

# Investigating the contribution of grown new particles to cloud condensation nuclei with largely varying pre-existing particles - Part 2: Modeling chemical drivers and 3-D NPF occurrence

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**Abstract.** In this study, we utilized a 20-bin WRF-Chem (Weather Research and Forecast coupled with Chemistry regional model) to investigate the contributions of chemical drivers to the growth of **new particles**, as well as to simulate the three-dimensional dynamics of new particle formation (NPF) events over the North China Plain during a summer campaign in 2019. **The model replicated the occurrence of NPF and the growth pattern of newly formed particles, and the performance to meet the benchmarks, i.e., absolute mean fractional bias  $\leq 50\%$  and mean fractional error  $\leq 75\%$ , in replicating number concentrations of particles in the 10–40 nm range in five events between June 29 and July 6, 2019.** Therefore, we further analyzed three NPF events with distinct particle growth characteristics. **In these instances, the model overpredicted daytime condensation of H<sub>2</sub>SO<sub>4</sub> vapor and nighttime formation of NH<sub>4</sub>NO<sub>3</sub>. These resulted in overestimation of the hygroscopicity parameter of nanometer particles.** Nevertheless, the model performance **met the benchmarks for** reproducing CCN at a **supersaturation (SS)** of 0.4 % **on NPF days**. This was because the overestimation **of inorganics** was offset by the **model underestimation of CCN originating from submicron particles**. Additionally, three-dimensional simulations of NPF events **demonstrated** some key findings. **First, NPF consistently begins in the upper parts of the planetary boundary layer (PBL) before expanding. Second, during daytime organics dominate growth of new particles in the PBL, whereas in the free troposphere the primary chemical drivers are inorganic species.** However, to confirm these findings, vertical observations are required.

**Keywords:** NPF; WRF-Chem; secondary organic aerosols; NH<sub>4</sub>NO<sub>3</sub>; spatial inhomogeneity

## 1 1 Introduction

2 In the atmosphere, gaseous precursors form critical nuclei that can grow. This new particle formation  
3 (NPF) has been extensively studied (Kulmala et al., 2004; Bzdek and Johnston, 2010; Zhang et al., 2012;  
4 Chu et al., 2019; Lee et al., 2019; Sellegri et al., 2019). NPF events cause a sharp increase in particle  
5 number concentrations (PNCs) and can potentially impact the global climate by acting as cloud  
6 condensation nuclei (CCN) (Huang et al., 2016; Gordon et al., 2017; Yu et al., 2017). Specifically, NPF  
7 has been estimated to contribute as much as 45 % to the global CCN budget (Spracklen et al., 2008;  
8 Merikanto et al., 2009; Williamson et al., 2019). Furthermore, NPF particles have been shown to continue  
9 growing for several days to make substantial contributions to atmospheric particle mass. This link  
10 between NPF and subsequent haze events has been observed in China (Zhang et al., 2012; Chu et al.,  
11 2021; Kulmala et al., 2022). In fact, if the newly formed particles grow to a sufficient size, they may  
12 have direct climate effects by altering atmospheric radiation.

13 Although the North China Plain (NCP) is one of the largest Asian plains, it still suffers from air  
14 pollution (Li et al., 2017; Jiang and Bai, 2018; Ma et al., 2019; Yang et al., 2021). Despite this, the NCP  
15 frequently experiences NPF events, due to the abundance of precursors such as sulfuric acid, ammonia,  
16 amines, and secondary gaseous organics, as well as dry weather (Wehner et al., 2004; Wu et al., 2007;  
17 Yue et al., 2010; Wang et al., 2015; Zhu et al., 2017; Ma et al., 2021; Chu et al., 2021). The significant  
18 decrease in NCP air pollutants, including SO<sub>2</sub> and NO<sub>x</sub>, over the past decade, as reported by Chen et al.  
19 (2019a), Wen et al. (2021), and Zhu et al. (2021a). This could affect CCN production (Dusek et al., 2006;  
20 Hudson, 2007; Zhu et al., 2021b). This situation makes it imperative and essential to conduct new studies  
21 that quantify chemical drivers for growth of new particles that may become CCN and modification of  
22 pre-existing particles simultaneously over NCP (Wei et al., 2023).

23 Commercial particle sizers, such as the Scanning Mobility Particle Sizer (SMPS), Wide-range  
24 Particle Sizer (WPS), and fast mobility particle sizer (FMPS) have limitations in detecting particles  
25 smaller than 3 or 5.6 nm and have low detection efficiency for particles smaller than 15 nm. As a result,  
26 newly formed particles are typically observed at initial sizes larger than 6–15 nm. Growth of clusters  
27 larger than 6–15 nm takes hours, during which time NPF can occur and move with the air mass. Although  
28 observations are important to characterize NPF events and explore related mechanisms, the use of one  
29 fixed-site to observe condensable vapors is not sufficient to explain NPF occurring downwind, and it is  
30 difficult to perform Lagrangian observations in moving air masses. Thus, three-dimensional (3-D)  
31 modeling studies are needed to determine where NPF events initially occurred. Furthermore, it has been  
32 suggested that long-duration NPF events can extend hundreds of kilometers horizontally (Wehner et al.,  
33 2007; Hussein et al., 2009; Crippa and Pryor, 2013; Pikridas et al., 2015; Kerminen et al., 2018). However,  
34 relying solely on data from one or two observation sites limits understanding of NPF regional scale spatial  
35 inhomogeneities. Factors such as NPF event duration, particle formation, and particle growth rates can  
36 differ significantly therein (Kim et al., 2016; Dai et al., 2017; Shen et al., 2018). Therefore, NPF modeling  
37 studies are critical to fully explore the 3-D dynamic evolution of NPF events.

38 Matsui et al. (2009) utilized the Weather Research and Forecasting (WRF)-Community Multiscale  
39 Air Quality (CMAQ) and WRF-Chem models to reasonably replicate PNCs and identify instances of  
40 NPF during the CARE-Beijing 2006 campaign. Similarly, Chen et al. (2014) implemented a few

1 simulations of PNCs in NCP using the Nested Air Quality Prediction Modeling System (NAQPMS) with  
2 an Advanced Particle Microphysics (APM) model (Chen et al., 2017; 2019b). The NPF-explicit WRF-  
3 Chem model has reportedly demonstrated a good performance in simulating some regional NPF events  
4 in East Asia and North America (Matsui et al., 2013; Dong et al., 2019; Yu et al., 2020), and Lai et al.  
5 (2022) investigated the vertical transport and distribution of particles using the WRF-Chem model.  
6 Therefore, it would be beneficial to utilize this model to examine the recently observed NPF events in  
7 NCP, especially in terms of their 3-D evolution and chemical drivers for growth of new particles in  
8 horizontal and vertical directions, as highlighted in the companion paper, which discusses the low  
9 probability of newly formed particles to grow to CCN sizes. It is important to note, however, that there  
10 may be significant uncertainties in PNCs emission factors and particle number size distributions (PNSDs)  
11 from primary sources in China due to the lack of such data (Yao et al., 2005; Shen et al., 2022).

12 In this study, the NPF-explicit WRF-Chem model is used to investigate NPF events observed at a  
13 mountain site in NCP from June 23 to July 14, 2019, focusing on chemical drivers to grow new particles  
14 to CCN, the uncertainty of estimated contributions of grown new particles to CCN, and 3-D NPF. Before  
15 the result presentation and discussion, a comprehensive model performance evaluation will be delivered  
16 in Section 2.4. Section 3.1 will provide an overview of modeling NPF events. The simulated chemical  
17 drivers to grow newly formed particles at ground level and different heights will be presented in Sections  
18 3.2 and 3.3, respectively. Section 3.4 will analyze 3-D evolution of NPF events and transport of new  
19 particles. Section 3.5 will address what happened to grown new particles after they disappeared from  
20 observations.

## 21 2 Methods

### 22 2.1 Observational information

23 The Beijing Forest Ecosystem Positioning Research Station is located at 1170 m elevation a.s.l.  
24 (above sea level, a.s.l.) and is surrounded by Yanshan mountain (39.96°N, 115.43°E; hereinafter referred  
25 to as the mountain station). This area is mainly covered by secondary forest vegetation, such as secondary  
26 shrub, oak, and birch. The mountain station is located at the western edge of Beijing (see Fig. 1), is far  
27 from industrial and urban areas. Strong air pollutant sources and heavily polluted cities are distributed to  
28 the southwest 200–500 km from the site (Ma et al., 2019). No strong air pollutant emission sources are  
29 found to the north where the playing fields of the 2022 Beijing Olympic Winter Games are located.  
30 Simulated NPF events from the north were significantly stronger than those from other directions, as will  
31 be explained later.

32 The performance of the model was assessed using a suite of observational data, which were detailed  
33 in the companion paper. These included both on-line and off-line measurements of total PNCs, PNSDs,  
34 and CCN, which were conducted using a condensation particle counter (CPC; TSI Model 3775), a fast  
35 mobility particle sizer (FMPS, TSI Model 3091), a scanning mobility particle sizer (SMPS, Grimm), and  
36 a continuous flow CCN counter (CCNC, DMT Model 100), respectively. The instruments were located  
37 on the third floor of the main station building and obtained ambient air through conductive silicone tubing  
38 (TSI Inc., product number 3001788, inner diameter 0.19 inch, outer diameter 0.375 inch) of ~ 2 m length.

1 In addition, a high-volume TSP sampler was used for off-line sampling to analyze water-soluble ions, as  
2 well as organic and elemental carbon. For measurements **took place** between June 14 and 30, 2019, both  
3 the SMPS and the additional CPC were used. The FMPS measured from June 23 to July **14, 2019**. **Thus**,  
4 it had a one-week overlap with **the** SMPS measurements. During laboratory tests conducted after the  
5 campaign, it was **determined** that the dryer caused significant particle diffusion losses when the SMPS  
6 and CPC were used with flow rates less than 1 L min<sup>-1</sup>. However, the FMPS with a flow rate of 10 L min<sup>-1</sup>  
7 did not suffer from this issue and was able to accurately capture rapid changes in PNCs from primary  
8 and secondary sources, as documented in previous studies (Yao et al., 2005; 2006; Man et al., 2015).  
9 Therefore, only the data collected between June 23 and July 14 were evaluated here. It is worth noting  
10 that the PNSDs were employed for a comparative analysis of NPF events with those measured at an  
11 urban site in Beijing, as presented in Zhou et al. (2020).

## 12 **2.2 NPF-explicit WRF-Chem Model**

13 The NPF-explicit WRF-Chem model (Grell et al., 2005; Fast et al., 2006) was employed to **simulate**  
14 **NPF events** in the NCP **between June 23 and July 14, 2019**. The model was equipped with a 20-bin  
15 MOSAIC module that covered particle **diameters from** 1 nm to 10 μm (Matsui et al., 2011; Matsui et al.,  
16 2013; Lupascu et al., 2015; Lai et al., 2022). The parameter settings used in the model are shown in Table  
17 S1. It should be noted **that anthropogenic** emissions in China **for 2019** were not publicly available.  
18 Therefore, custom-modified MEIC\_2019 emissions, which were based on MEIC\_2016 and assumed a  
19 linear downward trend in the total amounts of chemicals from 2016 to 2019, **were used**. The custom-  
20 modified MEIC\_2019 emissions were successfully applied to simulate PM<sub>2.5</sub> in the NCP, and more  
21 information can be found in Zhang et al. (2022).

## 22 **2.3 Selection of nucleation mechanism**

23 Based on previous research on the importance of H<sub>2</sub>SO<sub>4</sub> and organic vapors in modeling high-  
24 altitude NPF events, as well as the varied environmental conditions found in **forests** (Metzger et al., 2010;  
25 Schobesberger et al., 2013; Riccobono et al., 2014; Yu and Hallar, 2014; Bianchi et al., 2016; Dong et  
26 al., 2019), **this study selected** the empirical H<sub>2</sub>SO<sub>4</sub>-organic nucleation mechanism in the NPF-explicit  
27 WRF-Chem model for simulating NPF events. The mechanism can be expressed as:

$$28 J = K_{\text{ORG}} \times [\text{H}_2\text{SO}_4] \times [\text{NucORG}] \quad (1)$$

29 The variable J represents the rate of formation of activated clusters with a diameter of 1 nm  
30 (measured in cm<sup>-3</sup> s<sup>-1</sup>). K<sub>ORG</sub> (measured in cm<sup>-3</sup> s<sup>-1</sup>) is an empirical coefficient for nucleation, while  
31 [H<sub>2</sub>SO<sub>4</sub>] and [NucORG] represent the concentrations of gaseous sulfuric acid (measured in cm<sup>-3</sup>) and  
32 low-volatility organic compounds **with a saturation mass concentration of 10<sup>-5</sup>-10<sup>-8</sup> μg m<sup>-3</sup>**, respectively  
33 (Lupascu et al., 2015). The nucleation empirical coefficient is a key parameter **for** accurately simulating  
34 **NPF**, but its value can vary significantly **for** different atmospheric conditions (Sihto et al., 2006; Riipinen  
35 et al., 2007; Matsui et al., 2011; Cui et al., 2014; Sullivan et al., 2018). In this study, a series of sensitivity  
36 tests were conducted to identify the optimal value of K<sub>ORG</sub> for modeling NPF events in the NCP. The  
37 results showed that a value of K<sub>ORG</sub> = 6.2 × 10<sup>-18</sup> cm<sup>-3</sup> s<sup>-1</sup> produced the best performance, and was therefore  
38 used to replace the default value of K<sub>ORG</sub> = 1.00 × 10<sup>-15</sup> cm<sup>-3</sup> s<sup>-1</sup> for modeling purposes.

## 2.4 Model performance evaluation

To evaluate the performance of the model, we compared the modeled PNCs, mass concentrations of secondary ions, and PM<sub>2.5</sub> mass concentrations with the observations. Specifically, we evaluated the modeled CN<sub>10–40</sub>, which is the summed PNCs in the 10–40 nm range, by comparing it with the observations (Fig. 2a). The simulated CN<sub>10–40</sub> showed a higher agreement with the observations during June 29–July 6 (unshaded area in Fig. 2a) than that before June 29 or after July 6. To quantify the simulation performance of the CN<sub>10–40</sub>, we used three statistical parameters: mean fractional bias (MFB), mean fractional error (MFE), and correlation coefficient (R) (Fig. 2b–d). No benchmark is available regarding the statistical metrics for simulated atmospheric particle number concentrations but we adopt the benchmarks and the goal values widely used in air quality studies for PM<sub>2.5</sub> mass concentrations (US EPA, 2007, the benchmark with absolute MFB: ≤50 %; MFE: ≤75 %; the goal values with absolute MFB: ≤30 %; MFE: ≤50 %). During the NPF events period, the MFB of 24 % and the MFE of 66 % of June 29–July 6 met the benchmarks. This R was 0.61, which ranked high in 0.4–0.7 range reported in the literature (Matsui et al., 2013; Lupascu et al., 2015; Dong et al., 2019). However, the three parameters showed poor model reproduction of observations before June 29 and after July 6 (Fig 2b-d). The reasons for the poor simulations are not yet explained. In our recent study (Zhang et al., 2023), we tried to modify the model to improve its performance near the coast. However, those modifications are not applicable to this study and are therefore not applied.

The simulated mass concentrations of SO<sub>4</sub><sup>2-</sup> within PM<sub>10</sub>, shown in Fig. 3a–e, met the benchmarks, i.e., MFB = -28 %, MFE = 41 %, and R = 0.69, for reproducing the observations of total suspended particles collected at the mountain station. However, the model tended to overestimate organic mass concentrations (ORG in Fig. 3b), NO<sub>3</sub><sup>-</sup>, and NH<sub>4</sub><sup>+</sup> within PM<sub>10</sub> before June 29 and after July 6. Despite this, the model was able to reproduce the low mass concentrations of all three species within PM<sub>10</sub> between June 29 and July 6. Differences of simulated concentrations from observed values can be due to model weaknesses such as sometimes poorly predicting meteorology, poorly estimating air pollutant emissions, or lack of key mechanisms (US EPA, 2007; Matsui et al., 2011; Liu et al., 2021; Shen et al., 2022). However, the poor performance in overestimating observed organics, NO<sub>3</sub><sup>-</sup>, and NH<sub>4</sub><sup>+</sup> could also be attributed to sampling artifacts, given their higher volatility compared to ammoniated sulfate acid (Yao et al., 2002; Chow et al., 2010).

During this study period, the chemical composition measured by the ToF-ACSM in Beijing (39.98°N, 116.39°E) was used for evaluation. The simulated mass concentrations of SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, NH<sub>4</sub><sup>+</sup>, and organics within PM<sub>1.0</sub> were reasonably consistent with the June 29 to July 6 observations, which had frequent NPF events (unshaded area in Fig. S1a–d, referring to the frequent-NPF period in this study). However, this was not the case before June 29 and after July 6. Quantitatively, the model performed to meet the benchmarks for simulating SO<sub>4</sub><sup>2-</sup> (MFB = 7 %, MFE = 54 % in Fig. S1e) with R = 0.52, p < 0.05 and organics (MFB = -1 %, MFE = 40 % in Fig. S1h) with R = 0.58, p < 0.05 during the frequent-NPF period. On July 5, the model mistakenly predicted the daytime wind direction to be mainly from the northeast, while the on-site recorded wind direction swayed between southwest and northwest. When excluding the evidently overestimated concentrations of PM<sub>2.5</sub> and all ions on July 5, the model

1 performed better in reproducing  $\text{SO}_4^{2-}$  and organics. The MFB and MFE were below the goal values with  
2 MFB = 4 % and MFE = 49 %, and R largely increased. For the modeled  $\text{NH}_4^+$ , the MFB (-1%) and MFE  
3 (57%) met the benchmarks during the frequent-NPF period (Fig. S1g). Excluding the data on July 5 only  
4 slightly increased R from 0.41 to 0.45. On the other hand, for  $\text{NO}_3^-$ , the model largely overestimated the  
5 observations, and the MFB (-125 %) and MFE (178 %) did not meet the benchmarks during the frequent-  
6 NPF period (Fig. S1f). Excluding the data of July 5, the model performed even worse in predicting  $\text{NO}_3^-$ .  
7 Simulations of  $\text{NO}_3^-$  in the literature also showed significant overestimations (Zakoura and Pandis, 2018;  
8 Travis et al., 2022).

9 Model performance met the benchmarks in reproducing  $\text{PM}_{2.5}$  mass concentrations in both Beijing  
10 downtown ( $R = 0.49$ ,  $p < 0.05$ , MFB = 1 %, MFE = 49 %, as shown in Fig. S2c) and Beijing suburb ( $R$   
11 = 0.42,  $p < 0.05$ , MFB = -32 %, MFE = 69 %, as shown in Fig. S2d) during the frequent-NPF period  
12 (unshaded area in Fig. S2a–b). Notably, the model also met the goal values in simulating  $\text{PM}_{2.5}$  mass  
13 concentrations prior to June 29, with an  $R = 0.58$  and  $p < 0.05$  as well as MFB and MFE of -14 % and  
14 26 %, respectively, in Beijing downtown, and in suburban Beijing an R value of 0.54 with MFB and  
15 MFE values of -11 % and 33 % respectively. Overall, the model demonstrated performance that met the  
16 benchmarks or even met the goal values for simulating the variables of interest during June 29–July 6,  
17 except for  $\text{NO}_3^-$ . As such, we will focus our result analysis and discussion on the frequent-NPF period of  
18 better simulation performance.

## 19 3 Results and discussion

### 20 3.1 Overview of modeling NPF events

21 From June 29 to July 6, 2019, there were five NPF events, June 29 and 30, and July 1, 3, and 6, as  
22 shown in Fig. 4a–b. The high frequency of NPF events was associated with clean air masses from the  
23 north, and was favored by dry and sunny conditions, which is consistent with previous literature (Wu et al.,  
24 2007; Chu et al., 2021; Ma et al., 2021). During the NPF events on July 1 and 3, a typical banana-  
25 shaped growth pattern was observed, with the maximum median mode diameter of newly formed  
26 particles reaching above 60 nm and around 50 nm, respectively. The NPF events on June 29 and 30 also  
27 experienced rapid new particle growth during the initial 2–3 hours, but the maximum median mode  
28 diameters of the newly formed particles were smaller than 30 nm. Similarly, during the NPF event of  
29 July 6, the maximum median mode diameter of the newly formed particles was smaller than 30 nm before  
30 they disappeared from the observations. The details on the growth patterns can be found in the companion  
31 paper (Wei et al., 2023).

32 The modeling results effectively captured the occurrence characteristics of five NPF events,  
33 including their initial occurrence time and duration. Moreover, the model partially captured the growth  
34 characteristics of new particles while the low size resolution of modeled results cannot allow a  
35 quantitative comparison of the growth parameters. However, the model failed to predict the decreased  
36 median diameter of new particles at nighttime on June 29 and 30, because the related mechanisms were  
37 not included in the model and were poorly understood (Yao et al., 2010; Skrabalova et al., 2015; Alonso-



1 Blanco et al., 2017; Kamra et al., 2022). Furthermore, the modeling results performed to meet the  
2 benchmarks for reproducing the PNCs plumes occurring on the nights of July 2-3 and 4-5. However, the  
3 model overestimated PNCs plumes on July 5, which was consistent with the overestimation of PM<sub>2.5</sub>,  
4 SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup>, NH<sub>4</sub><sup>+</sup>, and organics, as mentioned above. This overestimation of PNCs was previously  
5 reported by Matsui et al. (2011) who argued that it was due to the underestimation of vertical mixing  
6 capacity at night and excessive ground chemical concentrations in the model. Similar arguments were  
7 also reported by Mckeen et al. (2007) and Matsui et al. (2009). More discussion of PNCs in the vertical  
8 can be found in Section 3.3.

9 When considering NPF and non-NPF days separately, the simulated N<sub>ccn</sub> at SS = 0.2 % met the goal  
10 values on non-NPF days, with R = 0.38, p < 0.05, MFB = 19 % and MFE = 48 % (Fig. 5a). However, on  
11 NPF days, the model substantially underestimated N<sub>ccn</sub> at SS = 0.2 % with R = 0.50, p < 0.05 (Fig. 5b).  
12 At SS = 0.4 %, the model performed to meet the benchmarks for reproducing N<sub>ccn</sub> on NPF days, with R  
13 = 0.51, p < 0.05, MFB = -46 % and MFE = 74 % (Fig. 5d). However, it substantially overestimated N<sub>ccn</sub>  
14 at SS = 0.4 % relative to the observations on non-NPF days (Fig. 5c). The underestimation or  
15 overestimation was determined not only by the estimated PNCs with sizes larger than 60–120 nm but  
16 also by the Kappa values. We will delve into this further later on.

17 Based on the reasonable model predictions of NPF events from June 29 to July 6, a comprehensive  
18 analysis of three distinct NPF events will be conducted in section 3.2–3.5. This analysis will include a  
19 detailed examination of the chemical drivers at ground level and their vertical profiles, 3-D growth  
20 patterns of new particles, contributions of grown new particles to CCN, and other relevant factors.

### 21 3.2 Chemical drivers to grow newly formed particles and subsequently contribute to CCN at the 22 ground level

23 Figure 6a–d presents time series data for modeled chemical components of particles within two size  
24 ranges (10–40 nm and 40–250 nm) on July 1–2. During daytime, ORG were found to be the dominant  
25 contributor to growth of 10–40 nm particles followed by (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>. In contrast, NH<sub>4</sub>NO<sub>3</sub> was the most  
26 significant chemical component driving growth in 40–250 nm particles during nighttime. An analysis of  
27 kappa presented later in this section confirms that this could be an overestimated effect. To determine the  
28 relative contribution of each species as examples, their fractions within particles within each size range  
29 were calculated at two time points: 15:00 on July 1 (Fig. 6e–f) and 03:00 on July 2 (Fig. 6g–h). To account  
30 for differences in the contributions of secondary organics (SOA) and primary organics (POA) to new  
31 particle growth in the model results, SOA and POA were analyzed separately. In this study, SOA was  
32 defined as the sum of secondarily generated particulate organics from anthropogenic and biogenic  
33 precursors, while POA was the sum of primary organics.

34 At 15:00 on July 1, the simulated SOA contributed 56 % of the total mass of particles ranging from  
35 10–40 nm, while (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> accounted for 36 %, and POA made up 8 % of the mass. The high mass  
36 fraction of (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> in 10–40 nm particles at 15:00 resulted in corresponding high hygroscopicity  
37 parameters ( $\kappa$ ) of up to 0.24. The model predicted that the fraction of (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> in 40–250 nm particles  
38 was larger than that in 10–40 nm particles at 15:00, with (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> contributing 49 % of the mass,  
39 followed by 44 % for SOA and 7 % for POA. This resulted in a corresponding  $\kappa$  value of up to 0.30.

1 However, the estimated  $\kappa$  values based on observations at SS=1.0 %, 0.4 %, and 0.2 % were only 0.08,  
2 0.08, and 0.16, respectively, at the same time, as reported in the companion paper. At 08:00, the model  
3 predicted that the concentration of H<sub>2</sub>SO<sub>4</sub> vapor was approximately 10<sup>8</sup> molecules cm<sup>-3</sup> (Fig. S3), which  
4 was substantially higher than previous observations in Beijing, where the maximum concentration was  
5 around 10<sup>7</sup> molecules cm<sup>-3</sup> (Wang et al., 2011; Lu et al., 2019). Other modeling studies (Matsui et al.,  
6 2011; 2013) have also reported an overestimation of H<sub>2</sub>SO<sub>4</sub> vapor similar to ours. In addition, **gas-to-**  
7 **particle condensation overwhelmingly** contributed to POA in 10–40 nm particles, which also aided in the  
8 growth of newly formed particles. According to our simulated mass fractions of 40–250 nm particles,  
9 **which are 68% in NH<sub>4</sub>NO<sub>3</sub> and (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 25% in SOA, and 7% in POA, led to the model-based  $\kappa$  value**  
10 **being estimated at 0.37.** However, at 03:00 on July 2, the observation-based estimated  $\kappa$  values at  
11 SS=1.0 %, 0.4 %, and 0.2 % were 0.10, 0.13, and 0.28, respectively, as reported in our companion paper.  
12 In this case, overestimation of NH<sub>4</sub>NO<sub>3</sub> is likely to have contributed to the overestimation of  $\kappa$  (Fig. S1b–  
13 c).

14 Similar to the NPF event on July 1–2, the modeling results on July 3–4 indicated that SOA was the  
15 dominant driver of new particle growth. Overestimation of H<sub>2</sub>SO<sub>4</sub> vapor **during daytime** resulted in  
16 overestimated  $\kappa$  values for particles ranging from 10–40 nm and 40–250 nm (Fig. S4 a–h). However, the  
17 model did not predict the formation of NH<sub>4</sub>NO<sub>3</sub> before 24:00 on July 3, which was consistent with  
18 observation-based estimated  $\kappa$  values of less than 0.1 at SS levels of 0.4 % or higher.

19 On July 6, the model predicted that new particle growth was dominantly driven by SOA, with higher  
20 contributions than those observed on July 1–2 and July 3 (Fig. S5a–j). Unlike the cases on July 1–2 and  
21 July 3, the  $\kappa$  values derived from the modeled mass fractions of 10–40 nm and 40–250 nm particles were  
22 reasonably consistent with the observation-based  $\kappa$  values on July 6. This suggests that there was no  
23 detectable evidence for overestimated H<sub>2</sub>SO<sub>4</sub> vapor condensation during daytime on July 6. Additionally,  
24 the model did not predict the formation of NH<sub>4</sub>NO<sub>3</sub> before 24:00 at nighttime.

25 Fig S6a–c presents the comparison of CCN **simulations with observations for 0.2 % SS** during the  
26 NPF events on July 1, 3, and 6, respectively. The comparison shows that the simulated N<sub>ccn</sub> at 0.2 % SS  
27 were clearly underestimated by several folds. This underestimation was mainly due to the underestimates  
28 of number concentrations of preexisting particles >100 nm, as the  $\kappa$  values of particles at different sizes  
29 during the NPF events had been overestimated to some extent. However, the model performed to meet  
30 the benchmark **for** reproducing N<sub>ccn</sub> at 0.4 % SS during the NPF events on July 1 and 3 (Fig S6d–e). In  
31 these cases, the overestimation of number concentrations of grown new particles and their  $\kappa$  values  
32 probably canceled out the effect of the underestimated preexisting particles (>100 nm). However, this  
33 was not the case for the NPF event on July 6 (Fig S6f), when the grown new particles were too small to  
34 be activated as CCN. On that day, the N<sub>ccn</sub> at 0.2 % SS were still underestimated to some extent.

### 35 **3.3 Chemical drivers to grow newly formed particles **in the vertical** direction**

36 Based on the fact that the model reproduced CN<sub>10–40</sub> and organic drivers, that lead to the growth of  
37 **new particles** at the ground level, **that met** the benchmark during June 29–July 6, this section delves  
38 deeper into the chemical drivers at different heights during three selected NPF events. Figure 7a–d shows  
39 the simulated chemical composition of 10–40 nm particles at three different heights (500 m, 1500 m, and



1 2500 m) above the observation site. These heights represent the lower part of the planetary boundary  
2 layer (PBL), the upper part of the PBL in the morning during the initial occurrence of NPF, and the top  
3 of the diurnal peak PBL on July 1–2, respectively. The results indicate that SOA dominated the growth  
4 of 10–40 nm particles at 500 m and 1500 m at 10:00, 15:00, and 22:00 on July 1. In contrast, inorganic  
5 species were found to control growth at 2500 m during the same time, i.e., the ammoniated sulfuric acid  
6 likely acted as the dominant driver at 10:00, while the dominant driver switched to  $\text{NH}_4\text{NO}_3$  at other  
7 times. This height-dependence of chemical drivers is consistent with previous findings in the literature,  
8 which attribute it to the low abundance of volatile organic compounds in the free troposphere (Sanchez  
9 et al., 2018; Williamson et al., 2019). On July 2 at 03:00,  $\text{NH}_4\text{NO}_3$  acted as the dominant driver at all  
10 heights. However, as mentioned earlier, the ammoniated sulfuric acid and  $\text{NH}_4\text{NO}_3$  in 10–40 nm particles  
11 may have been overestimated to some extent during daytime and nighttime at the ground level. The same  
12 overestimation could also occur at different heights, underscoring the urgent need for vertical  
13 observations of chemical composition in 10–40 nm particles.

14 On July 3, the simulation showed a similar height-dependence of chemical drivers for the growth of  
15 newly formed particles (see Fig. S7). However, the model did not predict any  $\text{NH}_4\text{NO}_3$  at a height of 500  
16 m before the new particle signal disappeared. On July 6, the model predicted that a combination of SOA  
17 and POA contributed to approximately 80–90 % of the  $\text{CN}_{10-40}$  mass concentrations at a height of 500 m  
18 during the event, as shown in Fig. S8. However, these percentages decreased with increasing height,  
19 dropping to approximately 35–60 % at a height of 2500 m. This indicates that inorganic species also  
20 played a significant role in the growth of newly formed particles, with an even greater contribution at  
21 higher altitudes.

### 22 3.4 3-D occurrence of NPF events and transport of newly formed particles

23 To investigate the 3-D evolution of NPF events, we utilized the  $\text{CN}_{10}$  (summed number  
24 concentrations of particles with diameter less than 10 nm) instead of  $\text{CN}_{10-40}$ . The simulated 3-D  
25 evolution of  $\text{CN}_{10}$  on July 1 is presented in Fig. 8a–b. The maximum  $\text{CN}_{10}$  value of approximately 8,000  
26  $\text{cm}^{-3}$  was predicted over the mountain station at around 1300 m a.s.l., starting from 08:00 on July 1. At  
27 this point, the PBL had risen to approximately 1260 m a.s.l. As reported in the literature, NPF events  
28 tend to occur initially in the residual layer, which was formed above the PBL at night, due to high  
29 oxidation capacity, low condensation sink, and abundant precursors (Stratmann et al., 2003; Wehner et  
30 al., 2010; Quan et al., 2017; Qi et al., 2019; Tröstl et al., 2016). From 08:00 to 09:00 on July 1, NPF  
31 rapidly extended to the ground level, leading to a sharp increase in  $\text{CN}_{10}$  at that altitude (Fig. 8b). The  
32 NPF event reached its maximum concentration at the ground level between 10:00–11:00 and  
33 subsequently weakened.

34 When examining the occurrence of NPF in the horizontal at an altitude of approximately 1300 m  
35 a.s.l., a significant spatial inhomogeneity was predicted across the NCP. Specifically, the simulated  $\text{CN}_{10}$   
36 revealed the presence of two stronger NPF regions located roughly 100–300 km away from the  
37 observation site at 08:00 (see Fig. 8a). From 09:00 to 12:00, these two stronger regions continued to  
38 expand and eventually connected with each other, forming a large zone approximately 270 km×135 km  
39 in size, located at 40.5–43 °N and 115–116.5 °E. When comparing the vertical distributions of the

1 mountain station and points A and B (which represent the high-value areas of the two strong NPF regions)  
2 from 10:00 to 12:00, it was found that the simulated  $CN_{10}$  across the stronger NPF zone were  
3 approximately 3–4 times larger than those observed over most of the weaker NPF zones. However, the  
4 simulation also showed that there was no time lag for the occurrence of NPF, whether it was at the priority  
5 nucleation height of approximately 1300 m a.s.l. or at the ground over the NCP, as shown in Fig. 8a.

6 Similar to the event on July 1, NPF also occurred widely over the NCP on July 3. However, the  
7 stronger NPF zone was situated far from the observation site (refer to Fig. S9). On July 6, NPF occurred  
8 over most parts of the NCP, as shown in Fig. S10. However, the areas of NPF occurrence were noticeably  
9 smaller compared to those on July 1 and 3, which could explain the shorter duration of NPF observed on  
10 July 6. Furthermore, the stronger NPF zone was located north of the observation site, and the strong  
11 northeast wind blew the new particle signal away from the observation site in the period.

### 12 3.5 What happened for grown new particles after the particles disappear from observations?

13 The total number concentration of particles with sizes between 40 and 250 nm ( $CN_{40-250}$ ) was used  
14 to characterize new particles that grew too large to observe. Figure 9a–b depicts the horizontal  
15 distribution of  $CN_{40-250}$  at ground level from 18:00 on July 1 to 07:00 on July 2 and the corresponding  
16 vertical profiles of  $CN_{40-250}$  over the observation zone and two stronger NPF zones. The simulated wind  
17 direction over the observation zone changed from northwest to southwest at 18:00 on July 1, coinciding  
18 with the decrease in observed new particle concentrations (as shown in Fig. 4a and Fig. 9a), due to weaker  
19 NPF events in the southwest direction. At that time, strong plumes were predicted in the southwest  
20 direction over a large area. By 24:00 on July 1, the modeling results indicated that these plumes had  
21 approached the observation zone. This intrusion likely led to an increase in both simulated and observed  
22  $CN_{40-250}$  from 24:00 on July 1 to 04:00 on July 2 (as shown in Fig. 4a), with the plume particle  
23 concentration eventually overwhelming the new particle concentration. The modeling results suggested  
24 that the new particles were mostly moved out of the observation zone. Consequently, the question of  
25 whether grown new particles can experience additional growth to become CCN was replaced by a new  
26 question: whether <20–50 nm preexisting particles, mainly composed of organics, can grow to become  
27 CCN. In this study, preexisting particle growth only occurred on July 5, as presented in the companion  
28 paper. Unfortunately, the model poorly reproduced the observations on July 5. However, the occurrence  
29 frequency of preexisting particle growth was much less than that of NPF events on the basis of  
30 observations in this study alone.

31 Similar to what occurred on July 1, the new particle concentration was also significantly diluted to  
32 a low level, and by July 3, the new particles had vanished over the observation zone (see Fig. S11).  
33 However, this was not the case on July 6, as demonstrated in Fig. 10a–b. On that day, the strong northeast  
34 wind carried the new particles out of the observation zone, rather than diluting them to normal  
35 levels prior to the event. It is still expected that the new particle concentrations will eventually be diluted  
36 to the ambient concentrations. Nevertheless, the model results need to be confirmed with Lagrangian  
37 observations that track moving air masses.

#### 4. Conclusion and uncertainties

To quantitatively understand 3-D evolution of NPF events and impacts on CCN, we used a 20-bin WRF-Chem model to simulate NPF events in the NCP during a three-week observational period in the summer of 2019. The model was able to reproduce the observations during June 29–July 6, which was characterized by a high frequency of NPF occurrence. Specifically, the model reproduced  $CN_{10-40}$ ,  $N_{ccn}$  at 0.4 % SS, mass concentrations of  $PM_{2.5}$ , mass concentrations of  $SO_4^{2-}$  in  $PM_{1.0}$  and TSP, ORG and  $NH_4^+$  in  $PM_{1.0}$ , and other variables, with the performance meeting the benchmark. However, the model consistently overestimated daytime  $H_2SO_4$  vapor by approximately one order of magnitude and frequently overestimated nighttime formation of  $NH_4NO_3$ . These overestimations led to an overestimation of the  $\kappa$  values of both grown new particles and pre-existing particles to some extent. The model also poorly reproduced most of the observed variables during the remaining two weeks, and we have yet to explain this poor simulation. Our modeling results indicated that the growth of newly formed particles from 10 nm to larger sizes was overwhelmingly determined by SOA, which is consistent with previous modeling studies in the literature. This implies that the critical challenge in modeling contributions of NPF events to CCN may be accurately reproducing those inorganic species, accounting for a small but appreciable fraction, rather than SOA.

The results of 3-D simulations of NPF events over the NCP, based on case studies, showed that NPF events occurred preferentially at the top of the PBL and then expanded vertically. In the horizontal direction, the NPF was predicted in a large regional scale with the stronger NPF zone located northeast of the observation site. The modeling results also suggested that SOA played a dominant role in controlling the growth of newly formed particles in the PBL. However, inorganic species likely replaced SOA as the dominant driver above the PBL. Additional observations are needed to confirm these findings.

The model performed to meet the goal values in reproducing CCN at SS = 0.2 % on non-NPF days, but it clearly overestimated CCN at SS = 0.4 % on those days. Conversely, the model performed to meet the benchmark in reproducing CCN at SS = 0.4 % on NPF days, but it noticeably underestimated CCN at SS = 0.2 %. This presents a significant challenge that must be urgently addressed, as it has a major impact on the accuracy of predicted contributions of NPF events to CCN budgets. Additionally, the disappearance of new particles from observations may simply be due to dilution effects or the movement of the particles elsewhere. In such cases, the issue of how newly formed particles grow into CCN becomes another important question: specifically, how do pre-existing particles with an organic-dominant composition of <20–50 nm grow into CCN?

The model poorly predicted the PNCs and CCN during the periods with low occurrence frequency of NPF events because of lack of local particle number size distributions (PNSDs) from various sources in China. The weakness may restrict the model application on various aspects and a localized PNSDs adaptor in China is urgently needed to be developed.

**Data availability.** The data of this paper are available upon contact with the authors, Yang Gao (yanggao@ouc.edu.cn), Xiaohong Yao (xyhao@ouc.edu.cn) and Ming Chu (cm5594@stu.ouc.edu.cn).

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5

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7

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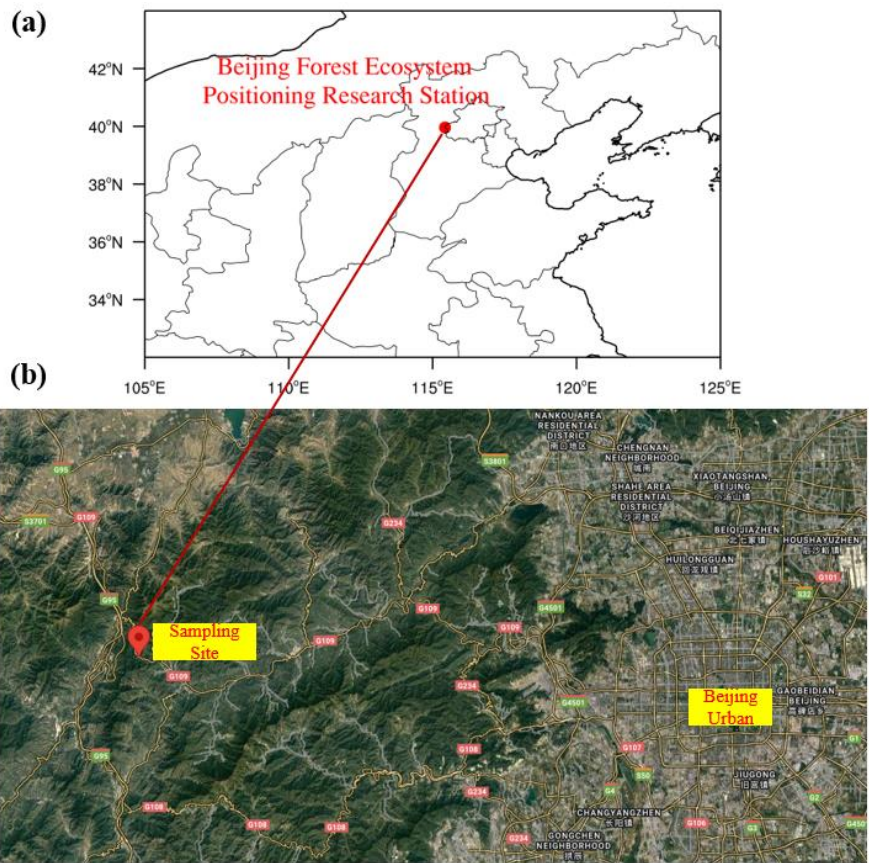


Figure 1. Map of sampling site (a) and 3-D view of sampling site (b) in summer (download from <https://www.google.com/maps/>).

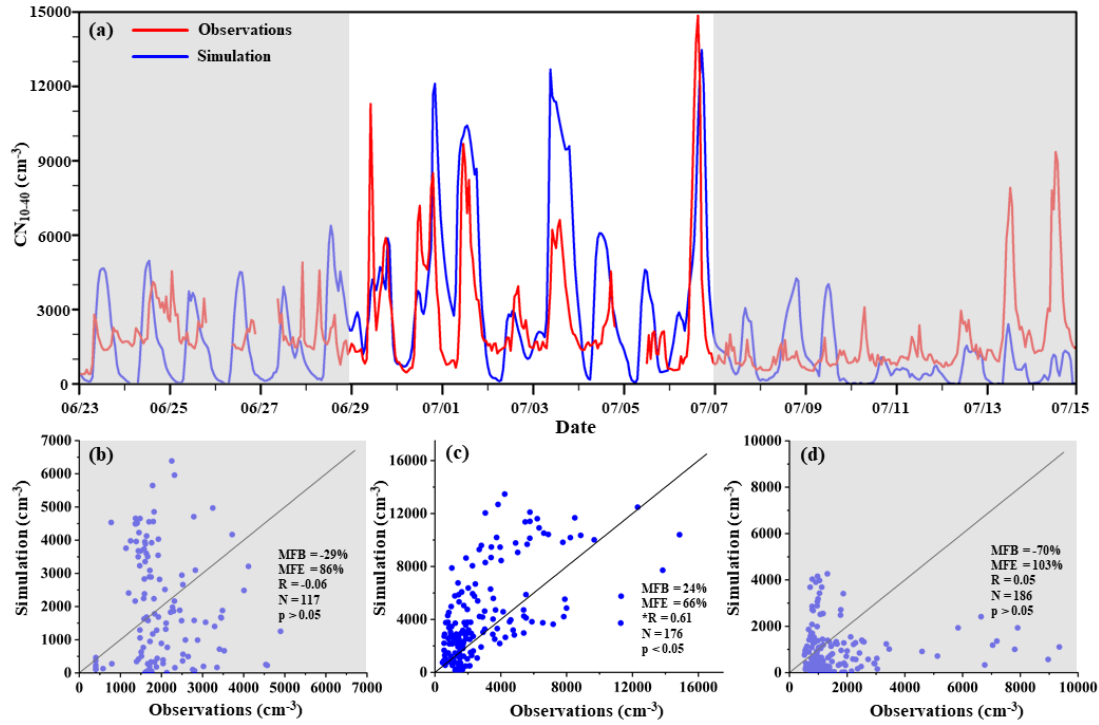


Figure 2. Time series of observed and modeled  $CN_{10-40}$  from June 23 to July 14 (a), the comparison of the modeled  $CN_{10-40}$  with the observations in June 23–28 (b), in June 29–July 6 (c) and in July 7–July 14 (d) (a: local time is used and the harsh marks with the data represent the beginning of each day at 00:00, and the same is applicable for all time plots presented later).

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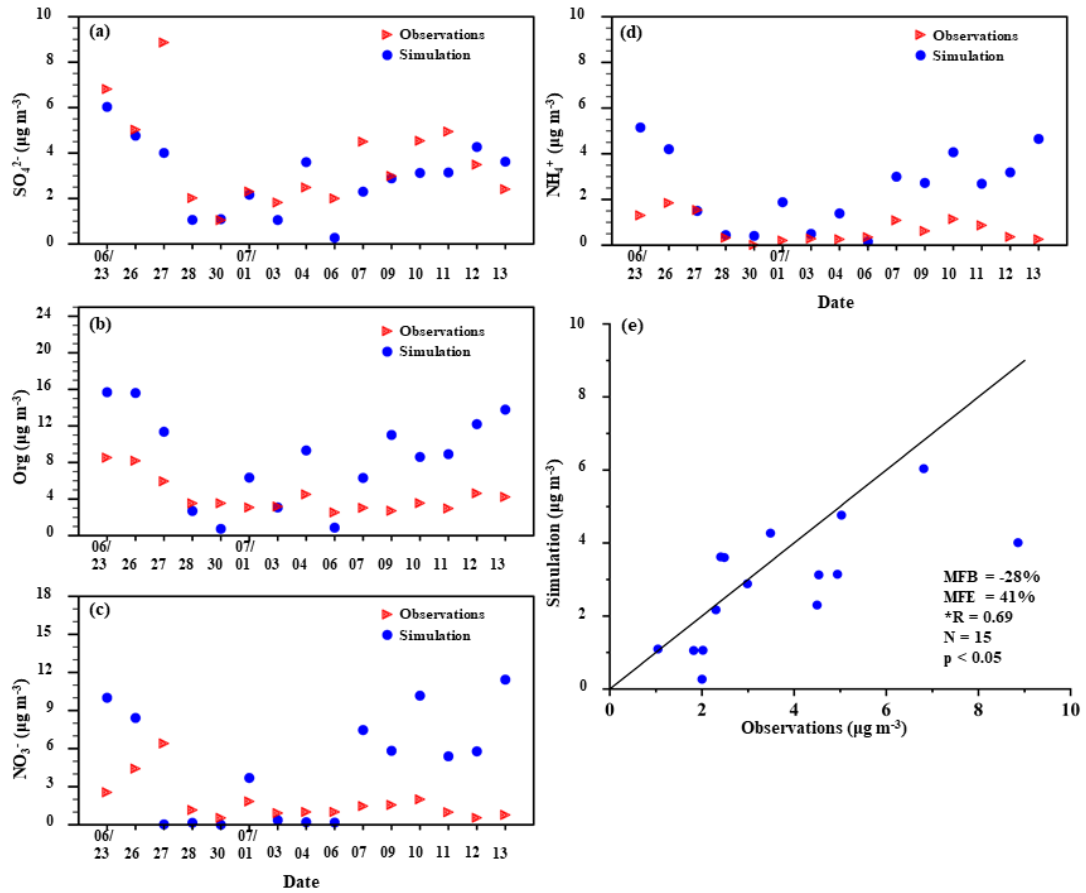


Figure 3. Time series of observed and modeled mass concentrations of  $\text{SO}_4^{2-}$  (a), organics (b),  $\text{NO}_3^-$  (c) and  $\text{NH}_4^+$  (d), and comparison of the modeled and observed  $\text{SO}_4^{2-}$  (e) in June 23–July 14.

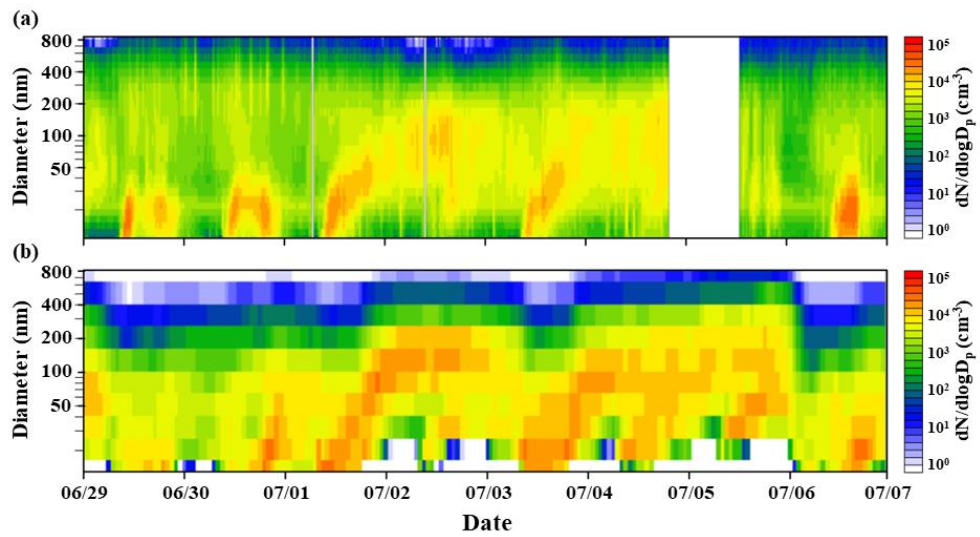


Figure 4. Contour plot of PNSDs from observations (a) and modeling (b) from June 29 to July 6, 2019.

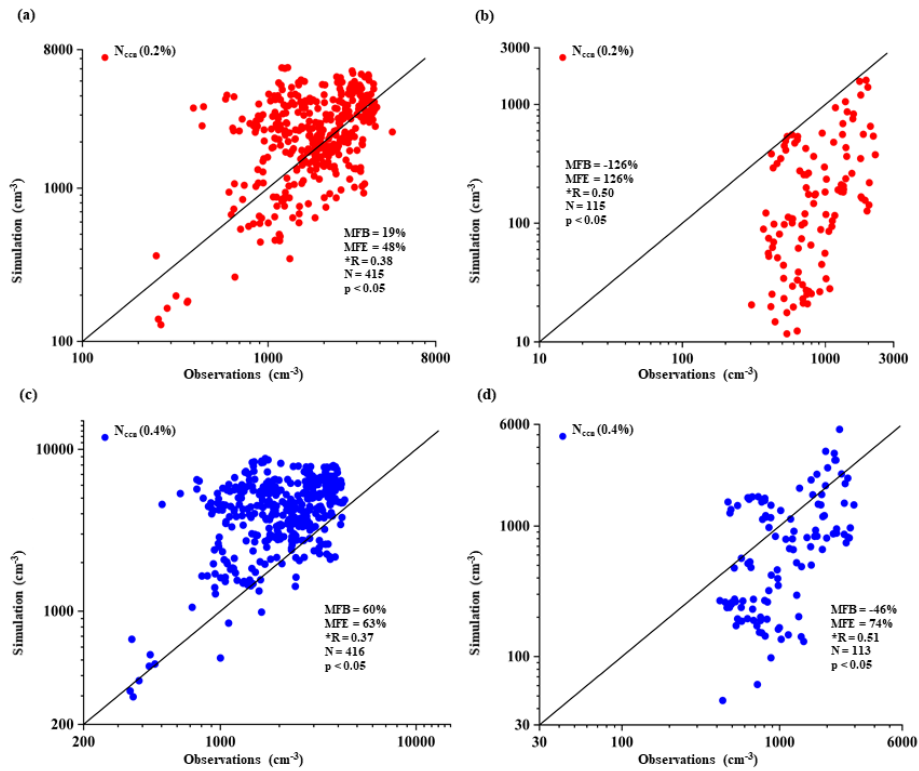


Figure 5. The simulated  $N_{ccn}$  against the observations at 0.2 % SS (a–b) and 0.4 % SS (c–d) on non-NPF days (a and c) and NPF days (b and d), respectively, during the frequent-NPF period.

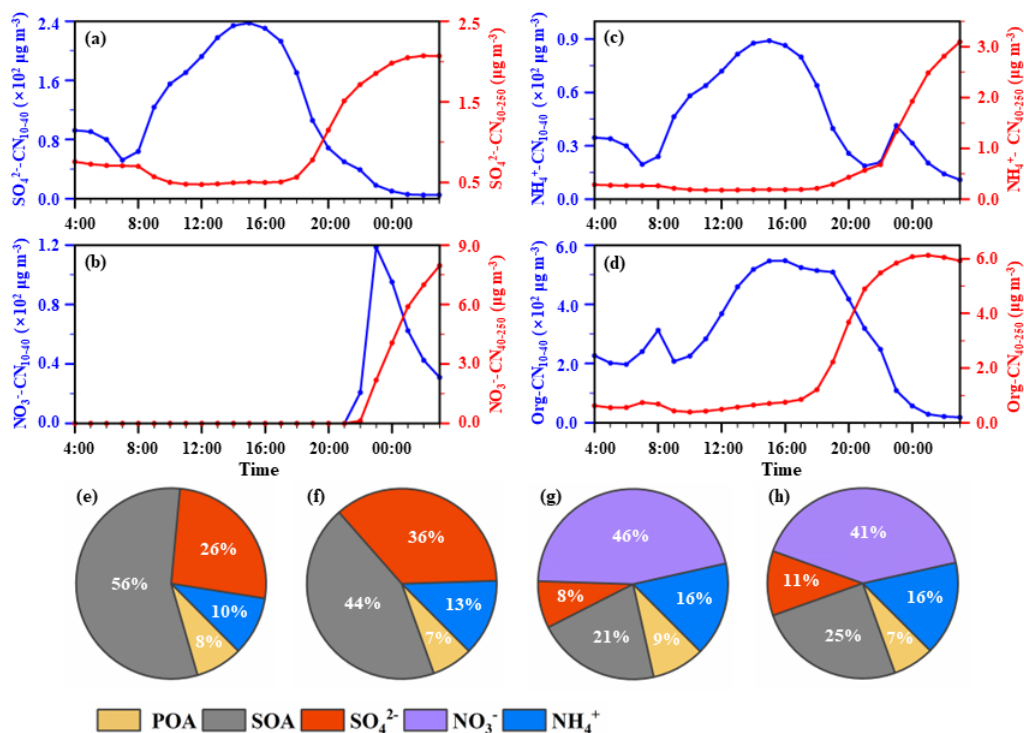


Figure 6. Diurnal variations in modeled chemical components in 10–40 nm particles and 40–250 nm particles:  $\text{SO}_4^{2-}$  (a),  $\text{NO}_3^-$  (b),  $\text{NH}_4^+$  (c), organics (d) on July 1–2; fractions of chemical species in 10–40 nm particles (e) and 40–250 nm particles (f) at 15:00 on July 1 and those at 3:00 on July 2 (g and h). POA and SOA stand for primary organics and secondary organics, respectively.

5



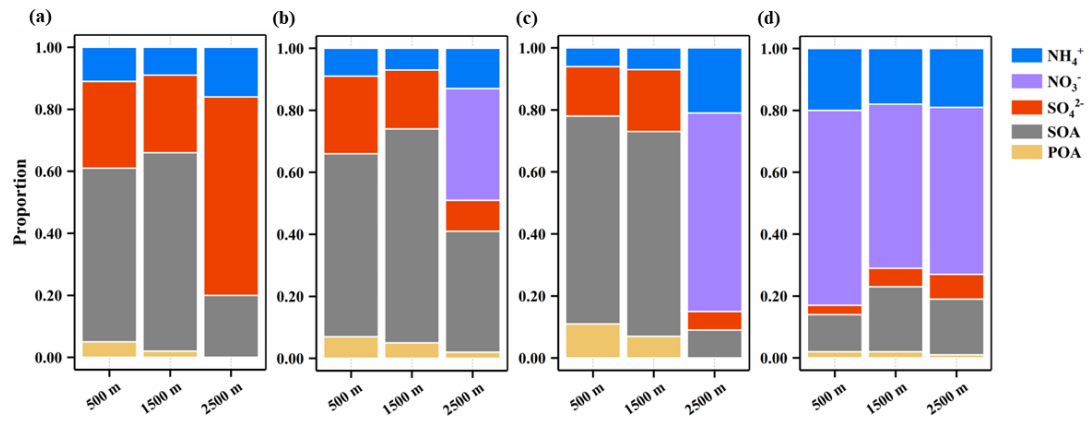
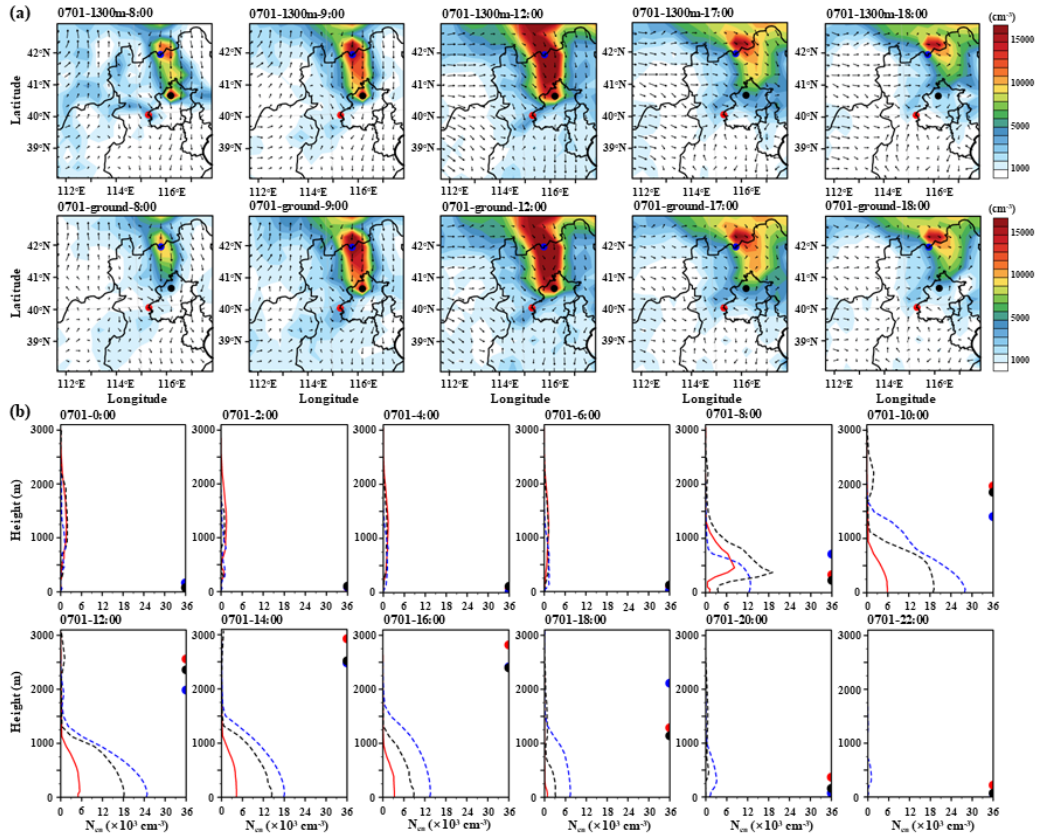
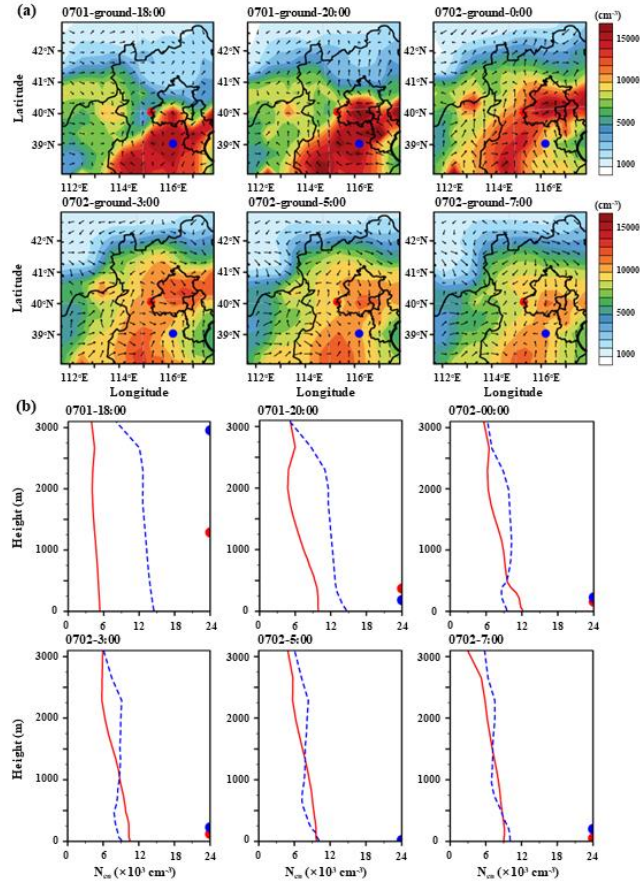


Figure 7. The simulated chemical components in 10–40 nm particles at 500 m, 1500 m and 2500 m above the ground respectively at 10:00 (a), 15:00 (b), 22:00 (c) on July 1 and 3:00 (d) on July 2.

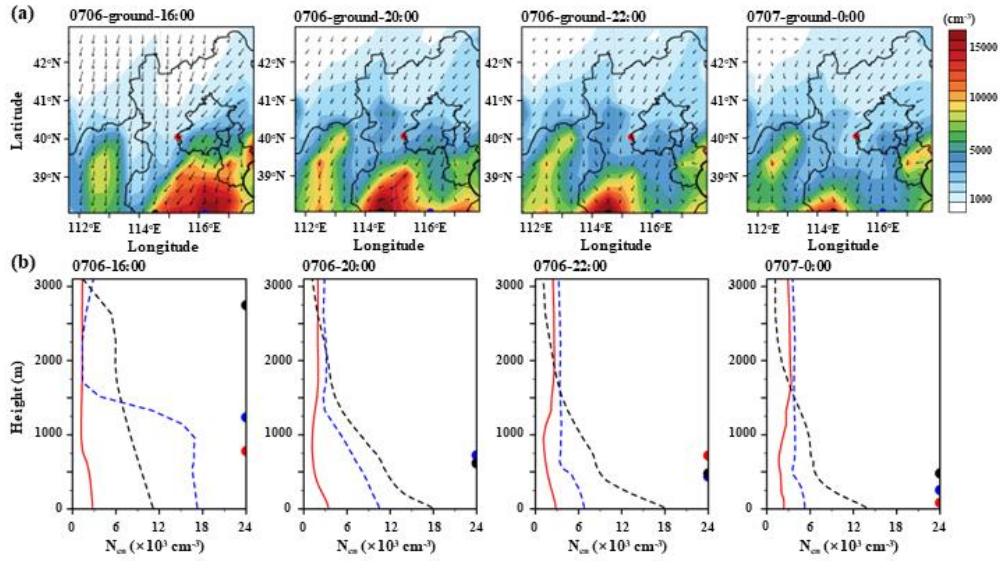


1

2 **Figure 8. Horizontal distribution of CN<sub>10</sub> at ~1300 m a.s.l. (a, the upper row) and on the ground level (a, the**  
 3 **bottom row) at 8:00, 9:00, 12:00, 17:00 and 18:00 on July 1, 2019 (the red, blue and black solid dots represent**  
 4 **the observation site, two centers of strong NPF zones (point A and point B), respectively; the direction and**  
 5 **length of the black arrows represent the wind direction and wind speed, respectively); Vertical profiles of CN<sub>10</sub>**  
 6 **above the observational site (red solid line), point A (blue dashed line) and point B (black dashed line) from**  
 7 **0:00 to 22:00 on July 1, 2019 (b, the Y-axis coordinate is the height above the ground; the red, blue and black**  
 8 **solid dots represent the height of the PBL over the observational site, point A and point B, and PBL exceeding**  
 9 **3000 meters above the ground are not shown in Figure).**



5 **Figure 9. Horizontal distribution of CN<sub>40-250</sub> on ground (a, the upper row) and vertical profiles of CN<sub>40-250</sub> over the observational site (red solid line) and point A (blue dashed line) from 18:00 on July 1 to 07:00 on July 2 (b, the Y-axis coordinate is the height above the ground; the red and blue solid dots represent the height of the PBL over the observational site and point A, and PBL exceeding 3000 meters above the ground are not shown in Figure).**



5 **Figure 10.** Horizontal distribution of  $CN_{40-250}$  on ground (a, the upper row) and vertical profile of  $CN_{40-250}$  in observation site (red solid line), point A (blue dashed line) and point B (black dashed line) in the NPF event occurred on July 6 from 16:00 on July 6 to 00:00 on July 7 (b, the Y-axis coordinate is the height above the ground; the red, blue and black solid dots represent the height of the PBL in observation site, point A and point B, and PBL exceeding 3000 meters above the ground are not shown in the figure).