Observations of particle number size distributions and new particle formation in six Indian locations

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Abstract. Atmospheric new particle formation (NPF) is a crucial process driving aerosol number concentrations in the atmosphere; it can significantly impact the evolution of atmospheric aerosol and cloud processes. This study analyses at least one year of asynchronous particle number size distributions at six different locations in India. We also analyze the frequency of NPF and its contribution to cloud condensation nuclei (CCN) concentrations. We found that the NPF frequency has a considerable seasonal variability. At the measurement sites analyzed in this study, NPF frequently occurs in March-May (pre-monsoon, about 21% of the days) and is the least common in October-November (post-monsoon, about 7% of the days). Considering the NPF events in all locations, the particle formation rate ($J_{NUC}$) varied by more than an order of magnitude (0.01 - 0.6)
cm$^{-3}$ s$^{-1}$) and the growth rate (GR$_{NUC}$) by about three orders of magnitude (0.2 - 17.2 nm h$^{-1}$). We found that J$_{NUC}$ was higher by nearly an order of magnitude during NPF events in urban areas than mountain sites. GR$_{NUC}$ did not show a systematic difference. Our results showed that NPF events could significantly modulate the shape of particle number size distributions and CCN concentrations in India. The contribution of a given NPF event to CCN concentrations was the highest in urban locations (4.3x10$^3$ cm$^{-3}$ per event and 1.2x10$^3$ cm$^{-3}$ per event for 50 nm and 100 nm, respectively) as compared to mountain-background sites (2.7x10$^3$ cm$^{-3}$ per event and 1.0x10$^3$ cm$^{-3}$ per event). To better understand atmospheric NPF and its contribution to CCN concentrations, we would need long-term observational data from various diverse environments in India, aided with regional model simulations to help interpret field observations.

**Keywords:** new particle formation, particle number size distribution, Aitken mode, accumulation mode, cloud condensation nuclei

1 Introduction

Cooling by atmospheric aerosols offset a significant fraction of the radiative forcing of the greenhouse gases (Paasonen et al., 2013) directly by scattering and absorbing solar radiation and indirectly by altering cloud microphysical properties via activation of cloud condensation nuclei (CCN) (Rosenfeld et al., 2014; Sarangi et al., 2018). New particle formation (NPF), as a result of the gas-to-particle conversion, is the largest source of the aerosol number to the terrestrial atmosphere (Kulmala et al., 2007; Zhang et al., 2012). While nucleated particles from NPF are initially very small molecular clusters (1-2 nm; Kerminen et al., 2012), these molecular clusters can grow to large sizes within a few hours to a few days and ultimately reach CCN-active sizes (>50-100 nm) (Pierce and Adams, 2007; Westervelt et al., 2013). Thus, CCN forms the direct microphysical link between aerosols and clouds and plays a vital role in the hydrological cycle and climate.

In India, several intensive field campaigns such as the Indian Ocean Experiment (INDOEX) (Ramanathan et al., 2001), Indian Space Research Organization (ISRO)-Geosphere-Biosphere Programme (GBP)- Land campaign II (Tripathi et al., 2006; Tare et al., 2006), and Integrated Campaign for Aerosols, gases, and Radiation Budget (ICARB) (Moorthy et al., 2008; Nair et al., 2020; Kompalli et al., 2020) measured sub-micron particle number size distributions.
There are also short- and long-term field observations of sub-micron PNSDs in a variety of diverse locations in India (Hyvärinen et al., 2010; Kanawade et al., 2014a; Shika et al., 2020; Tripathi et al., 1988; Komppula et al., 2009; Singh et al., 2004; Moorthy et al., 2011; Babu et al., 2016; Kompalli et al., 2018). But there are sparse studies in India characterizing seasonal variation in PNSDs and number concentrations (Kanawade et al., 2014a; Hyvärinen et al., 2010; Komppula et al., 2009; Hooda et al., 2018; Laj et al., 2020) and atmospheric NPF (Sebastian et al., 2021b; Siihgh et al., 2018; Neitola et al., 2011; Moorthy et al., 2011; Kanawade et al., 2014b; Kanawade et al., 2014c; Kanawade et al., 2020a). The characterization of PNSDs is critical because the PNSD is controlled by an evolving balance between NPF, condensation of vapor on pre-existing particles, evaporation of particles, coagulation and sedimentation (IPCC, 2013). Previous field measurements and modeling studies globally demonstrated a substantial enhancement in CCN number concentrations from nucleation (Yu et al., 2020; Wiedensohler et al., 2009; Sihhto et al., 2011; Rose et al., 2017; Tröstl et al., 2016; Kalivitis et al., 2015; Westervelt et al., 2013; Pierce et al., 2012; Pierce et al., 2014; Westervelt et al., 2014; Kerminen et al., 2012; Kerminen et al., 2018; Merikanto et al., 2009; Gordon et al., 2017). For instance, Merikanto et al. (2009) revealed that 45% of the global low-level CCN at 0.2% supersaturation originates from nucleation. Westervelt et al. (2014) also found that nucleation contributes to about half of the boundary layer CCN (at supersaturation of 0.2%), with an estimated uncertainty range of 49 to 78%, which is sensitive to the choice of nucleation scheme. In contrast, Reddington et al. (2011), using the global model GLOMAP against ground-based measurements at 15 European sites, found that CCN-sized particle number concentrations were driven by processes other than nucleation at more than ten sites. They explained that the weakened response of CCN-sized particles to boundary layer nucleation arises from an increase in coagulation and condensation sinks for ultrafine particles, thereby reducing the condensational growth of ultrafine particles to CCN-active sizes (Kuang et al., 2009; Pierce and Adams, 2007). Tröstl et al. (2016) also revealed that only a small fraction of total particles less than 50 nm grew beyond 90 nm (50-100 particles cm$^{-3}$), even on a timescale of several days. Therefore, to better understand atmospheric NPF and its contribution to the boundary layer CCN budget, we need highly-resolved spatiotemporal observational data in diverse environments globally, aided with aerosol model simulations, to help to interpret field observations.

Overall, studies pertinent to the impact of NPF on aerosol-cloud interactions are highly sparse in India. The sources of aerosols, and gaseous precursors required for secondary aerosol
formation, depict a considerable spatiotemporal heterogeneity over India. Therefore, observational aerosols and precursors data must be synthesized to understand the processes that govern NPF and its contribution to CCN concentrations in different settings of India. The primary objective of this study is to harmonize observational PNSDs data from six diverse locations in India to present analyses of PNSDs, atmospheric NPF, and the contribution of NPF to CCN concentrations.

2 Methods

2.1 Observation sites and aerosol sampling instrumentation

Figure 1 shows the geographical location of measurement sites on the surface elevation map. Table 1 provides details of measurement sites and particle data analyzed in this study.

Figure 1. The geographical location of measurement sites on the surface elevation map. Measurement sites such as Ranichauri (RNC), Mukteshwar (MUK), Mahabaleshwar (MBL), Hyderabad (HYD), Thiruvananthapuram (TVM), and Delhi (DEL) are shown by the plus sign. The global 1-arcsecond (30-m) SRTM digital surface elevation data is obtained from the United States Geological Survey (https://dds.cr.usgs.gov/srtm/version2_1/SRTM30/).
Table 1. Details of the measurement sites and particle number size distribution measurements analyzed in this study.

<table>
<thead>
<tr>
<th>Site Name</th>
<th>Site code</th>
<th>Site type</th>
<th>Instrument</th>
<th>Size range (nm)</th>
<th>Time resolution (minutes)</th>
<th>Time Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ranichauri</td>
<td>RNC</td>
<td>Mountain background</td>
<td>DMPS</td>
<td>10.1–757</td>
<td>10</td>
<td>12/2016 – 09/2018</td>
</tr>
<tr>
<td>Mukteshwar</td>
<td>MUK</td>
<td>Mountain background</td>
<td>DMPS</td>
<td>10.1–757</td>
<td>5</td>
<td>01/2012 – 12/2013</td>
</tr>
<tr>
<td>Mahabaleshwar</td>
<td>MBL</td>
<td>Mountain semi-rural</td>
<td>WRAS</td>
<td>5.14–1000</td>
<td>4</td>
<td>03/2015 – 03/2016</td>
</tr>
<tr>
<td>Hyderabad</td>
<td>HYD</td>
<td>Urban</td>
<td>SMPS</td>
<td>10.9–514</td>
<td>5</td>
<td>04/2019 – 03/2020</td>
</tr>
<tr>
<td>Thiruvananthapuram</td>
<td>TVM</td>
<td>Semi-urban coastal</td>
<td>SMPS</td>
<td>14.6–661.2</td>
<td>5</td>
<td>01/2013 – 01/2014</td>
</tr>
<tr>
<td>Delhi</td>
<td>DEL</td>
<td>Urban</td>
<td>WRAS</td>
<td>5.14–1000</td>
<td>5</td>
<td>11/2011 – 01/2013</td>
</tr>
</tbody>
</table>

DMPS: Differential Mobility Particle Sizer, WRAS: Wide-Range Aerosol Spectrometer, SMPS: Scanning Mobility Particle Sizer

Ranichauri observation site (RNC, 30.2°N, 78.25°E; ~1930 m above mean sea level, amsl) is located in Tehri–Garhwal district of Uttarakhand state in the southern slope of the Western Himalaya. The RNC site is situated on an isolated hilltop within the campus of the College of Forestry in the Ranichauri village. The RNC site is a Climate Monitoring station managed by the India Meteorological Department (IMD). It is a mountain background remote observatory (Sebastian et al., 2021b) and located about 70 km to the northeast of Rishikesh city, about 100 km to the northwest of the Srinagar city, and about 100 km to the east of Dehradun. Here, particle number size distributions in the size range from 10 nm to 757 nm (30 size bins) is measured using a differential mobility particle sizer (DMPS, Finnish Meteorological Institute assembled) from December 2016 – September 2018 are used (Sebastian et al., 2021b). The DMPS consisted of a Vienna-type differential mobility analyzer (DMA) that classifies the charged particles according to their electrical mobility and a TSI 3772 condensation particle counter (CPC) that counts particles of the selected mobility. The sample air was drawn inside through a stainless-steel inlet tube of about 2 meters in length and dried to less than 40% relative humidity with a Nafion dryer (Perma Pure model MD-700-48). Diffusion losses in the inlet and inside the DMPS instrument were considered in the data inversion. The inversion method was identical to that presented by Wiedensohler et al. (2012) for the Finnish Meteorological Institute (FMI) DMPS.
Mukteshwar observation site (MUK, 29.43° N, 79.62° E, 2180 m amsl) is located in the Nainital district of Uttarakhand state in the southern slope of the Central Himalaya. The Mukteshwar village is situated 3 km to the northeast of the measurement site at a similar altitude with ~800 inhabitants (Census of India, 2011). MUK can be considered a mountain background site, with the annual mean black carbon (BC) concentration of 0.9 µg m\(^{-3}\). The town of Almora (1650 m amsl, 34,000 inhabitants) is located at about 16 km to the north, Nainital (1960 m amsl, 41000 inhabitants) is located at about 25 km to the southwest, and the city of Haldwani (424 m amsl, 150,000 inhabitants) is located at about 32 km to the southwest to MUK. Delhi, the major metropolitan city (215 m amsl, 16.8 million inhabitants), is located approximately 250 km to the southwest.

Systematic measurements of aerosol properties have been conducted at MUK since 2005 in Indo-Finnish cooperation with the Finnish Meteorological Institute (Hooda et al., 2018 and references therein). Here, we used only two years (January 2012 to December 2013) of measurements of particle number size distributions in the size range of 10 nm to 757 nm (30 size bins). The air sampling procedure was similar to that of the RNC observation site.

Delhi observation site (DEL, 28.64° N, 77.17° E, 215 m amsl) is located at CSIR-National Physical Laboratory (NPL). Delhi, India’s national capital and largest metropolitan city in South Asia, is located in the northwestern Indo Gangetic Plain (IGP) in northern India. Delhi city has a population of 16.8 million, with a population density of 11,320 km\(^{-2}\) (Census of India, 2011). The Great Indian Desert (Thar Desert) of Rajasthan state is located to the southwest, hot central plains to the south, and hilly regions to the north and east of Delhi. Long-range transported air masses often influence Delhi’s air quality from the northwest (agricultural residue burning from Punjab and Haryana in October-November) and southwest (dust storms from Thar and Arabian Peninsula in April-June) (Kanawade et al., 2020b; Srivastava et al., 2014). Wide Range Aerosol Spectrometer (WRAS, manufactured by GRIMM, Germany), installed on the second floor of the NPL main building, was used to measure particle number size distributions. WRAS consists of a Scanning Mobility Particle Sizer (SMPS) and an Environmental Dust Monitor (EDM). GRIMM-SMPS system consists of a Vienna-type monodisperse differential mobility analyzer (M-DMA). DMA classifies the particle according to their electrical mobility, which is then counted using a CPC. EDM uses an Optical Particle Counter (OPC), which works on the light scattering technology for particle counting gives the particle number size distribution in the size range from 250 nm to 32 μm (Grimm and Eatough, 2009). Thus, the WRAS system gives the particle number...
size distribution in the size range from 5.5 nm to 32 μm (72 size bins). The detailed description and principle of the instrument are discussed elsewhere (Grimm and Eatough, 2009). In this study, we used particle number size distributions in the size range of 5.14 nm to 1000 nm from November 2011 to January 2013.

Mahabaleshwar observation site (MBL, 17.92° N, 73.65° E; 1378 m amsl) is located in the small town named Mahabaleshwar in the forested Western Ghats range in the Satara district of Maharashtra State. In MBL, a High-Altitude Cloud Physics Laboratory (HACPL) was established by the Indian Institute of Tropical Meteorology (IITM), Pune, in 2012, to study monsoon clouds in this region. HACPL site details are found elsewhere (Anil Kumar et al., 2021). Mahabaleshwar town is a tourist attraction consisting of dense vegetation, residential houses, hotels, and a rural market. Pune city is located on the leeward side of the Western Ghats about 100 km to the north, Mumbai city is located approximately 250 km on the northwest, and Satara city is located approximately 50 km to the southeast of Mahabaleshwar. Measurements of particle number size distributions were carried out using the GRIMM-WRAS system. The detailed description and principle of the instrument are discussed elsewhere (Grimm and Eatough, 2009). The sampling probe uses a Nafion dryer to reduce the relative humidity to ~40%. In this study, we used particle number size distributions in the size range of 5.14 nm to 1000 nm from March 2015 to March 2016.

Hyderabad observation site (HYD, 17.46° N, 78.32° E; 542 m amsl), University of Hyderabad, is located in the outskirts of Hyderabad urban city. HYD observation site details can be found in Sebastian et al. (2021a). Briefly, particle number size distributions in size range from 10.9 to 514 nm (108 size bins) were measured using TSI SMPS, which consists of an electrostatic classifier with a long differential mobility analyzer (TSI LDMA, model 3082) and a butanol CPC (TSI, model 3772), on the second floor of the Earth Sciences building located in the University of Hyderabad campus from April 2019 to March 2020. The scanning cycle of SMPS was 300 seconds, yielding a particle number size distribution every 5 minutes.

Thiruvananthapuram (Trivandrum) observation site (TVM, 8.55° N, 76.97°E, 3 m amsl) is a tropical semi-urban coastal city with a population of ~1 million (Census of India, 2011), located on the southwestern coast of the Indian peninsular. The observations were carried at the Space Physics Laboratory (SPL) within the Thumba Equatorial Rocket Launching Station, which is about 500 m due east of the Arabian Sea coast and 10 km northwest of the urban area of
Thiruvananthapuram. The experimental site is free from major industrial or urban activities (Babu et al., 2016). TVM station is a part of the Aerosol Radiative Forcing over India (ARFI) project network of the Indian Space Research Organisation - Geosphere-Biosphere Program (ISRO-GBP). Measurements of particle number size distributions in size range from 14.6 nm to 661.2 nm (108 size bins) were made using TSI SMPS, which consists of an electrostatic classifier with an LDMA (3081) and a water-based CPC (3786) from January 2013 to January 2014. More details about the site and prevailing meteorology are described in Babu et al. (2016).

Particle number size distributions are categorized by season. We have defined four seasons as indicated in Table 2. The overall particle number size distribution data coverage was adequate (>60%) at the RNC, MUK, MBL, and HYD sites (Fig. 2) for determining the main seasonal and annual features of particle number size distributions and NPF characteristics. The data coverage at TVM (34%) and DEL (47%) was lower. We also analyzed the number concentration of three sub-micron aerosol modes: Aitken mode (25-100 nm), accumulation mode (100-514 nm), and total particles (<514 nm).

Table 2. Seasons are defined in the analysis and average weather conditions.

<table>
<thead>
<tr>
<th>Season</th>
<th>Months</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Winter</td>
<td>December, January, February</td>
<td>Cold and dry</td>
</tr>
<tr>
<td>Pre-monsoon</td>
<td>March, April, May</td>
<td>Hot and dry</td>
</tr>
<tr>
<td>Monsoon</td>
<td>June, July, August, September</td>
<td>Warm, humid, and wet</td>
</tr>
<tr>
<td>Post-monsoon</td>
<td>October, November</td>
<td>Cool and humid</td>
</tr>
</tbody>
</table>
**Figure 2.** Particle number size distributions data coverage (% of days/month) at the sites. The values in the bracket indicate total data coverage. The blue, red, green, and grey colored thick lines indicate winter, pre-monsoon, monsoon, and post-monsoon months.

### 2.2 New particle formation event classification and features

We classified observation days into three types of events: NPF event day, non-event day, and undefined event day using visual inspection of the particle number size distributions following the methodology given by Dal Maso et al. (2005). A day was classified as an NPF event day by the presence of a distinctly new mode of particles with a diameter smaller than 25 nm and steady growth in diameter of this new mode such that the particle number size distributions display a noontime "banana" shaped aerosol growth. The particle mode diameter (i.e., the local maximum of the particle number size distribution) was obtained by fitting a log-normal distribution to the measured particle number size distribution. A day without any evidence of a distinctly new mode...
of particles diameter smaller than 25 nm was identified as a non-event day. Those days, which were difficult to be classified as any one of the above two event types, were identified as undefined event days. For NPF events, the particle growth rate (GR) was calculated by fitting a first-order polynomial line through growing particle mode diameter between the lowest detectable size (LDS) of the instrument (e.g., 10 nm for RNC) and 25 nm as a function of time and calculating its slope. The formation rate of a particle at the LDS (\(J_{LDS}\)) was also found using the simplified approximation of the General Dynamic Equation (GDE), describing the evolution of the particle number size distribution as given below;

\[
J_{LDS} = \frac{dN_{LDS-25}}{dt} + F_{\text{CoagS}} + F_{\text{growth}}
\]  

where the first term in Eq. (1) is the rate of the change of nucleation mode particle number concentrations, the second term is the coagulation loss of nucleation mode particles, and the third term is the flux out of the size range of LDS-25 nm, i.e., condensational growth (Dal Maso et al., 2005). A direct comparison of GR and J between all of the sites is not possible because of the different size ranges covered by the instruments.

2.3 Increase in CCN concentrations from NPF

The increase in CCN concentrations from any given NPF event can be estimated by comparing the CCN concentration before the event (\(N_{CCN_{\text{prior}}}\)) and the maximum CCN concentration during the event (\(N_{CCN_{\text{max}}}\)) following the methodology developed by Kerminen et al. (2012), which we modified further. In typical ambient in-cloud supersaturations, the total number of particles from 50 nm to >100 nm can be considered as a proxy for CCN concentrations (Westervelt et al., 2013; Kerminen et al., 2012). \(N_{CCN_{\text{prior}}}\) was chosen to be a one-hour average concentration of particles larger than 50 nm (and 100 nm) just before the start of the NPF event. \(N_{CCN_{\text{max}}}\) was taken as a maximum one-hour average concentration of particles larger than 50 nm (and 100 nm) during the event. The \(N_{CCN_{\text{max}}}\) is not the best representation of CCN concentration after the NPF event because it is not possible to estimate the end of an NPF event. But it gives a rough estimate of the observed maximum number of primary and secondary particles present in the atmosphere during an event (Kerminen et al., 2012). We calculated the seasonally averaged change in CCN-active particles on non-event days over the same time of day as the NPF events,
which would account for the CCN concentrations from processes other than NPF. Then, the absolute increase in CCN concentration from NPF is calculated as given below,

\[
\text{CCN increase} = (N_{\text{CCNmax}} - N_{\text{CCNprior}})_{\text{NPF event}} - (N_{\text{CCNmax}} - N_{\text{CCNprior}})_{\text{non-events}}
\]  

(2)

The first term on the right-hand side in Eq. (2) indicates the CCN increase during an NPF event, while the second term indicates the CCN increase during a non-event. This difference between them allows us distinguishing primary particles and particles formed originally from atmospheric nucleation and yields the best representation of CCN concentrations after the NPF event. But the atmospheric condition on non-event days is generally different from NPF event days; therefore, the calculated increase in CCN concentrations from NPF may be imprecise.

3. Results and discussion

3.1 Variability in particle number size distributions and number concentrations

Figure 3 shows the annual and seasonal median and 25th and 75th percentile values of particle number size distributions at all the sites. The thick line represents the median value, whereas the shaded area indicates particle number size distribution between 25th and 75th percentiles. The annual median particle number size distribution has the smallest mode diameter at DEL compared to the other sites. The smallest mode diameter necessarily indicates the significant near-surface anthropogenic sources at DEL as compared to other sites. The mountain sites (RNC, MUK, and MBL) all show similar mode diameters, with the lowest concentrations at RNC. Amongst urban areas (HYD, TVM, and DEL), TVM has the largest mode diameter, which is frequently influenced by the influx of marine air masses containing high moisture and coarser sea salt aerosols (Babu et al., 2016) (Fig. 3a). The peak number concentration of PNSDs is the highest in pre-monsoon (MAM) than in other seasons at RNC and MUK (Fig. 3b-c), while it was similar in winter and pre-monsoon at MBL (Fig. 3d). These elevated concentrations are accompanied by a smaller mode diameter of the Aitken mode particles. The highest number concentration is attributed to the frequent occurrence of NPF in these locations in pre-monsoon (Sebastian et al., 2021b; Neitola et al., 2011). The contribution of newly formed particles to total particles is also visible in the 75th percentile PNSDs at these sites. The number size distributions of particles were significantly the lowest in monsoon and post-monsoon.
The median number size distribution of particles at HYD is the highest in pre-monsoon and post-monsoon (Fig. 3e). The highest particle number concentrations in pre-monsoon and post-monsoon can be attributed to the frequent occurrence of NPF in these seasons at the site. The influence of NPF is also noticeable in the 75th percentile PNSDs. The PNSD is consistently the lowest in monsoon, attributed to the wet scavenging of particles. The concentrations of Aitken and accumulation mode particles are the highest in winter compared to the other seasons. The mode diameter of PNSDs at TVM is comparatively similar in all seasons (Fig. 3f). At DEL, the mode diameter of PNSDs is the highest in winter compared to the other seasons (Fig. 3g). The shallow boundary layer height, stagnant atmospheric conditions, and high emission rates of aerosol precursors in winter (Kanawade et al., 2020b) allow particles to stay close to the surface and grow larger under high relative humidity and high condensable vapor concentrations. The median PNSD is consistently the lowest in monsoon at TVM due to extensive wet scavenging. The strong seasonality in PNSDs is similar to those reported earlier in India (Hooda et al., 2018; Komppula et al., 2009; Gani et al., 2020; Kanawade et al., 2014a).
Figure 3. (a) Annual and (b-g) seasonal median particle number size distributions at all the sites. The solid line indicates the median, and the light-colored shading indicates 25th and 75th percentile distributions. The blue line and shading indicate winter (DJF), red line and shading indicate pre-monsoon (MAM), green line and shading indicate monsoon (JJAS), and grey line and shading indicate post-monsoon season (ON). Note that the y-axis scale is different for the DEL site.

Figure 4 shows the average observed PNSDs evolving over the day for each season, as a contour plot, at all the sites. For RNC and MUK, the average seasonal contour plot indicates daytime NPF in pre-monsoon. However, winter, monsoon, and post-monsoon had the lowest concentrations of smaller particles that are not associated with NPF. For MBL, NPF occurred in winter, pre-monsoon, and post-monsoon. For all urban sites (HYD, TVM, and DEL), the average seasonal contour plot indicates the highest concentration of particles in morning and evening peak traffic hours, in addition to daytime NPF. In Section 3.2, we investigate the frequency of occurrence of NPF and its contribution to CCN concentrations.
Figure 4. The diurnal-seasonal median particle number size distributions at all the sites; a) Ranichauri, b) Mukteshwar, c) Mahabaleshwar, d) Hyderabad, e) Thiruvananthapuram, and f) Delhi.
Figure 5 shows the box-whisker plot of the number concentrations of Aitken, accumulation, and total particles at all the sites. The median Aitken mode particle number concentrations are the lowest at RNC ($1.4 \times 10^3$ cm$^{-3}$) and the highest at DEL ($7.1 \times 10^3$ cm$^{-3}$). The median accumulation mode particle number concentrations are the lowest at MUK ($0.9 \times 10^3$ cm$^{-3}$) and the highest at DEL ($2.4 \times 10^3$ cm$^{-3}$). The total particle number concentrations are the lowest at MUK ($2.7 \times 10^3$ cm$^{-3}$) and the highest at DEL ($12.5 \times 10^3$ cm$^{-3}$). The median particle number concentrations are about 5-fold higher in urban locations (HYD, TVM, and DEL) than mountain sites (RNC, MUK, and MBL). Overall, the size-segregated particle number concentrations show strong spatial variability, with the lowest concentrations at the mountain sites and the highest at the urban sites. Further, the size-segregated particle number concentrations also show the large variability in each urban location than the mountain sites. Next, we discuss the seasonality in the number concentration of Aitken, accumulation, and total particles in all locations to understand space- and time-varying heterogeneity in particle number concentrations.

**Figure 5.** Box-whisker plot of the size-segregated particle number concentrations using the entire data. The filled square indicates the mean, the horizontal line indicates the median, the top and
bottom of the box indicate 25th and 75th percentile values, and the top and bottom whiskers indicate 10th and 90th percentile values.

The histograms of the relative occurrence of Aitken mode particle number concentrations at all the sites are presented in Figure 6(a-f). RNC and MUK show a similar seasonality in number concentration histograms of Aitken mode particles, with a reasonably log-normal shape and the highest concentrations in the pre-monsoon season. The lowest concentrations are observed in monsoon and post-monsoon due to increased removal of particles by wet-scavenging. MBL does not show notable seasonality in the number concentration histograms of Aitken mode particles. HYD, TVM, and DEL are urban environments but show different seasonality in the number concentration histograms of Aitken mode particles. DEL shows the highest Aitken mode particle number concentrations in winter, and post-monsoon, TVM show the highest concentrations in winter. In contrast, HYD shows comparable number concentrations in winter, pre-monsoon, and post-monsoon. The highest Aitken mode number concentrations in pre-monsoon at mountain-background sites are attributed to the high frequency of NPF occurrence in pre-monsoon (see Sect. 3.2.1). The highest Aitken mode number concentrations in winter at urban sites can be explained by the high pre-existing particle concentration. The difference in seasonality in the number concentration histograms of Aitken mode particles can be explained by the differences in the atmospheric conditions (e.g., prevailing synoptic air masses, mesoscale processes such as atmospheric boundary layer dynamics, and particle removal processes) and considerable heterogeneity in aerosol composition (natural versus anthropogenic aerosol emission sources); DEL is representative of a sub-tropical climate, HYD is representative of a tropical climate, and TVM is representative of a tropical-coastal climate.
Figure 6. Histogram of the relative occurrence of Aitken mode particle number concentrations at the sites. The concentration bins are logarithmically spaced in the x-axis, and the y-axis shows the relative occurrence of values in each bin compared to the total number of valid observations. The thick black line indicates all data. The red, blue, green, and grey lines indicate winter (DJF), pre-monsoon (MAM), monsoon (JJAS), and post-monsoon (ON) months. n indicates the number of 10 minutes averaged valid data points.
Similar histograms of accumulation mode particles are presented in Fig. 7(a-f). The seasonality in accumulation mode particles is slightly different as compared to Aitken mode particles at some sites. RNC shows similar number concentration histograms of accumulation mode particles in winter and pre-monsoon instead of dissimilar histograms for Aitken mode particles. The number concentration histograms of accumulation mode particles at MUK are similar to Aitken mode particles. MBL shows similar number concentration histograms in winter, pre-monsoon, and post-monsoon, with the lowest concentrations in monsoon due to wet scavenging. Among the urban sites, DEL shows the highest accumulation mode concentrations in post-monsoon and winter. TVM and HYD show the highest accumulation mode concentrations in winter and post-monsoon, respectively. The seasonality in total particles was also similar to Aitken mode particles, indicating that Aitken mode particles constituted the most considerable fraction of total particles at all the sites (Figure not shown). However, it is difficult to separate a fraction of Aitken or accumulation mode particles that originated from NPF from that of the primary emissions, especially in urban areas where the primary emission rates of aerosols are very high (Thomas et al., 2019). The survival probability of newly formed particles to >50-100 nm size depends on many factors such as the frequency and intensity of the NPF occurrence, availability of condensable vapors, pre-existing particles, and atmospheric conditions. In Sect. 3.2.3, we estimate the absolute increase of CCN concentrations from NPF following the methodology given by Kerminen et al. (2012) and modified to remove the possible contribution from the primary particles to CCN concentrations for any given NPF event.
3. New particle formation and its contribution to CCN concentrations

3.2.1 NPF event characteristics

The frequency of occurrence of NPF events, the particle formation rate of nucleation mode particles ($J_{NUC}$), and the particle growth rate of nucleation mode particles ($GR_{NUC}$) are typically derived to quantify the NPF (Kerminen et al., 2018; Nieminen et al., 2018; Kulmala et al., 2004).

**Figure 7.** Same as Fig. 6, but for accumulation mode particle number concentrations.
These NPF characteristics are closely associated with aerosol precursor concentrations, pre-existing aerosol particles, and atmospheric conditions. As a result, the frequency of occurrence of NPF events varies from one location to another as well as seasonally. NPF is thought to occur frequently during the spring (pre-monsoon) and rarely during the winter (Kanawade et al., 2012; Dal Maso et al., 2005; Nieminen et al., 2018). However, NPF events were also observed frequently during the thermal winter (Kulmala et al., 2004; Pikridas et al., 2012) and fall (September, October, and November) (Rodríguez et al., 2005). These studies indicate that there is no universal pattern in the occurrence of NPF events. Figure 8 shows the percentage of NPF, non-event, and undefined event days based on valid observation days at all the sites. Out of a total of 586 valid observation days at RNC, NPF events occurred on 21 days (3.9%), whereas 493 (83.7%) days were non-event days. Out of a total of 440 valid observation days at MUK, NPF events occurred on 13 days (2.9%), whereas 321 (73.1%) days were non-event days. Out of a total of 281 valid observation days at MBL, NPF events occurred on 16 days (5.9%), whereas 188 (66.1%) days were non-event days. Out of a total of 270 valid observation days at HYD, NPF events occurred on 38 days (16.3%), whereas 124 (44.8%) days were non-event days. Out of a total of 133 valid observation days at TVM, NPF events occurred on 23 days (16.6%), whereas 55 (41.4%) days were non-event days. Out of a total of 139 valid observation days at DEL, NPF events occurred on 39 days (28.1%), whereas 30 (21.1%) days were non-event days.
Figure 8. Monthly percentage of occurrence of NPF, non-event, and undefined events days based on total valid observations days at all the sites. The blue, red, green, and grey colored thick lines indicate winter, pre-monsoon, monsoon, and post-monsoon months.
3.2.2 Particle formation rate and growth rate

Overall, the frequency of occurrence of NPF is the highest in pre-monsoon as compared to other seasons. There is also an exception to this, with the highest frequency of NPF occurrence in the late monsoon (September) at TVM. Babu et al. (2016) have reported that NPF events over this site occurred due to a mixing of contrasting air masses due to the combined effect of mesoscale land-sea breeze circulation and local ABL dynamics. Though prevailing air masses are oceanic, the wind speeds and total rainfall were lower during September than other monsoonal months. A cleaner synoptic air mass (i.e., lower background concentrations and condensation sink), combined with the occurrence of well-defined mesoscale land-sea breeze transitions and horizontal convergence of contrasting air masses during September, was responsible for the highest NPF frequency. Amongst the sites, the mountain-background sites in the Western Himalaya (RNC and MUK) have the lowest annual mean frequency of occurrence of NPF (3.9% and 2.9%, respectively), with the highest seasonal frequency of occurrence of NPF in pre-monsoon. Previous studies also showed the infrequent occurrence of NPF at RNC (Sebastian et al., 2021b) and MUK (Neitola et al., 2011), with the highest frequency in pre-monsoon. The highest NPF frequency in pre-monsoon was connected to the planetary boundary layer uplifting to the measurement site elevation that appeared to transported aerosol precursors from nearby polluted lower-altitude regions (Hooda et al., 2018; Raatikainen et al., 2014). However, NPF occurred frequently (39%) at the Nepal Climate Observatory-Pyramid (NCO-P) site in the Eastern Himalaya (Venzac et al., 2008). A recent study also observed a very high NPF frequency (69%) at NCO-P from November to December when cleaner conditions prevailed, with little transportation from the polluted lower-altitude regions (Bianchi et al., 2021). They showed that up-valley winds bring gaseous aerosol precursors to higher altitudes. These precursors are oxidized into compounds of very low volatility and are subsequently converted into new particles during their transport to the site. The above discussion indicates that RNC and MUK mountain-background sites in the Western Himalayas are strikingly different from the NCO-P site in the Eastern Himalayas (Bianchi et al., 2021). The annual NPF frequency at RNC and MUK is lower than MBL and the high-altitude sites in Europe (Nieminen et al., 2018). DEL has the highest frequency of occurrence of NPF events in pre-monsoon (63.8%), followed by HYD (28.4%) and MBL (15.9%). TVM coastal semi-urban site witnesses frequent NPF events under the influence of continental air masses. As the air masses change from continental to mixed or marine origin, the NPF event frequency decreases (Babu et
NPF was also observed commonly at other urban sites in India (Kanpur and Pune) under a high source of aerosol precursors when pre-existing particle concentrations reduced sufficiently due to dilution (Kanawade et al., 2020a; Kanawade et al., 2014b). While the severe air pollution episode in Delhi in November 2016 suppressed the NPF, the co-condensation of vapors of anthropogenic origin along with water onto primary particles assisted the rapid particle growth (1.6 to 30.3 nm h\(^{-1}\)) (Kanawade et al., 2020b). The emission of precursor compounds from traffic and other sources in Beijing, China, also contributed significantly to the molecular cluster formation, particle growth and secondary aerosol mass formation, leading to haze formation under favorable meteorological conditions (Kulmala et al., 2021). In Europe, the atmospheric conditions (such as the solar radiation and relative humidity) appear to dictate the NPF occurrence at rural sites, whereas the increased concentrations of precursor gases are more important for the occurrence of NPF in urban areas (Bousiotis et al., 2021). This explains why NPF occurs more frequently in urban areas than rural, remote or high-altitude locations (Guo et al., 2020; Nieminen et al., 2018; Sellegri et al., 2019). This also indicates that the balance between the precursor concentration and pre-existing particles plays a vital role in the NPF occurrence. Owing to large spatial heterogeneity in aerosol precursor emissions and background aerosol concentrations in India, the chemical species contributing to aerosol nucleation and growth is unidentified (Kanawade et al., 2021). The atmospheric NPF can be quantified by calculating \( J_{\text{NUC}} \) and \( GR_{\text{NUC}} \) for the observed NPF events, which we discuss next.

Figure 9 shows the scatter plot of the \( J_{\text{NUC}} \) and the \( GR_{\text{NUC}} \) as a function of condensation sink at each site. A fairly good correlation between \( J_{\text{NUC}} \) and \( GR_{\text{NUC}} \) at each site (Pearson correlation coefficient of 0.48, 0.78, 0.85, 0.33, 0.68, and 0.18 at RNC, MUK, MBL, HYD, TVM, and DEL, respectively) indicates that \( J_{\text{NUC}} \) and \( GR_{\text{NUC}} \) are strongly coupled. The large scatter in data points is a result of important factors influencing the NPF, such as nucleation mechanisms (Dunne et al., 2016), the availability of other condensable vapors that are needed to stabilize molecular clusters containing sulfuric acid (Kirkby et al., 2011; Schobesberger et al., 2015), and atmospheric conditions (Bousiotis et al., 2021). A recent study showed that amines stabilize the nucleating cluster while organics contribute to higher concentrations of condensable vapors, particularly in urban areas (Xiao et al., 2021). The formation rate of 10 nm particles at mountain-background sites (RNC and MUK) varied from 0.01 to 0.1 cm\(^{-3}\) s\(^{-1}\), with a mean value of 0.08 cm\(^{-3}\) s\(^{-1}\). The formation rate of 5 nm particles at MBL varied from 0.02 to 0.1 cm\(^{-3}\) s\(^{-1}\), with a mean of
The formation rate of 10 nm particles at HYD varied from 0.01 to 0.56 cm$^{-3}$ s$^{-1}$, with a mean of 0.13 cm$^{-3}$ s$^{-1}$. The formation rate of 15 nm particles at TVM varied from 0.001 to 0.02 cm$^{-3}$ s$^{-1}$, with a mean of 0.07 cm$^{-3}$ s$^{-1}$. The formation rate of 5 nm particles at DEL varied from 0.01 to 0.5 cm$^{-3}$ s$^{-1}$, with a mean value of 0.12 cm$^{-3}$ s$^{-1}$. The mean growth rates of nucleation mode particles during NPF events were 6.3 nm h$^{-1}$, 2.5 nm h$^{-1}$, 4.7 nm h$^{-1}$, 5.7 nm h$^{-1}$, 1.1 nm h$^{-1}$, and 3.7 nm h$^{-1}$, at RNC, MUK, MBL, HYD, TVM, and DEL, respectively. Considering all the sites, GR$_{NUC}$ during NPF events varied from 0.2 to 17.2 nm h$^{-1}$. Overall, $J_{NUC}$ and GR$_{NUC}$ are within the observed large range of values in diverse environments in India and elsewhere (Nieminen et al., 2018; Kerminen et al., 2018; Kulmala et al., 2004). Expectedly, the condensation sink at the start of the NPF event is higher at urban sites than the mountain sites. The mean condensation sink at urban sites ($16.1 \times 10^{-3}$ s$^{-1}$) was twice as compared to mountain sites ($7.9 \times 10^{-3}$ s$^{-1}$). A previous study also showed that the higher pre-existing particles at Kanpur than at Pune suppressed the particle formation rate but favored the particle growth under high concentrations of condensable vapors (Kanawade et al., 2014b).

**Figure 9.** Scatter plot of the particle formation rate and the growth rate as a function of condensation sink at each site. The condensation sink is taken at the start of the NPF event. The lowest nucleation mode detectable size at each site is shown in the bracket.
3.2.3 Increase in CCN concentrations during NPF events

To reach climatologically relevant sizes, the newly formed particles must grow by condensation while avoiding coagulation removal by pre-existing particles because these freshly formed particles are small and highly diffusive (Vehkamäki and Riipinen, 2012). Based on the observed range of particle growth rates at all the sites (0.2 to 17.3 nm h⁻¹), newly formed particles may take from a few hours to 1-2 days to grow to CCN-active sizes (>50-100 nm). Over such time scales, it is observationally challenging to separate CCN originating from NPF from those emanating from the growth of small primary particles and direct emission of CCN-active sized particles. The increase in CCN concentrations during any given NPF event was estimated following the methodology developed by Kerminen et al. (2012), which we modified to remove CCN originating from the growth of small primary particles and direct emission of CCN-active sized particles based on non-event days.

Figure 10 shows the box-whisker plot of the absolute increase in CCN concentrations (50 and 100 nm) at all the sites. Considering all NPF events at mountain sites, increase in CCN₅₀ ranged from 168 cm⁻³ per event to 5.2×10³ cm⁻³ per event, with a median value of 2.7×10³ cm⁻³ per event, whereas the increase in CCN₁₀₀ ranged from 0.02×10³ cm⁻³ per event to 1.9×10³ cm⁻³ per event, with the median value of 1.0×10³ cm⁻³ per event. The increase in CCN₅₀ and CCN₁₀₀ is about two-fold lower than the free tropospheric site, Chacaltaya (5240 m amsl, Bolivia), for NPF events started in the boundary layer (5.1×10³ cm⁻³ per event and 1.5×10³ cm⁻³ per event for 50 and 100 nm, respectively) (Rose et al., 2017). The median increase in CCN₅₀ and CCN₁₀₀ at RNC (2.3×10³ cm⁻³ per event and 0.9×10³ cm⁻³ per event) and MUK (2.9×10³ cm⁻³ per event and 0.9×10³ cm⁻³ per event) are comparable to those reported at Botsalano (1420 m amsl, South Africa); 2.5×10³ cm⁻³ per event and 0.8×10³ cm⁻³ per event, respectively, but about three-fold higher than those reported at a remote continental site in Finland (1.0×10³ cm⁻³ per event and 0.2×10³ cm⁻³ per event for 50 nm and 100 nm, respectively) (Kerminen et al., 2012). Considering all NPF events at the urban sites, CCN₅₀ increase ranged from 0.08×10³ cm⁻³ per event to 9.4×10³ cm⁻³ per event, with a median value of 4.3×10³ cm⁻³ per event, whereas CCN₁₀₀ increase ranged from 0.03×10³ cm⁻³ per event to 4.9×10³ cm⁻³ per event, with a median value of 1.2×10³ cm⁻³ per event. These values are about two-folds lower as compared to values reported at the station of San Pietro Capofiume, in a polluted region of the Po Valley; 7.3×10³ cm⁻³ per event and 2.4×10³ cm⁻³ per event, respectively for 50 nm and 100 nm (Laaksonen et al., 2005). The overall effect of NPF
events on the CCN concentrations at DEL was the largest since the high background number concentrations of CCN$_{50}$ and CCN$_{100}$ resulted in a smaller relative increase, particularly in post-monsoon and winter seasons when compared to the other sites. In order to comprehensively investigate the atmospheric CCN budget and the contribution of NPF to it, Kerminen et al. (2012) pointed out that the analysis should include not only NPF events but also non-event days. Therefore, the modified methodology applied here following Kerminen et al. (2012) provides the best representative of the increase in CCN concentrations for an NPF event.

**Figure 10.** Box-whisker plot of absolute increase in CCN concentrations for (a) 50 nm and (b) 100 nm particles at all the sites based on the observed NPF and non-event events. The filled square indicates the mean, the horizontal line indicates the median, the top and bottom of the box indicate 25$^{th}$ and 75$^{th}$ percentile values, and the top and bottom whiskers indicate 10$^{th}$ and 90$^{th}$ percentile values.

The sites with low pre-existing particle concentrations (hence, low condensation sink values), high solar radiation, and cooler temperatures at high-altitude (or free tropospheric) (RNC, MUK, and MBL) should favor NPF with enhanced frequency as compared to near-surface urban environments (HYD, TVM, and DEL) wherein pre-existing particles concentration are high, leading to faster removal of nucleating vapors. However, NPF in polluted environments occurs more often than expected, with enhanced growth rates (Yu et al., 2017). Guo et al. (2014) also reported that NPF leads to winter-time haze formation in Beijing. Kulmala et al. (2021) recently showed that >65% of the number concentration of haze particles resulted from NPF in Beijing.
The observation sites at altitudes higher than 1000 m amsl also favored NPF at the high condensation sinks and linked precursor gases needed to initiate nucleation and early growth (Sellegri et al., 2019). Therefore, the low condensation sinks are not necessarily required to trigger nucleation and early growth, provided there are high vapor production rates. Because the higher pre-existing particle concentration is an indication of precursor-laden air, but when the condensation sink gets very high, it inhibits aerosol nucleation. Further, at Hyderabad, about half of the NPF events did not display aerosol nucleation (sub-3nm particle formation) with subsequent growth of these particles to larger sizes (>10 nm), perhaps due to lower organic vapor concentrations (Sebastian et al., 2021a). Rose et al. (2017) also reported a high frequency of NPF occurrence for boundary layer (48%) than free troposphere (39%) conditions at Chacaltaya mountain (5240 m amsl), Bolivia. Thus potential CCN formation was higher for NPF events initiated in the boundary layer (67%) than free troposphere (53%). Sellegri et al. (2019) reviewed the CCN concentrations from NPF events in the boundary layer and high-altitude locations. They revealed that the CCN production is the highest at San Pietro Capofiume, a polluted region of the Po Valley ($7.3 \times 10^3 \text{ cm}^{-3}$) (Laaksonen et al., 2005) as compared to high-altitude sites (Rose et al., 2017; Kerminen et al., 2012). Our findings are similar to these studies showing the highest increase in CCN concentrations in urban locations (HYD, TVM, and DEL) compared to mountain locations (RNC, MUK, and MBL) in India. It is not possible to track the nucleated particle until it becomes a CCN, and they are always mixed with CCN originating from primary sources. This makes it extremely difficult to estimate CCN arising from a given NPF event. In the light of the above discussion, these results offer some insights into potential CCN concentrations originating from NPF.

4 Conclusions

In this study, we used at least one year of asynchronous particle number size distribution measurements from six locations in India, consisting of mountain background sites (Ranichauri and Mukteshwar), mountain rural site (Mahabaleshwar), urban sites (Delhi and Hyderabad), and semi-urban coastal site (Thiruvananthapuram). The results from this study provide some insights into the processes influencing particle number size distributions and CCN concentrations in different environments (mountain and urban) of India.
We found that the regional NPF was most common in the pre-monsoon (spring) at all the measurement sites, with an exception at TVM where NPF occurred mostly in the late monsoon season (September), which was linked to the inflow of continental air masses that provided a source of low volatile vapors for nucleation. During pre-monsoon, DEL has the highest frequency of NPF occurrence (63.8%), followed by HYD (28.4%) and MBL (15.9%). NPF was the least common during winter at all the sites, particularly at the mountain-background sites (RNC and MUK) without a single NPF event. The high solar insolation (active photochemistry) and the elevated boundary layer (efficient ventilation leading to low pre-existing particles near the surface) explains the most common occurrence of NPF in the pre-monsoon (spring), but this is not a universal NPF frequency pattern in India and elsewhere globally. We found that the J_{NUC} during NPF events tends to increase with an increasing anthropogenic influence, with an order of magnitude higher in urban areas (0.12 cm$^{-3}$ s$^{-1}$) than mountain sites (0.06 cm$^{-3}$ s$^{-1}$). We did not find any systematic pattern in GR_{NUC}, with the highest GR_{NUC} at RNC (6.3 nm h$^{-1}$) and the lowest at TVM (1.1 nm h$^{-1}$). The observed values of the NPF frequency, J_{NUC}, and GR_{NUC} indicate that the regional NPF events can significantly influence the evolution of particles in the atmosphere. We found that NPF modulates the shape of the particle number size distributions significantly, especially at the mountain background sites (RNC and MUK), which are not directly influenced by the local direct emissions of aerosols (traffic and industries). The number size distribution of particles is higher in pre-monsoon at mountain-background sites, whereas it is higher in winter at urban sites, with the exception of HYD. All sites generally show lower concentrations of particles in monsoon due to the increased removal by wet-scavenging. The histograms of size-segregated particle number concentrations show large variability from one site to another, reflecting the varying contribution of different processes to the total aerosol loading. For instance, the Aitken mode particle concentrations were the highest in pre-monsoon at mountain-background sites (RNC and MUK), whereas they were the highest in winter at urban sites (HYD, TVM, and DEL). Amongst the sites, the lowest measured median total particle number concentration was found in MUK (2658 cm$^{-3}$) and the highest in DEL (12519 cm$^{-3}$).

We found that the increase in CCN concentrations during an NPF event is higher in urban locations ($4.3 \times 10^3$ cm$^{-3}$ per event and $1.2 \times 10^3$ cm$^{-3}$ per event for 50 nm and 100 nm, respectively) compared to mountain-background sites ($2.7 \times 10^3$ cm$^{-3}$ per event and $1.0 \times 10^3$ cm$^{-3}$ per event for 50 nm and 100 nm, respectively). We modified Kerminen and colleague’s approach for removing
the potential contribution of primary CCN-active particles to give the best possible estimate for
the increase in CCN concentrations during a given NPF event. Such analyses should be
supplemented by regional model simulations or high spatial resolution measurements of NPF and
CCN concentrations.

**Code availability**
Particle number size distributions data was analyzed in IGOR Pro 8.0. Figure 8 was created in
IGOR Pro 8.0, while all other figures were created in IDL 8.0.

**Data availability**
Particles data will be made available upon a reasonable request to the corresponding author.

**Author contribution:**
VPK conceived the idea and designed the research. MS and VPK carried out a comprehensive
data analysis. MS carried out CCN estimation analysis and interpretation with critical inputs
from JRP, VV, and VPK. MS, SKK, AK, and SJ performed particle size distribution
measurements and analysis. MS and VPK wrote the first draft, and MS edited with critical inputs
from all co-authors.

**Competing interests**
The authors declare that they have no conflict of interest.

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