

The Impact of aerosol–ice nuclei-cloud interactions on a Typical Spring Dust-Precipitation Event in China

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Abstract.

20 To investigate the impact of ice nuclei (IN) activated by dust aerosols on precipitation over China, this study uses regional Global/Regional Assimilation and Prediction System – China Meteorological Administration Unified Atmospheric Chemistry Environment (GRAPES/CUACE). The original temperature-dependent IN nucleation scheme in the Double-Moment 6-Class (WDM6) is improved by
25 incorporating an on-line aerosol–IN nucleation scheme to examine their effects during a typical dust affected precipitation event in East Asia.

Dust modifies the spatial distribution and number concentration of IN, affecting heterogeneous ice nucleation. Compared with the systematic underestimation in original WDM6, the peak values of nucleated INs reach 10^{-4} L^{-1} with the improved
30 scheme, which is closer to observations.

Dust inhibit the development of clouds. Above 7 km, dust suppresses both heterogeneous nucleation and deposition growth. Thus, the total production rate of cloud ice drops to less than 24% of that in the control test T_CTL, promoting snow

formation and reducing the total ice-phase hydrometeor content to 70–85% of T_CTL.

35 Between 4 and 7 km, dust enhances heterogeneous nucleation of cloud ice but suppresses deposition growth, leading to a decrease in the total ice-phase hydrometeor content to 85–91% of T_CTL. Below 4 km, dust suppresses the conversion of water vapor to cloud water, thereby reducing the liquid-phase hydrometeor content to 90–95% of T_CTL.

40 Dust modifies the precipitation distribution, bringing it closer to observations. It suppresses precipitation near dust source areas, where mean precipitation decreased by about 4.5 mm, while the downstream event-mean precipitation increased by about 1.1 mm.

45 **Keywords:** aerosol–IN–cloud–precipitation interactions; dust-precipitation event; on-line aerosol–IN nucleation scheme, CUACE

1 Introduction

The formation of cloud ice is one of the key processes in ice-phase precipitation, and ice nuclei (IN) associated with aerosols play a crucial role in the development of cloud ice, particularly in mid- to high-latitude areas and in the upper troposphere (Li et al., 2022; Chen et al., 2023; Knopf and Alpert, 2023). This is because homogeneous nucleation without IN occurs only below $-40\text{ }^{\circ}\text{C}$, which is relatively rare in natural atmospheric environments (Eastwood et al., 2008; Herbert et al., 2015; Kumar et al., 2020; Che et al., 2021). In contrast, heterogeneous nucleation mediated by IN can occur under ice-supersaturated conditions at much higher temperatures, making it the dominant pathway for cloud ice formation.

Aerosols can serve as IN, participating in cloud formation, altering cloud microphysical properties and lifetimes, and thereby affecting precipitation (Albrecht, 1989; Andreae and Crutzen, 1997). Among different species, mineral dust is recognized as one of the primary sources of atmospheric IN (Khain et al., 2000; Nenes et al., 2014; Tobo et al., 2019). Dust particles have unique surface structures that facilitate the adsorption and binding of water molecules, promoting the formation of cloud ice (Possner et al., 2017; Stevens et al., 2018). Stith et al. (2009) and DeMott et al. (2015) have found a strong correlation between IN number concentration and aerosols with diameters larger than $0.5\text{ }\mu\text{m}$, with mineral dust accounting for 33-50% of the total IN. Jiang et al. (2016) combined IN measurements during dust conditions at multiple sites in China, including Xinjiang (Jiang et al., 2016), Mt. Huangshan (Jiang et al., 2015), and Nanjing (Yang et al., 2013), and found that IN concentrations were significantly higher than those under non-dust conditions. Tobo et al. (2020) observed that IN concentrations increased remarkably during dust events in Tokyo when temperatures were above $-25\text{ }^{\circ}\text{C}$. In addition, aged dust aerosol has increased solubility, which can act as cloud condensation nuclei (CCN) and thereby further influencing precipitation (Trochkin et al., 2003).

Compared with the relatively well-understood impacts of aerosols as CCN, the role of dust as IN is considerably more complex and remains poorly understood, with substantial uncertainties (Kaufman et al., 2002; Eastwood et al., 2008; Pan et al., 2017;

Possner et al., 2017). Based on chemistry (WRF-Chem) model and multiple observational and reanalysis data, Wang et al. (2024) found that dust aerosols can suppress light precipitation by increasing atmospheric stability and inhibiting the conversion of cloud droplets into raindrops. In contrast, Naeger (2018) found that dust could enhance precipitation over Florida based on multi-sensor satellite observations and field campaigns. More recently, Hu et al. (2023) demonstrated that the impact of springtime dust on precipitation is strongly modulated by the presence of other aerosol types. Liu et al. (2024) analyzed the spatiotemporal patterns and trends of dust aerosols and precipitation and found that dust increases suppress precipitation over source regions such as the Gobi and Taklamakan deserts, but enhance precipitation in downwind areas like northern China. Overall, due to the multiple factors influencing precipitation beyond aerosols, it remains challenging to quantify the impact of dust on precipitation from observations alone (Zhou et al., 2016; Stier et al., 2024), highlighting the need for process-oriented numerical modeling studies with physically based aerosol–ice nucleation parameterizations.

Numerical model is a crucial approach for numerically studying the impact of dust on precipitation. In early cloud microphysics scheme, the ice nucleation scheme did not account for aerosols, with IN concentrations typically expressed as functions of temperature or supersaturation (DeMott et al., 2010). Moreover, many clouds ice microphysical schemes were single-moment, which only simulated the mass mixing ratio of cloud ice. Such single-moment schemes often led to large biases in cloud ice mass concentrations (Molthan and Colle, 2012; Igel et al., 2015). In contrast, double-moment ice schemes, which simulate both cloud ice mass and number concentrations, outperform the single-moment schemes in terms of the simulated structure, life cycle, cloud coverage, precipitation, and microphysical properties (Pu and Lin, 2015; Zhao et al., 2021). The double-moment ice schemes can provide more stable and improved precipitation simulations (Kang et al., 2018; Shen et al., 2022; Shen et al., 2024). Mascioli et al. (2021) used the Thompson aerosol-aware microphysics scheme, incorporating the IN nucleation scheme of DeMott et al. (2010), to study the sensitivity

of precipitation to different prescribed dust aerosol concentrations. Park and Lim (2023) develops the revised Weather Research and Forecasting Double-Moment 6-class (WDM6) scheme through the implementation of prognostic cloud ice number concentrations. The excess generation of cloud ice mixing ratio is considerably alleviated. However, these studies did not establish an explicit quantitative relationship between on-line aerosols and IN. Su and Fung (2018a) implemented the simplified Goddard Chemistry Aerosol Radiation and Transport aerosol model (GOCART) together with Shao's dust emission scheme (Kang et al., 2011; Shao et al., 2011) in WRF/Chem and incorporated the online IN nucleation scheme of DeMott et al. (2015) for producing real time IN into the double-moment Thompson–Eidhammer microphysics scheme. They analyzed the impact of dust on radiative forcing and temperature in East Asia, but only the sensitive impacts in terms of precipitation rate in March and April in 2012 (Su and Fung, 2018b). The spring of 2012 is not a typical dust season, most dust storm concentrated in Mongolia. Therefore, the microphysical pathways through which dust affects precipitation during typical dust events remain insufficiently studied. In this study, we focus on a representative spring dust–precipitation event and explicitly examine the cloud microphysical processes associated with dust-induced heterogeneous ice nucleation, together with direct comparisons to precipitation observations in the Global/Regional Assimilation and PrEdiction System, China Meteorological Administration (CMA) Unified Atmospheric Chemistry Environment (GRAPES/CUACE) model. GRAPES/CUACE provides on-line sectional aerosol concentrations with multi chemical composition information (Wang et al., 2010; Zhou et al., 2012). Zhou et al. (2016) introduced an on-line aerosol–CCN–cloud interaction scheme into the system, allowing the model to simulate real time CCN activation and their influence on precipitation. However, in the GRAPES/CUACE microphysics scheme WDM6, IN is a function of temperature only, and cloud ice is represented by a single-moment scheme only for the mass mixing ratio (Hong et al., 2004; Zhang et al., 2022). To address these limitations, this study implements a double-moment cloud ice scheme and incorporates an on-line aerosol–IN nucleation scheme to

135 explicitly represent heterogeneous processes. Using this improved framework, we then
investigate the impact of dust on precipitation by a typical dust affected precipitation
event in East Asia. This paper is organized as follows: Section 2 introduces the model
configuration, cloud microphysical processes, on-line aerosol–IN nucleation scheme,
study region, and observational datasets. Section 3 presents the evaluation of the
140 improved model's simulation performance and discusses the effects of dust on
precipitation. Section 4 summarizes the main conclusions of the study.

2 Model description and methodology

2.1 GRAPES/CUACE

The GRAPES is a fully compressible, non-hydrostatic numerical weather model
145 that adopts a semi-implicit and semi-Lagrangian discretization scheme (Chen et al.,
2008; Xu et al., 2008; Zhang and Shen, 2008; Wang et al., 2022a). The physical
packages include cumulus convective, single-moment cloud microphysics, radiative,
land surface, and boundary layer processes. CUACE is a regional chemical weather
forecasting system developed by Gong and Zhang (2008) coupled on-line with
150 GRAPES (Wang et al., 2010). It is capable of simulating on-line seven aerosol species
of sulfate, nitrate, ammonium, black carbon, organic carbon, sea-salt together with dust
(Zhou et al., 2008, 2012; Wang et al., 2015). The sectional dust emission scheme is by
Marticorena and Bergametti (1995) and Alfaro and Gomes (2001) which has been
improved by surface dust flux observations and desertification in East Asia (Gong et al.,
155 2003), and new desertification map and soil texture samples from Chinese deserts
(Zhou et al., 2019; Zhou et al., 2024). The aerosol size spectra have been divided into
12 size bins with a radius range of 0.005–0.01, 0.01–0.02, 0.02–0.04, 0.04–0.08, 0.08–
0.16, 0.16–0.32, 0.32–0.64, 0.64–1.28, 1.28–2.56, 2.56–5.12, 5.12–10.24, and 10.24–
20.48 μm . GRAPES/CUACE has a horizontal resolution of 0.15° and 31 vertical levels
160 extending to approximately 28.6 km in altitude.

2.2 WDM6 microphysics scheme

In this study, we select the WDM6 microphysics scheme in GRAPES for
simulating precipitation (Hong et al., 2004; Zhang et al., 2022) . The WDM6 scheme

simulates the mass mixing ratio of water vapor (Q_v), as well as the mass and number
165 concentrations of cloud water (Q_c) and rainwater (Q_r) in warm clouds. For icy clouds,
it includes the mass mixing ratios of cloud ice (Q_i), snow (Q_s), and graupel (Q_g). A
double-moment cloud ice scheme by Park and Lim (2023) is incorporated into the
WDM6 scheme, allowing for the explicit prediction of cloud ice number concentration.
A sectional CCN activated scheme has been introduced in WDM6 in GRAPES/CUACE,
170 connecting the multi-component multi-section aerosols from CUACE into the WDM6
microphysics and the sub-grid convective parameterization scheme by newly activated
CCN at each time step (Zhou et al., 2016).

2.3 On-line aerosol-IN nucleation scheme

In the original WDM6 scheme, when the temperature is below 0 °C, the increase
175 in cloud ice mass concentration arises from two processes: heterogeneous nucleation
(Pigen) and deposition–sublimation of cloud ice (Pidep, when positive). Both consume
water vapor to form ice clouds. The abbreviations for the remaining cloud
microphysical processes are listed in Table 2. The IN concentration, N_{ice} (m^{-3}), is
calculated by a classical ice nuclei nucleation scheme, which is an empirical function
180 of temperature and does not account for the influence of atmospheric aerosols (Hong et
al., 2004):

$$N_{ice} = 10^3 e^{0.1(T_0 - T_k)} \quad (1)$$

Where, T_k is atmospheric temperature, T_0 is the freezing point (273.15 K).

This study implements an on-line aerosol-IN nucleation scheme in
GRAPES/CUACE that accounts for heterogeneous ice nucleation processes influenced
185 by atmospheric aerosols. Heterogeneous nucleation mechanisms are generally
classified into immersion freezing, condensation freezing, deposition nucleation, and
contact freezing (Hiranuma et al., 2015; Ilotoviz et al., 2016; Lee et al., 2017). Among
these mechanisms, immersion freezing, condensation freezing, and deposition
nucleation are selected, as they are relatively well developed. This selection is based on
190 the fact that dust aerosols primarily affect ice nucleation at temperatures below 258.15
K through these three mechanisms (Cantrell et al., 2013; Patnaude et al., 2025), whereas

the efficiency of contact freezing by dust particles is relatively low (Niehaus et al., 2014).

Immersion freezing is a heterogeneous ice nucleation process with existence of liquid drops at temperatures between 233.15 K and 273.15 K, which ice nucleus immersed in supercooled liquid, triggering it freezing into an ice crystal (Boose et al., 2016). Immersion freezing consumes water vapor to form cloud ice. The initial size of the ice crystal is influenced by the size of the liquid droplet (Fan et al., 2014; Gibbons et al., 2018), therefore the cloud ice formation through this mechanism is relatively easier compared to other nucleation mechanisms. The selected immersion freezing nucleation scheme here is developed by DeMott et al. (2015), based on continuous flow diffusion chamber measurements. The number concentration of ice nuclei, N_{icenui} (m^{-3}), activated via immersion freezing is given by:

$$N_{icenui} = 3 * n_{aer,0.5}^{1.25} * e^{(0.46*(273.16-T_k)-11.6)} \quad (2)$$

Where, $n_{aer,0.5}^{1.25}$ is the number concentration of insoluble aerosol particles with diameters exceeding 0.5 μm such as dust, black carbon and part of organic carbon.

Deposition and condensation freezing are both heterogeneous ice nucleation processes that occur at temperatures between 248.15 K and 258.15 K (Chen et al., 2019). In condensation freezing, water vapor first condenses on the surface of IN and subsequently freezes to form an ice crystal, while in deposition nucleation, water vapor directly deposits onto the IN surface (Kanji et al., 2017). The the ice formation through these two pathways is generally harder than that through immersion freezing (DeMott et al., 2015). The parameterization used here follows the formulation of Chen et al. (2019). In Jiang et al. (2016), the ice-nucleating ability of dust aerosols was derived from measurements conducted at several sites in China (Yang et al., 2013; Jiang et al., 2015; Jiang et al., 2016), using a static vacuum vapor diffusion chamber based on the FRIDGE (Frankfurt Ice Nuclei Deposition Freezing Experiment) design. Chen et al. (2019) further refined the parameterization to explicitly represent deposition and condensation freezing processes within a specified temperature range. The number concentration of ice nuclei produced by deposition and condensation freezing, $N_{icenuid}$

220 (m^{-3}), is calculated as follows:

$$\begin{aligned} \text{Nicenud} = & 5.7 * 10^{-7} n_{\text{aer},0.5}^{0.018(273.16-Tk)-0.007S_i+0.342} \quad (3) \\ & * (273.16 - Tk)^{3.745} * S_i^{1.31} \end{aligned}$$

Where, S_i is supersaturation with respect to ice.

WDM6 uses the formula $\rho_{q_{I0}}(\text{kg m}^{-3}) = 4.92 \times 10^{-11} N_{\text{ice}}^{1.33}$ and $\text{Pigen}(\text{kg kg}^{-1} \text{ s}^{-1}) = \frac{(q_{I0} - q_I)}{\Delta t}$ to calculate newly nucleation of ice. Where, ρ denotes the newly-formed air density, and q_{I0} is the predicted ice mixing ratio (kg kg^{-1}). Δt is the
225 integration time step. Production rate for heterogeneous nucleation is calculated as the difference between q_{I0} and the current ice mixing ratio (q_I). However, it does not account for the influence of nucleated IN size or the specific characteristics of different heterogeneous ice nucleation mechanisms on ice crystal development.

Here, the mass production rate of cloud ice newly nucleated is calculated as the
230 following:

$$\begin{aligned} \text{Pinud} &= \frac{4}{3} \pi \frac{\rho_i}{\rho_a} (r_{\text{df}}^3 N_{\text{icenud}}) / \Delta t \quad (4) \\ \text{Pinui} &= \frac{4}{3} \pi \frac{\rho_i}{\rho_a} (r_{\text{if}}^3 N_{\text{icenui}}) / \Delta t \end{aligned}$$

Where, Pinud ($\text{kg kg}^{-1} \text{ s}^{-1}$) is mass production rate for deposition/condensation freezing, Pinui ($\text{kg kg}^{-1} \text{ s}^{-1}$) is for immersion freezing. Pinud depletes water vapor to form cloud ice, while Pinui depletes cloud water to form cloud ice. ρ_i is 500 kg m^{-3} (Park and Lim, 2023). r_{if} represents the initial radius of cloud ice formed via immersion freezing,
235 while r_{df} represent the initial radius of cloud ice formed through deposition and condensation freezing, respectively. Δt is the integration time step. In the new online scheme, production rate for nucleated IN number concentration (N_{igen}) is the sum of N_{icenui} and N_{icenud} .

The typical range of ice crystal radius in East Asia is about 10–100 μm (Chen et al.,
240 2021), droplet radius range is about 1~30 μm (Um et al., 2018; Yang et al., 2021). Considering ice crystals generally grow from smaller particles and the radius of initial ice crystal size are often smaller than observed values, and with reference to the bin sizes of aerosol particles in CUACE (Um et al., 2018; Chen et al., 2021; Yang et al.,

2021), this study assumes the characteristic radius of ice crystals of r_{df} and r_{if} to be:

$$\begin{cases} r_{df} = 10 \mu\text{m} (r_{aer} < 10 \mu\text{m}) \\ r_{df} = 30 \mu\text{m} (r_{aer} > 10 \mu\text{m}) \\ r_{if} = 30 \mu\text{m} (r_{aer} < 10 \mu\text{m}) \\ r_{if} = 50 \mu\text{m} (r_{aer} > 10 \mu\text{m}) \end{cases} \quad (5)$$

245 Then, the original production rate for nucleation of ice from vapor Pigen in WDM6 is replaced by the Pinud and Pinui described above.

2.4 Case description and test design

The typical dust affected precipitation event

The typical dust affected precipitation event is from 00:00 UTC on 9 April to 00:00
250 UTC on 15 April 2018, which contains two dust storms events in East Asia. One is from 9 to 11 April, originating in Mongolia and affected northern China. Lots of dust storm phenomena are observed in Mongolia, while blowing dust and floating dust phenomena are reported in central and western Inner Mongolia, central Gansu, Ningxia, northern Shaanxi, most parts of Shanxi, southern Hebei, northern Henan, and western Shandong
255 in China (see Fig. S1 for locations). Another event is from 13 to 14 April. It also gains with widespread dust storm phenomena in Mongolia and central Inner Mongolia, blowing or floating dust phenomena observed in central Inner Mongolia, northern Shanxi, Beijing, Tianjin, and northern Hebei in China. Between the two dust storm events, the precipitation occurred from west to east covering most of northern China
260 extending to the Yangtze River area, from 00:00 UTC on 12 April to 00:00 UTC on 15 April, concentrated in Shaanxi, Henan, southern Hebei, and along the Yangtze River in Sichuan, Hubei, Anhui, and the Jiangsu-Zhejiang-Shanghai area.

Figure 1a presents the dust-affected areas by dust phenomenon from Meteorological stations and PM_{10} from the National Environmental Monitoring
265 Network of the Ministry of Environmental Protection. Based on the distribution of dust in this event, the domain bounded by 90-135 °E and 20-54 °N is defined as the major dust-affected area (DA, region 1 in Figure 1). Together with the real precipitation distribution (Fig. 5a), the domain bounded by 103°–130.5°E and 27.5°–50°N is defined as the dust-affected precipitation (DP) area (DPA, region 2 in Figure 1). The whole

270 model domain covers 70° – 145° E and 15° – 64.5° N, containing the DA and DPA. To investigate the impact of dust on precipitation in regions distant from the dust source in Section 3.3, we calculate horizontal hydrometeor fluxes across 116° E (33° – 50° N) and 33° N (103° – 116° E) during 12:00 UTC on 12 April to 18:00 UTC on 13 April (Fig. 6). The area bounded by 103° – 116° E and 33° – 50° N is defined as the near-dust-source area
275 (NDSA, region 3 in Figure 1).

GRAPES/CUACE successfully reproduces both the spatial distribution and intensity of the dust events (Fig. 1b). Considering that many radar observations and model studies have indicated that dust mainly participates in heterogeneous ice nucleation as ice nuclei within the mid-tropospheric layer ($-20 - 0^{\circ}$ C) (Haarig et al.,
280 2019; He et al., 2021; He et al., 2023), which corresponds to altitudes between 4 and 7 km in the present case, Fig. 1c shows the simulated dust concentration within this layer.

Test design

As shown in Table 1, two tests are designed. The first test uses the on-line aerosol–CCN–cloud interaction scheme from Zhou et al. (2016), denoted as T_CTL. Based on
285 T_CTL, the second test adds the on-line aerosol-IN nucleation scheme described in Section 2.3, denoted as T_IN.

The successive integration is cut into several three-days-interval with a warm restart. It starts at 00:00 UTC on April 5, 2018 with 6 days spinning up for tracers in CUACE. As simulation time increases, integration errors tend to accumulate (Zhang et al.,
290 al., 2019), and to minimize the influence of initial conditions on precipitation, the simulations in this study were divided into several time segments: 5–8 April, 8–11 April, 11–14 April, and 13–16 April. Among these, the simulation results for 13 April were taken from the 11–14 April experiment to minimize the influence of initial conditions on precipitation development. Except for water vapor, all initial values of hydrometeors
295 are zero for each run. The model outputs 1-hourly precipitation data. To compare with the observed 6-hourly precipitation, the model outputs are temporally interpolated to the time stamps of the observations.

The initial and boundary meteorological conditions for GRAPES/CUACE are from

the Final Operational Global Analysis data produced jointly by the National Centers for
 300 Environmental Prediction (NCEP) and the National Center for Atmospheric Research
 (NCAR) at a temporal resolution of 6 hours and a spatial resolution of 0.15°. The
 anthropogenic emissions are from Multi-resolution Emission Inventory for China (Li
 et al., 2017).

2.5 Data and evaluation methodology

305 The initial and boundary meteorological conditions for GRAPES/CUACE are
 obtained from the NCEP/NCAR Final Operational Global Analysis (FNL) data, with a
 temporal resolution of 6 hours and a spatial resolution of 0.25°. Dust observations are
 obtained from two sources: weather phenomena from the CMA surface meteorological
 observation network with a temporal resolution of 3 hours, while PM₁₀ and PM_{2.5}
 310 concentration data from the national environmental monitoring network of the Ministry
 of Ecology and Environment of China, with a temporal resolution of 1 hour. 6-hour
 rainfall data are also from CMA surface meteorological observation network. As there
 are more than 2,000 precipitation stations in DA, only 63 stations of levels 1 and 2 are
 selected for evaluation, of which 43 stations are in DPA to avoid overfitting with the
 315 model outputs. Due to the complex sources of PM₁₀ and considering the relatively long
 atmospheric residence time of dust, we select precipitation stations where the
 PM_{2.5}/PM₁₀ ratio is less than 0.6 within 24 hours prior to the precipitation event as
 representative of dust-influenced precipitation (DP) stations (Wang and Yan, 2007;
 Filonchyk et al., 2019).

320 Model performance is evaluated using mean absolute error (MAE), root mean
 square error (RMSE), and symmetric mean absolute percentage error (sMAPE)
 (Shcherbakov et al., 2013):

$$\begin{aligned} \text{MAE} &= \frac{\sum_{i=1}^n (r_{mi} - r_{oi})^2}{n} & (8) \\ \text{RMSE} &= \sqrt{\frac{\sum_{i=1}^n (r_{mi} - r_{oi})^2}{n}} \\ \text{sMAPE} &= \frac{1}{n} \sum_{i=1}^n \frac{|r_{mi} - r_{oi}|}{|r_{mi}| + |r_{oi}|} \end{aligned}$$

$$\text{aMAPE} = \frac{r_{mi} - r_{oi}}{|r_{mi}| + |r_{oi}|}$$

where r_{mi} represents the simulated cumulative precipitation at station i , and r_{oi} denotes the observed precipitation. For MAE, RMSE and sMAPE, values closer to 0 indicate better simulation performance. The aMAPE is used to evaluate whether the simulated precipitation is overestimated or underestimated compared with the observation. When $\text{aMAPE} > 0$, the precipitation is overestimated; when $\text{aMAPE} < 0$, the precipitation is underestimated.

The horizontal hydrometeor fluxes shown in Section 3.3 are calculated using a grid-based mass transport formulation. For each model layer, the flux is computed as

$$F = \rho_{air} q_x V \Delta z \Delta s \quad (9)$$

where F is the hydrometeor flux (kg s^{-1}), ρ_{air} is the air density (kg m^{-3}), q_x is the mass mixing ratio of the hydrometeor species (kg kg^{-1}), V_n is the wind component normal to the cross section (m s^{-1}), Δz is the layer thickness (m), and Δs is the horizontal grid spacing along the cross section (m).

3 Results

3.1 Ice nuclei

During the DP event, the implemented on-line aerosol-IN nucleation scheme enables dust aerosols to modify the nucleated IN number concentration. Figures 2a and 2b show the horizontal distribution of the maximum nucleated IN number concentration between 4 and 7 km above ground level at DP stations during the time period from 00:00 UTC on 11 April to 00:00 UTC on 15 April 2018 for T_CTL and T_IN, respectively. Figure 2c presents the vertical distribution of DP-event-averaged production rate for Nigen for T_CTL (red line) and T_IN (blue line). Figure 2d presents the vertical distribution of cloud ice mass production rate for heterogeneous ice nucleation for T_CTL and T_IN. Based on the variation characteristics, the vertical layer is divided into three parts: layer A, above 7 km (temperature below $-18\text{ }^{\circ}\text{C}$); layer B, between 4 and 7 km (temperature approximately $-18\text{ }^{\circ}\text{C}$ to $-1.5\text{ }^{\circ}\text{C}$); and layer C, below 4 km (temperature approximately $-1.5\text{ }^{\circ}\text{C}$ to $18\text{ }^{\circ}\text{C}$).

The on-line aerosol-IN nucleation scheme can correct the systematic

350 underestimation of IN concentrations. The maximum nucleated IN number concentrations in T_CTL can reach 10^2 L^{-1} in layer B during the DP event (Fig. 2a), showing a relatively uniform horizontal pattern, which is much lower than observed IN concentrations (10^2 – 10^4 L^{-1}) during East Asian dust events (Bi et al., 2019; Tobo et al., 2019; Chen et al., 2021; Hu et al., 2023). For example, Chen et al. (2021) measured
355 immersion-mode INPs at Peking University Atmosphere Environment Monitoring Station during spring 2018–2019 and found that dust periods increased INP concentrations by approximately two orders of magnitude, reaching 10^2 L^{-1} between -15°C and -28°C . The DP-event-averaged production rate for nucleated IN number concentration ranges 0.005 – $0.01 \text{ L}^{-1} \text{ s}^{-1}$ in layer B (Fig. 2c). In T_CTL, the production
360 rate for nucleated IN number concentration increases with height (Fig. 2c), primarily due to the temperature-dependent nature of original WDM6 scheme. As a result, cloud ice mass production rate due to heterogeneous ice nucleation peak near the -40°C level (Fig. 2d). Above this layer, IN concentration continues to increase, but production rate of heterogeneously nucleated cloud ice begins to decline due to limited water vapor
365 (Fig. 2d). In the real atmosphere, the number concentration of effective ice-nucleating particles often reaches a maximum in the mid-troposphere rather than at the highest altitudes (He et al., 2023), suggesting that the continuous increase of IN at higher altitudes in T_CTL may inconsistent with typical observed . In T_IN, the maximum nucleated IN number concentrations can reach 10^4 L^{-1} in layer B during the DP event
370 (Fig. 2b), closer to those observed or simulated in other East Asian dust events (Bi et al., 2019; Tobo et al., 2019; Chen et al., 2021; Hu et al., 2023). The DP-event-averaged production rate for nucleated IN number concentration ranges from 0.2 to $3.7 \text{ L}^{-1} \text{ s}^{-1}$ in layer B (Fig. 2c), and the cloud ice mass production rate for heterogeneous ice nucleation also peaks in this layer, which is consistent with radar observations and other
375 modeling studies (Haarig et al., 2019; He et al., 2021; He et al., 2023). As immersion freezing is the dominant heterogeneous nucleation mechanism (DeMott et al., 2015; Hiranuma et al., 2015), this study compares the number concentration of ice-nucleating particles activated by immersion freezing with those activated by deposition and

condensation freezing. The DP-event-averaged results indicate that the activated IN
380 number concentration from immersion freezing exceeds that from deposition and
condensation freezing by approximately 4–5 orders of magnitude.

3.2 Hydrometeors

During the DP event, the introduction of the on-line aerosol-IN nucleation scheme
allows dust aerosols to alter the distribution of cloud hydrometeors. Figure 3 shows the
385 DP-event-averaged vertical distributions of hydrometeors in T_CTL and T_IN,
averaged over the dust–precipitation period (00 UTC 11 April–00 UTC 15 April 2018)
and over dust–precipitation stations, as well as their difference ($T_IN - T_CTL$), by
using budget analysis. Figure 4 shows the differences in the production rates of different
hydrometeors ($T_IN - T_CTL$). To further examine the thermodynamic conditions
390 responsible for the weakened production rate of cloud droplet activation from CCN in
T_IN in Figure 4, the vertical profiles of temperature and water vapor were analyzed,
averaged over the dust–precipitation stations in DPA and NDSA during the period when
the dust impact was most pronounced (18:00 UTC 11 April to 18:00 UTC 12 April)
(Figure 5).

395 Cloud ice

In layer A, when dust aerosols are considered, the IN number concentration
decreases in T_IN (Fig. 2c), resulting in cloud ice number concentrations in T_IN that
are approximately 5 L^{-1} lower than those in T_CTL, about 40% of T_CTL (Fig. 3d).
The cloud ice mass concentration is reduced to only 10% - 50% of T_CTL (Fig. 3a,3b).
400 Because the two primary processes contributing to cloud ice formation in this layer—
heterogeneous nucleation and deposition-sublimation of cloud ice—are both
suppressed (Fig. 4a), and the total production rate of cloud ice ($P_{igen}+P_{idep}-P_{saut}-$
 $P_{raci}-P_{saci}-P_{gaci}$) drops to less than 24% of that in T_CTL. On the one hand, the
nucleated IN number concentration decreases, weakening the P_{igen} in T_IN by 1–2
405 orders of magnitude relative to T_CTL. On the other hand, the reduction in cloud ice
number concentration allows the ice crystals to grow more efficiently, with their
effective particle size generally reaching 98%-135% of that in T_CTL. The combined

effect of these two factors ultimately limits the deposition of water vapor onto the ice crystals. Consequently, P_{dep} decreases to 20%–50% of T_CTL , with the maximum
410 suppression occurring at approximately 7–8 km (Fig. 4a).

In layer B, cloud ice number concentrations in T_IN range from 7 to 10 L^{-1} , approximately 120% of those in T_CTL . However, the cloud ice mass concentration in T_IN is reduced to only 70%–90% of T_CTL . The effective diameters of cloud ice also decrease to only 77%–97% of T_CTL , with occasional reductions exceeding 50%. This
415 reduction is mainly attributable to combined effects of enhanced heterogeneous nucleation and suppressed depositional growth, and the total production rate of cloud ice drops to less than 82% of that in T_CTL . Dust aerosols provide additional ice nuclei, leading to a substantial enhancement of heterogeneous nucleation in T_IN and the formation of a much larger number of newly formed small ice crystals, with P_{gen}
420 exceeding that in T_CTL by more than two orders of magnitude. However, the increase in cloud ice number concentration is accompanied by a reduction in individual particle size, which limits the deposition of water vapor onto ice crystals. This effect is combined with a decrease in relative humidity over dust–precipitation stations in the DPA (Fig. 5c), further inhibiting the deposition process. As a result, P_{dep} in T_IN is
425 reduced to about 30% of that in T_CTL , indicating that growth of cloud ice via depositional processes is inhibited.

Snow

In layer A, the total snow production rate in T_IN increases to approximately 88%–200% of that in T_CTL ($P_{sdep}+P_{aacw}+P_{saut}+P_{iacr}+P_{raci}+P_{saci}+P_{sacr}-P_{gaut}-P_{racs}$,
430 Fig. 4b), leading to an increase in snow mass concentration to 120%–200% of T_CTL (Fig. 3a, 3b). This increase results from the combined effects of enhanced production rate for deposition–sublimation of snow (P_{sdep}) and weakened production rate for aggregation of cloud ice to snow (P_{saut}) and production rate for accretion of cloud ice by snow (P_{saci}). The P_{sdep} can reach approximately 2–5 times that in T_CTL (Fig. 4b).
435 In WDM6, the deposition growth of ice-phase hydrometeors is constrained by the available water vapor, with cloud ice deposition given priority and snow deposition

consuming the remaining vapor. Because P_{idep} is reduced to about 20%–50% of that in T_{CTL} , more water vapor is allocated to snow deposition, P_{sdep} is then enhanced. Meanwhile, as cloud ice reduces, P_{saut} and P_{saci} are weakened in T_{IN} , with both
440 processes reduced to approximately 40%–60% of their values in T_{CTL} (Fig. 4a, 4b). Despite the suppression of these source terms, the substantial enhancement of snow deposition growth dominates the snow budget in layer A, resulting in a net increase in snow production and cloud-snow mass concentration.

Finally, the ratio of cloud ice to cloud snow changes from 1:1 to 1:3 in layer A,
445 more closely consistent with observation, which shows that cloud ice generally has higher number concentrations but lower mass concentrations than cloud snow (Gao et al., 2020; Yang et al., 2021; Feng et al., 2021; Fang et al., 2022). In the aircraft observations reported, small ice particles ($>50 \mu\text{m}$) reached concentrations of up to 300 L^{-1} , whereas large ice crystals ($>600 \mu\text{m}$) were only about 3 L^{-1} (Wang et al., 2023).
450 Despite their much lower number concentrations, the larger particles contributed more to the ice mass because of their substantially greater sizes. In the present simulation, the cloud ice number concentrations in T_{IN} reach on the order of 10^1 L^{-1} in the main mixed-phase layer, which is substantially higher than in T_{CTL} and closer to the observed magnitudes, although still lower than some aircraft measurements.

In layer B, the snow mass concentration shows relatively small changes, ranging
455 from approximately 90% to 100% of T_{CTL} . From the perspective of cloud microphysics, the mechanisms are similar to those in layer A. Despite the reduction of P_{idep} , the P_{sdep} increases to 130%–200% of T_{CTL} . At the same time, the decrease in cloud ice mass leads to the continued suppression of P_{saut} and P_{saci} , resulting in a
460 total snow production rate of about 95% of T_{CTL} .

In layer C, although the model diagnostics indicate an enhancement in cloud-snow production processes (production rate for accretion of rain by snow (P_{sacr}) and production rate for accretion of rain by cloud ice (P_{iacr})) and a reduction in the production rate for accretion of snow by rain (P_{racs}), newly formed cloud snow cannot
465 be maintained because the temperature is already above 0°C which makes it

instantaneously melt, rapidly converting to rain. As a result, there is no significant change in snow mass concentration in this layer.

Cloud water and rainwater

Cloud water and rainwater are mainly distributed in layer C (temperature
470 approximately $-2\text{ }^{\circ}\text{C}$ to $18\text{ }^{\circ}\text{C}$). In this layer, both cloud-water and rainwater mixing ratios in T_IN are about 90%-95% of those in T_CTL. This small reduction is primarily attributed to a weakening of the production rate for cloud droplet activation from CCN (P_{act}), which decreases by about 5% in T_IN relative to T_CTL, indicating a suppressed conversion of water vapor into liquid water. As a consequence of the
475 reduced cloud-water content, the production rate for accretion of rainwater by cloud water (P_{acw}) is also weakened, by 5%–10%. Meanwhile, the conversion of rainwater into ice-phase hydrometeors (P_{sacr}, P_{gacr}, and P_{iacr}) is enhanced. However, under the thermodynamic conditions of layer C, temperatures exceed the melting thresholds of ice-phase hydrometeors, the newly formed snow and graupel rapidly melt and are easily
480 converted back into rainwater. Consequently, these ice-phase conversion processes contribute only marginally to the net change in rainwater mixing ratio.

Fig 5 show that, over dust–precipitation stations in both the NDSA and the DPA, the introduction of the on-line aerosol–IN nucleation scheme leads to temperature increases below 4 km, with changes of about 0.16 to 0.52 K, while the water vapor
485 mixing ratio changes by -0.04 to 0.2 g kg^{-1} during this 18:00 UTC 11 April to 18:00 UTC 12 April. These changes lead to a decrease in relative humidity within the warm-cloud layer. The relative humidity averaged over the dust–precipitation stations decreases by up to about 3 percentage points in the DPA during this period. In the NDSA, it locally reaches reductions of 6–7 percentage points around 4 km at 06:00 UTC 12
490 April. The reduced relative humidity suppresses droplet activation and condensational growth, thereby inhibiting the development of warm clouds in T_IN compared to T_CTL.

Overall, dust suppresses cloud development, reducing the total ice-phase hydrometeor content in layer A to 70 – 85% of T_CTL, the total ice-phase hydrometeor

495 content in layer B to 85 – 91% of T_CTL, and the liquid-phase hydrometeor content in layer C to 90 – 95% of T_CTL. Our results indicate that dust aerosols tend to suppress cloud development in springtime dust-related precipitation over East Asia, where precipitation is predominantly stratiform. Similar suppression effects have also been reported in previous observational studies (Zhu et al., 2023).

500 3.3 Precipitation

The on-line aerosol–IN nucleation scheme can modulate the spatial distribution of precipitation. Figure 6a shows the observed event-accumulated precipitation of DPA stations, and Figure 6b shows the simulated event-accumulated precipitation of T_CTL. In T_CTL, 18 of 43 stations in DPA exhibit overestimated simulation precipitation compared to observations (overestimated stations), primarily located in areas near dust sources area such as Gansu, Ningxia, Shaanxi, and Inner Mongolia, as well as northeastern provinces including Shandong, Liaoning, Jilin, and Heilongjiang (Fig.6b). At these overestimated stations, the observed mean accumulated precipitation is 11.49 mm, while the simulated mean accumulated precipitation is 25.55 mm (Fig.7), 505 with an average sMAPE of 45 %. The other 25 stations show underestimated simulated precipitation compared to observations (underestimated stations), mainly distributed across Hebei, Beijing, Henan, and the Yangtze River Basin downwind area of the dust events (Fig.6b). At underestimated stations, the observed mean accumulated precipitation is 31.58 mm (Fig.7), while the simulated value is only 4.63 mm, with an 515 average sMAPE of –64 %.

In T_IN, the on-line aerosol–IN nucleation scheme does not alter the overall pattern of overestimation precipitation north of 35° N and underestimation precipitation to south of 35° N in T_CTL (Fig. 6d). However, compared to T_CTL, notable improvements are mainly observed primarily between 34° and 40° N. This is driven by 520 the process discussed in Section 3.2, where the presence of dust in layer C suppresses P_{act}, thereby reducing the overestimation of precipitation near the dust source areas. sMAPE is reduced by about 1–10 % in areas near the dust source area, resulting in more accurate forecasts compared to both T_CTL (Fig. 6e, f).

Rather than being removed by precipitation or evaporation, the suppressed cloud
525 hydrometeors are transported downstream in T_IN. We calculate horizontal
hydrometeor fluxes across 116°E, 33°–50°N and 33°N, 103°–116°E from 12:00 UTC
on 12 April to 18:00 UTC on 13 April (Fig. 7). Over the entire 0–12 km layer, the total
hydrometeor flux slightly increases to about 102% of that in T_CTL.

Within the temperature range from 0 to -40 °C, the total horizontal hydrometeor
530 flux decreases by about 11 %, primarily due to a substantial reduction in cloud ice flux,
accompanied by increases in snow and graupel fluxes. In Layer A, the total hydrometeor
flux is about $4.4 \times 10^{-5} \text{ kg s}^{-1}$, corresponding to about 75 % of T_CTL. Cloud ice flux
drops sharply to about 8 % of T_CTL, while snow and graupel fluxes increase markedly
to about 19.8 times and 7.8 times, respectively. In Layer B, the total hydrometeor flux
535 is about $2.6 \times 10^{-6} \text{ kg s}^{-1}$, corresponding to about 93 % of T_CTL, with cloud ice flux
reduced to about 28 % of T_CTL, and snow and graupel fluxes increased to about 2.3
times and about 1.8 times, respectively. At temperatures above 0 °C, the total horizontal
hydrometeor flux increases to about 106 % of T_CTL, with cloud water and rainwater
fluxes increasing to about 115 % and about 108 %, respectively.

540 These results indicate that although dust suppresses cold-cloud development in the
upper and mid-troposphere, it enhances the downstream transport of liquid-phase
hydrometeors near and below the melting layer, enhancing downstream precipitation.
Finally, for underestimation stations, the mean accumulated precipitation increases by
1.1 mm compared to T_CTL, and precipitation simulation improves by approximately
545 4 %, with little changes in MAE and RMSE (Fig. 8b). For overestimated stations, the
mean accumulated precipitation decreases by 4.5 mm compared to T_CTL, and
precipitation simulations improves by approximately 40%, with MAE reduced by 1.4
and RMSE reduced by 4.1 (Fig. 8a).

In summary, because the reduction in cloud water in the 0–4 km layer is relatively
550 small, the corresponding decrease in rainwater reaching the surface is also limited. As
a result, the on-line aerosol-IN nucleation scheme exerts only a weak influence on the
total precipitation amount. Nevertheless, it can modulate the spatial and temporal

distribution of precipitation, impressing overestimated and altering underestimation in a degree, which is consistent with the findings of Park and Lim (2023) and Su and Fung
555 (2018b).

4 Conclusions and discussion

In order to explore the impact of spring dust aerosols on precipitation, this study develop an on-line aerosol-IN nucleation scheme in the regional model GRAPES/CUACE. The model performance has been evaluated by a typical dust-
560 precipitation event from 00:00 UTC on 9 April to 00:00 UTC on 15 April 2018.

Dust provides ice nuclei by heterogeneous nucleation during dust-precipitation event. The on-line aerosol-IN nucleation scheme significantly modifies nucleated IN concentration distributions. The original WDM6 scheme exhibits a systematic underestimation of ice nuclei concentrations, with nucleated IN concentrations can
565 reach 10^2 L^{-1} between 4 and 7 km altitude during the dust-precipitation event, and abnormally increase with height due to the temperature-dependent formulation of original WDM6 scheme, peaking near the -40° C layer. With the on-line aerosol-IN nucleation scheme, IN concentrations can reach 10^4 L^{-1} , so for the cloud ice mass production rate concentrated peaking at about the layer between 4 and 7 km in height,
570 more closer to the observations

Dust can inhibit the development of clouds. Above 7 km, dust suppresses the growth rate of cloud ice (through both heterogeneous nucleation and deposition-sublimation rate of cloud ice), and the total production rate of cloud ice drops to less than 24% of that in T_CTL, promoting snow formation and ultimately reducing the
575 total ice-phase hydrometeor content to 70–85% of T_CTL. Meanwhile, the total snow production rate in T_IN increases to approximately 88% - 200% of that in T_CTL, reducing total ice-phase hydrometeor content to 70 - 85% of T_CTL. Between 4 and 7 km height, dust enhances heterogeneous nucleation of cloud ice, but the new smaller particles suppress cloud ice and reduces the deposition rate, resulting in the total ice-
580 phase hydrometeor content decreasing to 85–91% of T_CTL. Below 4 km in height, the relative humidity decreases by about 3 percentage points on average over the dust-

precipitation stations in T_IN. This decrease in relative humidity limits the conversion of water vapor to cloud water and of cloud water to rain, reducing the liquid-phase hydrometeor content to 90–95% of that in T_CTL.

585 The dust can also modulate the spatial distribution of precipitation even though the on-line aerosol-IN nucleation scheme cannot alter completely the overall pattern of overestimation precipitation north of 35° N and underestimation precipitation to the south of 35° N as seen in T_CTL. The on-line aerosol-IN nucleation scheme mitigates the overestimation of precipitation near dust source areas. For overestimated stations, 590 the event-mean accumulated precipitation decreases by about 4.5 mm relative to T_CTL, with the MAE reduces by 1.4 and the RMSE reduces by 4.1. Meanwhile, the cloud hydrometeors suppressed by dust IN are not removed from the atmosphere; instead, they remain in the weather system and transported downstream as the air mass moves, thereby alleviating the underestimation of precipitation in downstream areas. In 595 stations where precipitation is previously underestimated, the mean accumulated precipitation increases by about 1.1 mm relative to T_CTL.

This study shows improvements of dust as IN on cloud and precipitation simulation by a comprehensive online aerosol-IN-cloud interaction scheme. Considering both CCN and IN effects, rather than CCN alone, improves precipitation 600 simulations by up to approximately 40 %. Aerosol and clouds interactions are an old open question, but there are still many uncertainties, due to the complex mechanisms of both CCN and IN. Furthermore, the scarcity of real-time observations hinders the in-depth exploration of detailed microphysical processes and their underlying mechanisms. More cases in different seasons and different dusty cases are needed to perform in the 605 future with more observations.

Code/data availability

All source code and data can be accessed by contacting the corresponding author Chunhong Zhou (zhouch@cma.gov.cn).

Author contributions.

610 JZ developed the on-line aerosol-IN nucleation scheme, conducted the data analysis,

and wrote the original draft of this paper. CHZ developed the aerosol-CCN-cloud interaction scheme and the on-line aerosol-IN nucleation scheme, and reviewed and edited the manuscript, providing critical insights. XYS reviewed the manuscript. SLG reviewed the manuscript and provided general insight. HW reviewed the manuscript. 615 XYZ reviewed the manuscript and gave guidance on the data analysis. All authors have given approval to the final version of the paper.

Competing interests

The authors declare that they have no conflict of interest.

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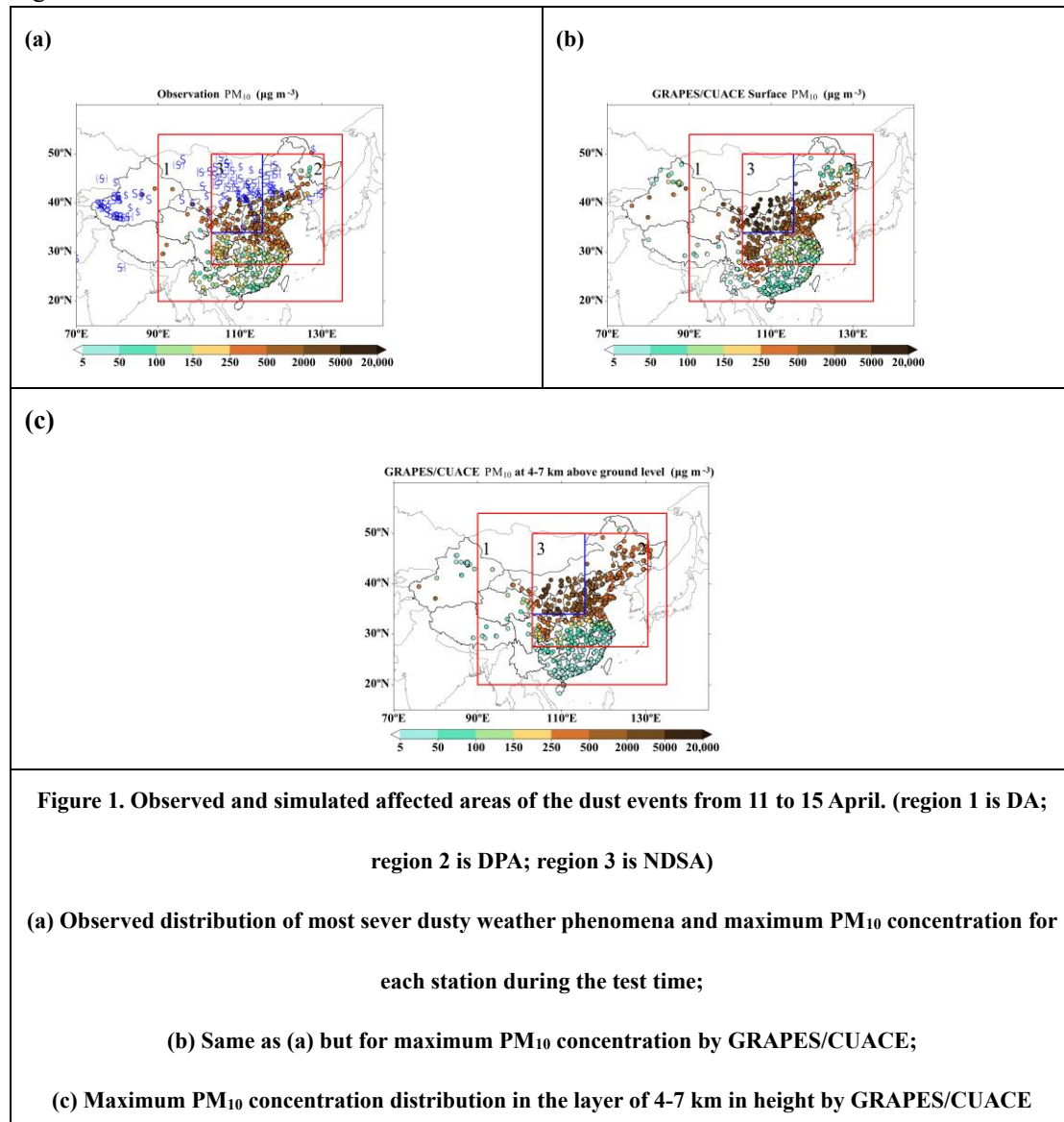
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Figure

Figure 1



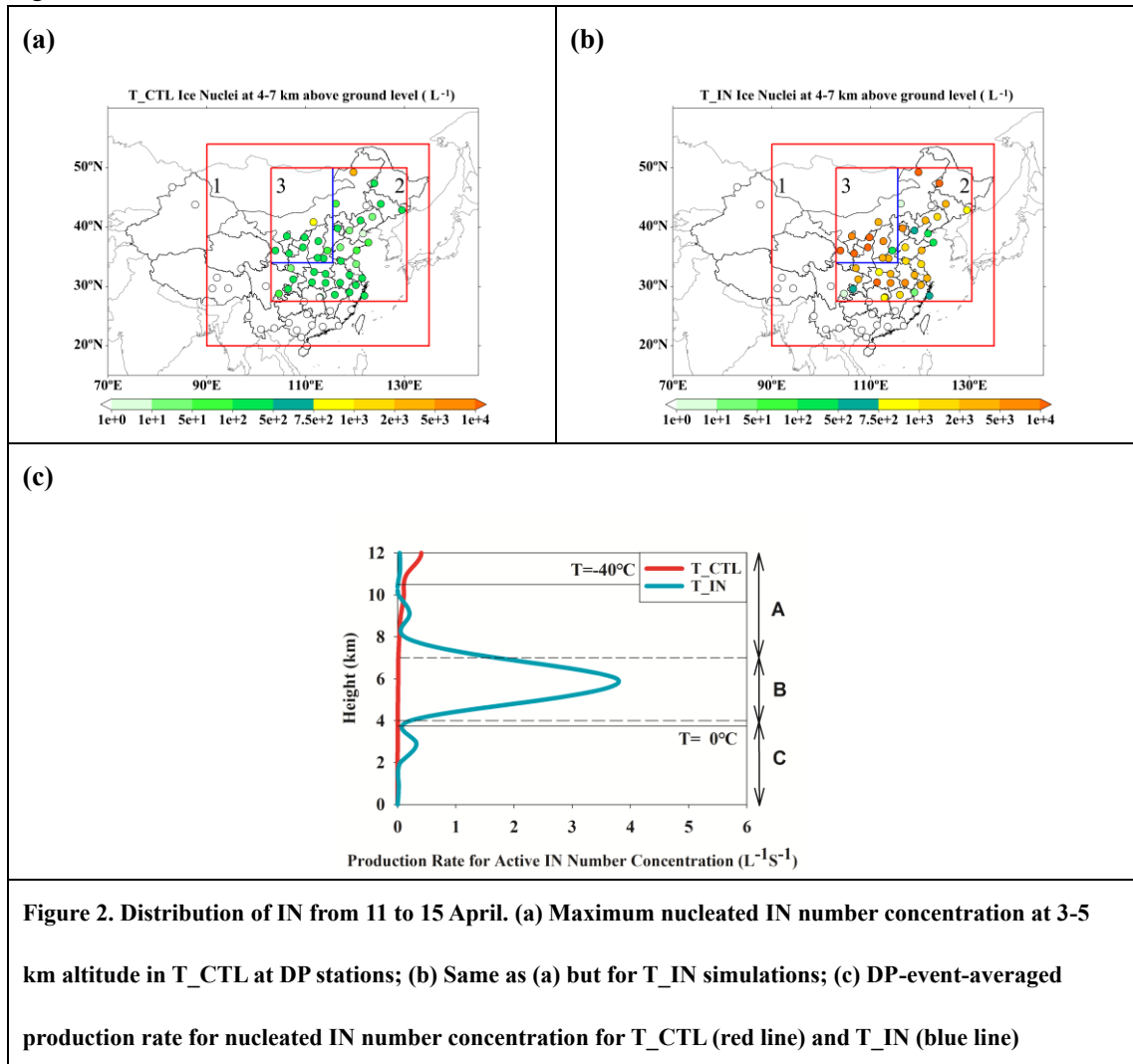


Figure 2. Distribution of IN from 11 to 15 April. (a) Maximum nucleated IN number concentration at 3-5 km altitude in T_CTL at DP stations; (b) Same as (a) but for T_IN simulations; (c) DP-event-averaged production rate for nucleated IN number concentration for T_CTL (red line) and T_IN (blue line)

Figure 3

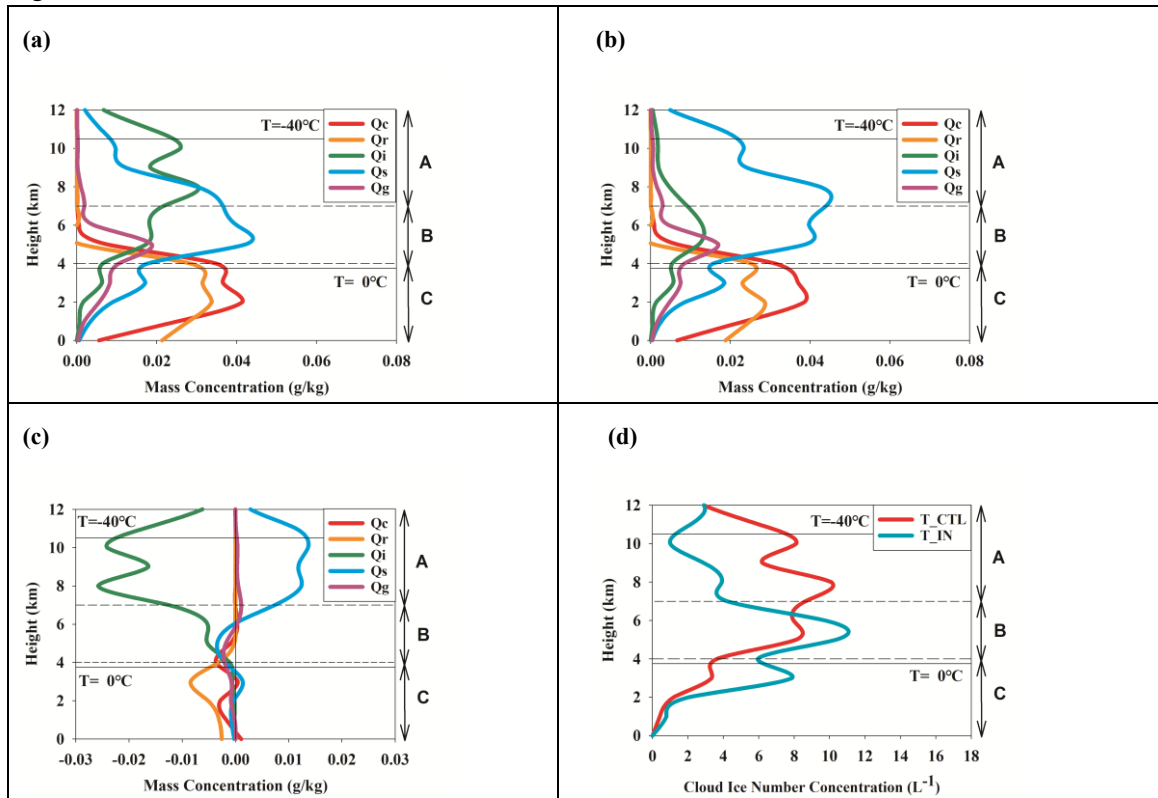


Figure 3. Distributions of hydrometeors, averaged over the dust–precipitation period (00 UTC 11 April–00

UTC 15 April 2018) and over dust–precipitation stations:

(a) hydrometeors simulated in T_CTL (b) hydrometeors simulated in T_IN

(c) hydrometeors difference by (T_IN- T_CTL);

(d) DP event-averaged vertical distributions of cloud ice number concentration

Figure 4

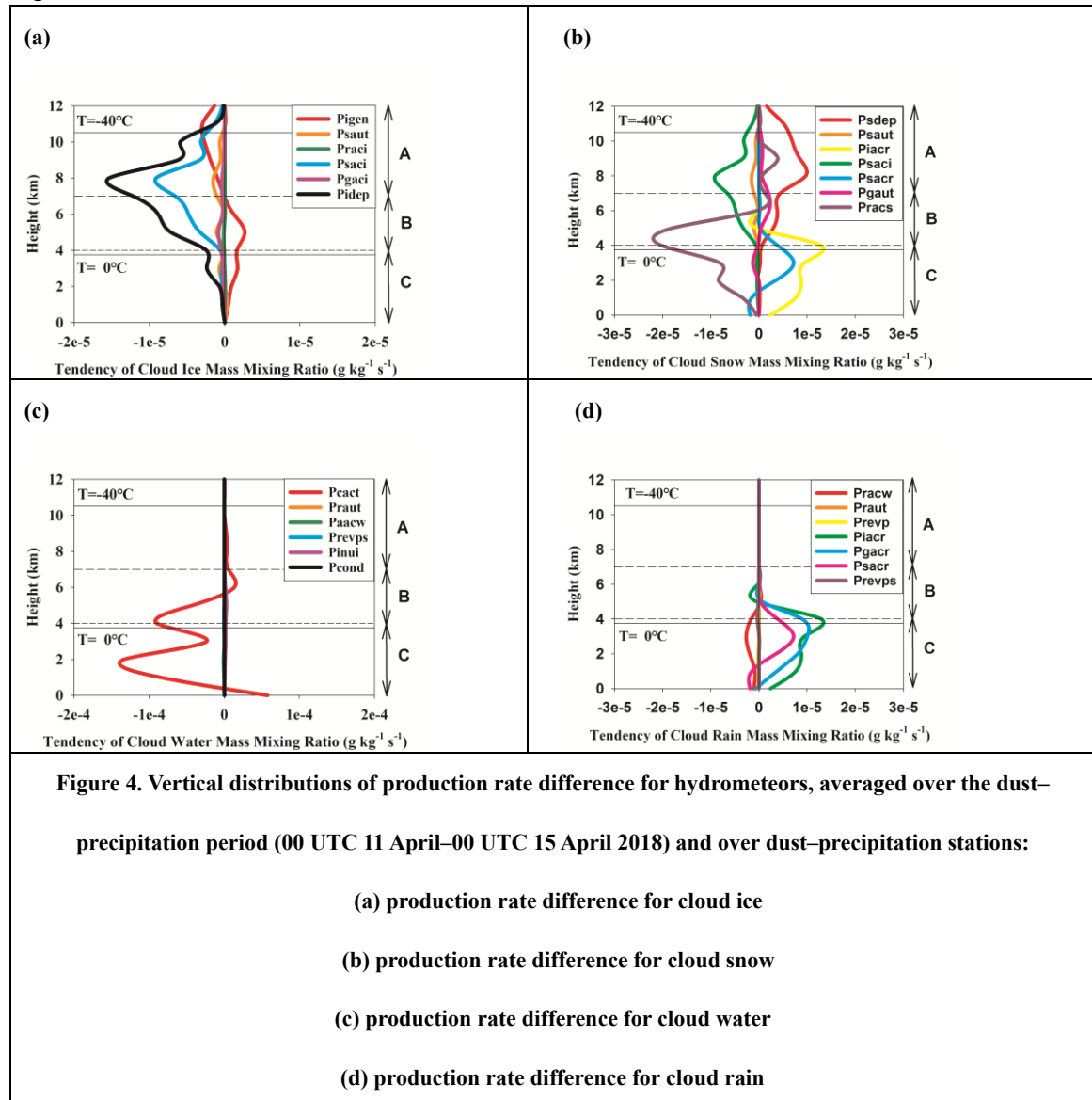


Figure 5

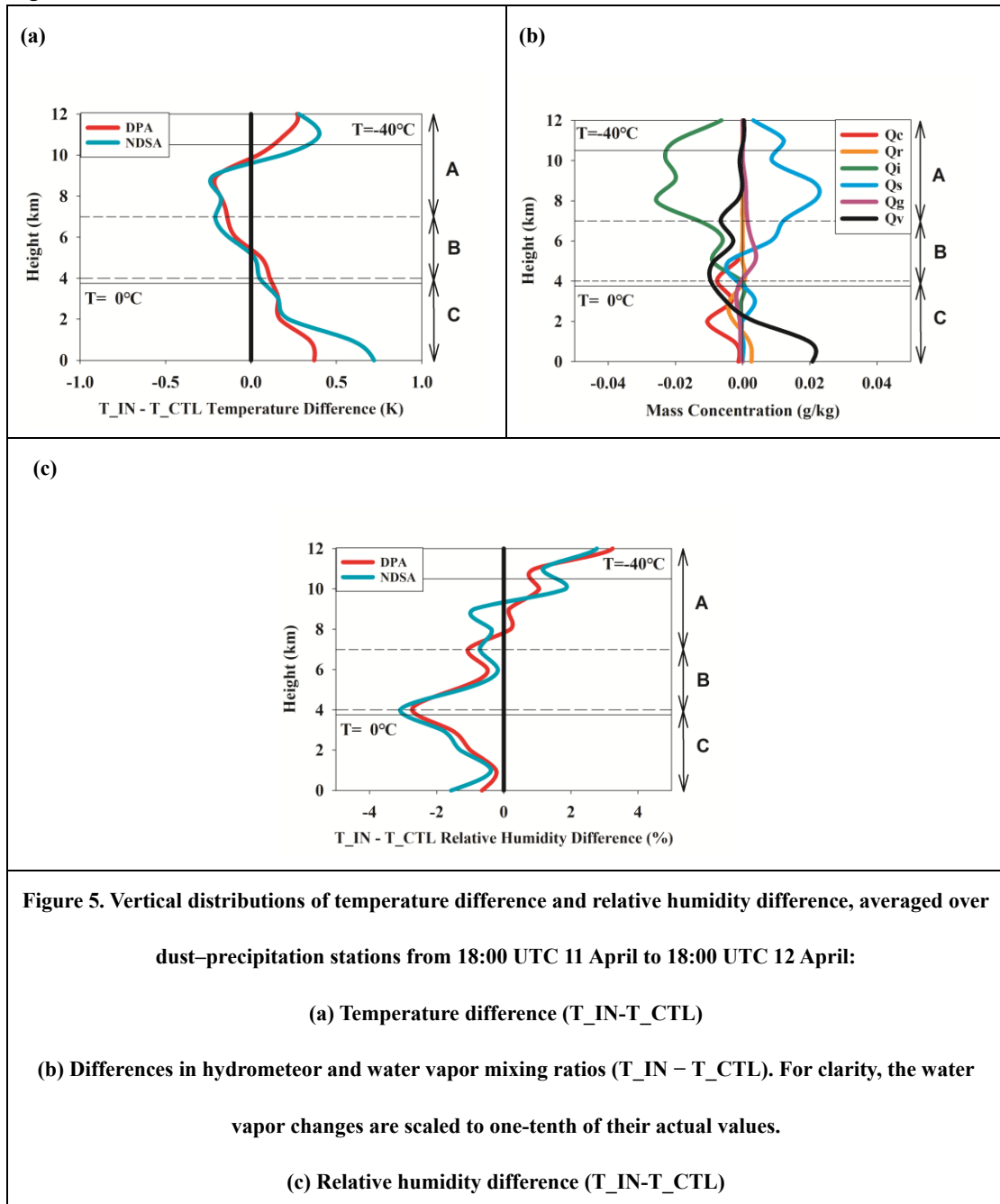


Figure 6

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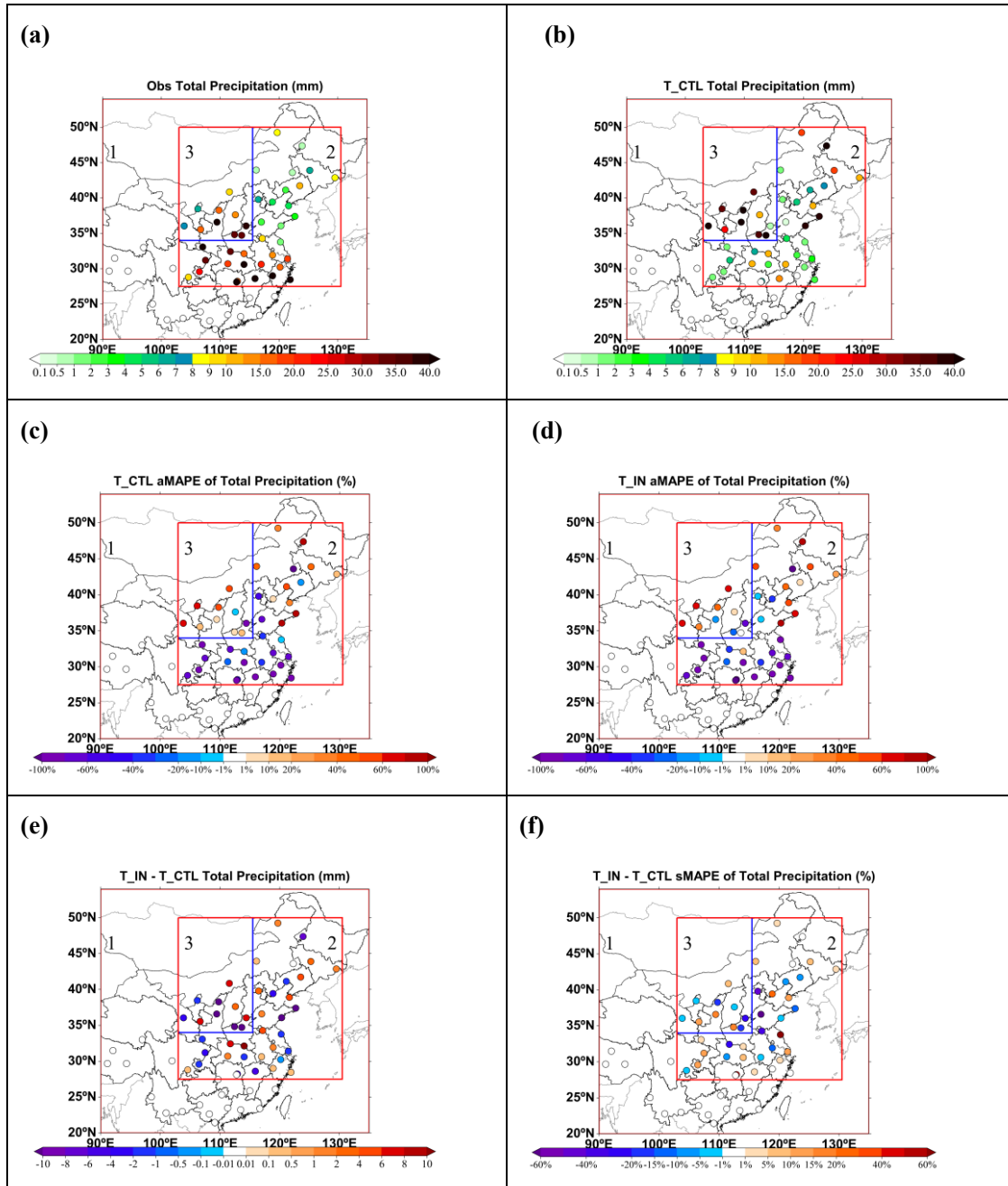


Figure 6. Comparison of observed and simulated accumulated precipitation at dust-precipitation stations:

(a) Observed accumulated precipitation from 11th 00:00 UTC to 15th 00:00 UTC;

(b) Same as a but for T_CTL;

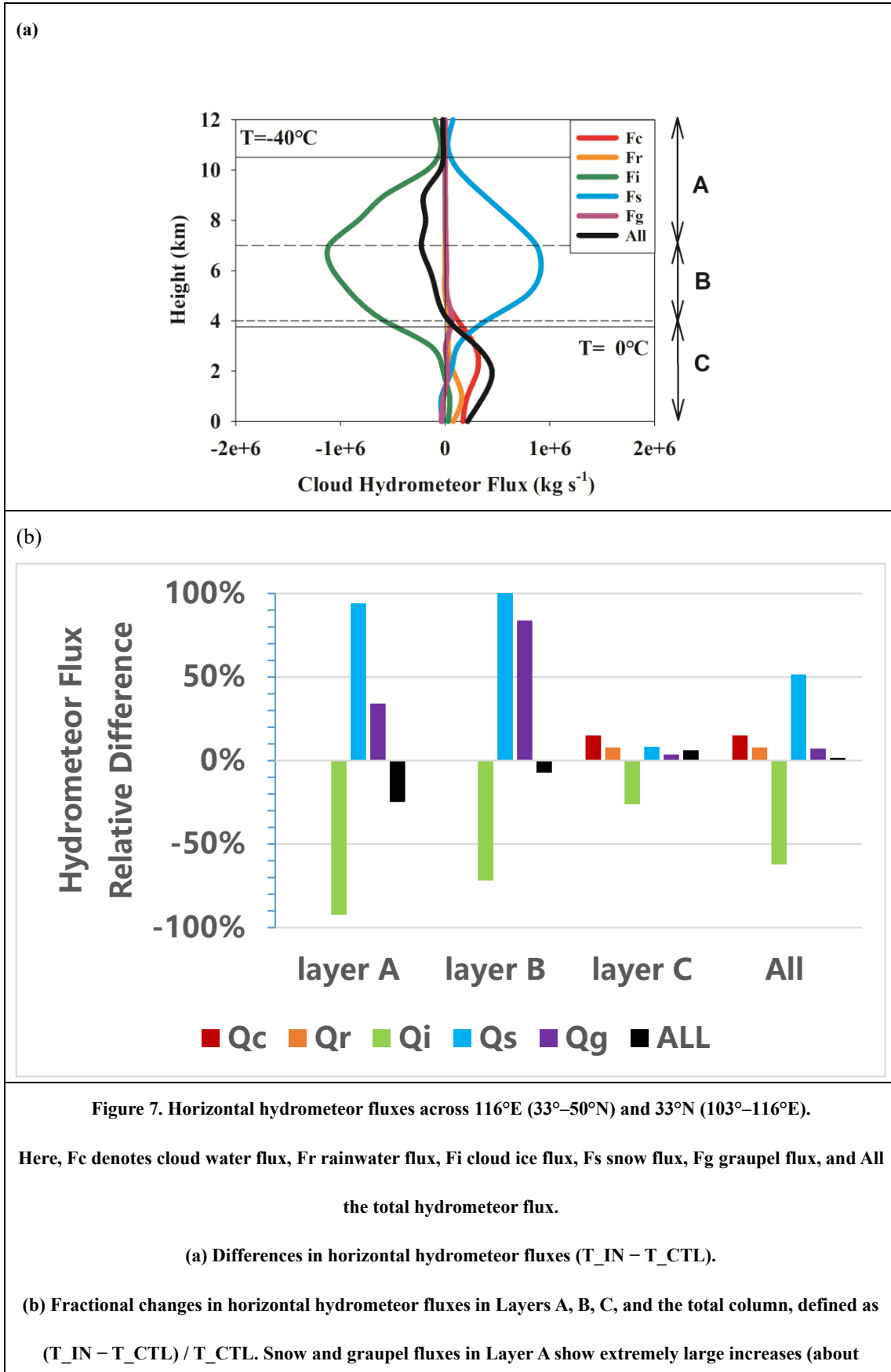
(c) aMAPE of simulated accumulated precipitation in T_CTL;

(d) aMAPE of simulated accumulated precipitation in T_IN;

(e) Difference in precipitation between T_IN and T_CTL;

(f) Difference in sMAPE between T_IN and T_CTL.

Figure 7



1883 % and 683 %, respectively); for better visualization, these values are scaled down by a factor of 20 in the figure.

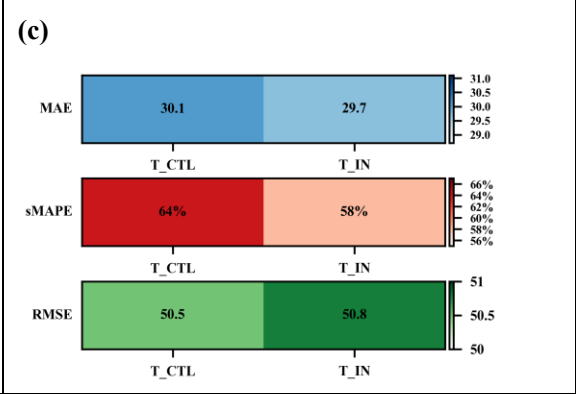
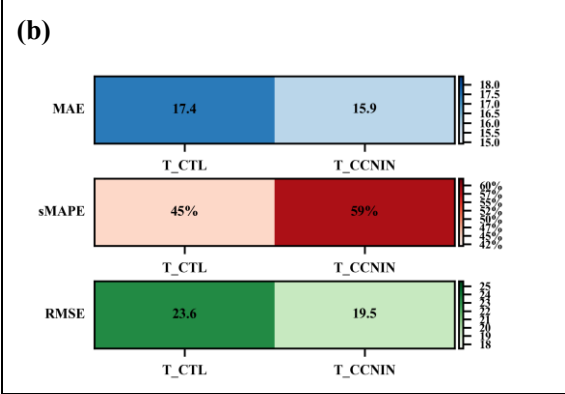
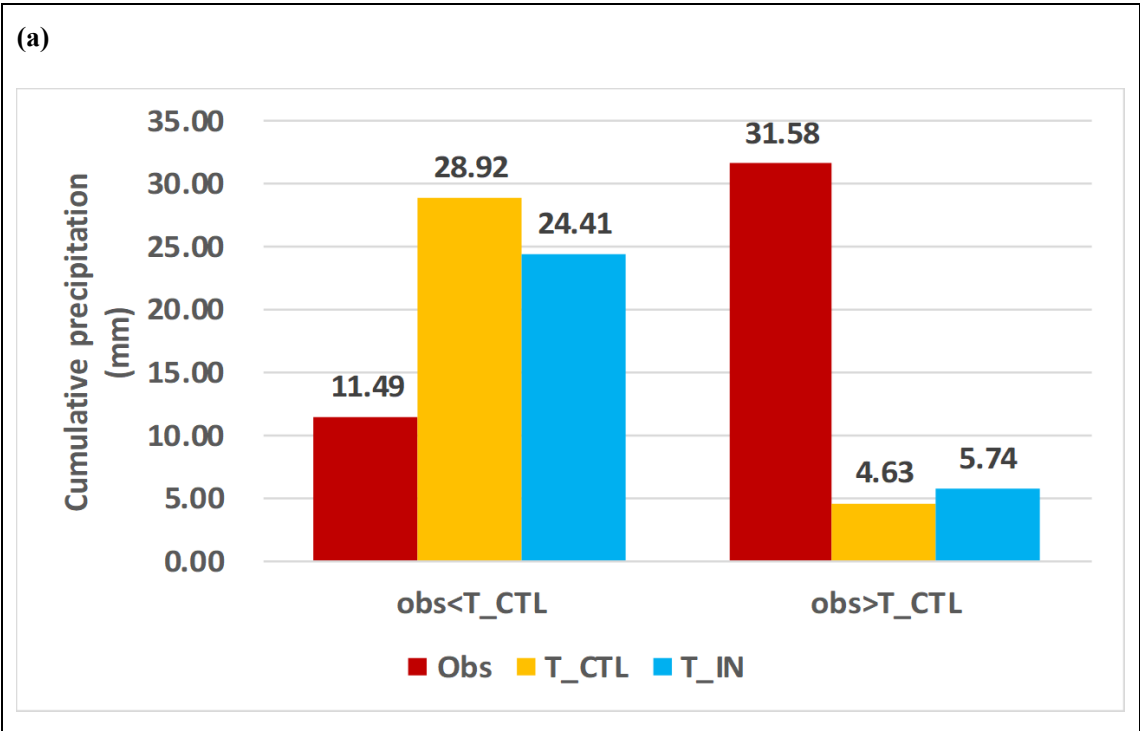


Figure 8.

(a) mean accumulated precipitation during DP event at overestimated stations and underestimated stations

(b) Statistical analysis of observed versus simulated accumulated precipitation at DPA stations for Overestimated stations

(c) same as (b), but for Underestimated stations

Table

Table 1. Three Tests designed for different types of precipitation

Test	Warm cloud	Cold cloud
T_CTL	on-line aerosol–CCN interaction scheme	original WDM6
T_IN	on-line aerosol–CCN interaction scheme	on-line aerosol-IN nucleation scheme

985

Table 2. List of Symbols

Symbol	Meaning
Paacw	Production rate for accretion of cloud water by averaged snow/graupel (Under subfreezing conditions ($T < 0$ °C), the collected droplets are typically supercooled and freeze onto the surface of snow or graupel particles. In this case, cloud water is converted into snow or graupel, and latent heat of fusion is released, contributing to local warming. Under above-freezing conditions ($T > 0$ °C), the accreted droplets do not freeze.)
Pcact	Production rate for cloud droplet activation from CCN
Pcond	Production rate for condensation rate of water vapor to cloud liquid water
Pgacr	Production rate for accretion of rain by graupel
Pgaci	Production rate for accretion of cloud ice by graupel
Pgaut	Production rate for aggregation form snow to graupel
Piacr	Production rate for accretions of rain by cloud ice
Pidep	Production rate for deposition- sublimation rate of cloud ice
Pigen	Production rate for heterogeneous nucleation
Pinud	Production rate for deposition/condensation freezing to form cloud ice
Pinui	Production rate for immersion freezing of cloud water to form cloud ice
Pracs	Production rate for accretions of cloud snow by rain
Pracw	Production rate for accretion of cloud water by rain
Praci	Production rate for accretion of cloud ice by rain
Praut	Production rate for aggregation form cloud water ice to form rain

Prevps	Production rate for evaporation/condensation rate of cloud water
Prevp	Production rate for evaporation/condensation rate of rain
Psacr	Production rate for accretions of rain by cloud snow
Psaci	Production rate for accretion of cloud ice by snow
Psaut	Production rate for aggregation form cloud ice to snow
Pscar	Production rate for accretion of rain by snow
Psdep	Production rate for deposition- sublimation rate of cloud snowFig 1b
