, South Korea. Research in Beijing indicated that NPF events could increase local  $N_{\text{CCN}}$  by 0.4–6 times (Yue et al., 2011). In polluted environments, additional condensable species, particularly ammonium nitrate and secondary organic aerosols, can accelerate growth and shorten the time needed for freshly formed particles to reach CCN-active sizes (Zhu et al., 2026). However, some studies suggest that an increase in hydrophobic organic components during subsequent particle growth may inhibit CCN generation. Therefore, understanding the role of different components during particle growth is crucial for assessing their subsequent climate effects.

Although NPF research in China is widespread, most studies focus on reporting occurrence frequencies, formation, and growth rates, or are limited to discussing nucleation mechanisms. For instance, NPF event frequencies at sites like Shangdianzi, Mount Tai, and Lin'an in eastern China range from 15 % to 29 % (Shen et al., 2018). Frequencies in Beijing, Jinan, and Shanghai are approximately 30 %, 40 %, and 21 %, respectively (Jayaratne et al., 2017; Lv et al., 2018; Xiao et al., 2015). Currently, understanding how regional variations in atmospheric oxidants and precursors affect the

growth of newly formed particles to CCN sizes, especially the quantification of their CCN efficiency, remains a challenge and frontier in current research (Cai et al., 2018; Kulmala et al., 2021; Rose et al., 2017; Xiao et al., 2015; Yao et al., 2018). Given that Fuzhou is a rapidly developing southeastern coastal city with unique sea-land breeze conditions, a high-temperature and high-humidity environment, and complex pollution emission characteristics, it may have unique NPF mechanisms. However, there is a lack of long-term comprehensive observation or systematically quantified NPF's CCN efficiency in this region. Therefore, this study conducted a one-year comprehensive observation in Fuzhou from June 2021 to May 2022, providing new insights and data support for understanding the CCN efficiency and potential climate relevance of NPF under China's complex atmospheric environment.

## 2 Data and Methods

### 2.1 Observation site

Observation data for this study were collected from 1 June 2021 to 30 May 2022, during comprehensive atmospheric environmental observations conducted at the Fujian Provincial Environmental Monitoring Center Station (26.11° N, 119.30° E, altitude 65 m) and the Fuzhou Meteorological Bureau Station (26.05° N, 119.26° E, altitude 18 m). Both stations are located within Fuzhou's urban area, approximately 8 km apart horizontally. The Fujian Provincial Environmental Monitoring Center Station is situated in Gulou District, the central urban area of Fuzhou, surrounded primarily by commercial, residential, and transportation land, representing areas heavily influenced by intense human activities. The Fuzhou Meteorological Bureau Station is located in Cangshan District, southern Fuzhou, approximately 1.5 km east of the Min River. Fuzhou is situated at the Min River estuary and along the East China Sea coast, characterized mainly by plains (average altitude 10–30 m) and a typical East Asian monsoon climate, significantly influenced by sea-land breeze circulation and marine air masses (Hu et al., 2024). Given the short distance (~ 8 km) and the regional background homogeneity, the local environments of the two sites are not expected to be drastically different, although each is influenced by its specific surroundings (urban vs. riverside). A comparison of air pollutant concentrations between the two sites using nearby national monitoring stations (Table S2 in the Supplement).

### 2.2 Measurement and instrumentation

A CCN counter (CCN-100; DMT, USA) equipped with a continuous flow of  $500 \text{ cm}^3 \text{ min}^{-1}$  and a thermal gradient was used to measure CCN concentrations at five SS levels. To maintain counting accuracy, the instrument was regularly calibrated for  $T$  gradient, flow rate, pressure, SS, and the op-

tical particle counter (OPC) using standard ammonium sulfate according to the method by Rose et al. (2008). Additionally, zero-point determination was performed before and after each observation to minimize instrumental error. During observations, the measurement interval for each SS level was 10 min, and a few minutes were required to stabilize after switching SS levels. Therefore, CCN data collected before reaching stable SS were excluded from subsequent analysis. The typical measurement uncertainty of the CCN-100 is approximately  $\pm 10 \%$  in Table S1.

An online organic carbon/elemental carbon analyzer (Sunset Laboratory semicontinuous OC/EC analyzer, Model-4, Sunset Laboratory Inc., USA) was used to determine organic carbon (OC) and elemental carbon (EC) content in atmospheric particulate matter samples. The instrument provides hourly averaged concentrations. Details of instrument operation can be found in Chang et al. (2017). Based on replicate analyses reported by Zhang et al. (2021), the within-model measurement uncertainties (relative standard deviation) for the Sunset Model-4 are  $\pm 3.6 \%$  for OC and  $\pm 6.8 \%$  for EC.

A Wide-Range Particle Spectrometer (WPS-1000, MSP) measured aerosol number size distributions (10–350 nm) with a time resolution of 6 min across 96 channels. To align with the 1 h meteorological data, the 6 min distributions were arithmetically averaged to hourly particle number size distributions. Instrument details and principles are described in Wang et al. (2014). The typical uncertainty for particle size distribution measurements in this size range is approximately  $\pm 10 \%$  in Table S1.

Black carbon (BC) mass concentration was measured using an Aethalometer (AE-33, Magee Scientific) with a time resolution of 1 h. BC data from the 880 nm wavelength (channel 6) were used (Kirchstetter et al., 2004). The measurement uncertainty is approximately  $\pm 10 \%$  (Table S1).

An online particle chromatograph (MARGA ADI-2080) continuously monitored mass concentrations of soluble aerosol ionic components ( $\text{SO}_4^{2-}$ ,  $\text{NO}_3^-$ ,  $\text{NH}_4^+$ ,  $\text{Na}^+$ ,  $\text{K}^+$ ,  $\text{Ca}^{2+}$ ,  $\text{Cl}^-$ ) and trace gases ( $\text{NH}_3$ ,  $\text{HNO}_2$ ,  $\text{HNO}_3$ ,  $\text{HCl}$ ,  $\text{SO}_2$ ). Sampling, operation, and internal calibration methods followed Du et al. (2010). The instrument provides hourly averaged concentrations. According to an independent verification study (Battelle, 2009), the measurement precision (median absolute relative percent difference between duplicate units) ranges from 5 % for  $\text{SO}_2$  to 20 % for  $\text{NH}_4^+$ , with data completeness > 90 % for all major ions.

Meteorological data (including wind speed (WS), wind direction (WD), temperature ( $T$ ), relative humidity (RH), and precipitation) with a time resolution of 1 h were obtained from the Fuzhou Olympic Sports Center Meteorological Station. Data on conventional air pollutants ( $\text{O}_3$ , CO,  $\text{NO}_2$ ,  $\text{PM}_{2.5}$ , and  $\text{PM}_{10}$ ) were sourced from the China National Environmental Monitoring Centre's real-time urban air quality release platform (<https://quotsoft.net/air/>, last access: 26 May 2026).

### 2.3 Analysis methods

The growth rate (GR) of new particles was calculated following (Kulmala et al., 2012):

$$\text{GR} = \frac{\Delta D_m}{\Delta t} \quad (1)$$

where  $D_m$  is the median diameter of the nucleation mode particles, obtained by fitting a log-normal distribution to the particle number size distribution. Statistics of the fitting results (see the Supplement) demonstrate that the log-normal distribution represents the observed particle size distributions well, with the majority of fits yielding high coefficients of determination ( $R^2 > 0.85$ ); only fits with  $R^2 > 0.7$  were used to avoid propagating poor-fitting uncertainties. GR describes how rapidly particles grow from the nucleation size to larger sizes. It is later used to estimate the formation rate of new particles (Eq. 4) and condensable vapor concentration (Eq. 5) and to evaluate the competition between condensation and coagulation during NPF events.

The condensation sink (CS) reflects the rate at which condensable vapor molecules condense onto the surface of pre-existing atmospheric particles and was calculated as follows (Kulmala et al., 2012):

$$\text{CS} = 4\pi D \sum_i \beta_{M,i} \cdot D_i \cdot N_i \quad (2)$$

where  $D$  is the diffusion coefficient of the vapor (typically assumed to be sulfuric acid),  $N_i$  is the number concentration of particles in a given size bin, and  $\beta_M$  is a correction factor.

The coagulation sink (CoagS) reflects the ability and rate of pre-existing atmospheric particles to remove newly formed nucleation particles via coagulation. For particles of size  $i$ , the coagulation sink can be expressed as:

$$\text{CoagS}_i = \sum_j K_{ij} N_j \quad (3)$$

where  $N_j$  is the number concentration of particles in size bin  $j$ , and  $K_{ij}$  is the Brownian coagulation coefficient between particles of size  $j$  and  $i$ .

The formation rate (FR) of new particles was calculated following Kulmala et al. (2012):

$$\text{FR} = \frac{dN_{\text{nuc}}}{dt} + \text{CoagS}_{\text{nuc}} \cdot N_{\text{nuc}} + \frac{\text{GR}}{\Delta dp} \cdot N_{\text{nuc}} + S_{\text{losses}} \quad (4)$$

where  $N_{\text{nuc}}$  is the number concentration of nucleation-mode particles. Following the definition by Kulmala et al. (2012), the nucleation-mode size range in this study was also limited to below 25 nm.  $\text{CoagS}_{\text{nuc}} \cdot N_{\text{nuc}}$  is the flux of particles lost due to coagulation with pre-existing particles, where  $\text{CoagS}_{\text{nuc}}$  is the coagulation sink for nucleation-mode particles.  $\text{GR}/\Delta dp$  represents the flux of particles growing out of the nucleation size range (exceeding 25 nm), which is generally negligible under typical atmospheric conditions (Dal

Maso et al., 2005). FR is later used to compare NPF event intensity under different meteorological and chemical conditions, and to identify periods with active nucleation. The additional loss term  $S_{\text{losses}}$  (e.g., dilution due to boundary layer growth, wall losses) was negligible under our field conditions (Dal Maso et al., 2005). For regional NPF events, transport losses can also be ignored. FR is a direct measure of NPF intensity and is later compared across different meteorological and chemical conditions.

Condensable vapor concentration ( $C$ ) and source rate ( $Q$ ). Assuming that particle growth is dominated by condensation of a low-volatility vapor (typically sulfuric acid), the vapor concentration can be estimated from the observed growth rate (Dal Maso et al., 2005; Kulmala et al., 2012):

$$C = A \cdot \frac{ddp}{dt} \quad (5)$$

where  $dp$  is the particle diameter, and  $A$  is a constant, which has the value  $1.37 \times 10^{-7} \text{ h cm}^{-3} \text{ nm}^{-1}$  for a vapor with molecular properties of sulfuric acid (Dal Maso et al., 2005). This provides an upper-limit estimate of the condensable vapor concentration, as it assumes growth is solely due to condensation of the vapor and neglects contributions from coagulation.

The CS (Eq. 2) quantifies the rate at which this vapor is removed by pre-existing particles. Under steady-state conditions ( $dC/dt = 0$ ), the vapor source rate  $Q$  can be derived as (Dal Maso et al., 2005):

$$Q = \text{CS} \cdot C \quad (6)$$

This source rate represents the net production of condensable vapor needed to maintain the observed growth and is later compared with precursor gas concentrations (e.g.,  $\text{SO}_2$ ) to infer the chemical pathways driving NPF.

Hygroscopicity parameter ( $\kappa_{\text{inorg}}$ ) for inorganic species. Due to the lack of organic composition measurements, we estimated the hygroscopicity of the inorganic fraction only. The measured water-soluble ions ( $\text{SO}_4^{2-}$ ,  $\text{NO}_3^-$ ,  $\text{NH}_4^+$ ,  $\text{Cl}^-$ ) were converted to mass concentrations of inorganic salts using the ion-pairing scheme described in Gysel et al. (2007) and Kuang et al. (2020). The following salts and their  $\kappa$  values (Kuang et al., 2020, Table S2) were considered:  $(\text{NH}_4)_2\text{SO}_4$  ( $\kappa = 0.48$ ),  $\text{NH}_4\text{NO}_3$  ( $\kappa = 0.58$ ),  $\text{NH}_4\text{HSO}_4$  ( $\kappa = 0.56$ ), and  $\text{NH}_4\text{Cl}$  ( $\kappa = 0.93$ ). The volume fraction of each salt was calculated using its density (also from Kuang et al., 2020). The overall inorganic hygroscopicity parameter  $\kappa_{\text{inorg}}$  was then obtained by volume-weighted mixing (Petters and Kreidenweis, 2007):

$$\kappa_{\text{inorg}} = \sum_i \varepsilon_i \kappa_i \quad (7)$$

where  $\kappa_i$  and  $\varepsilon_i$  represent the hygroscopicity parameter and volume fraction of component  $i$  in the mixture, respectively, and  $i$  denotes the number of components. This  $\kappa_{\text{inorg}}$  represents the hygroscopicity of the inorganic aerosol components

and is used as an upper limit estimate for the total particle hygroscopicity, as organic matter (typically less hygroscopic) was not included.

The concentrations of secondary organic carbon (SOC) and primary organic carbon (POC) were estimated following (Wu and Yu, 2016):

$$\text{POC} = (\text{OC}/\text{EC})_{\text{min}} \cdot \text{EC} \quad (8)$$

$$\text{SOC} = \text{OC}_{\text{total}} - (\text{OC}/\text{EC})_{\text{pri}} \cdot \text{EC} \quad (9)$$

where  $\text{OC}_{\text{total}}$  is the measured OC,  $(\text{OC}/\text{EC})_{\text{min}}$  is the minimum  $(\text{OC}/\text{EC})$  ratio during the observation period, POC is primary organic carbon, and SOC is secondary organic carbon.

The enhancement effect on cloud condensation nuclei number concentration ( $E_{N_{\text{CCN}}}$ ) was defined as the ratio of CCN number concentration after the NPF event to that before the event (Ren et al., 2021):

$$E_{N_{\text{CCN}}} = N_{\text{CCN, after}} / N_{\text{CCN, prior}} \quad (10)$$

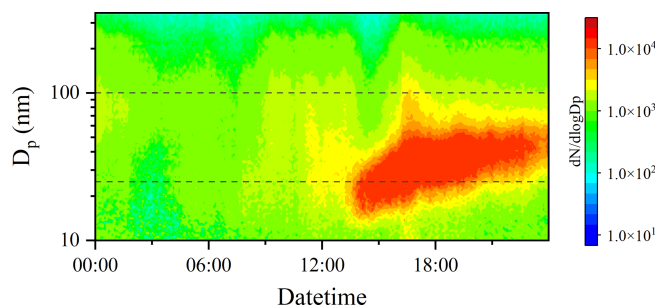
where  $N_{\text{CCN, after}}$  is the average CCN number concentration during the NPF event (from its start to end), and  $N_{\text{CCN, prior}}$  is the average CCN concentration during the 2 h before the event. This factor directly links NPF to potential cloud formation: a value  $> 1$  indicates that NPF increases CCN availability.

**XG Boost-SHAP framework.** To quantitatively evaluate the nonlinear effects of meteorological factors ( $T$ ,  $\text{RH}$ ) and precursor gases ( $\text{NH}_3$ ,  $\text{SO}_2$ ) on the particle formation rate (FR), we applied an interpretable machine learning framework combining XG Boost (Extreme Gradient Boosting) with SHAP (SHapley Additive exPlanations). A detailed description of the feature selection, model training, validation, and SHAP interpretation is provided in the Supplement (Sect. S1). The main quantitative thresholds and interaction strengths derived from this analysis are discussed in Sect. 3.4.

## 2.4 Identification of NPF events

NPF events were identified based on criteria from Kulmala et al. (2012): (1) significant increase in nucleation-mode number concentration ( $N_{\text{nuc}}$ ) (diameter 10–25 nm); (2) formation of a new mode lasting several hours; (3) growth of the newly formed mode over several hours. Additional criteria for NPF identification included: low pre-existing particle number concentration, a clear “banana-shaped” evolution in particle number concentration over time and size, and exclusion of interference from pre-existing particles (especially in urban environments) (Heintzenberg et al., 2007).

In this study, a day was defined as an effective NPF day if the nucleation-mode (10–25 nm) particle number concentration increased continuously for at least 2 h from its initial value to its maximum and showed clear growth to larger sizes (e.g., 12–50 nm) over several hours (Fig. 1). Other days were



**Figure 1.** An example NPF event observed on 21 April 2022.

defined as non-NPF days (Leng et al., 2014). During the one-year observation, a total of 46 NPF events and 319 non-NPF days were identified.

## 3 Results and Discussion

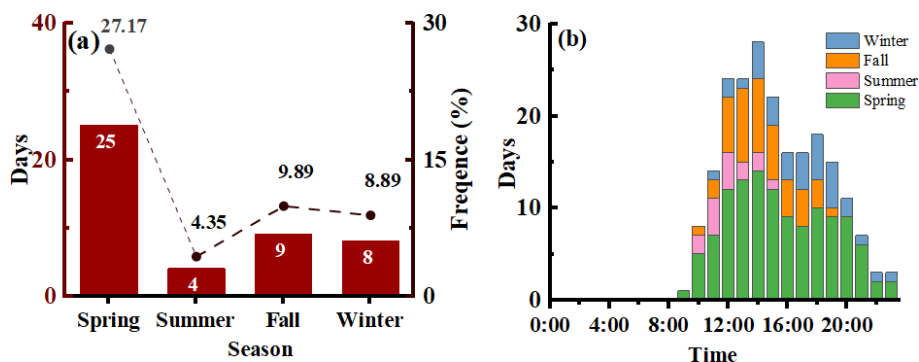
### 3.1 Overall characteristics of NPF occurrence

NPF events in Fuzhou exhibited a distinct seasonal preference. As shown in Fig. 2a, spring was the season with the highest NPF frequency (27.17 %), while summer had the lowest (4.35 %). Fall (9.89 %) and winter (8.89 %) showed intermediate to low frequencies. NPF events mainly occurred between 09:00 and 12:00 local time (LT).

NPF events led to significant increases in nucleation-mode ( $N_{\text{nuc}}$ ) and Aitken-mode ( $N_{\text{ait}}$ ) particle number concentrations (Fig. S1 in the Supplement). Spring showed the highest increase in  $N_{\text{nuc}}$  (196.7 %), while fall showed the highest increase in  $N_{\text{ait}}$  (70.5 %), indicating differences in new particle formation and subsequent growth across seasons.

NPF days typically corresponded to a lower background of  $\text{PM}_{2.5}$  and  $\text{PM}_{10}$  (Fig. S3), indicating that NPF occurred in relatively clean atmospheres with a weak condensation sink (CS). It should be noted that the absolute  $\text{PM}_{2.5}$  and  $\text{PM}_{10}$  in winter and spring were still relatively high compared to other seasons (Fig. S3), implying that even on NPF days, these seasons experienced higher pollution levels. In spring, summer, and winter, POC on NPF days were lower than on non-NPF days, corroborating this point. Meanwhile, in spring, summer, and fall, SOC was higher on NPF days, suggesting that secondary organic vapors may actively participate in particle formation and growth.

Although spring and winter had heavy background pollution, Fig. S4 shows that the gaseous precursor  $\text{SO}_2$  concentration on NPF days was significantly higher than on non-NPF days (winter:  $0.88 \mu\text{g m}^{-3}$  vs.  $0.76 \mu\text{g m}^{-3}$ ; spring:  $0.64 \mu\text{g m}^{-3}$  vs.  $0.53 \mu\text{g m}^{-3}$ ). This indicates that in polluted seasons, high gaseous precursors can overcome the inhibitory effect of a high CS and thus trigger nucleation. In contrast, NPF days in summer and Fall exhibited a distinctly clean background, with  $\text{NH}_3$  and  $\text{HNO}_2$  significantly lower than on non-NPF days (Fig. S4). In summer, fall,



**Figure 2.** The occurrence frequency and diurnal distribution of NPF events.

and winter, the concentrations of sulfate ( $\text{SO}_4^{2-}$ ), nitrate ( $\text{NO}_3^-$ ), and ammonium ( $\text{NH}_4^+$ ) on NPF days were significantly lower than on non-NPF days (Fig. S5). This suggests that in these seasons, the lower pre-existing particles and the reduced CS lessen the consumption of gaseous precursors, thereby favouring new particle formation and subsequent growth. However, spring presents a special case in that the SNA (sulfate, nitrate, and ammonium) concentrations on NPF days were comparable to those on non-NPF days ( $\text{SO}_4^{2-}$ :  $4.35 \mu\text{g m}^{-3}$  vs.  $4.27 \mu\text{g m}^{-3}$ ). This indicates that spring NPF events are driven by high precursor concentrations. Even when the background particle level is high, the abundant supply of gaseous precursors (Fig. S4) can still overcome the inhibition and trigger NPF events.

According to the hygroscopicity parameter ( $\kappa_{\text{inorg}}$ ) shown in Fig. S6, the  $\kappa_{\text{inorg}}$  values on NPF days were generally lower than on non-NPF days in all seasons except summer, which is generally attributed to an increased proportion of weakly hygroscopic components in newly formed particles. However, the  $\kappa_{\text{inorg}}$  on NPF days was significantly higher than on non-NPF days in summer. Despite reduced hygroscopicity, NPF events effectively increased  $N_{\text{CCN}}$ , which was also combined with higher  $\text{Cl}^-$  concentration on summer NPF days in Fig. S5 ( $0.59 \mu\text{g m}^{-3}$  vs.  $0.28 \mu\text{g m}^{-3}$ ). At SS above 0.4 %,  $N_{\text{CCN}}$  on NPF days was higher than on non-NPF days in all seasons except spring, with the most significant increases observed in winter and fall (Fig. S7).

Figure 3a shows that in spring, particle number concentrations in the 10–20 nm range are generally elevated. NPF typically occurs from 09:00 to 20:00 LT, with the peak concentration of 10–15 nm particles reaching  $29\,498 \text{ cm}^{-3}$ . Concurrently, the concentration of particles  $> 20 \text{ nm}$  increases significantly, with some growing beyond 100 nm. The aerosol size distributions on NPF days in summer, fall, and winter all exhibit an NPF process pattern similar to that shown in Fig. 1, typically occurring in 09:00–20:00 LT. During summer NPF events, the maximum particle number concentration reaches approximately  $11\,410 \text{ cm}^{-3}$ , with particles growing up to around 50 nm. In fall and winter, peak particle number concentrations are lower than in summer ( $10\,110$

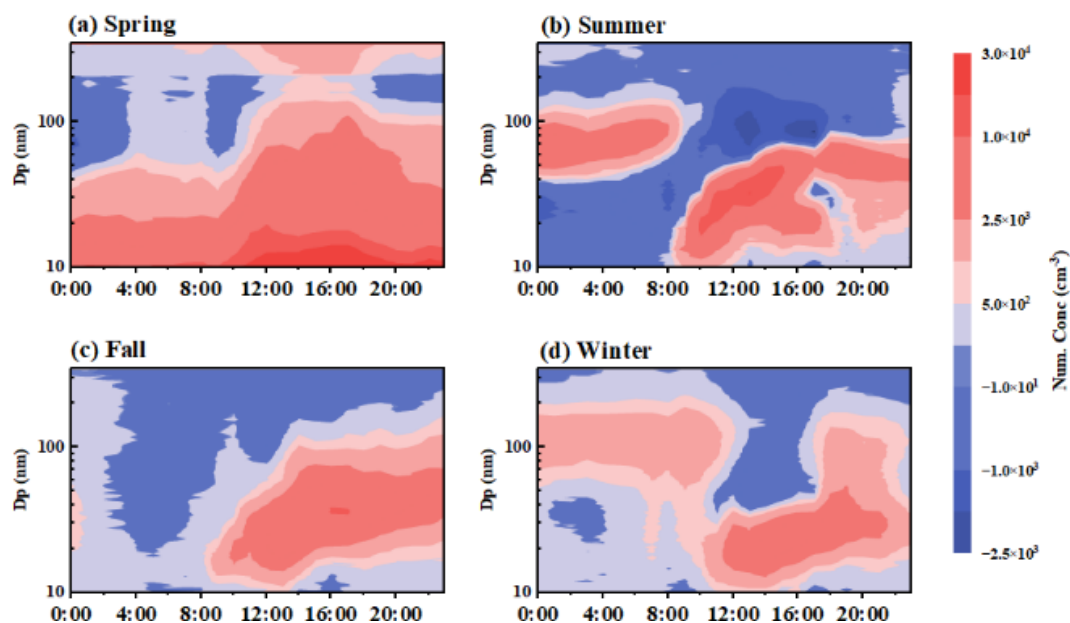
and  $5276 \text{ cm}^{-3}$ ), indicating weaker NPF intensity, and the maximum particle growth can extend up to 100 nm in these seasons.

### 3.2 Evolution characteristics and key parameters of NPF events

Spring NPF events had the highest FR ( $7.13 \text{ cm}^{-3} \text{ s}^{-1}$ ) and the highest CS ( $4.1 \times 10^{-2} \text{ s}^{-1}$ ) among all seasons (Table 1). Before NPF events ( $-4$  to  $0 \text{ h}$ ),  $N_{\text{nuc}}$  increased from  $5425$  to  $7701 \text{ cm}^{-3}$ , peaking at  $1 \text{ h}$  ( $8840 \text{ cm}^{-3}$ ), then gradually declining. This trend was consistent with changes in the FR (Fig. 4a). FR rose sharply to  $6.31 \text{ cm}^{-3} \text{ s}^{-1}$  at the onset of NPF ( $0 \text{ h}$ ) and peaked at  $7.21 \text{ cm}^{-3} \text{ s}^{-1}$  at  $1 \text{ h}$ , demonstrating strong new particle formation capability. However, intense competition under a high CS background significantly suppressed subsequent growth. The growth rate (GR) exhibited large fluctuations, reverting to a negative value at  $3 \text{ h}$  after an initial peak.  $N_{\text{ait}}$  increased by only 32 % from  $0 \text{ h}$  to its peak at  $2 \text{ h}$ , much lower than in other seasons. Spring NPF events were characterized by strong formation but suppressed growth under high CS.

Despite the low average FR ( $0.40 \text{ cm}^{-3} \text{ s}^{-1}$ , Table 1), summer exhibited the highest average GR ( $4.20 \text{ nm h}^{-1}$ ), and the lowest average CS ( $1.8 \times 10^{-2} \text{ s}^{-1}$ ). In summer NPF events (Fig. 4b),  $N_{\text{nuc}}$  peaked at  $1 \text{ h}$  ( $3459 \text{ cm}^{-3}$ ). FR relatively high from  $0$ – $2 \text{ h}$  ( $0.97$ – $0.44 \text{ cm}^{-3} \text{ s}^{-1}$ ). The most prominent feature of summer was the high growth efficiency (GR) under the lowest CS, with a maximum peak of  $11.68 \text{ nm h}^{-1}$  at  $2 \text{ h}$ . After NPF onset in summer,  $N_{\text{ait}}$  peaked at  $3 \text{ h}$  ( $6461.6 \text{ cm}^{-3}$ ), with the highest increase of 202.91 %. This indicates that the growth process of new particles in summer NPF events was far stronger than particle formation.

Winter had the lowest average FR ( $0.23 \text{ cm}^{-3} \text{ s}^{-1}$ ) and GR ( $1.87 \text{ nm h}^{-1}$ , Table 1). Winter NPF events were characterized by a low FR and delayed growth under a low condensation sink (CS), as shown in Table 1. In Fig. 4d, the FR peak observed at  $0 \text{ h}$  ( $0.43 \text{ cm}^{-3} \text{ s}^{-1}$ ) was the lowest among all seasons, indicating weak nucleation. In contrast, the growth rate (GR) displayed a distinct multi-peak pattern, with an ini-



**Figure 3.** The differences in particle number size distributions between NPF days and non-NPF days for each season.

**Table 1.** Seasonal variations in key parameters of NPF events: particle growth rate (GR), formation rate (FR), coagulation sink (CoagS), condensable vapor concentration ( $C$ ) and its production rate ( $Q$ ), condensation sink (CS), and hygroscopicity parameter ( $\kappa_{\text{inorg}}$ ).

Average	FR ( $\text{m}^{-3} \text{s}^{-1}$ )	GR ( $\text{nm h}^{-1}$ )	CS ( $\times 10^{-2} \text{s}^{-1}$ )	CoagS ( $\times 10^{-4} \text{s}^{-1}$ )	$C$ ( $\times 10^7 \text{cm}^{-3}$ )	$\kappa_{\text{inorg}}$
Total	3.94	5.20	3.1	3.9	16.7	0.56
Spring	7.13	3.69	4.1	5.2	20.1	0.54
Summer	0.40	4.20	1.8	2.0	10.9	0.59
Fall	0.28	2.35	2.0	1.8	9.5	0.55
Winter	0.23	1.87	2.1	1.9	4.7	0.58

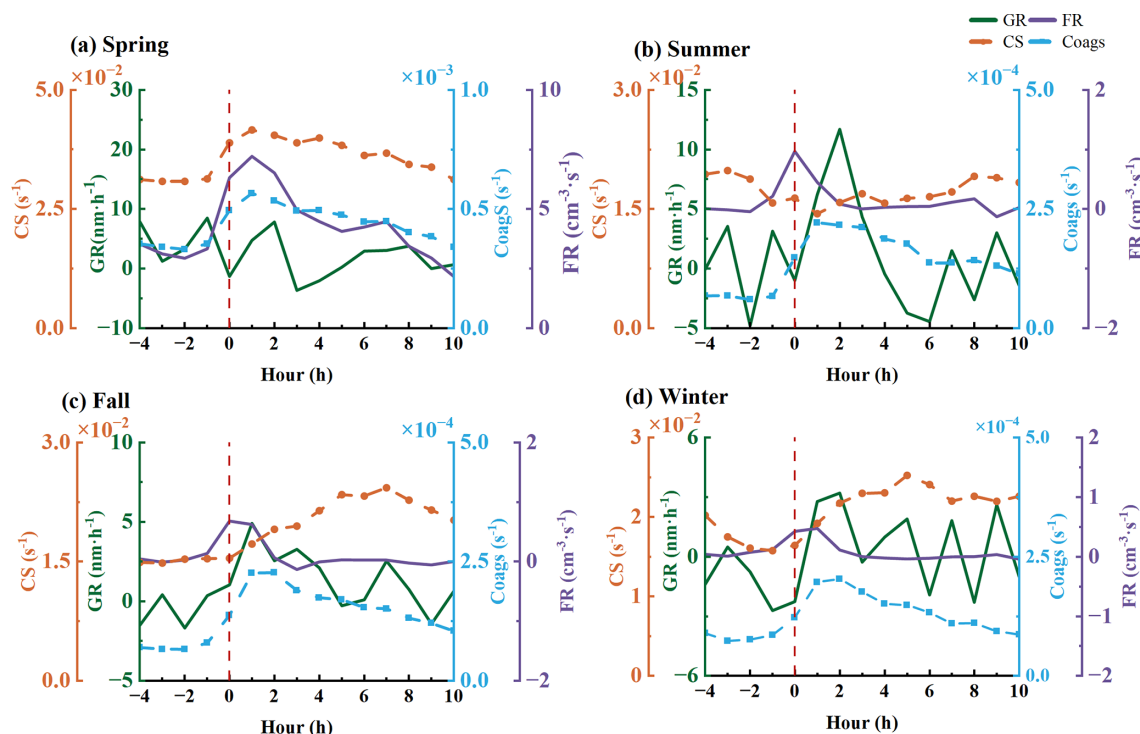
tial peak at 2 h ( $3.20 \text{ nm h}^{-1}$ ) and subsequent peaks occurring between 5 and 9 h, suggesting that different mechanisms may have driven particle growth at different stages. Correspondingly,  $N_{\text{ait}}$  reached a maximum at 3 h ( $4794.2 \text{ cm}^{-3}$ ), which was the lowest seasonal peak (Fig. S8). Nevertheless,  $N_{\text{ait}}$  remained at relatively high levels ( $4500\text{--}4800 \text{ cm}^{-3}$ ) over an extended period from 2 to 6 h, reflecting sustained particle growth throughout the event.

Fall presented transitional characteristics, with average FR ( $0.28 \text{ cm}^{-3} \text{ s}^{-1}$ ) and GR ( $2.35 \text{ nm h}^{-1}$ ) higher than winter but lower than spring and summer (Table 1). Fall NPF process parameters showed transitional characteristics between summer and winter, generally similar to winter (Fig. 4c). Its FR peak ( $0.68 \text{ cm}^{-3} \text{ s}^{-1}$ ) and GR peak ( $4.90 \text{ nm h}^{-1}$ ) were higher than winter but much lower than spring and summer. The increase in  $N_{\text{ait}}$  after NPF onset was 165 %, significantly stronger than in winter (Fig. S8). The  $N_{\text{ait}}$  peak ( $6240.9 \text{ cm}^{-3}$ ) occurred latest (4 h) and remained above  $5600 \text{ cm}^{-3}$  from 5–7 h, higher than winter.

In summary, spring shows the highest FR ( $7.13 \text{ cm}^{-3} \text{ s}^{-1}$ ) but the lowest GR ( $3.69 \text{ nm h}^{-1}$ ) due to a large CS, indicating strong nucleation yet suppressed growth. Summer achieves the highest GR (peak  $11.68 \text{ nm h}^{-1}$ ) and the cleanest background (lowest CS), where growth dominates over formation. Fall and winter exhibit low FR and GR with delayed growth, reflecting weaker NPF intensity.

### 3.3 Influence of chemical composition on NPF events

Spring NPF days were characterized by the highest CS ( $4.1 \times 10^{-2} \text{ s}^{-1}$ ) and were predominantly influenced by secondary pollution. Spring NPF events were mainly driven by the continuous transport from the southeast coastal pathway (CL2, 51.33 %, Fig. S9a), which supplied high levels of  $\text{NH}_3$  ( $6.76 \pm 2.09 \mu\text{g m}^{-3}$ ) and  $\text{HNO}_2$  ( $2.81 \pm 2.16 \mu\text{g m}^{-3}$ ) (Table S4). Consistently, during the NPF events, the frequency of SE/S winds continuously increased and became dominant in the later stage (maximum 38.1 %), reflecting the influence of this coastal air mass at the local scale. Before NPF



**Figure 4.** Key NPF parameters: growth rate (GR), formation rate (FR), condensation sink (CS), and coagulation sink (CoagS) for each season. The NPF events in each season have been normalized. The  $x$  axis represents time relative to the NPF event start, where  $t = 0$  h is the onset time (defined as the time when the nucleation-mode particle number concentration begins a sustained increase). Negative times indicate hours before the start, and positive times indicate hours after the start. The time axis ranges from  $-4$  to  $10$  h, showing the evolution from  $4$  h before to  $10$  h after the event.

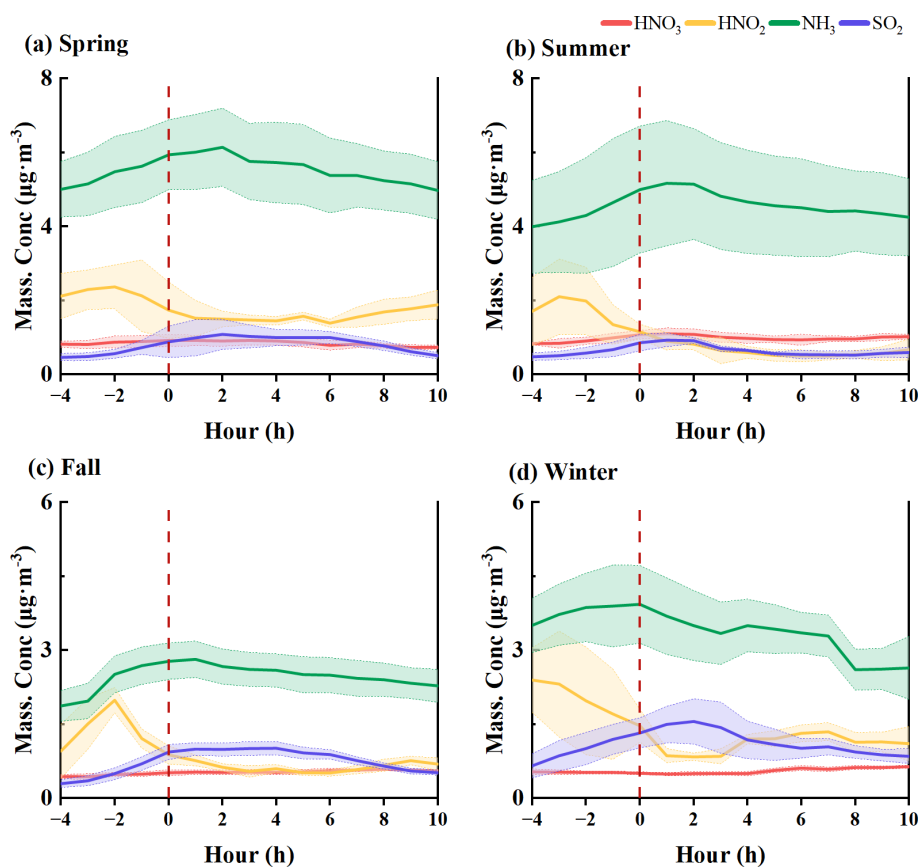
events, gaseous  $\text{SO}_2$  and  $\text{NH}_3$  increased from  $0.47$  and  $5.01$  to  $0.89$  and  $5.95 \mu\text{g m}^{-3}$ , respectively (Fig. 5a). Two hours after NPF onset, their concentrations began to decline continuously, indicating substantial consumption. Mass concentrations of secondary inorganic salts ( $\text{SO}_4^{2-}$  and  $\text{NO}_3^-$ ) fluctuated between  $4.5$ – $4.8$  and  $4$ – $6 \mu\text{g m}^{-3}$ , respectively. Although  $\text{NO}_3^-$  showed minor fluctuations, it remained at relatively high levels (Fig. 6a). NPF events occurred when the temperature was between  $18.9$ – $23.2^\circ\text{C}$  (Fig. S12a), favoring accelerated photochemical reactions and gas-particle conversion of semi-volatile gases (Chen et al., 2023). Wind speeds were generally low ( $< 1.9 \text{ m s}^{-1}$ ), and the atmospheric stratification was stable. After NPF onset,  $\text{O}_3$  increased significantly (from  $81.1$  to  $98.8 \mu\text{g m}^{-3}$ ; Fig. S14a), indicating enhanced atmospheric oxidizability. Precursor gases ( $\text{SO}_2$  and  $\text{NO}_2$ ) were oxidized via photochemistry to form  $\text{NO}_3^-$  and  $\text{SO}_4^{2-}$ , promoting the generation of secondary inorganic salts and secondary aerosols, including secondary organic aerosol (Fig. S13a). The high CS background, spring had high-frequency and high-FR NPF events, primarily attributed to higher precursor gas and strong photochemistry.

However, high CS competed for condensable vapors and scavenged newly formed particles, suppressing the growth stage of new particles. The high particle hygroscopicity

( $0.56$ ) was observed at  $-4$  h before the event, and it decreased to  $0.53$  during the NPF event ( $1$ – $4$  h) (Fig. S11).

CS in summer, fall, and winter were relatively low (around  $2.0 \times 10^{-2} \text{ s}^{-1}$ ), indicating fewer surfaces available for condensation in the atmosphere.

Summer NPF events occurred under a high-temperature environment ( $> 32^\circ\text{C}$ ) (Fig. S12b), which inhibits nucleation (Yu et al., 2017). Two hours after NPF onset ( $2$  h), the frequency of northeasterly (NE) winds surged from  $0\%$  (at  $-4$  h) to  $50\%$ , corresponding to marine air mass cluster CL2 ( $60\%$ ), which had a low average CS of  $1.08 \pm 0.31 \times 10^{-2} \text{ s}^{-1}$  (Table S3). The arrival of this air mass diluted local pollutants, causing  $\text{PM}_{10}$  to drop from  $14.25 \mu\text{g m}^{-3}$  at  $0$  h to  $8.75 \mu\text{g m}^{-3}$  at  $2$  h; meanwhile, it supplied abundant gaseous precursors or enhanced local photochemical activity, leading to continuous increases in  $\text{SO}_2$  and  $\text{NH}_3$  from  $0.48$  and  $4.00 \mu\text{g m}^{-3}$  to peak values of  $0.94$  and  $5.18 \mu\text{g m}^{-3}$ , respectively, at  $2$  h (Fig. 5b). Therefore, despite favorable conditions of low CS and ample sunlight, summer NPF frequency and formation rate were low. As ozone continued to rise ( $0$ – $6$  h), secondary ions ( $\text{NO}_3^-$ ,  $\text{SO}_4^{2-}$ ) and SOC increased (Figs. 6b and S13b). These efficiently condensed onto particle surfaces under a low CS background, achieving extremely high growth efficiency (GR peak of  $11.59 \text{ nm h}^{-1}$ ).  $\kappa_{\text{inorg}}$  in the range of  $0.61$ – $0.75$  was higher than in other sea-



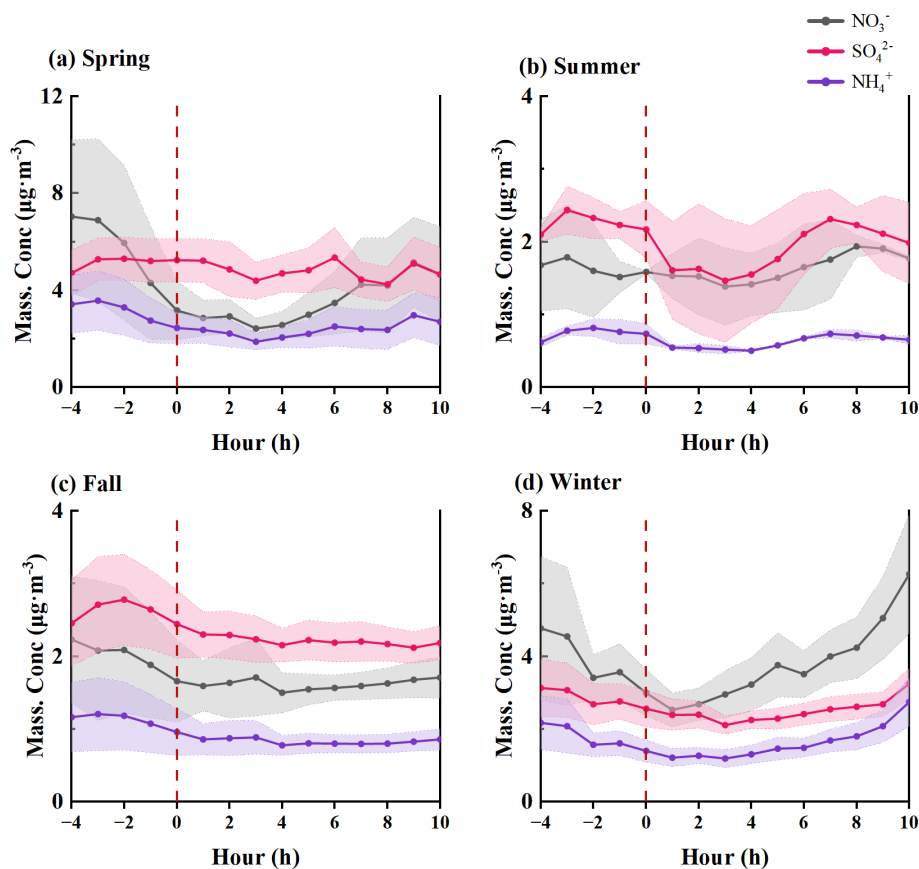
**Figure 5.** The evolution of relevant trace gases ( $\text{SO}_2$ ,  $\text{NH}_3$ ,  $\text{NO}_2$ ) for (a) spring, (b) summer, (c) fall, and (d) winter. The  $x$  axis follows the same normalized time scale as defined in Fig. 4 ( $t = 0$  h represents NPF event start). Shaded bands indicate  $\pm 1\sigma$  standard deviation.

sons, surging to 0.75 at 3 h after NPF onset. This increase was primarily driven by prevailing easterly and southerly winds, corresponding to air mass cluster CL3, which had a high  $\kappa_{\text{inorg}}$  (0.71). This indicates that inorganic salts of marine origin facilitated the sustained growth of particles.

Before Fall NPF events,  $\text{SO}_2$  and  $\text{NH}_3$  increased from 0.36 and 1.99 to 2.83 and 2.83  $\mu\text{g m}^{-3}$ , respectively (Fig. 5c). This stage was dominated by northwesterly winds (NW: 62 %) and strongly influenced by continental air masses from CL2 and CL3, which contributed to the initial accumulation of precursor gases ( $\text{SO}_2$  and  $\text{NH}_3$ ). The lower temperature (around 22 °C) and high humidity ( $\text{RH} > 71\%$ ) environment (Fig. S12c) favored the combination of sulfuric acid molecules, promoting nucleation (Lehtipalo et al., 2018; Tröstl et al., 2016; Yue and Hamill, 1979). After NPF onset, although atmospheric oxidizability increased continuously ( $\text{O}_3$  from 36 to 94  $\mu\text{g m}^{-3}$ ), it was short-lived. Within 1–3 h after NPF onset, the local wind direction shifted notably to southerly (S: 44.4 %, Fig. S10c), corresponding on a larger scale to the marine/coastal air mass CL1 (22.22 %, Fig. S9c). CL1 served as the most important transport pathway for  $\text{NH}_3$  in the fall, with an average concentration as high as 3.13  $\mu\text{g m}^{-3}$  (Table S4), substantially higher than that

of continental pathways. After 4 h, as  $\text{O}_3$  gradually decreased and primary emissions increased, pollutant accumulation occurred, and BC and POC rebounded (Fig. S13c). The contribution of primary emissions (e.g., BC, POC) to aerosols was significantly enhanced, and overall  $\kappa_{\text{inorg}}$  (0.55) was low.

Winter NPF events were also preceded by an accumulation of gaseous precursors (Fig. 5d). However, elevated emissions from sources such as heating led to high BC and POC during the initial NPF stage (1.86 and 1.02  $\mu\text{g m}^{-3}$  at  $-4$  h, Fig. S13d). These abundant primary particles strongly suppressed new particle formation via intense coagulation scavenging, resulting in persistently low formation rates ( $\text{FR} \leq 0.47 \text{ cm}^{-3} \text{ s}^{-1}$ ). Photochemical activity was limited under low winter temperatures (Fig. S12d). As temperatures continued to drop later in the event (after 6 h), condensation-driven conversion of gaseous precursors to particles increased (Jokinen et al., 2018). During the 6–10 h period, the high concentrations of precursors reached supersaturation under the low-temperature conditions, promoting the rapid formation of particulate ammonium nitrate (nitrate concentration increased from 4.21 to 5.14  $\mu\text{g m}^{-3}$ , Fig. 6d). In winter, the air masses were mainly dominated by the northwesterly continental air mass originating from Siberia (CL1,



**Figure 6.** The evolution of major secondary inorganic ions ( $\text{SO}_4^{2-}$ ,  $\text{NO}_3^-$ ,  $\text{NH}_4^+$ ) for each season. The  $x$  axis follows the same normalized time scale as defined in Fig. 4 ( $t = 0$  h represents NPF event start). Shaded bands indicate  $\pm 1\sigma$  standard deviation.

Fig. S9d). Particles within this air mass exhibited relatively high  $\kappa_{\text{inorg}}$  (0.59, Table S3), and the introduction of these hygroscopic components effectively offset the dilution of overall hygroscopicity caused by BC and POC. During the 8–10 h period, northwesterly winds persisted, and their frequency gradually increased, which also explains the gradual increase in  $\kappa_{\text{inorg}}$  values (from 0.57 to 0.61, Fig. S11).

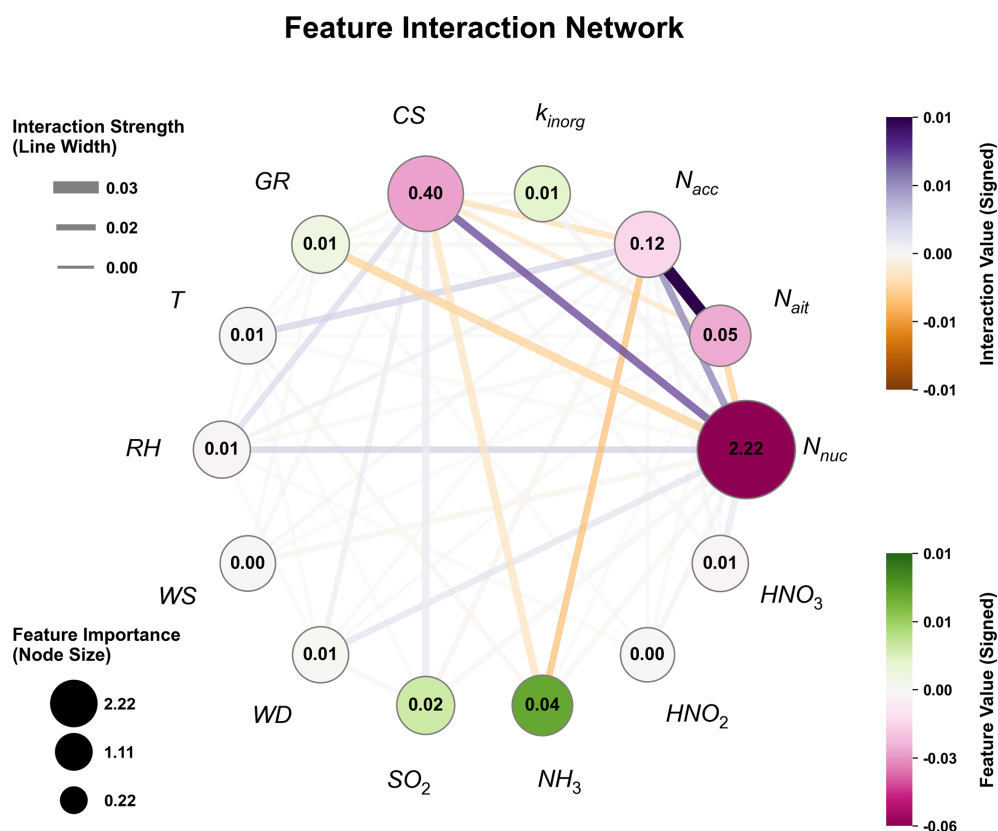
In summary, in spring, high  $\text{NH}_3$  and photochemistry drive strong nucleation, but high CS and secondary inorganic salts suppress growth. Summer marine air masses provide low CS and high hygroscopicity ( $\kappa_{\text{inorg}} > 0.6$ ), favoring efficient growth, while fall and winter continental emissions (BC, POC) lower  $\kappa_{\text{inorg}}$  ( $\leq 0.55$ ) and limit NPF intensity.

### 3.4 Attribution analysis of particle formation and growth

According to SHAP analysis (Fig. S15), the main contributors to FR in Fuzhou are nucleation mode (76.2%), CS (13.8%),  $\text{NH}_3$  (1.3%), and  $\text{SO}_2$  (0.6%). Physical processes explain about 90% of the variance. A prominent feature is a sharp  $\text{NH}_3$  threshold at  $4 \mu\text{g m}^{-3}$ . Below this value, SHAP is approximately  $-0.025$ ; above it, SHAP becomes positive and rises to approximately 0.2 at  $10 \mu\text{g m}^{-3}$ . CS shows a clear

inhibition onset at  $0.03 \text{ s}^{-1}$ , with SHAP dropping from 0 to  $-0.5$  as CS increases to  $0.08 \text{ s}^{-1}$ . Temperature has a positive SHAP only within 20–25 °C (peak 0.01), decreasing by 40% at 35 °C. As shown in Fig. 7, the strongest interaction is between nucleation mode and CS (0.40), far exceeding other pairs ( $\text{NH}_3$ -nucleation mode: 0.04), highlighting that the net FR is governed by the competition between particle formation and scavenging.

For the Aitken mode, the driving forces shift notably. CS remains the top contributor, followed by accumulation mode (13.8%) and nucleation mode (7.1%). As shown in Fig. S16, a key distinction from FR is the temperature response. Instead of a bell shape, SHAP increases linearly from 0 to 0.1 over 20–35 °C, indicating that high temperatures accelerate particle growth, enabling a rapid transition from nucleation to the Aitken mode. The  $\text{NH}_3$  threshold at  $4 \mu\text{g m}^{-3}$  persists, showing its continued role in particle growth. RH exerts a linear negative effect, turning SHAP negative above 60%, suggesting hygroscopic growth or coagulation loss. In Fig. S17, the CS-nucleation mode interaction strength is 0.12, lower than in FR but still dominant, implying that strong nucleation can offset high CS losses.



**Figure 7.** Non-linear response relationships and interaction analysis between core environmental factors and FR. Node size and inner numbers denote feature importance (mean |SHAP|). Node color indicates contribution direction (green: positive; pink: negative). Edge width reflects interaction strength; edge color indicates interaction direction (purple: synergistic; orange: antagonistic).

In summary, the SHAP attribution reveals two distinct regimes. FR is dominated by CS, accompanied by a sharp chemical trigger ( $NH_3 > 4 \mu g m^{-3}$ ) and a narrow temperature window (20–25 °C). The Aitken mode, while still influenced by CS, is primarily driven by temperature-accelerated growth (linearly increasing above 20 °C). Ammonia acts as a persistent enhancer in both stages, whereas high RH (> 60 %) consistently suppresses Aitken mode concentrations.

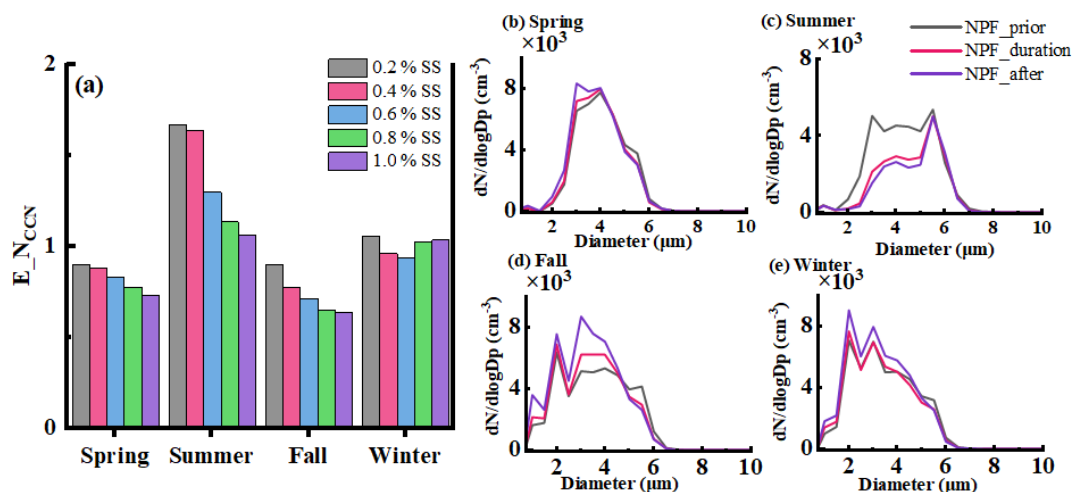
### 3.5 Particle growth controls CCN formation from NPF events

In summer, although NPF frequency was the lowest, it had a significant enhancing effect on  $N_{CCN}$ .  $E_{N_{CCN}}$  at 0.4 % SS was as high as 1.64 (Fig. 8a). In the initial stage of NPF events (0–2 h),  $N_{CCN}$  showed a sharp decline. At 0.4 % SS, it decreased from about 2078 to 1187  $cm^{-3}$  (Fig. S18b). However, after 2 h,  $N_{CCN}$  recovered noticeably and later returned to or even exceeded initial levels. The CCN size distribution (Fig. 8c) showed that during the event, smaller CCN (1–3  $\mu m$ ) decreased, while CCN of 5.5–6.5  $\mu m$  increased. For instance, CCN at 6  $\mu m$  increased from 2607.7 to 3147.8  $cm^{-3}$ . Summer NPF promoted CCN growth to larger sizes. However, the daily average  $N_{CCN}$  on summer NPF days was lower

than on non-NPF days, possibly due to the extremely low NPF frequency (4.35 %) and the sharp decline in CCN at the beginning of events, affecting the daily average.

Spring NPF events' impact on  $N_{CCN}$  showed significant suppression. Throughout the event,  $N_{CCN}$  at various supersaturation levels showed only weak and slow increases.  $N_{CCN}$  (0.4 % SS, the same as below) increased from a pre-event average (–4 to –1 h) of about 2636 to 3192  $cm^{-3}$  at 4 h, then decreased slowly from 4 to 6 h, with a larger decline at higher supersaturations (Fig. S18a).  $E_{N_{CCN}}$  at 0.4 % SS was only 0.88. This phenomenon corresponds to the “high formation, suppressed growth” characteristic of spring NPF. Despite the explosive generation of nucleation-mode particles, severe competition under high CS severely hindered subsequent growth of new particles, preventing them from effectively growing to CCN activation sizes, resulting in a weak or even negative contribution to CCN.

In fall events, from 2 h onward,  $N_{CCN}$  began to increase, rising from 3471  $cm^{-3}$  to a maximum of 4752  $cm^{-3}$ , forming a high-value plateau lasting from 3 to 8 h (Fig. S18d). After the NPF event,  $N_{CCN}$  increased significantly across the 1–5  $\mu m$  size range (Fig. 8d). Fall NPF events most effectively and broadly increased the number of particles avail-



**Figure 8.** The contribution of NPF events to cloud condensation nuclei (CCN) across seasons: (a) CCN enhancement ( $E_{NCCN}$ ), and (b–e) particle number size distributions at 2 h before, during, and 5 h after the NPF event for seasons.

able for cloud droplet activation in the atmosphere. Although  $E_{NCCN}$  was lower due to the modest initial stage, the post-event enhancement effect was significant. This evolution process is consistent with the sustained particle growth process in fall NPF, allowing new particles to grow steadily to CCN activation sizes.

In winter NPF events,  $N_{CCN}$  was low in the early stage (0–4 h), with insignificant growth. However, from 5 h onward,  $N_{CCN}$  growth became significant, increasing from 4483 to 6173  $\text{cm}^{-3}$  and remaining stable at high levels (Fig. S18d). During the event, changes in the CCN size distribution were not obvious. However, after the NPF event, CCN in the 2–5  $\mu\text{m}$  size range showed the most significant growth (e.g., at 2  $\mu\text{m}$ , from 7073.0 to 9045.1  $\text{cm}^{-3}$ ; Fig. 8e).

#### 4 Conclusions

This year-long observational study in coastal Fuzhou revealed distinct seasonal patterns in new particle formation (NPF) and its impact on cloud condensation nuclei (CCN). We identified 46 NPF events, which predominantly occurred between 09:00 and 12:00 LT. Key quantitative results include: the highest seasonal formation rate (FR) in spring ( $5.56 \text{ cm}^{-3} \text{ s}^{-1}$ ), the highest growth rate (GR) in summer (peak at  $11.68 \text{ nm h}^{-1}$ ), and the strongest CCN enhancement in summer ( $E_{NCCN} = 1.64$  at 0.4 % SS). In fall and winter, CCN increases (13 %–65 %) lag NPF events by 3–5 h.

A total of 46 NPF events occurred during the observation period, with a frequency of 12.7 %. NPF event start times were mainly concentrated between 08:00 and 13:00 LT (accounting for 85 % of all events), with an average duration of 4 h. The annual averages for formation rate (FR), growth rate (GR), condensation sink (CS), coagulation sink (CoagS), and condensable vapor concentration (C) were

$3.94 \pm 8.26 \text{ cm}^{-3} \text{ s}^{-1}$ ,  $5.20 \pm 1.78 \text{ nm h}^{-1}$ ,  $4.2 \times 10^{-2} \text{ s}^{-1}$ ,  $5.6 \times 10^{-4} \text{ s}^{-1}$ , and  $16.7 \times 10^7 \text{ cm}^{-3}$ , respectively.

The seasonal contrast is mechanistically driven by the interplay between precursor availability, CS, and aerosol chemistry. Spring conditions favor high FR due to strong photochemistry and abundant precursors, but high CS suppresses subsequent growth. In summer, the occurrence of NPF events is often accompanied by a transition in sea-land breeze circulation. The northerly land breeze shifts to northeasterly/easterly sea breezes, effectively transporting clean marine air masses to the observation site. This reduces the CS to its seasonal minimum ( $1.08 \pm 0.31 \times 10^{-2} \text{ s}^{-1}$ ), favoring efficient particle growth and thereby sustaining the highest growth rate (GR) observed across all seasons. The chemical composition shifts from being dominated by sulfate and sea salt with high hygroscopicity in summer ( $\kappa_{\text{inorg}} > 0.6$ ) to low hygroscopicity in fall ( $\kappa_{\text{inorg}} = 0.55$ ), directly modulating particle hygroscopicity and cloud condensation nuclei (CCN) activation potential.

Using XGBoost-SHAP, FR is dominated by nucleation mode (76.2 %) and CS (13.8 %).  $\text{NH}_3 > 4 \mu\text{g m}^{-3}$  enhances FR,  $\text{CS} > 0.03 \text{ s}^{-1}$  inhibits FR (SHAP drops to  $-0.5$  at  $0.08 \text{ s}^{-1}$ ), and Temperature benefits FR only within 20–25 °C. For Aitken mode, temperature linearly promotes growth above 20 °C, while  $\text{RH} > 60 \%$  suppresses it. The strong nucleation-CS interaction (0.40) highlights the competition between formation and scavenging.

This work demonstrates that the climatic impact of NPF in coastal urban areas is not simply a function of its occurrence frequency or formation strength. Instead, it is seasonally modulated by a competition between formation and growth-sink processes, and by the resulting aerosol chemical composition. The decoupling of high FR from effective CCN production (as in spring) implies that climate models using NPF frequency as a proxy for aerosol indirect effects may

overestimate the impact in high-CS coastal regions. Conversely, the efficient growth and CCN enhancement in summer suggest that even infrequent NPF events can substantially influence cloud properties in such environments. These insights are crucial for developing more accurate parameterizations of aerosol-cloud-climate interactions in rapidly developing coastal zones.

**Data availability.** Data will be made available on request.

**Supplement.** The supplement related to this article is available online at <https://doi.org/10.5194/acp-26-7539-2026-supplement>.

**Author contributions.** Conceptualization was completed by ZW and HW. Formal analysis and software modification were carried out by ZW and HW. The initial draft was written by ZW, incorporating tables provided by HW. The first draft was revised by ZW, HW, and YB, with input from HW, YB, FZ, WL, JH, LS, and ZX; subsequent versions of the manuscript were reviewed and edited by all authors. Data curation and collection were conducted by all authors. Funding acquisition was secured by HW.

**Competing interests.** The contact author has declared that none of the authors has any competing interests.

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