Substantially positive contributions of new particle formation to Cloud Condensation Nuclei under low supersaturation in China based on numerical model improvements

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Abstract

New particle formation (NPF) and subsequent particle growth are important sources of condensation nuclei (CN) and cloud condensation nuclei (CCN). While a number of observations have shown positive contributions of NPF to CCN at low supersaturation, negative NPF contributions were often simulated. Using the observations in a typical coastal city of Qingdao, we thoroughly evaluate the simulated number concentrations of CN and CCN using a NPF-explicit parameterization embedded in WRF-Chem model. In terms of CN, the initial simulation shows large biases of particle number concentrations at 10–40 nm (CN_{10-40}) and 40–100 nm (CN_{40-100}). By adjusting the process of gas-particle partitioning, including mass accommodation coefficient of sulfuric acid, the phase changes of primary organic aerosol emissions and the condensational amount of nitric acid, the concomitant improvement of the particle growth process yields a substantial reduction of overestimates of CN_{10-40} and CN_{40-100}.

Regarding CCN, SOA formed from the oxidation of semi-volatile and intermediate volatility organic vapors (SI-SOA) yield is an important contributor. In the original WRF-Chem model with 20 size bins setting, the yield of SI-SOA is too high without considering the differences in oxidation rates of the precursors. Lowering the SI-SOA yield results in much improved simulations of the observed CCN concentrations. On the basis of the bias-corrected model, we find substantial positive contributions of NPF to CCN at low supersaturation (~0.2%) in Qingdao and over the broad areas of China, primarily due to the competing effects of increasing particle hygroscopicity surpassing that of particle size decrease. This study highlights the potentially much larger NPF contributions to CCN on a regional and even global basis.
1. Introduction

New particle formation (NPF) is a process in which gaseous vapors nucleate and form critical molecular clusters, followed by subsequent growth to larger sizes through condensation and coagulation (Kulmala et al., 2004; Kulmala et al., 2013; Lee et al., 2019). Newly formed particles could effectively grow into the size of cloud condensation nuclei (CCN) under certain supersaturation (SS), which exerts an impact on the cloud microphysical process and global radiation balance (Merikanto et al., 2009; Gordon et al., 2017; Kerminen et al., 2018; Ren et al., 2021). In addition, the high-efficiency nucleation and explosive growth of particles may contribute to the formation of haze (Guo et al., 2014), affecting air quality and human health (Yuan et al., 2015; Chu et al., 2019; Kulmala et al., 2021).

The overestimate of condensation nuclei (CN) in numerical models are commonly seen, despite the attempt to correctify the bias (Matsui et al., 2013; Arghavani et al., 2022). It is a common way to reduce the nucleation rate which may reduce the particle number concentration in proportion (Matsui et al., 2013). For instance, in the study of NPF in East Asia in the spring of 2009, even after lowering the nucleation rate in a regional model of WRF-Chem applied in their study, the reduced number concentration of particles at 10–130 nm remained to be overestimated (Matsui et al., 2013). Using the same regional model and a similar method to reduce the nucleation rate, Arghavani et al. (2022) found particle number concentration at 10–100 nm was still overestimated by nearly one order of magnitude, despite the effectiveness to reduce the overestimates for the smaller particles such as 2.5–10 nm. In addition to the rate of NPF, the growth process of particles also has a crucial effect on particle number concentration and size distribution. In this process, the condensation of some chemical species such as sulfuric acid, nitrate and organic gases on particles plays a major role in particle growth (Yao et al., 2018; Lee et al., 2019; Li et al., 2022), and the uncertainty of their condensation amount may lead to the bias of CN simulation.

In addition to CN, there are large discrepancies in the predicted CCN between the numerical models and observational results. Furthermore, as an important source of CCN (Merikanto et al., 2009), the contribution of nucleation to CCN quantified by
numerical models is also highly uncertain. For example, in terms of predicting CCN, Fanourgakis et al. (2019) evaluated the CCN concentrations simulated by 16 global aerosol–climate and chemistry transport models with observations at 9 sites in Europe and Japan from 2011 to 2015, and found that all models underestimated CCN concentrations with a mean normalized mean bias (NMB) of -36% at low supersaturation (SS=0.2%). Models also tend to underestimate the contribution of NPF on CCN, especially at low supersaturation. The continuous observation of CCN concentrations throughout the year (July 2008–June 2009) carried out in Hyytiälä, Finland, showed that under low SS, nucleation enhanced the CCN by 106% and 110% at SS=0.1% and 0.2% respectively (Sihto et al., 2011). Observations acquired in Beijing from July 12 to September 25, 2008, also suggested that nucleation significantly increases CCN at all supersaturations, even when supersaturation is low (i.e., 0.07% and 0.26%). Thus, the occurrence of NPF enhanced CCN by a factor of 1.7 and 2.2, respectively (Yue et al., 2011).

However, previous numerical experiments behave oppositely. For instance, Matsui et al. (2011) quantified the contribution of nucleation to CCN using WRF-chem in Beijing in August and September 2006 and found reduced CCN under low SS, e.g., when SS=0.02%, the concentration of CCN is reduced by up to ~50%. They attributed this to the fact that the small particles produced by nucleation may inhibit the growth of the preexisting particles (Matsui et al., 2011). Similarly, Dong et al. (2019) conducted NPF simulations with the WRF-Chem for the summer of 2008 focusing on the Midwest of the United States, and found that the nucleation resulted in decreased CCN at low supersaturation (SS=0.1%). Besides, a study carried out for East Asia in 2009 also indicated that at low supersaturation (e.g. SS=0.1%), nucleation has little impact on CCN (Matsui et al., 2013). The contrasting effects of nucleation on CCN at low supersaturations in model and observations is not explained in these previous studies.

At the stage of particle growth, secondary organic aerosol (SOA) formed by atmospheric oxidation of organic vapors is a major contributor to particle growth to CCN-related sizes (Liu and Matsui, 2022; Qiao et al., 2021). SOA formed by multi-generational gas-phase oxidation of semi-volatile and intermediate volatility organic
compounds (S/IVOC) is called SI-SOA (Jimenez et al., 2009; Zhang et al., 2007). Zhao et al. (2016) made a comprehensive assessment of the roles of various SOA precursors in SOA formation in real atmosphere in China in 2010, and the results demonstrated that evaporated POA and IVOC (i.e. S/IVOC) made a significant contribution to SOA, contributing up to 82% to the average SOA concentration in eastern China. However, the effect of SI-SOA on CCN has not been fully studied.

In this paper, WRF-Chem was applied to simulate the effect of the NPF on CCN in Qingdao in February 2017. The simulated results from the WRF-Chem model are firstly compared with observations in Qingdao, exhibiting large biases in CN. This is followed by an improvement through a few processes. At the end, the impact of SI-SOA yield and nucleation on CCN is investigated.

2. Data and methods

2.1 Observations

The measurements used in this study were carried out over the sampling site from February 5 to 24, 2017 at the campus of Ocean University of China (36°09'37″N, 120°29'44″E) in Qingdao, which is surrounded by residential buildings and is situated about 10 km away from the city center. A fast mobility particle sizer (FMPS, TSI Model 3091) was applied to measure the aerosol particle size distribution for the size range of 5.6 nm to 560 nm (Liu et al., 2014b). The size-resolved CCN activity is measured by a cloud condensation nuclei counter (CCNe) at three different supersaturations (0.2%, 0.4% and 0.6%) and each supersaturation lasts for 5 minutes. More information about the CCN measurement can be found in Li et al. (2015).

2.2 Model configurations

WRF-Chem version 3.9 is used to simulate NPF events, with the main physical and chemical parameterization settings summarized in Table 1. The spatial resolution is 36 km with 35 vertical layers and a model top at 50 hPa. A continuous run from February 1 to 25, 2017, was conducted, with the first five-day results as the spin-up and discarded in the analysis.
Table 1 WRF-Chem model configurations used in this work

<table>
<thead>
<tr>
<th>Model configuration</th>
<th>Microphysics</th>
<th>Planetary Boundary Layer (PBL)</th>
<th>Longwave and Shortwave Radiation</th>
<th>Land model</th>
<th>Cumulus</th>
<th>Aerosol module</th>
<th>Gas-phase Chemistry</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microphysics</td>
<td>Morrison 2-moment microphysics scheme (Morrison et al., 2009)</td>
<td>YSU boundary layer scheme (Hong et al., 2006)</td>
<td>RRTMG longwave and shortwave radiation (Iacono et al., 2008)</td>
<td>Unified Noah Land Surface scheme (Chen and Dudhia, 2000; Tewari et al., 2016)</td>
<td>Grell-3D cumulus parameterization scheme (Grell, 1993)</td>
<td>MOSAIC module (Zaveri et al., 2008; Matsui et al., 2011)</td>
<td>SAPRC-99 gas-phase chemistry scheme (Carter, 2000)</td>
</tr>
</tbody>
</table>

The meteorological initial and boundary conditions are driven by Climate Forecast System model version 2 (CFSv2; (Saha et al., 2014)) reanalysis developed by National Centre for Environmental Prediction (NCEP). The initial and boundary chemical conditions of WRF-Chem are provided by Community Atmosphere Model with Chemistry (CAM-Chem; (Buchholz et al., 2019)). Anthropogenic emissions for the year of 2017 are obtained from the Multiresolution Emission Inventory for China (MEIC, http://www.meicmodel.org/) emission dataset (Li et al., 2017; Zheng et al., 2018).

The Model for Simulating Aerosol Interactions and Chemistry (MOSAIC) was used to delineate dynamic gas-particle mass transfer to represent the condensation growth of aerosol (Zaveri et al., 2008). The gas-particle partitioning of gas species on particles is regulated by the mass transfer rate, which is related to mass accommodation coefficient (α), a parameter involved in the model representing the probability of gas molecules entering the bulk liquid phase (Pöschl et al., 1998). The original setting of α for all condensing species for all size bins a in MOSAIC is 0.1 (Zaveri et al., 2008).
the default release of WRF-Chem, MOSAIC was implemented in the sectional framework with aerosol size distributions divided into 4 or 8 size bins spanning 39 nm to 10 µm in diameter. To explicitly express the nucleation and the growth of newly formed particles, the aerosol size range in the MOSAIC module was extended from 1 nm to 10 µm, with the number of aerosol size bins increased to 20 (Matsui et al., 2011; Matsui et al., 2013; Lupascu et al., 2015; Lai et al., 2022).

The chemical aging process of organic aerosols (OA) is modeled by the volatility basis set (VBS) approach, which was widely used in air quality models to represent complex mixtures of thousands of organic species (Donahue et al., 2006; Shrivastava et al., 2011; Chrit et al., 2018). The VBS method classifies compounds according to the effective saturation concentration (C*), which represents the proportion of the component in the gas phase to the particle phase (Donahue et al., 2006), and species with higher C* values are more volatile. The oxidation of highly volatile precursors to form relatively low volatile components represents the aging process of OA. OA consists of directly emitted primary organic aerosols and photochemically produced secondary organic aerosols (SOA) (Shrivastava et al., 2011). In this study, the simplified 2-species VBS mechanism was applied to the simulation of SOA, during which primary organic aerosol was represented by two species based on volatility with effective saturation concentration C* values (at 298 K and 1 atm) of 10⁻² and 10⁵ µg m⁻³ (Shrivastava et al., 2011). Primary organic aerosols with C* of 10⁵ µg m⁻³ refers to S/IVOC, which is in the gas phase under most atmospheric conditions due to its high volatility, while for those primary organic aerosols with C* of 10⁻² µg m⁻³, it is treated as gas phase as well in the original model. The SOA formed by photochemical oxidation of S/IVOC precursors is called SI-SOA and the SOA formed by oxidation of VOC precursors is named V-SOA. In the simplified 2-species VBS mechanism, SI-SOA (C* of 10⁻²µg m⁻³) is formed by the oxidation reaction of S/IVOC precursors (C* of 10⁵µg m⁻³) and OH with an oxidation rate constant of 4 × 10¹¹ cm³ molec⁻¹ s⁻¹. A detailed description of 2-species VBS mechanism can be found in Shrivastava et al. (2011).

2.3 Model sensitivity formulations
Three sets of sensitivity tests are designed and listed in Table 2. The purposes of the three sets of experiments are as follows: (1) Adjust the condensation growth process of ultrafine particles in WRF-Chem model (Base, MAC, POA, NOCD, RACD, with details in Table 2); (2) Explore the effect of SI-SOA yield on CCN (Low-Yield and High-Yield); (3) Study the effect of nucleation process on CCN under the change of SI-SOA yield (Low-Yield and High-Yield and their corresponding cases without nucleation parameterization, i.e., Low_nucoff and High_nucoff). Each scenario will be explained in conjunctions with the results.

<table>
<thead>
<tr>
<th>Purposes</th>
<th>Simulation scenarios</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adjust the condensation growth process of ultrafine particles</td>
<td>Base</td>
<td>Simulation with the default setting with nucleation coefficient set as $2 \times 10^{-6}$ s$^{-1}$, the same as Lai et al. (2022)</td>
</tr>
<tr>
<td>Mass accommodation coefficient (MAC)</td>
<td>It is the same as Base except that the mass adjustment coefficient ($\alpha$) of gaseous sulfuric acid is adjusted from 0.1 to 0.65.</td>
<td></td>
</tr>
<tr>
<td>POA emission phase (PEP)</td>
<td>It is the same as MAC except that the phase of POA is changed from gas phase to particle phase.</td>
<td></td>
</tr>
<tr>
<td>No condensation (NOCD)</td>
<td>It is the same as PEP except that no NH$_4$NO$_3$ condenses on particles below 40 nm.</td>
<td></td>
</tr>
<tr>
<td>Ratio method for condensation (RACD)</td>
<td>It is the same as PEP except that the condensation of NH$_4$NO$_3$ on particles below 40 nm is reduced according to the ratio of acid particles to total matter.</td>
<td></td>
</tr>
</tbody>
</table>
Explore the effect of SI-SOA yield on CCN (Explore the effect of nucleation process on CCN under the change of SI-SOA yield)

<table>
<thead>
<tr>
<th>Simulation Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>High-Yield</td>
<td>Simulation with high oxidation rate of SI-SOA formation with reaction rate constant of $5 \times 10^{-11}$ cm$^3$ molec$^{-1}$ s$^{-1}$</td>
</tr>
<tr>
<td>Low-Yield</td>
<td>Simulation with low oxidation rate of SI-SOA formation with reaction rate constant of $2 \times 10^{-11}$ cm$^3$ molec$^{-1}$ s$^{-1}$</td>
</tr>
</tbody>
</table>

Explore the effect of nucleation process on CCN under the change of SI-SOA yield

<table>
<thead>
<tr>
<th>Parameterization Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>High_NUCOFF</td>
<td>Simulations without nucleation parameterizations based on High-Yield</td>
</tr>
<tr>
<td>Low_NUCOFF</td>
<td>Simulations without nucleation parameterizations based on Low-Yield</td>
</tr>
</tbody>
</table>

3. Results

3.1 Observational analysis

Based on the criteria (Dal Maso et al., 2005; Kulmala et al., 2012), NPF is defined as an event with the emergence of a nucleation mode with particle diameters smaller than 25 nm, lasting for 2 hours or more, followed in general by a continuous particle growth. Six NPF events were identified in February 2017 in Qingdao, on the days of 6, 9, 10, 17, 20 and 23 (Fig. 1a), yielding a frequency of ~30% and displaying a typical banana-shaped growth of particles in the particle number size distribution. Compared to a few other studies on NPF frequency in Qingdao, the results in this study are to a large extent consistent with that in the fall of 2012–2013 (30%; Zhu et al., 2019), slightly higher than that in summer 2016 (22%; Zhu et al., 2019) and lower than that in spring of 2010 (41%; Liu et al., 2014c). The higher frequency in spring in Qingdao is consistent with the observational results at different stations in the Northern Hemisphere in Nieminen et al. (2018).
Fig. 1 Distribution of particle number concentration. (a) Temporal evolution of particle size distributions (colored shading) and geometric median diameter (GMD; dots in black) in Qingdao on February 5-24, 2017. (b) The mean diurnal variation of CN_{10-40} (blue), CN_{40-100} (orange) and CN_{100-1000} (green) composited during the NPF (solid lines) and non-NPF (dashed lines) days on February 5-24, 2017. All times are local times (LT).

During the six NPF events identified in February in Qingdao, the mean diurnal cycle of CN_{10-40} (10–40 nm) particles exhibits triple peaks (solid blue in Fig. 1b), in the morning (8:00 LT), noon (12:00–14:00 LT) and evening (19:00 LT), respectively. A comparable three-peak feature was also observed in earlier years during 2016-2018 in Qingdao (Zhu et al., 2021). The morning and evening peaks of CN_{10-40}, with values of ~5300 cm^{-3} and ~12000 cm^{-3}, respectively, are likely caused by the primary emissions from traffic and cooking activities (Wu et al., 2021a; Wang et al., 2022; Cai et al., 2020). The occurrence of NPF starts approximately at 9:00 am LT, accompanied by a substantial increase in CN_{10-40} compared with non-NPF days (solid vs. dashed lines, in blue), yielding a peak around noon (20000 cm^{-3} during 12:00–14:00 LT). In addition,
larger particles (e.g., CN_{40–100} and CN_{100–1000}) displayed a slow or no increase in the afternoon.

3.2 Model improvement in particle number concentration simulations

Particle number concentrations, primarily in two ranges of 10–40 nm and 40–100 nm, are commonly simulated with large biases. In the smaller size range (10–40 nm), the particle number concentration is associated with NPF and particles growth. During NPF, despite differences among the formation mechanisms, H$_2$SO$_4$ is considered the common species (Yu, 2005; Lovejoy et al., 2004), which often suffer large biases (Cai et al., 2016; Matsui et al., 2011). In the size range of 40–100 nm, the particle number concentration is primarily affected by the condensation growth of particles below 40 nm, which is closely related to chemical components such as SOA and nitrate.

3.2.1 Bias correction of particle number concentration at 10–40 nm

In this study, as shown in Fig. 2, comparisons of CN_{10–40} between simulations (red line in Fig. 2a) and observations (black line in Fig. 2a) results of the six NPF events mentioned in the previous section in Qingdao in February 2017 indicate that model overestimates CN_{10–40} with mean fractional bias (MFB) of 48%. As one of the major processes affecting the particle number concentration of 10–40 nm, nucleation is governed by the particle nucleation rate of 1 nm particles (cm$^{-3}$ s$^{-1}$), which is closely associated with the concentration of H$_2$SO$_4$. For instance, in a commonly applied activation mechanism, the nucleation rate calculated by $J^* = K_{ACT} \times [H_2SO_4]$. Note that $K_{ACT}$ is the nucleation coefficient considering the physical properties and chemical species of nucleation process under different environments, indicating that a lumped chemical species are included in the scheme reflected primarily in the nucleation coefficient $k$, set as $2 \times 10^{-6}$ s$^{-1}$ based on previous studies (Sihto et al., 2006; Riipinen et al., 2007). Dong et al. (2019) simulated NPF occurring in the summer of 2008 in the United States using the NFP-explicit WRF-Chem based on the activation mechanism, which overestimated the particle number concentration at 10–63 nm by nearly doubled, even when the $K_{ACT}$ decreased by one order of magnitude (set at a very low value of $10^{-7}$ s$^{-1}$). Therefore, it is likely that the overestimation of particle number concentration...
in the smaller particle size segment is probably due to the bias of simulated sulfuric acid. 

![Figure 2 Time series of (a) CN_{10-40} on NPF days, where red and blue represent Base and MAC simulation results respectively, and black represents observation results, and (b) sulfuric acid gas concentration obtained by simulation and by proxies (dark grey: Eq. 1; green: Eq. 2; purple: Eq. 3; brown: Eq. 4; pink: Eq. 5; yellow: Eq. 6). All times are in local times.]

Measurement of sulfuric acid gases in the lower troposphere is challenging due to the generally low ambient concentration of sulfuric acid ($10^6$–$10^7$ molecule cm$^{-3}$). Different methods have been proposed to estimate ambient sulfuric acid concentrations based on observations such as SO$_2$ (Petäjä et al., 2009; Lu et al., 2019; Mikkonen et al., 2011). For instance, Petäjä et al. (2009) proposed a linear method to approximate observed H$_2$SO$_4$ concentration in Hyytiala, southern Finland. Moreover, a recent study by Lu et al. (2019) proposed a nonlinear method to construct a number of proxies for gaseous sulfuric acid concentration (Eq. 1–5), indicating that compared to the linear method in Petäjä et al. (2009), the nonlinear relationship can provide more accurate H$_2$SO$_4$ concentration in Beijing during February–March 2018 period. In addition, we also used another sulfuric acid nonlinear proxy (Eq. 6) based on long-term observations in Germany, Finland, the United States, etc. (Mikkonen et al., 2011). In this study, we adopt the above six nonlinear proxy methods (referred as proxy1 to proxy6) to estimate H$_2$SO$_4$ in Qingdao.
\[ [H_2SO_4] = 515.74 \times [SO_2]^{0.38} \times \text{Radiation}^{0.14} \times CS^{0.03} \] (1)

\[ [H_2SO_4] = 280.05 \times \text{Radiation}^{0.14} [SO_2]^{10.40} \] (2)

\[ [H_2SO_4] = 9.95 \times [SO_2]^{0.39} \times \text{Radiation}^{0.13} \times CS^{-0.01} \times [O_3]^{0.14} \] (3)

\[ [H_2SO_4] = 14.38 \times [SO_2]^{0.38} \times \text{Radiation}^{0.13} \times [O_3]^{10.14} \] (4)

\[ [H_2SO_4] = 0.0013 \times [SO_2]^{0.38} \times \text{Radiation}^{0.13} \times CS^{-0.17} \times ([O_3]^{10.14} + [NO_x]^{0.41}) \] (5)

\[ [H_2SO_4] = 8.21 \times 10^{-3} \times [SO_2]^{0.62} \times \text{Radiation} \times (CS \times RH)^{-0.13} \] (6)

where \([SO_2], [O_3]\) and \([NO_x]\) (molecule cm\(^{-3}\)) represents concentration of observed \(SO_2, O_3\) and \(NO_x\), respectively. “Radiation” (W m\(^{-2}\)) is global radiation. RH (%), is the relative humidity, and CS (s\(^{-1}\)) is the condensation sink, which is calculated based on observed particle distribution.

The simulated \(H_2SO_4\) concentration from the Base simulation (dots in Fig. 2b) is compared with observations obtained by proxies (see Fig. 2b), indicating that Base simulations apparently overestimate by one order of magnitude compared to the \(H_2SO_4\) estimated by proxies. The overestimation has been frequently reported previously, i.e., over Beijing (Matsui et al., 2011), which ascribes the bias to the overestimation of the \(SO_2\) concentration. In a more recent study, the sensitivity of \(H_2SO_4\) to \(SO_2\) is tested, and the result shows that even when \(SO_2\) is reduced to an unrealistically low level, the simulated \(H_2SO_4\) is still more than one order of magnitude higher than the observed value (Lai et al., 2022), suggesting that the \(SO_2\) concentration cannot fully explain the overestimates.

In addition to the precursor of \(H_2SO_4\), the mass accommodation coefficient \((\alpha)\), representing the probability of impaction of a gaseous molecule on a liquid surface and entering the bulk liquid phase, is another important factor affecting the concentration of sulfuric acid gas. In the public release of WRF-Chem, \(\alpha\) is typically set to a low value of 0.1 for all gas species under different volatility during the condensation process, including \(H_2SO_4\) (Davidovits et al., 2004; Zaveri et al., 2008). Recent studies indicate that the low \(\alpha\) value may not be applicable to the low volatile gases, which tend to have a mean \(\alpha\) value of 0.7 and close to the unity (Krechmer et al., 2017). In fact, an earlier study has indicated based on experimental determination, the mass accommodation coefficient of \(H_2SO_4\) vapor in sulfuric acid aqueous solution was measured, and the best
fit value was 0.65 (Pöschl et al., 1998). Accordingly, a sensitivity simulation was conducted by adjusting the mass accommodation coefficient of H$_2$SO$_4$ from 0.1 to 0.65, referred to as MAC.

This simulation brought the H$_2$SO$_4$ concentration (see Fig. 2b) much closer to the calculated results from proxies, and the corresponding biases reduced by approximately an order of magnitude. Notably, the MAC simulation decreases the overestimate of sulfuric acid gas concentration, resulting in a lower particle formation rate. The MAC simulation also significantly reduces overestimate of CN$_{10-40}$ (Fig. 2b), and MFB compared to observations decreases from 48% to 1%.

### 3.2.2 Improvement of particle number concentration simulations at 40–100 nm

The number concentration of particles in the 40–100 nm range is mainly affected by the coagulation and condensation processes. While the coagulation process tends to largely affect ultrafine particles below 10 nm than those with larger sizes (Wu et al., 2011), the condensation growth of particles during gas-particle partitioning at sizes of 10–40 nm, to a large extent, governs the variations in number concentration of 40–100 nm particles. The condensation process is primarily controlled by gas-particle partitioning of chemical species, which may change the chemical composition of particles, such as organic compounds and inorganics including sulfate, nitrate and ammonium.

Among the species contributing to the condensation growth of particles at 10–40 nm, the organic compounds with with C$_\ast$ of 10$^{-2}$ µg m$^{-3}$ play the dominant role (Pierce et al., 2011). In the current model setting, there is a total of two volatility sets, including 10$^{-2}$ µg m$^{-3}$ and 10$^{5}$ µg m$^{-3}$. The low volatile organic matter of 10$^{-2}$ µg m$^{-3}$ comes from two gas-phase sources, including the direct emission of primary organic aerosol (POA) and SOA formed from S/IVOC (SI-SOA), conducive to condensation on particles. While the condensation of gaseous SOA is in general reasonable, the gas phase emissions of POA may be problematic. For instance, previous studies suggested that POA is in gas phase close to the emissions source, but with rapid dilution and cooling in the atmosphere away from the source, most POA condenses to particle-phase (Roldin...
et al., 2011b; Roldin et al., 2011a; Shrivastava et al., 2008). Therefore, away from the emissions source POA, being in the particle phase, will not be involved in the growth of newly formed particles. Therefore, we assume that low volatility POA is emitted in the particle phase rather than the gas-phase, which caused different size distributions of POA compared to when it was emitted in the gas-phase (Fig. S1a vs. Fig. S1b). Emitting low volatility POA in the particle phase eliminates the unreasonable quasi-banana shape pattern exhibiting concomitant growth of newly formed particles with increasing mass concentration of POA.

The composition analysis (Fig. S1c) in the 10–40 nm particles mass from the model results indicates that organic compounds mentioned above only account for 21% of total mass (sulfates, nitrates, ammonium salts and organics) in this size range and the dominant species is nitrate which accounts for 51% of total mass, exhibiting inconsistencies with the previous studies which in general indicates a much smaller contribution of nitrate. For instance, Liu et al. (2014a) suggested that over North China Plain in summer 2009, organic matter accounted for 77% of particles around 30 nm, while the sum of SO$_4^{2-}$, NO$_3^-$ and NH$_4^+$ only accounted for 18%. Another study showed that nitrate accounted for 7–8% at urban sites and 17% at rural sites for particles mass in 7–30 nm in the United States in 2007 (Bzdek et al., 2012). Therefore, the potentially too high modeled nitrate fraction in 10–40 nm in this study is tightly associated with the condensation process, with the specific reasons explained below.

The condensation of nitric acid on particles is highly constrained by the particle acidity. The acidity in smaller particles (i.e., 10–40 nm) tends to be higher than that in large particles, primarily due to the larger condensation of H$_2$SO$_4$ (Lu et al., 2022), and particles with sizes greater than 40 nm have a much weaker acidity or are nearly neutral. For example, observed evidence has shown that acidic ultrafine particles account for a large proportion of ultrafine particles from 22 December 2010 to 15 January 2011 in Hong Kong, e.g., 65% for particles within 5.5–30 nm (Wang et al., 2014).

In the model, a particle is determined to be in solid phase when the ambient relative humidity is lower than the mutual deliquescence relative humidity of the particles (Zaveri et al., 2005; Zaveri et al., 2008), which is in general suitable for particles...
dominated by inorganics. In the study area, the results indicate that at most conditions relative humidity are relatively low and the particles are in solid phase, in which the condensation process is not affected by particle acidity and the condensation of nitric acid on particles is directly calculated based on the gas-particle equilibrium concentration (Zaveri et al., 2008). However, for particles below 40 nm, the main compositions are likely to be organic matter (Zhu et al., 2014; Ehn et al., 2014), which tends to be in liquid phase (Virtanen et al., 2011; Cheng et al., 2015), under which the condensation of nitric acid is strongly constrained by acidity. Therefore, the phase misrepresentation ignores the weakening effect of acidity on nitric acid condensation, resulting in too high nitrate therein.

To overcome this issue, we propose a ratio method for condensation (RACD) to partition the condensation of nitric acid on particles under 40 nm, by applying a ratio of the number concentration of non-acidic particles to ultrafine particles. The method is based on two assumptions, including: 1) little condensation of nitric acid on particles with strong acidity (Lu et al., 2022); 2) the condensation of nitric acid on particles is proportional to the ratio of the number concentration of non-acidic ultrafine particles to the total particles, despite the existence of uncertainties. Fig. S2 depicts the average particle number concentration and acid particle in the 1 to 40 nm range, calculated based on Wang et al. (2014). The ratio of non-acidic particles is 8% for particles below 10 nm, 18% for particles at 10–15.8 nm, 30% for particles at 15.8–25.1 nm, and 55% for particles at 25.1–39.8 nm (Fig. S2). Note that the ratio is based on measurements acquired at a single site in Hong Kong, therefore more observational studies are needed to warrant the robustness of the method. Alternatively, the condensation of nitric acid on particles in bins from 1 nm to 40 nm is completely suppressed, referred to as NOCD.

The simulation results based on the two methods (RACD and NOCD) are shown in Fig. 3. Compared to MAC, RACD simulations reduce previously noted overestimation of PNC in the 40–100 nm size range (Fig. 3b), with the MFB decreases from 83% to 63%. In addition to the amount of nitrate condensation during particle growth mentioned above, the overestimation of particle number concentrations in the 40–100 nm range may be attributed to nucleation process. More specifically, in the
H$_2$SO$_4$-H$_2$O binary nucleation mechanism used in this study, when the concentration of sulfuric acid gas is reduced (Section 3.2.1), the resulting decrease in nucleation rate leads to a slight decrease in particle number concentration at 40–100 nm relative to Base (MFB from 98% to 83%). Apart from that, it may also be related to the choice of nucleation parameterization scheme. For example, using a global chemical transport model GEOS-Chem, Yu et al. (2015) overestimated the concentration of particles in the 10–100 nm range by 161% at nine sites in the summer in North America. A possible explanation for this overestimation was given by the uncertainty of the predicted concentration of organic compounds involved in organics-mediated nucleation parameterization. After they switched to another scheme of the ion-mediated nucleation mechanism without organic matter, the number becomes 27% lower than the observations (Yu et al., 2015). The test based on different schemes is beyond the scope of the study, which is therefore not investigated.

Moreover, the overestimation of particles over 100 nm (CN$_{100-1000}$; Fig. 3c), which have a strong influence on CCN, also decrease in the RACD simulation. Thus, the MFB decreases from 25% (MAC) to 1%. Note that the slight increase of CN$_{10-40}$ through the application of RACD, can be linked to the decrease of nitrate condensation, and leads to weakened particle growth and enhanced particle number concentration at 10–40 nm (Fig. 3a). The alternative method by completely removing the nitrate condensation (NOCD) yields even better performance in particle number concentration of 40–100 nm (MFB of 34%), indicating the feasibility by reducing the nitrate condensation. The proportion of nitrate simulated by RACD is 23%, closer to values reported in past observations (Bzdek et al., 2011; Bzdek et al., 2012), while the nitrate (1%) in the scenario of NOCD seems to be too low. Considering the limited observational information obtained based on previous studies, RACD is applied in this study.
Figure 3 The time series of (a) CN\textsubscript{10–40}, (b) CN\textsubscript{40–100} and (c) CN\textsubscript{100–1000} on NPF days in Qingdao on February 5-24 simulated MAC (marked in red), NOCD (marked in blue) and RACD (marked in green) as well as from observations (OBS) (marked in black). All times are local time.

3.3 Substantial contributions of SI-SOA to CCN

Compared with the original model setting, after adjusting the growth process of ultrafine particles (RACD), the number concentration of particles tend to decrease, especially for particles above 40 nm. Ultrafine particles above 40 nm are important sources of CCN (Dusek et al., 2006), in this way, the number concentration of CCN also tends to decline. In addition, in the Base case, we found that the model overestimated CCN\textsubscript{0.4\%} and CCN\textsubscript{0.6\%}, with MFB being 64\% and 87\%, respectively. After adjusting the condensation growth process of ultrafine particles, under high supersaturation (i.e., CCN\textsubscript{0.4\%} and CCN\textsubscript{0.6\%}), the capability of the model in reproducing the CCN is improved. RACD reduces the overestimation of CCN\textsubscript{0.4\%} and CCN\textsubscript{0.6\%}, with MFB reduced to 30\% and 56\%, respectively, although the overestimates still exist (Figs. S3b, c). However, for low supersaturation (i.e., CCN\textsubscript{0.2\%}), the decrease of number concentration of CCN is too large, and MFB decreases from 7\% to -45\% (Figs. S3a), therefore the bias will be further adjusted later.
In addition to the growth process, the remaining overestimate of CCN under high SS and underestimate of CCN over low SS is likely to be influenced by the chemical compositions involved in the activation of ultrafine particles into CCN. Specifically, ultrafine particles can grow up to CCN size under certain SS (Pierce and Adams, 2007). This process is influenced by both particle size and hygroscopicity, and hygroscopicity is closely related to the chemical composition of particles (Petters and Kreidenweis, 2007). In particular, inorganic compounds generally increase particle hygroscopicity, increasing CCN. SOA has dual effects on CCN since it decreases particle hygroscopicity but also promotes growth of particles, and these two effects are competitive with each other (Wu et al., 2015; Zaveri et al., 2021). Ultrafine particles must grow to a critical size to be activated into CCN (Dusek et al., 2006). SOA act as a major contributor in promoting the condensational growth of ultrafine particles to the critical size, facilitating particles activation into CCN. In contrast, SOA tends to reduce the hygroscopicity of particles, leading to a diminished ability of activation to CCN (Wu et al., 2015). These two competing effects work together and modulate the number of CCN. Moreover, considering that SI-SOA is the main SOA component on ultrafine particles (Fig. S3d), the effect of SI-SOA on CCN is therefore explored in this study.

Considering SI-SOA is a product of S/IVOC oxidation, the oxidation rate of S/IVOC is tightly associated with CCN, which likely affects the bias of CCN. In the original model setup, the oxidation rate is set to be a constant of $4 \times 10^{-11}$ cm$^3$ molec$^{-1}$ s$^{-1}$ for all S/IVOC. However, a recent study (Wu et al., 2021b) proposed that the oxidation rate can be as high as $5 \times 10^{-11}$ cm$^3$ molec$^{-1}$ s$^{-1}$ such as for polycyclic aromatic hydrocarbons (PAHs), close to the original model value, but can be as low as half (i.e., $2 \times 10^{-11}$ cm$^3$ molec$^{-1}$ s$^{-1}$) of the original modeling setting for S/IVOC species except PAHs (O-S/IVOCs). It is noteworthy that the oxidation rates of $5 \times 10^{-11}$ and $2 \times 10^{-11}$ in general represent the upper and lower bounds (Zhao et al., 2016; Wu et al., 2021b).

To delve into how oxidation rates affect CCN, we set up a few numerical experiments (Table 2) to investigate the response of CCN to the oxidation rate of S/IVOC at three supersaturations (0.6%, 0.4%, 0.2%), including cases of High_Yield and Low_Yield. As it is shown in Fig. 4, decreasing the oxidation rate (Low_Yield)
leads to a reduction of ~10% of CCN at high supersaturation (i.e., CCN0.6%) as compared to the High_Yield simulation. This behaviour is a consequence of the decrease of particle number concentrations associated with Low_Yield, particular of the particles close to the critical diameter (40–100 nm). In this case, the effect of particle size dominates the hygroscopicity. In contrast, at a lower supersaturation (CCN0.2%), CCN increases by 42% when the oxidation rate is switched from a high to a low value, which is due to the smaller fraction of SI-SOA contributing to particulate mass when the oxidate rate is low. In this case, relative to SOA, a larger fraction of other particle constituents such as inorganics, increase the volume weighted particle hygroscopicity (Dusek et al., 2006) which causes the increase of CCN number. This means that the effect of hygroscopicity on CCN surpasses the influence on particle size at low supersaturations. This conclusion is consistent with the observation conducted by Ma et al. (2016) in the North China Plain in 2013, which suggested that along with the decrease of SS, the particles that can be activated to CCN is more sensitive to changes of particle hygroscopicity.

Figure 4: Average diurnal variation of (a) CCN0.2%, (b) CCN0.4% and (c) CCN0.6% and (d) CN10–40, (e) CN40–100, (f) CN100–1000 on NPF days in Qingdao on February 5–24, 2017, in Low-yield and High-yield simulations, shown as blue and brown lines, and black lines represent observation results.

Furthermore, compared to the high yield of SI-SOA, the low SI-SOA yield results in a high CCN concentration under low SS and low CCN concentration under high SS. Therefore, both the underestimates of CCN0.2% (MFB of -45%) and overestimates of
CCN$_{0.6\%}$ (MFB of 56%) mentioned above are improved, with MFB of CCN$_{0.2\%}$ and CCN$_{0.6\%}$ reaching 7% and 43%, respectively (Figs. 4a,c). This result suggests that the oxidation rate of S/IVOC is possibly closer to the low value, which is understandable based on Wu et al. (2021b), who found that the amount of O-S/IVOCs, which corresponds to a low oxidation rate, is in general much larger (i.e., 20 times) than that of PAHs with a high oxidation rate.

In addition to the single site of Qingdao, we further explore the impact of SI-SOA yield on CCN from a larger spatial coverage (Fig. 5). Consistent with the mechanism revealed over Qingdao, even from a larger spatial perspective, a lower oxidation rate of S/IVOC essentially enhances CCN at a lower SS (e.g., CCN$_{0.2\%}$; Fig. 5a) with the highest increase over north China Plain area (Figs. 5a), and weakens CCN (i.e., by 10–20% over Beijing-Tianjin-Hebei) at a higher SS (Figs. 5c), particularly over the dense emission area (Fig. S4). It is worth noting that in the 2-species VBS mechanism used in our study, all S/IVOC in the inventory is calculated based on a constant emission ratio of S/IVOC to POA from all source categories (Shrivastava et al., 2011), which may miss part of S/IVOC due to different emission ratios of POA from different source categories (Chang et al., 2022).
Figure 5. Spatial distributions of CCN concentrations at different supersaturations (SS), (a) and (d) are CCN$_{0.2\%}$, (b) and (e) are CCN$_{0.4\%}$, and (c) and (f) are CCN$_{0.6\%}$. The top panels exhibit the results from the High_Yield simulation, and the bottom panels shows the difference between the Low_Yield and High_Yield simulations.

3.4 Contribution of nucleation to CCN under different SI-SOA yields

Considering the importance of nucleated particles on CCN (Yu et al., 2020; Westervelt et al., 2013), we further investigate the influence of nucleation on CCN under different SI-SOA yield conditions discussed above.

As shown in Fig. 6, in simulations close to the original model setting (High_Yield), when SS is low (i.e., SS=0.2%), the nucleation process tends to reduce the CCN by ~10–50%. In contrast, when the SS is high (0.6%), the nucleation results in a significant increase in CCN in most regions of China. When the yield of SI-SOA is adjusted to a lower level, the nucleation process has a positive contribution to CCN under both low and high SS. Especially, when SS is low (0.2%), the sign reversal, i.e., from negative (Fig. 6a) to positive (Fig. 6d) contribution of NPF to CCN along with the decrease of SI-SOA yield, i.e., the increase is concentrated in the eastern China with an average of...
10–20%. The primary mechanism lies in that along with the decrease of SI-SOA yield, the smaller fraction of SI-SOA yields an increase in hygroscopicity, which surpasses the suppression effect on particle growth due to reduced SI-SOA formation. In the real atmosphere, when the supersaturation is usually low, e.g. about ~0.1% in polluted areas (Kalkavouras et al., 2019; Hudson and Noble, 2014), CCN will likely reduce with increasing oxidation rate of S/IVOC and corresponding SI-SOA formation.

Figure 6. Spatial distribution of contribution of nucleation to CCN calculated by the ratio of the difference between the parameterization with and without nucleation to the parameterization with nucleation under different SI-SOA yields in China in February 2017. (a), (d) is CCN$_{0.2\%}$, (b), (e) is CCN$_{0.4\%}$, (c), (f) is CCN$_{0.6\%}$. The upper panel and lower panel represent High_Yield and Low_Yield simulation respectively.

Conclusions

In this study, WRF-Chem explicit-NPF simulations are used to investigate the observed wintertime NPF events and their contribution to CCN in China. Based on
observations in a typical coastal city of Qingdao, we identify high biases of the model simulated CN and CCN concentrations. Therefore, we updated and improved the parameterization setting on particle growth in the model, mainly including: (1) adjusting the mass accommodation coefficient ($\alpha$) to from the default value of 0.1 to 0.65, an important parameter for sulfuric acid condensation; (2) proportionally reducing the condensation amount of nitric acid on particles below 40 nm, (3) changing the emitted low-volatility POA from gas to particle. Through these adjustments, the capability of the model in reproducing CN and CCN is substantially improved, leading to better agreement with the observed results, which significantly reduces the overestimation of CN$_{10-40}$ (MFB decreases from 48% to 1%) and CN$_{40-100}$ (MFB decreases from 98% to 63%).

For CCN, due to the crucial role of SI-SOA in promoting the growth of ultrafine particles, on the basis of previous studies, we lower the oxidation rate of S/IVOC and hence the production rate of SI-SOA, which weakens the growth of particles to reach the critical size of CCN activation, but enhances particulate hygroscopicity favoring the activation to CCN. When the yield of SI-SOA is adjusted to the lower bound of literature value, CCN$_{0.6\%}$ is reduced by $\sim$10% and is closer to observations. At low SS (CCN$_{0.2\%}$), the decrease of SI-SOA yield has greater effects on the increase of particle hygroscopicity compared to the effect of the reduction of particle size due to the decrease of condensation growth. It results in an increase of CCN (as large as $\sim$42%) in better agreement with observations. Under low SS conditions, common in the atmosphere, a 2.5-fold reduction in SI-SOA yield results in a substantial increase of CCN that switches from a negative contribution of new particle formation to CCN from $-50\%$ to $-10\%$ to a positive contribution of 10$\%$--20$\%$.

**Competing interests.** At least one of the (co-)authors is a member of the editorial board of Atmospheric Chemistry and Physics.
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