Title

A comprehensive characterization of ice nucleation by three different types of cellulose particles immersed in water

Authors

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Abstract

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We present the laboratory results of immersion freezing efficiencies of cellulose particles at supercooled temperature (T) conditions. Three types of chemically homogeneous cellulose samples are used as surrogates that represent supermicron and submicron ice nucleating plant structural polymers. These samples include micro-crystalline cellulose (MCC), fibrous cellulose (FC) and nano-crystalline cellulose (NCC). Our immersion freezing dataset includes data from various ice nucleation measurement techniques available at seventeen different institutions, including nine dry dispersion and eleven aqueous suspension techniques. With a total of twenty methods, we performed systematic accuracy and precision analysis of measurements from all twenty measurement techniques by evaluating T-binned (1 °C) data over a wide T range (-36 °C < T < -4 °C). Specifically, we inter-compared the geometric surface area-based ice nucleation active surface-site (INAS) density data derived from our measurements as a function of T, $n_{s,geo}(T)$. Additionally, we also compared the $n_{s,geo}(T)$ values and the freezing spectral slope parameter $(\Delta \log(n_{s,geo})/\Delta T)$ from our measurements to previous literature results. Results show all three cellulose materials are reasonably ice active. The freezing efficiencies of NCC samples agree reasonably well, whereas the diversity for the other two samples spans for ≈10 °C. Despite given uncertainties within each instrument technique, the overall trend of the $n_{s,qeo}(T)$ spectrum traced by the T-binned average of measurements suggest that predominantly supermicron-sized cellulose particles (MCC and FC) generally act as more efficient ice-nucleating particles (INPs) than NCC with about one order of magnitude higher $n_{s,geo}(T)$.

1. Introduction

1.1 Background

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Glaciation of supercooled clouds through immersion freezing induced by ice-nucleating particles (INPs) is an important atmospheric process affecting the formation of precipitation and the Earth's energy budget (*Boucher et al.*, 2013; *Vergara-Temprado et al.*, 2018). Currently, the climatic impact of INPs is, however, uncertain due to our insufficient knowledge regarding their diversity and abundance in the atmosphere (e.g., *Hoose and Möhler*, 2012; *Murray et al.*, 2012; *Kanji et al.*, 2017; *Knopf et al.*, 2018). Recently, micro-crystalline cellulose (MCC) particles of <16 µm in diameter, extracted from natural wood pulps (Aldrich, 435236), have been identified as an efficient INP (*Hiranuma et al.* 2015a, H15a hereafter). Experiments with this surrogate may provide useful information to understand the role of biological INPs in the troposphere as presented in H15a. Conspicuously, the H15a modeling results suggest that the tropospheric concentration of ice-nucleating cellulose becomes substantial (>0.1 L⁻¹) below about -21 °C.

Cellulose is a linear polymer of 1–4 linked β -d-anhydroglucopyranose molecules, deriving from plant fragments, leaf litter, wood fiber, non-wood fiber and/or even microbes (Quiroz-Castañeda & Folch-Mallol, 2013; Thakur and Thakur, 2014; Chawla et al., 2009). The composition and structure of cellulose-containing bio-fiber depends on the source and several different factors, summarized in Khalil et al. (2012) and Dittenber and GangaRao (2012). In general, airborne cellulose particles are prevalent ($>0.05 \mu g \, m^{-3}$) throughout the year even at remote and elevated locations as reported in Sánchez-Ochoa et al. (2007). More recent study of carbonaceous aerosol composition in Switzerland over two years showed that ambient cellulose represents approximately 36-60% of primary biological organic aerosols, and the ambient cellulose concentration exceeded a few µg m⁻³ (Figs. 6 and 7d of *Vlachou et al.*, 2018). Their water insoluble, hydrolysis resistant and heat resistive features (Fernández et al., 1997; Quiroz-Castañeda & Folch-Mallol, 2013) may in part explain the long-range transport and high concentrations of cellulose even at geographically dispersed sites. Another unique characteristic of ambient cellulose is its wide range of physical size available for freezing. For example, the size distribution measurements of ambient cellulose particles by Puxbaum and Tenze-Kunit (2003) indicate the presence of particulate cellulose in the range from 10 nm to >20 µm. The presence of supermicron particles, possessing larger surfaces as compared to submicron ones, is remarkable since they can potentially act as supermicron-sized INPs since large surfaces may promote efficient formation of ice embryos (Pruppacher and Klett, 2010; Schnell and Vali, 1972 and 1973). Nevertheless, more comprehensive characterization of icenucleating properties of various cellulose-containing particles is necessary to examine if the ice-nucleating activity is specific to MCC or generally relevant to all cellulose materials in the atmosphere.

1.2 Previous INUIT Activities

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In 2012, the German research consortium-led INUIT (Ice Nuclei research UnIT) project was commenced to comprehensively study heterogeneous ice nucleation processes in the atmosphere. Throughout the period since, this project has provided a trans-national platform to bolster collaborative research activities between meticulous groups who study atmospheric INPs. In turn, INUIT has accelerated ice nucleation research in a wide range of study scales from nanoscopic microphysics (e.g., *Kiselev et al.*, 2017) to cloud scale modeling (e.g., *Diehl and Mitra*, 2015; *Paukert and Hoose*, 2014) in cross- and inter-disciplinary manners.

Formerly, several INUIT studies addressed quantitative validations of ice nucleation (IN) instruments using test proxies of atmospheric particles (Wex et al., 2015; Hiranuma et al., 2015b; Burkert-Kohn et al., 2017). Some studies focused on identifying potential reasons of the data diversity (e.g., different experimental methods and sample preparation methods). For example, Burkert-Kohn et al. (2017) conducted the inter-comparison workshop by codeploying instruments with a uniform aerosol dispersion procedure and size segregation method to minimize the diversity in ice nucleation results. Hiranuma et al. (2015b), H15b henceforth, took a different approach to perform an inter-comparison of INP measurement techniques. The authors demonstrated the collaborative multi-institutional laboratory work with a total of fourteen institutions (seven from Germany, four from U.S., one from U.K., one from Switzerland and one from Japan) by distributing a test particulate sample to partners and allowing measurements at their home laboratories. The authors discussed the potential effect of sampling of the dust, agglomeration, flocculation, surface estimation methods, multiple nucleation modes and chemical aging on the observed data deviation amongst seventeen different IN instruments. This study suggested that a combination of above-listed factors may be responsible for ~8 °C diversity in terms of temperature and up to three orders of magnitude difference with respect to the ice nucleation active surface-site (INAS) density, $n_s(T)$, parameters. Further, two follow-up studies on potential effects of aggregation upon IN were performed in Emersic et al. (2015) and Beydoun et al. (2016). The former study presented the potential role of aggregation and sedimentation of mineral particles, altering their IN efficiency in aqueous suspension, by combining experimental and modeling approaches. The latter study presented a subset of cellulose data used in this study, and the authors postulated that the widening of the frozen fractions and enhanced ice activity towards high *T* was attributable to increased diversity in ice nucleating activity for lower concentrations and particle surfaces. In other words, there is a distribution of active sites between individual droplets depending on the total surface area. Nevertheless, our understanding of overall consistency of current INP measurement techniques and dominant mechanisms that may be responsible for diversity among measurements is still insufficient.

1.3 Goals

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The measurement strategy for this study was formulated in 2015 to further augment our understanding of the sensitivity of various ice nucleation instruments with respect to immersion freezing efficiencies. Beyond official INUIT-participating institutes, including Bielefeld University (BU), Goethe University Frankfurt (GUF), Johannes Gutenberg University of Mainz (JGU), Karlsruhe Institute of Technology (KIT), the Max Planck Institute for Chemistry (MPIC), the Leibniz Institute for Tropospheric Research (TROPOS), the Technical University of Darmstadt (TUD) and the Weizmann Institute of Science (WIS), ten associated institutes (five from U.S., three from E.U. and two from Japan) are involved in this study. These associated partners include Carnegie Melon University (CMU), Colorado State University (CSU), North Carolina State University (NC State), the Pacific Northwest National Laboratory (PNNL), West Texas A&M University (WTAMU), the Institute of Atmospheric Sciences and Climate-National Research Council (ISAC-CNR), the University of Basel, the University of Leeds, the Meteorological Research Institute (MRI) and the National Institute of Polar Research (NIPR). In this study, we have used three cellulose samples: micro-crystalline cellulose (MCC, Aldrich, 435236), fibrous cellulose (FC, Sigma, C6288) and nano-crystaline cellulose (NCC, Melodea, WS1) as atmospheric surrogates for non-proteinaceous biological particles. These samples were shared with all collaborators, and immersion freezing experiments were conducted individually at each institution to obtain immersion freezing data as a function of multiexperimental parameters (see Sect. 3.1). The motivation of using multiple types of cellulose was to (1) examine the immersion freezing abilities of both predominantly supermicron (MCC and FC) and submicron (NCC) cellulose particles to assess a wide size range of chemically uniform biological particles and (2) look into diverse surface structure (Table 1)

A total of twenty measurement techniques are used in this study to compile a comprehensive dataset for evaluating immersion freezing properties of cellulose samples. The dataset is analyzed to understand functional dependence of various experimental parameters and of cellulose particle characteristics. In this work, eleven instruments test samples used aqueous suspensions, while nine examined aerosolized powders dispersed in synthetic air

with a low RH or atomized/nebulized-suspensions containing cellulose samples followed by diffusion drying process, referred to as dry dispersion methods henceforth. The basic experimental methods and parameterization approaches used to interpret the data are discussed in **Sects. 3.1 and 3.2**.

This work extends a previous proof-of-principle experiment that demonstrated the importance of cellulose-containing particles in the atmosphere (H15a). Due to increasing and diverse awareness of presence of atmospheric cellulose (e.g., *Vlachou et al.*, 2018; *Schütze et al.*, 2017; *Legrand et al.*, 2007; *Yttri et al.*, 2018; *Samake et al.*, 2018) – not as levogulcosan (the pyrolysis product of cellulose), the main objective of this study is to comprehensively examine the immersion freezing efficiency of cellulose that could be important in an atmospheric context. Besides, the comprehensive ice nucleation data of cellulose materials presented in this work can be used to elucidate the role of airborne biological ice-nucleating aerosols derived from leaf litters and their emissions over natural surfaces (e.g., *Schnell and Vali*, 1976) and harvest regions, which certainly contained populations of plant matter in the air (*Suski et al.*, 2018).

2. Sample Characterization

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All of our samples are linear polymers of glucosyl derivatives, mechanically extracted through <200 °C heat application and catalytic oxidation (e.g., Battista et al., 1962; Brinchi et al., 2013). In particular, MCC is extracted from hardwoods (e.g., oak, personal communication with the manufacturer, Aldrich). A summary of major properties of three samples is provided in Table 1. Briefly, these highly stable biopolymers, whose bulk density ranges between 1.0-1.5 g cm⁻³, exhibit different physical dimensions depending on sample processing and treatments. As seen in Table 1, the geometric size of dispersed particles are more than ten-fold smaller than the size of bulk materials measured by electron microscopy without any exception, suggesting the presence of super aggregates in non-dispersed bulk samples. We note that the powder size of MCC reported by the manufacturer (~50 μm) is in good agreement with our Scanning Electron Microscope (SEM)-measured size. In contrast, the particle size of NCC reported on the manufacturer's material data sheet (TEM-based data) is more comparable to the dispersed particle diameter of ~0.2 μm than the SEM-based size. In this manuscript, the NCC size by SEM represents the size of NCC residuals (i.e., leftover particles after evaporating water content) from 5 μL suspension droplet of 0.03 wt%. Due to the high viscosity of the gelatinous form of NCC (4,665 ± 200 cP at 25 °C), aggregation may have occurred while evaporating water. Even after the 15 minute ultrasonic bath treatment of the suspension, aggregates seem to remain unelucidated, which is reflected in its SEM-based diameter of >2.5 μm. A more detailed discussion of particle and residual size distributions are available in the Supplemental Sect. **S.1.**

The average aspect ratios (ARs) of each cellulose material in **Table 1** were estimated with an identical procedure employed in our previous H15a study. We evaluated a total of 4,976 MCC, 371 FC and 764 NCC particles. The Everhart-Thornley Detector (ETD) of a scanning electron microscope (SEM, FEI, Quanta 650 FEG) was used to acquire the below-the-lens micrograph image and measure two dimensional axis length of particles deposited on membrane filters. The degree of elongation appears to be higher for NCC (average AR up to 2.93) when compared to MCC and FC (average AR of <2.30). Nonetheless, all sample types show that particles are elongated with an aspect ratio varying from ~2 to 3, which is similar to our previous measurement on MCC particles (i.e., 2.1).

Three different measurements of the unit surface area per unit mass (specific surface area, SSA), namely geometric SSA, SEM-based SSA and BET-SSA, for each system are also shown in **Table 1**. These measurements correspond to SSA of (1) mechanically aerosolized particles (<10 μ m in diameter) in the Aerosol Interaction and Dynamics in the Atmosphere

(AIDA) chamber, (2) droplet residuals obtained after evaporating water content of 5 μ L droplet of 0.03 wt% aqueous suspension and (3) bulk samples, respectively. Our intention of using different SSA metrics is to provide the most adequate parameter for the $n_{s,geo}(T)$ estimation of individual techniques based on their characteristics (e.g., geometric SSA for dry dispersion techniques and SEM-based SSA for aqueous suspension techniques). As demonstrated in our previous H15b comparison effort, when a reduced SSA value is observed for a same sample, it indicates the presence of agglomeration. Hence, the degree of aggregation of cellulose fibers is presumably responsible for the observed differences in SEM-based SSA values for residuals obtained from suspensions from geometric SSA of the mechanically aerosolized particles (Table 1). Alternatively, a loss of larger particles from the sample which may happen in airborne aerosols due to settling or impaction in the particle generation set-up may also lead to different SSA values if the surface properties of the cellulose particles differ with the particle size. A more detailed discussion of chemical composition and impurity analyses of our sample materials, including single particle aerosol spectrometry and scanning electron microscopy, are discussed in the **Supplemental Sects. S.2. and S.3.**, respectively.

3. Methods

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3.1. Ice Nucleation Measurements

Twenty techniques were used to investigate the ice-nucleating properties, in particular immersion freezing (*Vali et al.*, 2015), of cellulose particles (**Table 2**). In this study, nine techniques employed dry dispersion methods that refer to experiments employing water vapor condensation onto dry dispersed particles followed by droplet freezing, and another set of eleven techniques used aqueous suspension methods that denote the experiments started with the test sample pre-suspended in water before cooling. Detailed information of individual methods and their applications to study atmospherically relevant INPs are provided in references given in **Table 2** and elsewhere (e.g., *DeMott et al.*, 2017). More detailed quantitative and nominal descriptions of both dry dispersion and aqueous suspension methods used in this study are available in the **Supplemental Sect. S.4**.

Note that only non-mandatory guidelines were provided as an experimental protocol by INUIT to those who employed aqueous suspension techniques, and the experimental protocol for the wet suspension techniques was decided by each investigator. The intention was not to introduce limitations and constrains to participants. For MCC and FC, the INUIT protocol recommended the following procedures:

- 1. Measurements with <0.05 wt% suspension,
- 2. Idle time of ~30 min without stirring for large particles to settle out,
- Prepare droplets out of the quasi-steady state suspension (i.e., the upper layer of the suspension),
 - 4. Storage of the sample in the chemically inert container at ambient temperature.

In a similar way, for NCC, the INUIT protocol suggested:

- 1. One minute sonication of the original sample for initial homogenization,
- 2. Dilution to the desired final concentration using deionized water (18.2 M Ω cm⁻¹),
- Mixing the suspension vigorously for 3 minutes using high shear mechanical stirrer, homogenizer or probe sonicator to get homogenous suspension; alternatively, using an ultrasonic bath for 30 minutes in the case of sample volume <10 ml,
- 4. Measurements with <0.03wt% in order to diminish particle aggregation,
- 30 5. Storage of the sample in dry and cool (4 °C) environment.

More detailed discussion regarding nominal parameters is given in the **Supplemental Sect. S.4**.

3.2. Ice Nucleation Parameterization

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In this section, we describe a procedure to parameterize immersion freezing abilities for both dry dispersion methods and aqueous suspension techniques. The immersion freezing data of cellulose particles in a wide range of temperatures is then discussed by comparing $n_{s,geo}(T)$ spectra from all twenty instruments. Please note that using the scaled metrics for the validation (e.g., $n_{s,geo}(T)$ scaling with the technique specific *SSA* value) is indispensable in this study because the changes or uncertainties in surface area amongst groups are an issue as described in the **Supplemental Sect. S.4**. The INP concentration per volume of air ($n_{INP}(T)$, e.g., *DeMott et al.*, 2017; *Vali*, 1971) is a useful parameter for instrumental evaluation when utilizing identical samples at a single location with known sampling flows, but is not applicable in this work.

The majority of dry dispersion methods employs the approximation of *Niemand et al.* (2012). If the activated ice fraction is small (< 0.1), the Taylor series approximation can be applied, and we can estimate $n_{s,geo}(T)$:

$$n_{s,\text{geo}}(T) = -\ln\left(1 - \frac{N_{\text{ice}}(T)}{N_{\text{total}}}\right) \left(\frac{1}{S_{\text{ve}}}\right) \approx \frac{N_{\text{ice}}(T)}{N_{\text{total}}S_{\text{ve}}} = \frac{N_{\text{ice}}(T)}{S_{\text{total}}} , \tag{1}$$

in which $N_{\text{ice}}(T)$ is the cumulative number concentration of formed ice crystals at T (cm⁻³), N_{total} is the total number concentration of particles prior to any freezing event (cm⁻³), S_{ve} is the volume equivalent surface area of an individual particle (m²), and S_{total} is the total surface area (m²). For the LACIS data, the left part of Eqn. (1) was used without any approximation.

One distinct exception is the electrodynamic balance (EDB) method, in which the probability of contact freezing on a single collision, e_c , is first inverted from frozen fraction (*FF*) to take into account the rate of collision and, then, scaled to surface area of a single INP to estimate $n_{s,geo}(T)$ (*Hoffmann et al.*, 2013a; 2013b):

$$n_{s,geo}(T) = \frac{e_c(T)}{k_{imm} \cdot S_{re}} \tag{2}$$

Note that the INP particle colliding with the supercooled droplet is only partially submersed in water, and therefore the surface available for nucleation is corrected by a dimensionless factor k_{imm} . The value of this factor depends on the wettability of the particle surface and is generally unknown. In this work, $k_{imm}=1$ has been assumed. The effective surface area of MCC particles has been derived from the scanning electron microscope images of the particles collected on the Nuclepore® membrane filters placed inline to the EDB, as described in the Supplemental Information (Sect. S.1).

The results of eleven aqueous suspension methods are interpreted in terms of the frozen fraction (FF), INP concentration per volume of liquid (c_{INP} , Vali, 1971) and geometric

size-based ice nucleation active surface-site density ($n_{s,geo}(T)$, Connolly et al., 2009; H15b). The cumulative FF at T is:

$$FF(T) = 1 - \frac{N_u}{N},\tag{3}$$

where N_u is the number of unfrozen droplets and N is the total number of originally liquid entities. Following Eqn. 1 in *DeMott et al.* (2017), conversion to c_{INP} at T is expressed by

$$c_{INP}(T) = -\frac{1}{V_d} ln\left(\frac{N_u(T)}{N}\right),\tag{4}$$

where V_d represents the individual droplet volume. Finally, the $n_{s,geo}(T)$ value as a function of T can be estimated by

$$n_{s,geo}(T) = \frac{c_{INP}(T)}{\rho_w \omega \theta},\tag{5}$$

where ρ_w is the water density (= 997.1 g L⁻¹), ω is the mass ratio of analyte and water (unitless) and θ is the SSA value (m² g⁻¹), provided in **Tables 2 and, S1 and S2**.

Accordingly, we compare the $n_{s,geo}(T)$ and $\Delta \log(n_{s,geo})/\Delta T$ (i.e., the freezing spectral slope parameter, H15b) data from our measurements to five literature results. These reference results include previously reported $n_{s,geo}(T)$ curves of illite NX particles from H15b (hereafter H15NX), MCC particles from H15a (hereafter H15MCC), Snomax (*Wex et al.*, 2015, hereafter W15), desert dusts (*Ullrich et al.*, 2017, hereafter U17) and K-feldspar (*Atkinson et al.*, 2013, hereafter A13). The $n_{s,geo}(T)$ (m⁻² as a function of °C) fits from the reference literature are:

$$n_{s,\text{geo}}^{H15NX,dry} = \exp((27.92 \times \exp(-\exp(0.05 \times (T+13.25)))) + 6.32),$$

$$T \in [-37, -18]; \Delta \log(n_{s,\text{geo}})/\Delta T = 0.18,$$
(6)

 $n_{s,\text{geo}}^{H15NX,wet} = \exp((22.64 \times \exp(-\exp(0.16 \times (T + 20.93)))) + 5.92),$

$$T \in [-34, -11]; \Delta \log(n_{s,geo})/\Delta T = 0.37,$$
 (7)

 $n_{s,\text{geo}}^{H15MCC,dry} = \exp(-0.56 \times T + 7.50),$

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$$T \in [-30, -15]; \Delta \log(n_{s,geo})/\Delta T = 0.24,$$
 (8)

 $n_{s,\text{geo}}^{H15MCC,wet} = \frac{2.57 \times 10^7 + \frac{-2.84 \times 10^7}{1 + \exp(\frac{-25.19 - T}{1.45})}}{SEM - based SSA_{MCC}},$

$$T \in [-28, -22]; \Delta \log(n_{s,geo})/\Delta T = 0.35,$$
 (9)

$$n_{s,\text{geo}}^{W15} = \frac{(1.40 \times 10^{12}) \times (1 - (\exp((-2.00 \times 10^{-10}) \exp(-2.34 \times T))))}{geometric\,SSA_{Snomax}},$$

$$T \in [-38, -2]; \Delta \log(n_{s,geo})/\Delta T = 0.88 (-2 \text{ °C} < T < -10.7 \text{ °C}),$$
 (10)

 $n_{s,\text{geo}}^{U17} = \exp(150.577 - (0.517 \times (T + 273.150))),$

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$$T \in [-30, -14]; \Delta \log(n_{s,geo})/\Delta T = 0.22,$$
 (11)

$$n_{s,geo}^{A13} = 10^4 \times \exp(-1.038(T + 273.150) + 275.260) \times \frac{BET - SSA_{K-feldspar}}{geo - SSA_{K-feldspar}}$$

$$T \in [-25, -5]; \Delta \log(n_{s,geo})/\Delta T = 0.45.$$
 (12)

For H15MCC (wet), the $n_m(T)$ to $n_{s,geo}(T)$ conversion was performed using SEM-based *SSA* constants of 0.068 m² g⁻¹. The geometric *SSA* value of 7.99 m² g⁻¹ was used for W15. This *SSA* value was derived from the polydisperse particle size distribution measurements of Snomax obtained during AIDA studies, whose IN data are included to compute immersion freezing results reported in *Wex et al.* (2015). For microcline (K-feldspar), the $n_{s,geo}(T)$ to $n_{s,BET}(T)$ conversion was performed using a laser diffraction-based geometric *SSA* of 0.89 m² g⁻¹ and an N₂ BET-SSA of 3.2 m² g⁻¹ reported in *Atkinson et al.* (2013). Please note that laser diffraction tends to be sensitive to the larger particles in a distribution, so it may miss the smaller particles and underestimate surface area.

3.3. Temperature Binning

A consistent data interpolation method is important to systematically compare different ice nucleation measurement methodologies as demonstrated in H15b. In this study, we present *T*-binned average ice nucleation data (i.e., 1 °C bins for -36 °C < T < -4 °C). Unless the data were originally provided in 1 °C binned-data (i.e., weighted-average or cumulative counts) [i.e., BINARY, DFPC-ISAC, FRIDGE-CS (MCC portion), LINDA, NC State-CS, NIPR-CRAFT, WISDOM and WT-CRAFT], all data are binned in a consistent manner using either a moving average (where original data points are finer than 1 °C) or a Piecewise Cubic Hermite Interpolating Polynomial function (where original data points are equivalent or coarser than 1 °C). For the former case, the default span for the moving average is 3 (i.e., centered moving average for a 0.5 °C resolution data). If the temperature resolution is finer than 0.5 °C, the number of moving average span is equal to the number of data points in each temperature bin (an even span is reduced by 1). The comparison of *T*-binned immersion freezing spectra from particle dispersion methods and aqueous suspension methods is discussed in **Sect. 4.1**.

4. Results and Discussions

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4.1. Dry dispersion vs. aqueous suspension methods

Temperature-binned ensemble $n_{s,geo}(T)$ spectra of MCC, FC and NCC in a temperature range between -4 and -38 °C are presented in **Fig. 1.** Different columns (**a-c**) correspond to different sample types: (a) MCC, (b) FC and (c) NCC. The top panels show a comparison between dry dispersion type measurements and aqueous suspension measurements of cellulose samples with previous parameterizations of other reference samples (**panels i**). The $n_{s,geo}(T)$ spectra from each subgroup of techniques are independently summarized in **panels ii and iii**. More detailed representations of $n_{s,geo}(T)$ spectra from individual techniques are discussed in **Sect. 4.3**. Lastly, the bottom panels (**panels iv**) show the overall deviation between maxima and minima of $n_{s,geo}(T)$ as pink shaded areas. As inferred from the first three panels (**i, ii and iii**), dry particle-dispersed measurements generally show higher $n_{s,geo}(T)$ values than aqueous suspension measurements above -24 °C regardless of sample types. Furthermore, as apparent in **panels iv**, the $n_{s,geo}(T)$ differences among measurements can extend up to three orders of magnitude at -20 °C (for MCC and FC) and -15 °C (for NCC), where the results from particle dispersion measurements and a majority of suspension measurements coexist.

The observed divergence in $n_{s,geo}(T)$ is most significant at temperature higher than -24 °C, where the slope in the aqueous suspension spectra is steeper (i.e., $\Delta \log(n_s)/\Delta T > 0.34$). Most aqueous suspension methods capture the abruptly increasing segment of the $n_{s,geo}(T)$ spectral slopes at -20 °C > T > -25 °C. In this T region, the slope is virtually identical to the slopes of wet H15NX and H15MCC spectra (0.35-0.37, **Eqns. 7 and 9**) and is also closely parallel to the A13 parameterization (0.45, **Eqn. 12**), suggesting the number of active sites are different. Likewise, our T-binned data from dry dispersion methods exhibit similar $n_{s,geo}(T)$ values when compared to the previous parameterizations. For instance, our dry dispersed cellulose spectra (i.e., $\Delta \log(n_s)/\Delta T$ of 0.20, 0.28 and 0.22 for MCC, FC and NCC) present comparable trends to the dry H15 curves (0.18-0.24, **Eqns. 7 and 9**) and U17 parameterization (0.22, **Eqn. 11**).

It is interesting that a similar difference between dry dispersion and aqueous suspension results (i.e., $n_{s,geo}(T)$ of dry dispersed particle > $n_{s,geo}(T)$ of suspension results) is made by previous inter-comparison activities with mineralogically heterogeneous dust particles (*Emersic et al.*, 2015; H15b). In brief, *Emersic et al.* (2015) reports the dry dispersion chamber-measured $n_{s,geo}(T)$ can be up to a factor of 1000 larger than the cold stage results for multiple mineral dust samples, including illite NX, Kaolinite and K-feldspar. Our previous study also shows that $n_{s,geo}(T)$ of illite NX increases sharply at colder temperatures in the T range from -18 °C to -27 °C, followed by the leveling off segment at the low temperature region. It is

certainly common for the $n_{s,geo}(T)$ spectrum to level off at the $n_{s,geo}(T)$ maxima. As mentioned in **Sect. 1.2.**, several studies (*Emersic et al.*, 2015; *Beydoun et al.*, 2016) reported the mechanism of the observed divergence between two subsets of methods. Nonetheless, the reduction in the slope of $n_{s,geo}(T)$ spectrum may be a plausible contributor to the higher reported $n_{s,geo}(T)$ values in some aqueous suspension measurement results (WISDOM, CMU-CS in **Sect. 4.3**), which are comparable to the dry dispersion results (i.e., data of freezing of individual droplets containing a single aerosol particle) for illite NX and cellulose (*Beydoun et al.*, 2016).

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Next, **Fig. 2** depicts the $n_{s,geo}(T)$ diversity in $log(n_{s,ind.})/log(n_{s,avg})$, which represents the ratio of the log of individual measurements ($n_{s,ind}$) to the log average of $n_{s,geo}(T)$ expressed as $n_{s,avg}$ at given temperatures. In other words, this figure provides an overview of the $n_{s,geo}(T)$ deviations across the various techniques employed in this work. These n_s ratios are shown for the temperature range covered by at least two measurement techniques used in the present study. In this figure, different panels show three different $n_{s,avg}$ values as denominators, including the average based on all bulk data (All, panels i, ii and iii), dry dispersion subgroup (Dry, panels iv), or aqueous suspension subgroup (Sus, panels v). As for numerators ($n_{s,ind}$), the interpolated T-binned data (1 °C) from **Fig. 1** are used. A total of five panels are presented. First, a summary comparison of two method categories (dry dispersion and aqueous suspension) in a temperature range of -33 $^{\circ}$ C < T < -15 $^{\circ}$ C is given in the top panels (panels i). As shown in these panels, data deviation (i.e., scatter from the average $log(n_{s,ind.})/log(n_{s,avg}) =$ 1 line) can be seen in both dry dispersion and aqueous suspension measurements. Other panels provide more evidence on the measurement diversity. In short, while the $log(n_{s,ind.})/log(n_{s,avg})$ values range within 0.8-1.2 for Dry Dispersion (DD) and Aqueous Suspension (AS) cases (panels iv and v), more prominent scatter of the $log(n_{s,ind.})/log(n_{s,avg})$ values (0.6-1.4) is seen when All is used as $n_{s,avg}$ values (panels i, ii and iii). Thus, the observed deviation is the largest with $n_{s,avg}$ of All (i.e., both AS and DD). Furthermore, the deviation becomes more apparent towards higher temperatures. This trend persists regardless of sample type. Further discussion on the observed deviations and diversity between dry and aqueous suspension measurement techniques is beyond the scope of this study. Some discussions regarding potential sources and explanations of deviations, which warrant future studies, are given in the Supplemental Sects. S.4, S.5, S.9 and S.10.

4.2. Comparison of three cellulose sample types

The multiple exponential distribution fits (also known as the Gumbel cumulative distribution function) for *T*-binned data of all three cellulose samples are summarized in **Table**

3. Fit parameters as well as $\Delta \log(n_s)/\Delta T$ for each category are given in this table. As can be inferred from the table, the overall $\Delta \log(n_s)/\Delta T$ value is almost identical for all three sample types (0.31-0.33) in spite of some deviations observed for min-max (0.26-0.40). The observed consistency in the spectral slopes suggests cellulose material exhibits relatively similar ice nucleation above examined temperatures (>-36 °C).

For all cellulose types, a reasonable correlation coefficient (r) is found for each technique (i.e., DD and AS), suggesting reasonable agreement and consistency for the results from a similar group of immersion freezing techniques. However, we must reiterate the discrepancy between DD and AS. For instance, our observation of lower values of DD slopes (0.20-0.29) as compared to those of AS slopes (0.29-0.37) in the similar temperature range suggests distinct differences between the two subsets of methods. Moreover, the dry dispersed-MCC shows relatively lower $\Delta \log(n_s)/\Delta T$ of 0.20 than FC and NCC (note not all instruments delivered FC and NCC measurements, see **Table 2**). This exception potentially indicates a fundamental difference of dry dispersed-MCC from other sample types.

Table 4 provides the log average of T-binned $n_{s,geo}(T)$ values for all of the cellulose samples, representing detailed comparisons of MCC, FC and NCC. The **Supplemental Sect. S.6** (**Fig. S7**) also summarizes the comparison between the averages for each material (see Supplemental Information for details). As seen in the table and figure, there exists a discrepancy between this study and previous work for MCC. At -28 °C, for example, our log average $n_{s,geo}(T)$ of MCC (3.25 x 10^9 m⁻², **Table 4**) is smaller than the previous MCC result at the same T (1.18 x 10^{10} m⁻², H15a). This difference possibly reflects the fact that our average $n_{s,geo}(T)$ includes the results from a multitude of aqueous suspension measurements, which typically fall in the lower range of DD measurements (**Sect. 4.1**), while H15MCC (**Eqn. 9**) is derived from a dry dispersion method only. Note that the $n_{s,geo}(T)$ maxima from **Table 4** reasonably overlap with the H15MCC parameterization.

The highest $n_{s,geo}(T)$ value of the FC experiments (3.6 x 10^{10} m⁻² at -29 °C from AIDA) is somewhat lower than that of MCC. Similarly, the highest $n_{s,geo}(T)$ value of the NCC experiments (1.5 x 10^{10} m⁻² at -35 °C from WISDOM) is an order magnitude lower than that of MCC as well as W15.

Table 4 also implies that MCC possesses higher ice nucleation efficiency relative to the other two types. First, at above -25 °C, the immersion freezing ability of MCC typically exceed that of NCC. Second, at -22 to -24 °C, where more than seven instruments are involved to calculate the average T-binned $n_{s,geo}(T)$, MCC's $n_{s,geo}(T)$ is consistently one order magnitude higher than FC and NCC. Third, when compared to FC, MCC generally possess slightly higher $n_{s,geo}(T)$ at T below -16 °C. Likewise, a similar trend holds true when we compare MCC to NCC

at T below -17 °C. The observed difference is up to two orders of magnitude at -20 °C. Please note that, at the high T region (> -17 °C), dry dispersion techniques are not sensitive enough to detect INPs with their experimental parameters used in this study (**Supplemental Tables S1 and S3**). In contrast, detecting rare INPs by increasing the concentration of the aqueous particle suspension is advantageous yet also challenging. In other words, the measurement uncertainties generally propagate towards high temperatures because the confidence interval is relatively wider when there exists only a few frozen droplets. Hence, our observation of less immersion freezing ability of MCC at this T range (up to a factor of ~20 at -16 °C) may not be conclusive. Particle sedimentation, aggregation and the concentrations effects identified by *Beydoun et al.* (2016) are also more prominent at higher concentration, especially for cellulose samples.

4.3. Individual immersion freezing measurements

All individual $n_{s,geo}(T)$ spectra of MCC, FC and NCC from each technique are shown in **Figs. 3, 4** and **5**, respectively. Only brief remarks regarding each technique are summarized below. Several special experiments were carried out using seven techniques to complement our understanding of cellulose ice nucleation. The results from these unique experiments are first described (Sects. 4.3.1-4.3.7) followed by the other remarks (Sects. 4.3.8-4.3.19).

4.3.1. CSU-CFDC

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Immersion freezing ability of both polydisperse and quasi-monodisperse dry dispersed MCC particles were characterized by CSU-CFDC. In short, ice-nucleating efficiencies of DMA size-selected MCC particles (500 nm mobility diameter) were compared to that of the polydisperse population for immersion freezing experiments.

As seen in the **Fig. 3b**, the discrepancy between the results from two populations is substantial. Similar to the LACIS result, a weak temperature dependence of $n_{s,geo}(T)$ of monodisperse MCC particles is observed within defined experimental uncertainties (see the **Supplemental Table S1**). Observed quasi-flat $\Delta \log(n_{s,geo})/\Delta T$ of the monodisperse case suggests a week T-dependent immersion freezing ability of given specific size of MCC particles for the investigated temperature range. Conversely, a polydisperse spectrum, which represents the result of an ensemble of different MCC particle sizes, shows a stronger trend of the slope towards low T segment, suggesting a non-uniform distribution of active sites over the available S_{total} of cellulose in this study. Some previous INUIT studies demonstrated the size independence of the $n_{s,geo}(T)$ value using submicron hematite and illite NX particles based

on AIDA ice nucleation experiments (*Hiranuma et al.*, 2014 and 2015b). Such a characteristic may not remain true for the immersion mode freezing of supermicron-sized fiber particles.

For all sample types, as seen in **Figs 3b, 4b and 5b**, the CSU-CFDC results do not agree well with H15a (MCC_dry, **Eqn. 8**). Instead, they virtually agree with the wet generation results. This is especially true for the results with polydisperse population. Note that formerly observed agreement within a factor of three in $n_{s,geo}(T)$ estimation (cloud simulation chamber INAS > CSU-CFDC INAS; *DeMott et al.*, 2015) is seen only at -30 °C. The observed discrepancy may be due to non-uniform active site density for different sizes. Another possible explanation may be due to the alternation of cellulose physico-chemical properties perhaps upon humidification during shipping, causing behaviour more like aqueous suspended particles. One thing that we need to keep in mind is that the CFDC uses a 2.4 μ m particle impactor at its inlet (**Supplemental Table S3**). Because of the impactor, there is loss of larger particles. Thus, the $n_{s,geo}(T)$ results may vary, possibly due to the difference in the size of cellulose samples examined. At -23 °C, where the data of size-selected measurements exist for all three cellulose samples, CSU-CFDC show $n_{s,geo,MCC} \approx n_{s,geo,NCC}$ (**Figs. 3b, 4b and 5b**).

4.3.2. DFPC-ISAC

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DFPC-ISAC instrument (Santachiara al. 2010) provided et data condensation/immersion freezing. The use of 103% RHw in this investigation was optimized to count statistically significant amount of INPs in this system for examined cellulose particles (i.e., MCC and FC). With this system, we assessed the IN efficiencies of different sizes of MCC and FC particles generated by means of different cyclone cut-sizes (0.5, 1.0, 7.0 μm or none). Further, both dry dispersed (Dry) and nebulizer-generated particles (Wet) were systematically assessed for their INP activities. Without an exception, INP concentrations were measured at -22°C for all specimens. For the case of particles (<0.5 μm cyclone-selected), we additionally measured INP concentrations at -18 °C to assess the general trend of the INP activates as a function of T. This particular case was selected for the extended study due to the similarity of their geometric SSAs to those of the AIDA cloud parcel simulation measurements. In addition, while collecting the cellulose particles on nitrate membrane filters (Millipore, 0.47 um pore size) used for IN assessment, parallel measurements of particle size distributions using an optical particle counter (Grimm, 1.108) were carried out. The results of size distributions, represented by the SSA values, are summarized in **Table 5**.

For Dry, increasing the cut-size tends to decrease the *SSA* value, implying large particles come through, and the dominance of the mass relative to the surface becomes significant. This observation is valid as the cyclone is used to remove particles larger than the

designated cut-size. Regardless of whether using the cyclone or not, particle sizes out of the nebulizer-generation is somehow comparable to that of Dry dispersion with a cyclone of 1 μ m cut-size. The observed difference between Wet and Dry is indicative of the changes in particle size and morphology while drying atomized particles from a suspension of the powder in water as described in the **Supplemental Sect. S.2**.

Figures 3c and 4c show all the results of INP measurements by DFPC-ISAC. For MCC, the interpolated DFPC results of the immersed particles (<0.5 μm cyclone-selected) falls in the middle of FRIDGE results of two different modes for -22 °C < T < -18 °C. More interestingly, the slope of the DFPC $n_{s,geo}(T)$ spectrum ($\Delta log(n_{s,geo})/\Delta T = 0.24$) represents the median of the slopes of FRIDGE measurements (i.e., 0.17 for default mode and 0.31 for immersion mode). This observation is consistent with other results of (1) size-selected particles tend to exhibit a gentle slope (similar to the observations from CFDC and LACIS) and (2) nebulizer-generated techniques tend to result in a deteriorated INP activity (H15b).

Another important implication of the DFPC results is the fact that submicron dry particles show the highest INP efficiencies, practically lie on $n_{s,geo}(T)$ data points of H15a parameterization at given T for both MCC and FC. Moreover, inclusion of supermicron sizes (no cyclone or 7 μ m) seems reducing IN efficiencies of both MCC and FC. Further investigation is required to interpret these results.

Over the temperature range of -18 to -22 °C, the DFPC results of immersed particles (<0.5 µm cyclone-selected), show $n_{s,\text{geo},FC} \approx n_{s,\text{geo},MCC}$ (Figs. 3c and 4c). Note that $n_{s,\text{geo},FC}$ appears to be slightly higher than $n_{s,\text{geo},MCC}$. This observation is not consistent with the general trend of $n_{s,\text{geo},MCC} > n_{s,\text{geo},FC}$ (Sect. 4.2). However, the observed difference is only a factor of <2 on average.

4.3.3. FRIDGE

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The FRIDGE data were derived from both default mode (a combination of deposition, condensation ice nucleation and immersion freezing at $RH_{\rm w}$ of 101%) and immersion mode operation for MCC. With these two different operational modes, FRIDGE investigated the ice nucleation ability of both dry and droplet suspended particles deposited on a substrate. Particularly, the default mode operation of FRIDGE provided data from -16 to -30 °C (MCC) by scanning $RH_{\rm ice}$ and $RH_{\rm w}$ (low to high) at a constant temperature. Accordingly, ice crystals formed at the highest $RH_{\rm w}$ of 101% were considered as a measure of immersion $N_{\rm ice}$ from dry dispersed particle measurements. Likewise, the immersion mode operation of FRIDGE provided data from -19 to -28 °C (MCC) and from -13 to -23 °C (NCC). As demonstrated in H15b, this immersion mode counts immersion freezing of suspended particles in which the particles

are first washed into droplets and then placed on the substrate to be comparable to the dry dispersion method. Hence, this method is advantageous to collect a filter sample of cellulose, prepared the same way as in the dry dispersion experiment, and then run it on a cold-stage.

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Figure 3e shows the comparison of $n_{s,geo}(T)$ derived from the two different operation modes of FRIDGE. There are a few important implications from the FRIDGE results. First, on average, the measurements with dry particles in the 'default' setting showed more than an order of magnitude higher $n_{s,geo}(T)$ in comparison to the immersed particles in FRIDGE experiments at T > -22 °C. As shown in **Fig. 3e**, the deposition mode data suggest that $n_{s,geo}(T)$ values for -22 °C < T < -19 °C are close (within a factor of two) to those from MRI-DCECC, in which experiments were carried out with a high degree of particle agglomeration. In comparison to the default mode result, FRIDGE experiments in the pure immersion mode showed much lower $n_{s,geo}(T)$ than that with the default setting, but agreed with other immersion datasets. Second, a steeper $\Delta \log(n_{s,geo})/\Delta T$ of 0.31 was found for the measurements with immersed particles at T > -24 °C when compared to the slope of the deposition mode data (i.e., 0.17). As a temperature shift (i.e., shifting the data a few °C) does not offset the discrepancy, other mechanistic interpretations might be plausible causes of this discrepancy. For instance, this difference may be a consequence of the different IN efficiencies of nucleation modes of both experimental approaches (e.g., deposition + condensation + immersion vs. immersion alone) in the examined temperature range, the different sample preparation processes, effects of agglomeration or a combination of the three. The divergence of default-mode and CS-mode becomes notable T > -24 °C, perhaps suggesting the effect of agglomeration. Specifically agglomeration may take place inside the pipetted droplets. While pipetting agglomeration and separation is avoided by shaking the sample, but during cooling it lasts 15-30min until a droplet freezes.

Figure 5c presents the summary of FRIDGE-CS measurements for NCC. The $n_{s,geo}(T)$ spectrum nearly overlaps with the H15b (illite NX wet) reference spectrum. It also agrees well with the other droplet freezing instruments CMU-CS, NIPR-CRAFT, NCS-CS, BINARY and WISDOM. Similar $n_{s,geo}(T)$ values were obtained although the methods analysed droplets of different volumes. In particular FRIDGE and WISDOM $n_{s,geo}(T)$ attach to each other better than 0.3 °C. By comparing NCC to MCC at -23 °C < T < -19 °C, the FRIDGE-CS results show $n_{s,geo,MCC}$ > $n_{s,geo,NCC}$ for >one order magnitude throughout this overlapping T range. Note that the $\Delta \log(n_{s,geo})/\Delta T$ value of NCC (0.40) is somewhat higher than the average slope parameters listed in **Table 3**.

4.3.4. LACIS

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With LACIS, we examined immersion mode freezing of both atomized and dry dispersed MCC particles separately. For atomized particle generation, particles were dried in a diffusion dryer directly after spraying the suspension. Succinctly, LACIS measured immersion ice nucleation ability of atomizer-generated MCC particles for 700 nm mobility diameters in the temperature range of -35 °C < T < -30 °C. The selection of this relatively large size was necessary to get a signal above the limit of detection in the system. The experiments with dry dispersed MCC were performed with polydisperse MCC particles for -36 °C < T < -27 °C. Note that a cyclone was used in the air stream of LACIS (see **Supplemental Table S3**).

Generally, LACIS measurements with dry dispersed MCC particles are in agreement with that from H15a as apparent in **Fig. 3g** ($n_{\rm s,geo}$ (-30 °C) ~ 1.5 x 10¹⁰ m⁻²). Furthermore, LACIS measurements down to -36 °C with dry polydisperse MCC particles show that $\Delta \log(n_{\rm s,geo})/\Delta T$ (= 0.17, Supplemental Table S1) is identical to MRI-DCECC for -28 $^{\circ}$ C < T < -16 $^{\circ}$ C. Contrastively, the slope of the spectrum for 700 nm size-segregated MCC particles (= 0.05) is considerably lower than that of the polydisperse case. This slope of the LACIS $n_{s,geo}(T)$ spectrum is parallel to that of the CSU-CFDC spectrum (dry dispersed 500 nm case, slope = 0.05 for -30 $^{\circ}$ C < T < -24 °C; Fig. 3b). Thus, though we cannot certainly define the relative importance of the aerosol generation method (e.g., the changes in physico-chemical properties of particles occurred during atomization as prescribed in the Supplemental Sect. S.2), the aerosol size might have a non-negligible impact on the variation in spectral slopes. Therefore, the immersion freezing efficiency of MCC particles likely is different for differently sized MCC particles, meaning that a single $n_{s,geo}(T)$ curve cannot be reported for MCC. With this, the method of accounting for differences in surface area between different groups/methods becomes questionable for a complex system like cellulose. Furthermore, its complex morphology (see the Supplemental Sect. S.5) causes that the determination of the surface area is quite prone to errors which can be a reason for the observed differences in $n_{s,geo}(T)$. The n_s framework must be rigorously tested with more empirical data. Nevertheless, for LACIS, both polydisperse and quasimonodisperse MCC particles exhibit similar $n_{s,geo}(T)$ values above -30 °C (e.g., $n_{s,geo}(-30 °C)$ ~ 1.5 x 10^{10} m⁻² in **Fig. 3g**), suggesting a negligible size dependency of $n_{\rm s,geo}(T)$ for MCC particles in this temperature range.

4.3.5. LINDA

This vial-based immersion freezing assay was utilized to compare the freezing activity of bulk suspension (0.1 wt% cellulose in NaCl solvent) to that of dry powders individually suspended in each vial (sus vs. pow henceforth). Such comparison was carried out to ensure that

employing different methods of vial preparation did not impede ice nucleation of cellulose samples, including MCC and FC. For the latter procedure (pow), pre-weighed cellulose powders (0.2 mg) were directly poured into 200 mg (199.8 μ L) of 0.1% NaCl solution to realize the concentration of cellulose in each vial to be equivalent to 0.1 wt%, such that two procedures became comparable. We note that all vials, regardless of the procedure, were sonicated (46 kHz) for 5 minutes prior to each LINDA measurement. Note that we used non-sterile NCC (NCC01) for the IN characterization with LINDA.

The results of MCC and FC are shown in **Figs. 3m and 4g**. The results suggest similarity of $n_{s,geo}(T)$ within the experimental uncertainties of LINDA (*Stopelli et al.*, 2014) for the range of examined temperatures (-7 °C to -18 °C). Further, the slope of LINDA $n_{s,geo}(T)$ spectra $(\Delta \log(n_{s,geo})/\Delta T)$ of 0.29 is identical for both scenario cases (i.e., sus and pow). Hence, for given mass concentration of 0.1 wt%, both vial preparation procedures seem valid. Nonetheless, suspended cellulose powders settle rapidly in both cases, implying the necessity of taking a great care when measuring INP activity of supermicron-sized particles with the ~200 μ L vial-based assay.

For -18 °C < T < -12 °C, the LINDA results (bulk suspension) show $n_{s,\text{geo},\text{MCC}} > n_{s,\text{geo},\text{FC}}$ with similar $\Delta \log(n_{s,\text{geo}})/\Delta T$ (0.29-0.30), verifying comparable performance of this vial-based technique to other suspension methods (**Figs. 3m and 4g**).

Figure 5f shows the freezing spectrum of NCC01 with the slope parameter $(\Delta \log(n_{s,geo})/\Delta T)$ of 0.21. The observation of higher activity of NCC01 compared to MCC and FC implies possible inclusion of INA materials in the original 3% solution of NCC01. The source is not known, and the source identification is beyond the scope of this work. The sample stability of another NCC sample from another batch, NCC02, is discussed in **Sect. 4.3.6**.

4.3.6. NIPR-CRAFT

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This suite of cold stage instruments offered the immersion freezing measurements of all three cellulose samples using droplets with volumes of 5 μ L. This microliter range volume was the largest amongst all aqueous suspension techniques employed within this work. Such a large drop volume advantageously enables high resolution immersion freezing analysis for a wide range of temperatures (-31 °C < T < -17 °C). The highest freezing temperatures are attained with the largest droplets, which contain the largest surface area of cellulose.

By means of Stokes-law gravity differential settling (Tobo, 2016), <10 μ m MCC and FC particles of were extracted to generate droplets containing size-segregated cellulose samples. These droplets were subsequently assessed on NIPR-CRAFT, estimating an immersion freezing ability of MCC and FC with SSA of 3.35 m^2 g^{-1} (The AIDA-derived geometric SSA value,

accounting for only <10 μ m particles). Afterwards, the obtained results of <10 μ m were compared to those of bulk (SEM-based SSA of 0.068-0.087 m² g⁻¹). Furthermore, we facilitated NIPR-CRAFT for the quality check of the NCC sample over time. Expressly, we stored NCC02 at 4 °C for 9 months and made follow-up measurements to examine the potential decay of the samples, potentially altering its immersion freezing.

Figures 3q and 4k show the NIPR-CRAFT results for MCC and FC. In general, the NIPR-CRAFT data represent the lower boundary of compiled $n_{s,geo}(T)$ spectrum defined by the bulk of the instruments (**Figs. 2.a.iii and 2.b.iii**). Constant offset between NIPR-CRAFT and the log average of AS methods in $n_{s,geo}(T)$ is seen at -28 °C < T < -21 °C for on average a factor of >9 for MCC and >2.7 for FC. Immersion freezing abilities of bulk and size-segregated samples are in agreement within the measurement uncertainties. The spectral slopes for bulk MCC and FC are 0.41 and 0.39, respectively, and are in agreements with WT-CRAFT (measurements with 3 μL sonicated samples), indicating the presence of systematic error (e.g., temperature shift towards the low end). The spectral slopes for size-segregated MCC and FC are 0.43 and 0.34, respectively, and are in agreement with bulk NIPR-CRAFT.

Figure 5i shows time-trials of NCC02 and similarity in IN activity over 9 months. As inferred from the overlapped spectra, the influence of the decay over time is negligible. Over the time, the spectral slopes and $n_{s,geo}(T)$ remain similar, indicating high stability of NCC02.

For investigated temperatures listed in **Table 2**, the bulk NIPR-CRAFT results show $n_{s,\text{geo},\text{MCC}} > n_{s,\text{geo},\text{FC}}$ (**Figs. 3q and 4k**). Corresponding $\Delta \log(n_{s,\text{geo}})/\Delta T$ values are similar (0.41 for MCC and 0.39 for FC) but notably higher than any averaged slope parameters listed in **Table 3**. With even higher slope value of 0.50, the $n_{s,\text{geo},\text{NCC}}$ values exceed both $n_{s,\text{geo},\text{MCC}}$ and $n_{s,\text{geo},\text{FC}}$ at T below -20 °C (**Fig. 5i**).

4.3.7. WT-CRAFT

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The WT-CRAFT system, which is a replica of NIPR-CRAFT (*Tobo*, 2016), measured the freezing abilities of droplets containing 0.05-0.0005 wt% MCC and FC at *T* > -26 °C. WT-CRAFT also examined if the pre-treatment of aqueous suspension (i.e., sonication of 50 mL falcon tube for 15 min) has any influences on IN efficiency of MCC and FC. More specifically, we compared the IN efficiency of 49 drops made out of the sonicated-suspension containing given wt% of MCC and FC to those of non-sonicated suspension left idle for at least 60 min.

The results are shown in **Figs. 3s and 4l**. As seen in these figures, early freezer only appears in the case of pre-application of sonication. This trend is especially notable for the MCC case. As a result, the difference of the spectral slope for MCC deviates from 0.36 (sonicated-case) to 0.52 non-sonicated case). Importantly, our results suggest that MCC may

suffer more from the particle settling in the suspension when compared to FC for examined ranges of temperature and wt%. Nevertheless, the difference in $n_{s,geo}(T)$ is within a factor of four at the most, which is well within our experimental uncertainty (see **Supplemental Table S2**).

Below -22 °C, WT-CRAFT shows $n_{s,geo,MCC} > n_{s,geo,FC}$ (**Figs. 3s and 4I**). The MCC result exhibits sharper increase in $n_{s,geo}(T)$ within the limited temperature range with $\Delta \log(n_{s,geo})/\Delta T$ of 0.36 than FC ($\Delta \log(n_{s,geo})/\Delta T = 0.30$).

4.3.8. AIDA

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The AIDA facility at KIT represents the world's foremost facility for studying ice clouds in a controlled setting. As shown in **Fig. 2**, for all cellulose types, the AIDA data hover in the upper bound of comprehensive $n_{s,geo}(T)$ spectrum defined by the bulk of the instruments. The corresponding $log(n_{s,ind.})/log(n_{s,avg})$ is within 1.2. The spectral slope for immersion freezing of cellulose from AIDA varies depending on the sample type. For MCC, $\Delta log(n_{s,geo})/\Delta T$ is 0.24 and equivalent to that of H15a (MCC, dry, **Eqn. 8**). The larger slope value is found for FC (0.47), which is practically parallel to A13 (0.45), and deviating from other DD instrument ($\Delta log(n_{s,geo})/\Delta T$ of 0.28). But, the $n_{s,geo}(T)$ data of FC form AIDA are in fair agreement with the $log n_{s,geo}(T)$ average for examined T. Finally, the NCC02 results agree well with CSU-CFDC and WISDOM. Observed quasi-flat $\Delta log(n_{s,geo})/\Delta T$ of NCC02 (0.04) suggests a week T-dependent immersion freezing ability for the investigated temperature range. In addition, similar to the observation made by LINDA, higher activity of NCC01 compared to NCC02 is seen in **Fig. 5a**. This difference suggests the inclusion of INA materials in the original 3% solution of NCC01 (the source is not known). For investigated temperatures listed in **Table 2**, AIDA show $n_{s,geo,MCC} > n_{s,geo,MCC} > n_{s,geo,NCC}$ (**Figs. 3a, 4a and 5a; see also the Supplemental Sect. S.7**).

4.3.9. EDB

The contact freezing experiments have been performed with MCC particles preselected in DMA at two electrical mobility diameters: 320 nm and 800 nm. Due to the low concentration (typically less than 30 cm⁻³) of the MCC particles produced by the dry dispersion method (a turbulent flow disperser, see **Supplemental Table S3**), and relatively low IN efficacy of MCC particles, the measurements of e_c were possible only in a limited temperature range between -29°C and -32 °C. A strong asphericity of the MCC particles contributes to the uncertainty of $n_{\rm s,geo}(T)$ determination, which differs by two orders of magnitude for particles with mobility diameters of 320 nm and 800 nm. Additional uncertainty factor is the unknown portion of the MCC particle submersed in water upon contact with the supercooled droplet (k_{imm} , see **Eqn. 2**). We set $k_{imm} = 1$ thus giving a lower estimate of the possible $n_{\rm s,geo}(T)$

value. On the whole, the contact INAS density falls nicely within the range of $n_{s,geo}(T)$ values measured by other instruments, but does not exceed H15MCC parametrization for dry NCC particles. This is not very surprising given the experimental uncertainties of the EDB-based method.

4.3.10. INKA

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INKA (Ice Nucleation Instrument of the Karlsruhe Institute of Technology; *Schiebel*, 2017) is a cylindrical continuous flow diffusion chamber built after the design of the CSU-CFDC (*Richardson*, 2010), but with a prolonged residence time of the sample (*Chen et al.*, 2000). Using INKA, we studied the condensation / immersion freezing of MCC, which was dry dispersed into a 4 m³ stainless steel tank using the same procedure as for the AIDA experiments. No additional impactor was used at the INKA inlet.

The aerosol freezing ability was measured from -32.5 °C to -25 °C for increasing relative humidity from well below liquid water saturation to about 110% RH in a total of eight scans. Data reported in this paper was interpolated at a relative humidity of 107% RH, taking into account that the nominal relative humidity for CFDCs has to be above 100% in order to enable full aerosol activation (*DeMott et al.*, 2015; *Garimella et al.*, 2017). INKA measured ice nucleation surface site densities which are close to the average of all measured data (see **Fig.** 2). The results match the data measured by the CSU-CDFDC for polydisperse aerosol, with slightly less pronounced temperature dependence.

20 **4.3.11. MRI**

MRI cloud simulation chamber experiments were conducted to demonstrate that MCC particles can act as efficient immersion freezing nuclei in simulated supercooled clouds. The evacuation rate was correspondent to the updraft velocity of 5 m s⁻¹. Dry MCC powders were dispersed by a rotating brush generator (PALAS, RBG1000) and injected into the ventilated 1.4 m³ chamber vessel. Using the data from six experiments, we calculated the ice nucleation active surface-site densities of aerosolized cellulose in the temperature range from -15 °C to -30 °C. The regression line for the experimental data is $n_{s,geo}(T) = \exp(-0.56T + 7.50)$ with a correlation coefficient of 0.84. As shown in **Figs. 2 and 3h**, for dry MCC type, the MRI cloud simulation chamber data exist in the upper bound of comprehensive $n_{s,geo}(T)$ spectrum.

30 **4.3.12. PNNL-CIC**

Immersion freezing properties of size-selected MCC samples at a temperature ranging from -20 to -28 °C were investigated. The chamber was operated at RH $_{\rm w}$ = 106 \pm 3%, and the evaporation section of the chamber was maintained at aerosol lamina temperature. The uncertainty (\pm 0.5 °C) in the aerosol lamina temperature was calculated based on aerosol

lamina profile calculations. $n_{s,geo}(T)$ calculations were performed using immersion freezing frozen fraction and surface area of MCC particles. The $n_{s,geo}(T)$ values varied from 1 x 10⁸ to 1 x 10⁹ m⁻². $\Delta \log(n_{s,geo})/\Delta T$ (= 0.13, **Fig. 3i**) agreed well with that of the U17 dust parameterization in the same temperature range.

4.3.13. BINARY

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The three different cellulose types were investigated with the BINARY setup ($Budke\ and\ Koop$, 2015), and their sample preparation is described in **Supplemental Table S4**. We note that the MCC and FC original data are those published in H15a, i.e., before the recommended suspension preparation procedure was developed. As described in H15a these bulk suspensions suffered from sedimentation and, hence, are not predestined for a $n_{s,geo}(T)$ intercomparison. The original raw data from H15a were re-analyzed here in order to have the same 1 °C binning and averaging as other techniques. Moreover, a different background correction was applied, also to the NCC samples: the first 5% and last 5% of nucleation data points in a given frozen fraction curve (i.e. the data smaller than 0.05 and greater than 0.95 in FF) were excluded, in order to account for a concentration variation between individual droplets due to sedimentation and for nucleation events triggered by the glass substrate or impurities in the "pure" water background.

For -25 °C < T < -22 °C, the bulk BINARY data for the different cellulose samples are in a similar active site range, i.e. the results show $n_{s,geo,MCC} > n_{s,geo,FC} \approx n_{s,geo,NCC}$ (**Figs. 3j, 4d and 5d**). At -25 °C the MCC and FC data show a rapid change in slope and at lower temperature they level off at a $n_{s,geo}(T)$ value of about 10^8 m⁻², which may be due to the sedimentation of cellulose particles with lower ice nucleation activity as discussed above. In contrast, no such change in slope is observed for NCC (which did not suffer from apparent sedimentation), thus being consistent with higher $n_{s,geo,NCC}$ values observed below -25 °C in small-droplet experiments and dry suspension techniques. Moreover, above -25 °C the NCC data agree well with other large-volume droplet experiments such as NIPR-CRAFT and NC-State CS as well as with small-droplet techniques such as WISDOM. In summary, these observations imply that techniques using large droplets may suffer from sedimentation if the suspended material consists of particles with a wide size distribution. However, if smaller and homogeneous particles are suspended they give results similar to small droplet techniques.

4.3.14. CMU-CS

The immersion freezing ability of wide range of aqueous suspension concentrations and immersion freezing temperatures was measured by CMU-CS (*Polen et al., 2016*; *Beydoun et al., 2017*; *Polen et al., 2018*). This cold stage device facilitates the sampling of drops within a

squalene oil matrix that allows for experiments using varied wt% of the cellulose test samples (0.001 to 0.15 wt%) for this study. Drops containing MCC, FC and NCC02 were studied at a cooling rate of 1 °C min⁻¹ to determine the immersion freezing temperature spectrum.

A total of 10 immersion mode freezing experiments with a droplet volume of 0.1 μ L were performed. Using this instrument, a wide range of temperatures was investigated (T > -30 °C) yielding $n_{s,geo}(T)$ values ranging from 10^5 to 10^{10} m⁻². The data from the ten individual runs collapsed into a single $n_{s,geo}(T)$ spectrum suggesting that the mass loading of dust in the droplet did not affect the measurements for the wt% values investigated. For MCC, the data are in fair quantitative agreement with the H15a (Dry MCC) parameterization at temperatures below -25 °C. The $n_{s,geo}(T)$ values of both FC and NCC are about one order magnitude lower than the MCC $n_{s,geo}(T)$ values, agreeing with a general trend and overlapping with the Wet MCC reference curve.

Remarkably, the CMU-CS data show that the value of $\Delta \log(n_{s,geo})/\Delta T$ for MCC (= 0.20, **Supplemental Table S2**) is the least amongst the aqueous suspension techniques and the closet to the results of the bulk dry techniques (the DD slope = 0.20, **Table 3**), potentially suggesting a similar and more atmospherically representative experimental condition (less particle inclusion in a single droplet) when compared to other aqueous methods.

At -25 °C, where the immersion freezing abilities of all three cellulose samples were assessed, the CMU-CS result shows $n_{s,\text{geo},\text{MCC}} > n_{s,\text{geo},\text{NCC}} > n_{s,\text{geo},\text{FC}}$ (**Figs. 3k, 4e and 5e**). Note that MCC and FC exhibit broad $n_{s,\text{geo}}(T)$ spectra with the $\Delta \log(n_{s,\text{geo}})/\Delta T$ values of 0.20 (MCC) and 0.34 (FC), detecting ice nucleation at <-29 °C, whereas the NCC spectrum spans for limited T range (-25 °C < T < -22 °C) with the $\Delta \log(n_{s,\text{geo}})/\Delta T$ value of 0.51. The observed widening of the spectra and detection temperature sensitivity suggests that supermicron-sized particles have increased diversity in immersion freezing as compared to submicron particles.

25 **4.3.15. Leeds-NIPI**

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 μ L-NIPI is a droplet freezing device which controls the temperature of 1 μ L water droplets supported on a hydrophobic glass slide and monitors freezing in those droplets (Whale et al. 2015). For this study, 0.1 wt% suspensions of FC and MCC cellulose were made up in Milli-Q water by stirring for 30 minutes in glass vials. The suspensions were then stirred continuously while 1 μ L droplets were pipetted onto a hydrophobic glass slide using an electronic pipette. Droplets were then cooled from room temperature (~18 °C) at a rate of 1 °C min⁻¹ until they froze, freezing being monitored by a digital camera. A gentle flow of dry nitrogen was passed over the droplets to ensure that ice did not grow across the hydrophobic slide and cause unwanted droplet freezing. Temperature error for the instrument has been estimated at \pm 0.4

°C and $n_{s,geo}(T)$ error bars were calculated by propagating the uncertainties from droplet volume and weighing of the cellulose and water. The instrument has a freezing background, likely caused by minor impurities in the Milli-Q water or on the hydrophobic slide. A background subtraction is performed to account for any freezing caused by this background (O'Sullivan et al. 2015) however the freezing reported here occurred at sufficiently warm temperatures such that they did not overlap with the background freezing. For investigated temperatures listed in **Table 2**, Leeds-NIPI show $n_{s,geo,FC} \approx n_{s,geo,MCC}$, but the $n_{s,geo,FC}$ values are on average a factor of two higher than $n_{s,geo,MCC}$ across the investigated T range (**Figs. 31 and 4f**). The $\Delta \log(n_{s,geo})/\Delta T$ values for MCC and FC are 0.47 and 0.57, respectively.

4.3.16. M-AL

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For investigating the immersion freezing of droplets containing cellulose particles we have utilized two independent contact-free drop levitation methods in our laboratory at the Johannes Gutenberg University of Mainz, Germany. One of them is the Mainz Acoustic Levitator (M-AL) which was placed inside a walk-in cold room where the ambient temperature was set to be -30 °C. After introducing single drops into M-AL the drops were cooling down (at a continuously varying cooling rate) adapting their surface temperature to the ambient temperature. The size of the levitated drops was approx. 2 mm which was determined for each drop from the images captured by a digital video camera attached to the M-AL. Such large droplet size enabled the direct measurement of the surface temperature during the experiments with means of an infrared thermometer, therefore reducing the error in temperature originating from indirect determination of droplet temperature. The onset of freezing was characterized by a sudden increase in the surface temperature caused by the latent heat released during nucleation. The freezing temperatures of 100 drops was measured for each cellulose samples (MCC, FC and NCC) at two distinct concentrations, 1.0 and 0.1 wt%. Due to the relatively large droplet size a wide range of temperatures was covered (-13 to -23 °C) yielding $n_{s,geo}(T)$ values ranging from 10^4 to 10^7 m⁻². The NCC sample we got for investigation was contaminated by mold therefore the $n_{s,geo}(T)$ deviates significantly from other techniques at temperature above -20 °C (see Fig. 1c. iii). For investigated temperatures listed in Table 2, M-AL shows $n_{s,geo,MCC} > n_{s,geo,FC}$ and $n_{s,geo,MCC} > n_{s,geo,NCC}$ (Figs. 3n, 4h and 5g). For example, at -17 °C, the $n_{\rm s,geo}(T)$ values of MCC, FC and NCC are 2.54 x 10⁵, 2.48 x 10⁵ and 8.28 x 10⁴ m⁻². The $\Delta \log(n_{\rm s,geo})/\Delta T$ values vary for 0.28 (FC)-0.40 (MCC) with the spectral parameter of NCC (0.31) falling around the middle.

4.3.17. M-WT

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The main facility of our laboratory at the JGU Mainz is a vertical wind tunnel (M-WT) in which atmospheric hydrometeors can be freely suspended in the updraft of the tunnel at temperatures down to -30 °C. Since all hydrometeors (from cloud droplets of few tens of μ m to large hailstones with sizes of several centimeters) can be freely floated at their terminal falling velocities the relevant physical quantities, as for instance the Reynolds number and the ventilation coefficient, are equal to those in the real atmosphere.

The immersion freezing measurements in the M-WT have been conducted under isothermal conditions. The air was cooled down to a certain temperature between -20 and -25 °C and at that temperature the frozen fraction of water droplets containing MCC or FC was measured by investigating typically 50 droplets a day. The drop temperatures were determined from the continuously recorded air temperature and humidity (*Diehl et al., 2014*; *Pruppacher and Klett, 2010*). The size of the droplets was calculated from the vertical air speed which can be measured by high accuracy in the M-WT (*Diehl et al., 2014*). Due to the small droplet size and the applied INP concentration (0.1 wt%) a relatively narrow temperature range could be investigated yielding $n_{s,geo}(T)$ values ranging from 10^6 to 10^8 m⁻². Over -23 °C < T < -22 °C, M-WT shows $n_{s,geo,MCC} > n_{s,geo,FC}$ (**Figs. 3o and 4i**). Corresponding $\Delta \log(n_{s,geo})/\Delta T$ values are 0.26 for MCC and 0.48 for FC.

4.3.18. NC State-CS

Across investigated temperatures ($T \in [-23, -16]$ °C), results from the NC State CS show that INAS is indistinguishable between FC, MCC, and NCC for all temperatures within experimental uncertainty, except for T > -18 °C where $n_{s,geo,NCC}$ is less than that of FC and MCC. Overall, the NCC spectrum is narrower than the FC and MCC spectra, suggesting that the distribution of active sites for NCC is slightly more homogenous. The data connect with the $n_{s,geo}^{H15MCC,wet}$ parameterization at T = -22 °C, but falls below by ~ 1 order of magnitude at T = -23 °C. The data intersect with the $n_{s,geo}^{H15NX,wet}$ parameterization in the -20 < T < -18 °C range. However, the $n_{s,geo}^{H15NX,wet}$ has a steeper slope with temperature and thus overpredicts and underpredicts $n_{s,geo,cellulose}$ at colder and warmer temperatures, respectively (see also the **Supplemental Sect. S.8**).

4.3.19. WISDOM

Over the investigated temperature range given in **Table 2**, WISDOM shows $n_{s,\text{geo},\text{MCC}} > n_{s,\text{geo},\text{NCC}}$ (**Figs. 3r and 5j**). The MCC result exhibits broader spectrum with $\Delta \log(n_{s,\text{geo}})/\Delta T$ of 0.26 than NCC ($\Delta \log(n_{s,\text{geo}})/\Delta T = 0.31$). The observed relation between widening of spectra and increased $n_{s,\text{geo}}(T)$ suggests that supermicron-sized particles have increased diversity in immersion

freezing as compared to submicron particles. Looking at the overall NCC data (Fig. 1.c.iii), nearly all aqueous suspension techniques, independently of the drop volume, agree with the WISDOM data and all point towards the AIDA data. We remark that the WISDOM team followed the suggested sample handling details described in Sect. 3.1.

5. Conclusion and Future Outlook

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This paper presents the immersion freezing efficiencies of cellulose particles of three different types evaluated by a total of twenty IN instruments at supercooled temperature conditions. Three cellulose samples examined in this study showed a propensity to nucleate ice, and their ice nucleation activity are comparable to another test system (i.e., illite NX) that we have previously evaluated. On average, supermicron-sized cellulose samples are more ice active than the nano cellulose one at *T* lower than -20 °C although the difference is not apparent for all temperatures when considering experimental uncertainty.

Our work also provides a comprehensive dataset of experimental variables in INP measurement techniques to complement our insufficient knowledge regarding inter-method diversity that, when filled, will enhance the credibility of our experiments to evaluate INP abundance in the atmosphere. Strikingly, our results indicate that the overall diversity derived from comparing techniques is significant when compared to the individual uncertainties of each instrument.

The observed diversity amongst measurement techniques for cellulose is larger than that observed for a mineralogically heterogeneous illite NX sample described in our previous inter-comparison study (H15b). For illite NX, the deviations in temperature (-36 °C < T < -4 °C) are within 8 °C (H15b) while they span 10 °C for cellulose. For $n_{s,geo}(T)$, while the span in results covers a maximum of three orders of magnitude for illite NX, they span 4 orders of magnitude for cellulose. These diversities suggest the complex surface structure and compositional heterogeneity may play a substantial role to explain the diversity. This also implies that the cellulose system might not be suitable as a calibrant at this stage unless we completely understand the complex properties of different cellulose materials.

In conclusion, we have shown that several types of cellulose have the capacity to nucleate ice as efficiently as some mineral dust samples. Given cellulose within plant residue is present in the atmosphere, it represents a poorly characterised non-proteinaceous INP type. While the diverse instruments employed in this study agree in that cellulose has the capacity to nucleate ice, their quantitative agreement is poor. Unfortunately, it is not possible as yet to say what the cause of this disagreement is. We suggest a number of topics that future studies could address in order to better understand and resolve this discrepancy (see the **Supplemental Sects. S.4, S.5, S.9 and S.10**). Nevertheless, we show that cellulose has the potential to be an important atmospheric ice nucleating particle and more work is warranted. Our knowledge of non-proteinaceous biological INPs is still limited. Thus, it is important to further conduct comprehensive studies on the ice nucleation activity of other important plant

structural materials, such as cellulose polymorphs, lignin materials, lipids, carbohydrates and other macromolecule saccharides (e.g., *Pummer et al.*, 2012; *Dreischmeier et al.*, 2017; *Suski et al.*, 2018), as well as natural plant debris in simulated supercooled clouds of the lower and middle troposphere. Such additional studies are especially important for assessing the overall role of non-proteinaceous bio-INPs in clouds and the climate system.

References

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- Atkinson, J. D., Murray, B. J., Woodhouse, M. T., Carslaw, K., Whale, T. F., Baustian, K., Dobbie, S., O'Sullivan, D., and Malkin, T. L.: The importance of feldspar for ice nucleation by mineral dust in mixed-phase clouds, Nature, 498, 355–358, doi:10.1038/nature12278, 2013.
 - Aulin, C., Ahola, S., Josefsson, P., Nishino, T., Hirose, Y., Österberg, M., Wågberg, L.: Nanoscale cellulose films with different crystallinities and mesostructures their surface properties and interaction with water, Langmuir, 25, 7675–7685, doi: 10.1021/la900323n, 2009.
- Battista, O. A. and Smith, P. A.: Microcrystalline cellulose, Ind. Eng. Chem., 54, 20–29, doi: 10.1021/ie50633a003, 1962.
- Belosi, F., and Santachiara, G.: Ice-formation nuclei in Antarctica: new and past measurements
 Atmos. Res., 145–146, 105–111, doi: https://doi.org/10.1016/j.atmosres.2014.03.030, 2014.
 - Beydoun, H., Polen, M., and Sullivan, R. C.: Effect of particle surface area on ice active site densities retrieved from droplet freezing spectra, Atmos. Chem. Phys., 16, 13359–13378, doi: https://doi.org/10.5194/acp-16-13359-2016, 2016.
- Beydoun, H., Polen, M., and Sullivan, R. C.: A new multicomponent heterogeneous ice nucleation model and its application to Snomax bacterial particles and a Snomax–illite mineral particle mixture, Atmos. Chem. Phys., 17, 13545–13557, doi: https://doi.org/10.5194/acp-17-13545-2017, 2017.
- Boucher, O., Randall, D., Artaxo, P., Bretherton, C., Feingold, G., Forster, P., Kerminen, V.-M., Kondo, Y., Liao, H., Lohmann, U., Rasch, P., Satheesh, S. K., Sherwood, S., Stevens, B., and Zhang, X. Y.: Clouds and Aerosols, in: Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, edited by: Stocker, T. F., Qin, D., Plattner, G.-K., Tignor, M., Allen, S. K., Boschung, J., Nauels, A., Xia, Y., Bex, V., and Midgley, P. M., Cambridge University Press, Cambridge, United
 Kingdom and New York, NY, USA, 571–657, 2013.
 - Brinchi, L., Cotana, F., Fortunati, E., and Kenny, J. M.: Production of nanocrystalline cellulose from lignocellulosic biomass: technology and applications, Carbohydrate Polymers, 94, 154–169, doi: https://doi.org/10.1016/j.carbpol.2013.01.033, 2013.
 - Brunauer, S., Emmett, P. H., and Teller, E.: Adsorption of gases in multimolecular layers, J. Am. Chem. Soc., 60, 309–319, doi: 10.1021/ja01269a023, 1938.
- Budke, C. and Koop, T.: BINARY: an optical freezing array for assessing temperature and time dependence of heterogeneous ice nucleation, Atmos. Meas. Tech., 8, 689–703, doi: https://doi.org/10.5194/amt-8-689-2015, 2015.
- Burkert-Kohn, M., Wex, H., Welti, A., Hartmann, S., Grawe, S., Hellner, L., Herenz, P., Atkinson, J. D., Stratmann, F., and Kanji, Z. A.: Leipzig Ice Nucleation chamber Comparison (LINC): intercomparison of four online ice nucleation counters, Atmos. Chem. Phys., 17, 11683–11705, doi: https://doi.org/10.5194/acp-17-11683-2017, 2017.
 - Chawla, P. R., Bajaj, I. B., Survase, S. A., Singhal, R.S.: Microbial cellulose: Fermentative production and applications, Food Technology and Biotechnology, 47, 107–124, 2009.
- Chen, Y., DeMott, P. J., Kreidenweis, S. M., Rogers, D. C., and Sherman, D. E.: Ice formation by sulfate and sulfuric acid aerosol particles under upper-tropospheric conditions. J. Atmos. Sci., 57, 3752–3766, doi: https://doi.org/10.1175/1520-0469(2000)057<3752:IFBSAS>2.0.CO;2, 2000.
- Connolly, P. J., Emersic, C., and Field, P. R.: A laboratory investigation into the aggregation efficiency of small ice crystals, Atmos. Chem. Phys., 12, 2055–2076, doi: https://doi.org/10.5194/acp-12-2055-2012, 2012.
 - DeMott, P. J., Prenni, A. J., McMeeking, G. R., Sullivan, R. C., Petters, M. D., Tobo, Y., Niemand, M., Möhler, O., Snider, J. R., Wang, Z., and Kreidenweis, S. M.: Integrating laboratory and field data to quantify the immersion freezing ice nucleation activity of mineral dust particles, Atmos. Chem. Phys., 15, 393–409, https://doi.org/10.5194/acp-15-393-2015, 2015.

- DeMott, P. J., Hill, T. C. J., Petters, M. D., Bertram, A. K., Tobo, Y., Mason, R. H., Suski, K. J., McCluskey, C. S., Levin, E. J. T., Schill, G. P., Boose, Y., Rauker, A. M., Miller, A. J., Zaragoza, J., Rocci, K., Rothfuss, N. E., Taylor, H. P., Hader, J. D., Chou, C., Huffman, J. A., Pöschl, U., Prenni, A. J., and Kreidenweis, S. M.: Comparative measurements of ambient atmospheric concentrations of ice nucleating particles using multiple immersion freezing methods and a continuous flow diffusion chamber, Atmos. Chem. Phys., 17, 11227–11245, https://doi.org/10.5194/acp-17-11227-2017, 2017.
- Diehl, K., Mitra, S. K., Szakáll, M., Blohn, N. v., Borrmann, S., and Pruppacher, H.R.: Chapter 2. Wind Tunnels: Aerodynamics, Models, and Experiments. In: The Mainz Vertical Wind Tunnel Facility: A Review of 25 Years of Laboratory Experiments on Cloud Physics and Chemistry [Pereira, J. D. (eds.)], Nova Science Publishers, Inc., Hauppauge, NY, USA, 2011.

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35

- Diehl, K., Debertshäuser, M., Eppers, O., Schmithüsen, H., Mitra, S.K., and Borrmann, S.: Particle-area dependence of mineral dust in the immersion mode: investigations with freely suspended drops in an acoustic levitator. Atmos. Chem. Phys., 14, 12343–12355, doi: https://doi.org/10.5194/acp-14-12343-2014, 2014.
 - Diehl, K. and Mitra, S. K.: New particle-dependent parameterizations of heterogeneous freezing processes: sensitivity studies of convective clouds with an air parcel model, Atmos. Chem. Phys., 15, 12741-12763, https://doi.org/10.5194/acp-15-12741-2015, 2015.
- Dittenber, D. B., and GangaRao, H. V. S. Critical review of recent publications on use of natural composites in infrastructure, Composites Part A: Applied Science and Manufacturing, 43, 1419–1429, doi: https://doi.org/10.1016/j.compositesa.2011.11.019, 2012.
- Dreischmeier, K., Budke, C., Wiehemeier, L., Kottke, T., and Koop, T.: Boreal pollen contain ice-nucleating as well as ice-binding 'anti-freeze' polysaccharides, Sci. Rep., 7, 41890, 2017.
- Emersic, C., Connolly, P. J., Boult, S., Campana, M., and Li, Z.: Investigating the discrepancy between wet-suspension- and dry-dispersion-derived ice nucleation efficiency of mineral particles, Atmos. Chem. Phys., 15, 11311–11326, doi: https://doi.org/10.5194/acp-15-11311-2015, 2015.
 - Fernández, I., Cabaneiro, A., and Carballas, T.: Organic matter changes immediately after a wildfire in an Atlantic forest soil and comparison with laboratory soil heating, Soil Biol. Biochem., 29, 1–11, doi: https://doi.org/10.1016/S0038-0717(96)00289-1, 1997.
 - Friedman, B., Kulkarni, G., Beránek, J., Zelenyuk, A., Thornton, J. A., and Cziczo, D. J.: Ice nucleation and droplet formation by bare and coated soot particles, J. Geophys. Res., 116, D17203, doi: https://doi.org/10.1029/2011JD015999, 2011.
- Garimella, S., Rothenberg, D. A., Wolf, M. J., David, R. O., Kanji, Z. A., Wang, C., Rösch, M., and Cziczo, D. J.: Uncertainty in counting ice nucleating particles with continuous flow diffusion chambers, Atmos. Chem. Phys., 17, 10855–10864, doi: https://doi.org/10.5194/acp-17-10855-2017, 2017.
- Grawe, S., Augustin-Bauditz, S., Hartmann, S., Hellner, L., Pettersson, J. B. C., Prager, A., Stratmann, F., and Wex,
 H.: The immersion freezing behavior of ash particles from wood and brown coal burning, Atmos. Chem. Phys., 16,
 13911–13928, doi: https://doi.org/10.5194/acp-16-13911-2016, 2016.
- Hartmann, S., Niedermeier, D., Voigtländer, J., Clauss, T., Shaw, R. A., Wex, H., Kiselev, A., and Stratmann, F.: Homogeneous and heterogeneous ice nucleation at LACIS: operating principle and theoretical studies, Atmos. Chem. Phys., 11, 1753–1767, doi: https://doi.org/10.5194/acp-11-1753-2011, 2011.
 - Hiranuma, N., Hoffmann, N., Kiselev, A., Dreyer, A., Zhang, K., Kulkarni, G., Koop, T., and Möhler, O.: Influence of surface morphology on the immersion mode ice nucleation efficiency of hematite particles, Atmos. Chem. Phys., 14, 2315-2324, https://doi.org/10.5194/acp-14-2315-2014, 2014.
 - Hiranuma, N., Möhler, O., Yamashita, K., Tajiri, T., Saito, A., Kiselev, A., Hoffmann, N., Hoose, C., Jantsch, E., Koop, T., and Murakami, M.: Ice nucleation by cellulose and its potential contribution to ice formation in clouds, Nat. Geosci., 8, 273–277, doi: https://doi.org/10.1038/ngeo2374, 2015a.
- Hiranuma, N., Augustin-Bauditz, S., Bingemer, H., Budke, C., Curtius, J., Danielczok, A., Diehl, K., Dreischmeier, K., Ebert, M., Frank, F., Hoffmann, N., Kandler, K., Kiselev, A., Koop, T., Leisner, T., Möhler, O., Nillius, B., Peckhaus, A., Rose, D., Weinbruch, S., Wex, H., Boose, Y., DeMott, P. J., Hader, J. D., Hill, T. C. J., Kanji, Z. A., Kulkarni, G., Levin, E. J. T., McCluskey, C. S., Murakami, M., Murray, B. J., Niedermeier, D., Petters, M. D., O'Sullivan, D., Saito, A., Schill, G. P., Tajiri, T., Tolbert, M. A., Welti, A., Whale, T. F., Wright, T. P., and Yamashita, K.: A comprehensive laboratory

- study on the immersion freezing behavior of illite NX particles: a comparison of 17 ice nucleation measurement techniques, Atmos. Chem. Phys., 15, 2489–2518, doi: https://doi.org/10.5194/acp-15-2489-2015, 2015b.
- Hoffmann, N., Duft, D., Kiselev, A., and Leisner, T.: Contact freezing efficiency of mineral dust aerosols studied in an electrodynamic balance: quantitative size and temperature dependence for illite particles, Faraday Discuss., 165, 383–390, doi:10.1039/C3FD00033H, 2013a.
 - Hoffmann, N., Kiselev, A., Rzesanke, D., Duft, D., and Leisner, T.: Experimental quantification of contact freezing in an electrodynamic balance, Atmos. Meas. Tech., 6, 2373–2382, doi:10.5194/amt-6-2373-2013, 2013b.
 - Hoose, C. and Möhler, O.: Heterogeneous ice nucleation on atmospheric aerosols: a review of results from laboratory experiments, Atmos. Chem. Phys., 12, 9817–9854, doi: https://doi.org/10.5194/acp-12-9817-2012, 2012.

25

35

40

- Kanji, Z. A., Ladino, L. A., Wex, H., Boose, Y., Burkert-Kohn, M., Cziczo, D. J., and Krämer, M.: Ice formation and evolution in clouds and precipitation: Measurement and modeling challenges, Chapter 1: Overview of ice nucleating particles, Meteor. Mon., 58, 1.1–1.33, doi: https://doi.org/10.1175/AMSMONOGRAPHS-D-16-0006.1, 2017.
- Khalil, H. P. S. A., Bhat, A. H., and Yusra, A. F. I.: Green composites from sustainable cellulose nanofibrils: A review, Carbohydrate Polymers, 87, 963–979, doi: https://doi.org/10.1016/j.carbpol.2011.08.078, 2012.
 - Kiselev, A., Bachmann, F., Pedevilla, P., Cox, S. J., Michaelides, A., Gerthsen, D., and Leisner, T.: Active sites in heterogeneous ice nucleation—the example of K-rich feldspars, Science, 355, 367–371, doi: https://doi.org/10.1126/science.aai8034, 2017.
 - Knopf, D. A., Alpert, P. A., and Wang, B.: The role of organic aerosol in atmospheric ice nucleation: A review, ACS Earth Space Chem., 2, 168–202, doi: 10.1021/acsearthspacechem.7b00120, 2018.
- Knopf, D. A., Alpert, P. A., and Wang, B.: The role of organic aerosol in atmospheric ice nucleation: A review, ACS Earth Space Chem., 2, 168–202, doi: 10.1021/acsearthspacechem.7b00120, 2018.
 - Legrand, M., Preunkert, S., Schock, M., Cerqueira, M., Kasper-Giebl, A., Afonso, J., Pio, C., Gelencser, A., and Dombrowski-Etchevers, L.: Major 20th century changes of carbonaceous aerosol components (EC, WinOC, DOC, HULIS, carboxylic acids, and cellulose) derived from Alpine ice cores, J. Geophys. Res.-Atmos., 112, D23S11, doi:10.1029/2006JD008080, 2007.
 - Möhler, O., Stetzer, O., Schaefers, S., Linke, C., Schnaiter, M., Tiede, R., Saathoff, H., Krämer, M., Mangold, A., Budz, P., Zink, P., Schreiner, J., Mauersberger, K., Haag, W., Kärcher, B., and Schurath, U.: Experimental investigation of homogeneous freezing of sulphuric acid particles in the aerosol chamber AIDA, Atmos. Chem. Phys., 3, 211–223, doi: https://doi.org/10.5194/acp-3-211-2003, 2003.
 - Murray, B. J., O'Sullivan, D., Atkinson, J. D., and Webb, M. E.: Ice nucleation by particles immersed in supercooled cloud droplets, Chem. Soc. Rev., 41, 6519–6554, doi:10.1039/c2cs35200a, 2012.
- Niemand, M., Möhler, O., Vogel, B., Vogel, H., Hoose, C., Connolly, P., Klein, H., Bingemer, H., DeMott, P., and Skrotzki, J.: A particle-surface-area-based parameterization of immersion freezing on desert dust particles, J. Atmos. Sci., 69, 3077–3092, doi: https://doi.org/10.1175/JAS-D-11-0249.1, 2012.
- Nishiyama, Y., Langan, P. and Chanzy, H.: Crystal structure and hydrogen-bonding system in Cellulose Iβ from synchrotron X-ray and neutron fiber diffraction, J. Am. Chem. Soc. 124, 9074–9082, doi: 10.1021/ja0257319, 2002.
 - O'Sullivan, D., Murray, B. J., Ross, J. F., Whale, T. F., Price, H. C., Atkinson, J. D., Umo, N. S., and Webb, M. E.: The relevance of nanoscale biological fragments for ice nucleation in clouds, Sci. Rep., 5, 8082, doi: 10.1038/srep08082, 2015.
 - Paukert, M. and Hoose, C.: Modeling immersion freezing with aerosol-dependent prognostic ice nuclei in Arctic mixed-phase clouds, J. Geophys. Res. Atmos., 119, 9073–9092, doi:10.1002/2014JD021917, 2014.
- Polen, M., Lawlis, E., and Sullivan, R. C.: The unstable ice nucleation properties of Snomax® bacterial particles, J. Geophys. Res., 121, 11666–11678, doi: 10.1002/2016JD025251., 2016.
 - Polen, M., Brubaker, T., Somers, J., and Sullivan, R. C.: Cleaning up our water: reducing interferences from nonhomogeneous freezing of "pure" water in droplet freezing assays of ice-nucleating particles, Atmos. Meas. Tech., 11, 5315–5334, https://doi.org/10.5194/amt-11-5315-2018, 2018.

- Pruppacher, H. R. and Klett, J. D.: Microphysics of Clouds and Precipitation, Kluwer Acad., Norwell, Mass., 954 pp, doi: 10.1007/978-0-306-48100-0, 2010.
- 5 Pummer, B. G., Bauer, H., Bernardi, J., Bleicher, S., and Grothe, H.: Suspendable macromolecules are responsible for ice nucleation activity of birch and conifer pollen, Atmos. Chem. Phys., 12, 2541–2550, https://doi.org/10.5194/acp-12-2541-2012, 2012.
- Puxbaum, H. and Tenze-Kunit, M.: Size distribution and seasonal variation of atmospheric cellulose. Atmos. Environ., 37, 3693–3699, doi:10.1016/S1352-2310(03)00451-5, 2003.
 - Quiroz-Castañeda, R. E. and Folch-Mallol, J. L.: Hydrolysis of Biomass Mediated by Cellulases for the Production of Sugars. In: Sustainable Degradation of Lignocellulosic Biomass Techniques, Applications and Commercialization, edited by: Chandel. A., ISBN: 978-953-51-1119-1, InTech, doi: 10.5772/53719, 2013.
- Reicher, N., Segev, L., and Rudich, Y.: The Welzmann Supercooled Droplets Observation on a Microarray (WISDOM) and application for ambient dust, Atmos. Meas. Tech., 11, 233-248, doi: https://doi.org/10.5194/amt-11-233-2018, 2018.
- Richardson, M. S., DeMott, P. J., Kreidenweis, S. M., Petters, M. D., and Carrico, C. M.: Observations of ice nucleation by ambient aerosol in the homogeneous freezing regime, Geophys. Res. Lett., 37, L04806, doi: https://doi.org/10.1029/2009GL041912, 2010.
- Samake, A., Jaffrezo, J.-L., Favez, O., Weber, S., Jacob, V., Albinet, A., Riffault, V., Perdrix, E., Waked, A., Golly, B., Salameh, D., Chevrier, F., Oliveira, D. M., Besombes, J.-L., Martins, J. M. F., Conil, S., Guillaud, G., Meshba, B., Rocq, B., Robic, P.-Y., Hulin, A., Le Meur, S., Descheemaecker, M., Chretien, E., and Uzu, G.: Polyols and glucose particulate species as tracers of primary biogenic organic aerosols at 28 french sites, Atmos. Chem. Phys. Discuss., https://doi.org/10.5194/acp-2018-773, in review, 2018.
- Sánchez-Ochoa, A., Kasper-Giebl, A., Puxbaum, H., Gelencserm, A., Legrand, M., and Pio, C.: Concentration of atmospheric cellulose: a proxy for plant debris across a west-east transect over Europe, J. Geophys. Res., 112, D23S08, doi:10.1029/2006JD008180, 2007.
- Santachiara, G., Di Matteo, L., Prodi, F., and Belosi, F.: Atmospheric particles acting as ice forming nuclei in different size ranges, Atmos. Res., 96, 266–272, https://doi.org/10.1016/j.atmosres.2009.08.004, 2010.
 - Schiebel, T.: Ice Nucleation Activity of Soil Dust Aerosols, Thesis, Karlsruhe Institute of Technology, October 20th, doi: 10.5445/IR/1000076327, 2017.
- Schnell, R. and Vali, G.: Atmospheric ice nuclei from decomposing vegetation, Nature, 236, 163–165, doi: https://doi.org/10.1038/236163a0, 1972.

- Schnell, R. and Vali, G.: World-wide source of leaf-derived freezing nuclei, Nature, 246, 212–213, doi: https://doi.org/10.1038/246212a0, 1973.
- Schnell, R. and Vali, G.: Biogenic Ice Nuclei: Part I, Terrestrial and Marine Sources, J. Atmos. Sci., 33, 1554–1564, 1976.
- Schrod, J., Danielczok, A., Weber, D., Ebert, M., Thomson, E. S., and Bingemer, H. G.: Re-evaluating the Frankfurt isothermal static diffusion chamber for ice nucleation, Atmos. Meas. Tech., 9, 1313–1324, doi: https://doi.org/10.5194/amt-9-1313-2016, 2016.
- Schütze, K., Wilson, J. C., Weinbruch, S., Benker, N., Ebert, M., Günther, G., Weigel, R., and Borrmann, S.: Submicrometer refractory carbonaceous particles in the polar stratosphere, Atmos. Chem. Phys., 17, 12475-12493, https://doi.org/10.5194/acp-17-12475-2017, 2017.
 - Stopelli, E., Conen, F., Zimmermann, L., Alewell, C., and Morris, C. E.: Freezing nucleation apparatus puts new slant on study of biological ice nucleators in precipitation, Atmos. Meas. Tech., 7, 129–134, doi: https://doi.org/10.5194/amt-7-129-2014, 2014.
- Suski, K. J., Hill, T. C. J., Levin, E. J. T., Miller, A., DeMott, P. J., and Kreidenweis, S. M.: Agricultural harvesting emissions of ice-nucleating particles, Atmos. Chem. Phys., 18, 13755–13771, https://doi.org/10.5194/acp-18-13755-2018, 2018.

- Szakáll, M., Mitra, S. K., Diehl, K., and Borrmann, S.: Shapes and oscillations of falling raindrops: A review, Atmos. Res., 97, 416–425, doi: https://doi.org/10.1016/j.atmosres.2010.03.024, 2010.
- Tajiri, T., Yamashita, K., Murakami, M., Orikasa, N., Saito, A., Kusunoki, K., and Lilie, L.: A novel adiabatic-expansion-type cloud simulation chamber, J. Meteor. Soc. Japan, 91, 5, 687–704, doi: https://doi.org/10.2151/jmsj.2013-509, 2013.

25

- Thakur, V. K., and Thakur, M. K.: Processing and characterization of natural cellulose fibers/thermoset polymer composites, Carbohydr. Polym., 109, 102–117, doi: https://doi.org/10.1016/j.carbpol.2014.03.039, 2014.
- Timko, M. T., Yu, Z., Kroll, J., Jayne, J. T., Worsnop, D. R., Miake-Lye, R. C., Onasch, T. B., Liscinsky, D., Kirchstetter, T. W., Destaillats, H., Holder, A. L., Smith, J. D., and Wilson, K. R.: Sampling artifacts from conductive silicone tubing, Aerosol. Sci. Tech., 43, 855–865, 2009.
- Tobo, Y.: An improved approach for measuring immersion freezing in large droplets over a wide temperature range, Sci. Rep., 6, 32930, doi: 10.1038/srep32930, 2016.

 Ullrich, R., Hoose, C., Möhler, O., Niemand, M., Wagner, R., Höhler, K., Hiranuma, N., Saathoff, H., and Leisner, T.:
- A new ice nucleation active site parametrization for desert dust and soot, J. Atmos. Sci., 74, 699–717, doi: https://doi.org/10.1175/JAS-D-16-0074.1, 2017.
- Vali, G.: Quantitative evaluation of experimental results on the heterogeneous freezing nucleation of supercooled liquids, J. Atmos. Sci., 28, 402–409, doi: https://doi.org/10.1175/1520-0469(1971)028<0402:QEOERA>2.0.CO;2, 1971.
 - Vali, G., DeMott, P. J., Möhler, O., and Whale, T. F.: Technical Note: A proposal for ice nucleation terminology, Atmos. Chem. Phys., 15, 10263–10270, doi: https://doi.org/10.5194/acp-15-10263-2015, 2015.
- Vergara-Temprado, J., Miltenberger, A. K., Furtado, K., Grosvenor, D. P., Shipway, B. J., Hill, A. A., Wilkinson, J. M., Field, P. R., Murray, B. J., and Carslaw, K. S.: Strong control of Southern Ocean cloud reflectivity by ice-nucleating particles, Proc. Natl. Acad. Sci. U.S.A., 201721627, doi: 10.1073/pnas.1721627115, 2018.
- Vlachou, A., Daellenbach, K. R., Bozzetti, C., Chazeau, B., Salazar, G. A., Szidat, S., Jaffrezo, J.-L., Hueglin, C.,
 Baltensperger, U., Haddad, I. E., and Prévôt, A. S. H.: Advanced source apportionment of carbonaceous aerosols by coupling offline AMS and radiocarbon size-segregated measurements over a nearly 2-year period, Atmos. Chem. Phys., 18, 6187–6206, https://doi.org/10.5194/acp-18-6187-2018, 2018.
- Wex, H., DeMott, P. J., Tobo, Y., Hartmann, S., Rösch, M., Clauss, T., Tomsche, L., Niedermeier, D., and Stratmann,
 40 F.: Kaolinite particles as ice nuclei: learning from the use of different kaolinite samples and different coatings,
 Atmos. Chem. Phys., 14, 5529–5546, doi: https://doi.org/10.5194/acp-14-5529-2014, 2014.
- Wex, H., Augustin-Bauditz, S., Boose, Y., Budke, C., Curtius, J., Diehl, K., Dreyer, A., Frank, F., Hartmann, S., Hiranuma, N., Jantsch, E., Kanji, Z. A., Kiselev, A., Koop, T., Möhler, O., Niedermeier, D., Nillius, B., Rösch, M., Rose,
 D., Schmidt, C., Steinke, I., and Stratmann, F.: Intercomparing different devices for the investigation of ice nucleating particles using Snomax* as test substance, Atmos. Chem. Phys., 15, 1463–1485, doi: https://doi.org/10.5194/acp-15-1463-2015, 2015.
- Whale, T. F., Murray, B. J., O'Sullivan, D., Wilson, T. W., Umo, N. S., Baustian, K. J., Atkinson, J. D., Workneh, D. A., and Morris, G. J.: A technique for quantifying heterogeneous ice nucleation in microlitre supercooled water droplets, Atmos. Meas. Tech., 8, 2437–2447, https://doi.org/10.5194/amt-8-2437-2015, 2015.
 - Wright, T. P. and Petters, M. D.: The role of time in heterogeneous freezing nucleation, J. Geophys. Res.-Atmos., 118, 3731–3743, doi: 10.1002/jgrd.50365, 2013.
 - Yttri, K. E., Simpson, D., Bergström, R., Kiss, G., Szidat, S., Ceburnis, D., Eckhardt, S., Hueglin, C., Nøjgaard, J. K., Perrino, C., Pisso, I., Prevot, A. S. H., Putaud, J.-P., Spindler, G., Vana, M., Zhang, Y.-L., and Aas, W.: The EMEP Intensive Measurement Period campaign, 2008–2009: Characterizing the carbonaceous aerosol at nine rural sites in Europe, Atmos. Chem. Phys. Discuss., https://doi.org/10.5194/acp-2018-1151, in review, 2018.

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Author contributions

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J. Curtius and O. Möhler proposed the framework of this collaborative multi-institutional laboratory work. The overall manuscript, coordinated and led by N. Hiranuma, was a collaborative effort of the partners of the INUIT group and associated partners. N. Hiranuma and O. Möhler conceived the AIDA experiments, analyzed and discussed the results and contributed to the AIDA text. K. Suski and G Schill performed CFDC experiments and K. Suski, G. Schill and P. DeMott analyzed CFDC experimental data. M. Piazza performed the DFPC experiments and analysis with the support of A. Nicosia. He also contributed to the data elaboration. F. Belosi and G. Santachiara contributed to the design of the cellulose aerosol

generation systems and data elaboration. F. Belosi oversaw the DFPC ice nucleation measurements. D. Weber performed the FRIDGE experiments, and contributed to the associated data analysis with a support of H. Bingemer. T. Schiebel and K. Höhler performed and analyzed INKA experiments. S. Grawe performed the LACIS measurements and data evaluation which were coordinated and overlooked by H. Wex. K. Yamashita, T. Tajiri, A. Saito and M. Murakami performed MRI cloud simulation chamber experiments and contributed to the associated data analysis in collaboration with O. Möhler and N. Hiranuma. G. Kulkarni carried out the PNNL-CIC experiments and data analysis. E. Jantsch and T. Koop designed the BINARY experiments, which were performed by E. Jantsch. C. Budke, E. Jantsch and T. Koop analyzed and discussed the data, and E. Jantsch and T. Koop contributed the associated text. M. Polen, H. Beydoun and R. Sullivan performed the CMU-CS experiments and their analysis. Y. Tobo and K. Cory performed the NIPR-CRAFT experiments. T. Whale performed μl-NIPI experiments. T. Whale and B. Murray analysed and discussed the data and contributed the associated text. E. Stopelli conduced the measurements with LINDA with a support of F. Conen. M. Szakáll, O. Eppers and A. Mayer performed the M-AL and M-WT measurements and evaluated and discussed the experimental results, and contributed to the associated text. H. Taylor and M. Petters coordinated and carried out the NC State CS experiments. M. Taylor helped with initial data processing. M. Petters performed the final data processing and contributed the associated text. N. Reicher, Y. Rudich and B. Bhadori performed and analyzed WISDOM experiments, L. Segev designed freezing detection program and helped with the data analysis. C. Whiteside and K. Cory designed performed the WT-CRAFT experiments, and contributed to the associated data analysis and text under the guidance of N. Hiranuma. J. Schneider and H.-C. Clemen conducted single particle mass spectrometric laboratory measurements. A. Zelenyuk, K. Suski and D. Bell performed the miniSPLAT and multidimensional characterization experiments, and accompanying data analysis. For offline microscopy, K. Adachi performed electron microscopy and image analyses. R. Ullrich contributed to the database work. All authors discussed the results and contributed to the final version of the manuscript.

Competing interests

30 The authors declare no competing financial interests.

Data Records

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Within the framework of INUIT, we established a new community database including the laboratory results on ice nucleation with access for registered users. The tabulated data are available in a publically accessible MySQL portal at

https://imk-aaf-s1.imk-aaf.kit.edu:8081/inuit/. This database helps the users to evaluate and interpret the comprehensive laboratory ice nucleation results measured over the past years. It also provides a good basis for a database with a wider public access.

Usage Notes

All data associated with this study will be made available without any barriers to the user. Any disputes about the use of other groups' data, particularly with respect to publications, will be resolved by the INUIT coordinators.

Figures

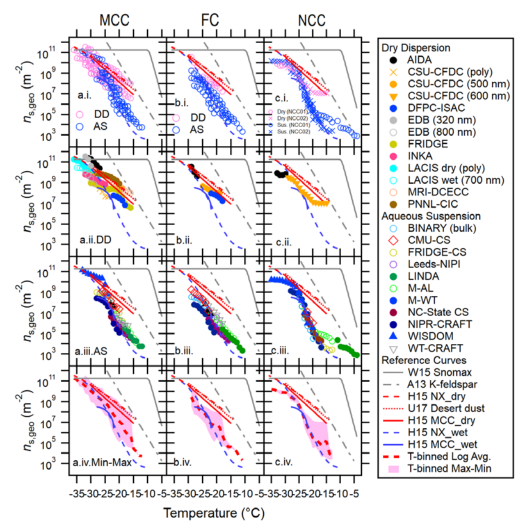


Figure 1. Immersion freezing $n_{s,geo}(T)$ spectra for MCC (a), FC (b) and NCC (c) from different techniques. Dry dispersion results (DD, pink markers) and aqueous suspension results (AS, blue markers) are shown in (i) to highlight the difference between these two subsets. Intercomparisons of DD and AS for each cellulose sample type using T-binned $n_{s,geo}$ are presented in (ii) and (iii), respectively. The log average of all results as well as the deviation between maxima and minima of $n_{s,geo}(T)$ are shown in (iv). Reference immersion freezing $n_s(T)$ spectra for MCC (H15a) illite NX (H15b), Snomax ($Wex \ et \ al.$, 2015), desert dusts (U17; $UIIrich \ et \ al.$, 2017) and K-feldspar (A13; $Atkinson \ et \ al.$, 2013) are also shown (See **Sect. 4.1**). For NCC, the results from two different batches (NCC01 from Dec 2014 and NCC02 from May 2015) are shown.

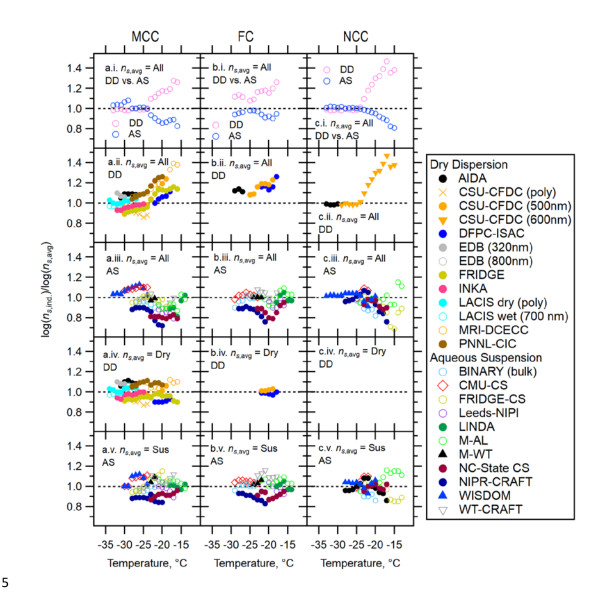


Figure 2. *T*-binned ratios of the interpolated individual measurements to the average of the data, $\log(n_{s,\text{ind.}})/\log(n_{s,\text{avg}})$, based on the geometric surface area $(n_{s,\text{geo}})$ for MCC (a), FC (b) and NCC (c). *T*-binned $\log(n_{s,\text{ind.}})/\log(n_{s,\text{avg}})$ are presented for (i) ratios of the log average to dry dispersion measurements (DD) or aqueous suspension measurements (AS) to the log average to all the data (All), (ii) ratios of the individual DD measurements to All, (iii) ratios of the individual AS measurements to All, (iv) ratios of the individual particle dispersion measurements to DD and (v) ratios of the individual aqueous suspension measurements to AS. The black dotted line represents $\log(n_{s,\text{ind.}})/\log(n_{s,\text{avg}}) = 1$. Panel c.iv is left blank since only one dataset is available at each temperature; thereby, no differences can arise.

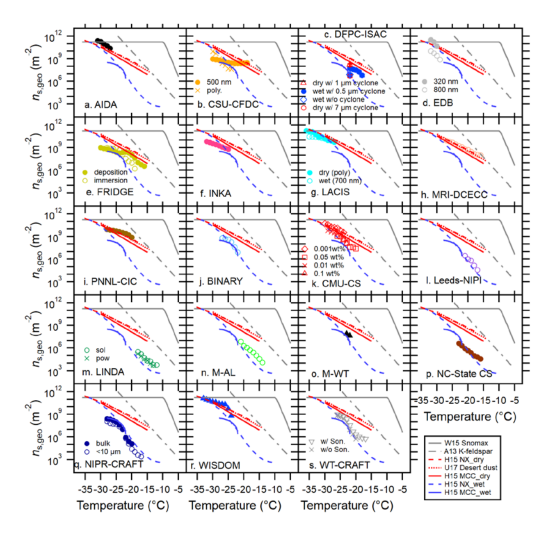


Figure 3. Inter-comparison of 20 INP measurement methods for MCC using T-binned $n_{s,geo}$. FRIDGE results of default (solid square) and imm.mode (open diamond) measurements are both presented in (e). Reference immersion freezing $n_s(T)$ spectra for MCC (H15a) illite NX (H15b), Snomax ($Wex\ et\ al.$, 2015), desert dusts (U17; $Ullrich\ et\ al.$, 2017) and K-feldspar (A13; $Atkinson\ et\ al.$, 2013), ATD and are also shown (See Sect. 3.2). Both aqueous suspension and dry dispersion results of FRIDGE are presented in panel e.

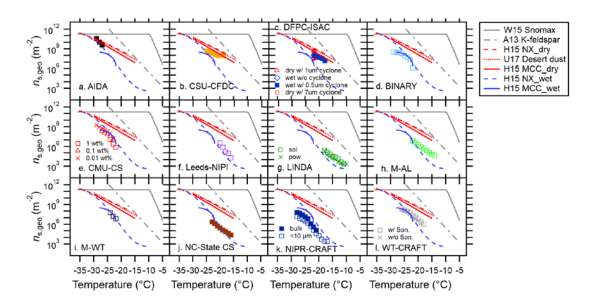


Figure 4. Inter-comparison of 12 INP measurement methods for FC using T-binned $n_{s,geo}$. Reference immersion freezing $n_s(T)$ spectra are provided as in **Fig. 3**.

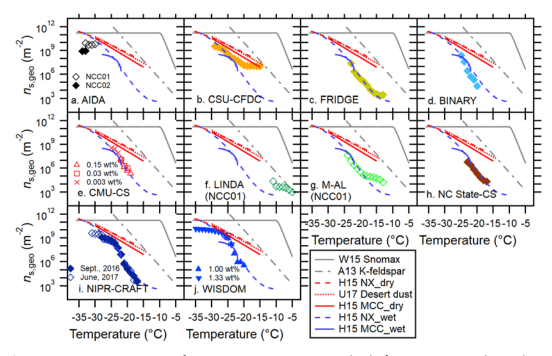


Figure 5. Inter-comparison of 11 INP measurement methods for NCC using T-binned $n_{s,geo}$. Reference immersion freezing $n_s(T)$ spectra are provided as in **Fig. 3**. Note: unless otherwise specified, the data are for NCC02.

Tables

Table 1. Properties of micro-crystalline cellulose (MCC), fibrous cellulose (FC) and nano crystalline cellulose (NCC).

System	MCC (Aldrich, 435236)	FC (Sigma, C6288)	NCC (Melodea, WS1)*
Chemical Formula	(C ₆ H ₁₀ O ₅) _n	(C ₆ H ₁₀ O ₅) _n	(C ₆ H ₉ O ₅) _n (SO ₃ Na) _x
Product Form	Powder	Powder	3wt% thixotropic gel (viscosity $^{\sim}4,665 \pm 200$ cP at 25 $^{\circ}$ C) in deionized water
¹ Density, g m ⁻³	~1.5	~1.5	~1.0-1.1
² Geometric Mode Diameter (± standard deviation) of dispersed particles, μm	1.22 ± <0.1 ^{3,4}	1.13 ± <0.1 ^{5, 4}	0.21 ± <0.1 ^{6,7}
SEM-based Mode Diameter of bulk materials (± standard deviation), µm	54.24 ± 6.2	>65	2.68 ± 0.38
Manufacturer-reported Diameter	51 μm	N/A	5-20 nm width, 100-500 nm length
Aspect Ratio	1.80-2.30 (4976/3) ⁹	~2.03 (371/1)	2.30-2.93 (764/2)
¹⁰ Geometric <i>SSA</i> , $m^2 g^{-1}$ 3.35 ± 0.1		3.35 ± 0.5	18.59 ± 2.5
¹¹ SEM-based <i>SSA</i> of residuals in 0.03wt% of 5 μL droplet, m ² g ⁻¹ 0.068		0.087	1.24
¹² BET-based <i>SSA</i> , m ² g ⁻¹	1.44 ± 0.10	1.31 ± 0.10	8.00 ± 1.00
Crystallinity	~80% (Cellulose Iβ crystallographic structure) ¹³	N/A	87% (Cellulose Iβ crystallographic structures) ¹⁴

^{*}Two NCC samples from different batches, namely non-sterile NCC (NCC01) and freshly generated NCC (NCC02), were used for the IN characterization.

5

¹Bulk density values according to manufacturers

 $^{^2}$ Based on $\Delta S/\Delta log D_{ve}$ from ADIA measurements

³Measured by a combination of SMPS and APS at AIDA (INUIT06_1, 17, 31, 42, 43, 44, 45, 46, 54)

⁴Dry particles were dispersed into the AIDA chamber using a rotating brush generator (RBG1000, PALAS).

⁵Measured by a combination of SMPS and APS at AIDA (INUIT06_6, 14)

⁶Measured by a combination of SMPS and APS at AIDA (INUIT08_6, 7, 9, 10)

Water-suspended NCC was aerosolized using the customized-atomizer (*Wex et al.*, 2015).

 $^{^8} The$ SEM-based mode diameter of atomized NCC is 0.28 \pm <0.1 μm , which is similar to that of bulk NCC.

⁹Average aspect ratio per substrate: the numbers in bracket represent a total number of particles/substrate(s) analyzed under SEM for each subset.

 $^{^{10}}$ Geometric SSA is derived from ADIA measurements (i.e., fraction of total surface area concentration to total mass concentration estimated from a combination of SMPS and APS; See **Fig. S1**). The particles in AIDA were all <10 μ m in diameter.

 $^{^{11}}$ Measured using droplet residuals derived from 5 μ L of 0.03wt% suspension. Uncertainty is not given because all individual particle counts were compiled to calculate the SSA value of each sample.

^{20 &}lt;sup>12</sup>Brunauer et al., 1938

¹³Nishiyama et al., 2002

¹⁴Aulin et al., 2009

 Table 2.
 Summary of INUIT measurement techniques and instruments. Their acronyms are available in the Supplementary Information. ID 1-9 and ID 10-20 represent dry dispersion measurements and suspension techniques, respectively (Alphabetical order).

_										
9	1	,	0.00	90	Ce	Cellulose type	уре	Investigable 7-	Investigated T-range for	CCA (202 2-1)**
⊇		Descripción	Moder	verei eiice	MCC	5	NCC	range	this study	8-III) HCC
Н	*AIDA	CECC	ON	Möhler et al., 2003; Niemand et al., 2012	×	×	×	-100°C< <i>T</i> <-5°C	MCC: -31 °C < T < -27 °C FC: -29 °C < T < -27 °C	MCC (poly): 3.35 FC (poly): 3.35
									NCC: -33 °C < T < -30 °C	NCC (poly): 18.59
2	CSU-CFDC	Cylindrical-walled CFDC	Yes	DeMott et al., 2015	×	×	×	-34 °C < <i>T</i> < -9 °C	MCC: -30 °C < 7 < -23 °C	MCC (poly): 2.09
									FC: -25 °C < <i>T</i> < -19 °C	MCC/FC (500 nm): 8.00
									NCC: -29 °C < 7 < -25 °C	NCC (600 nm): 6.67
æ	DFPC-ISAC	Substrate-supported diffusion	o N	Santachiara et al., 2010; Belosi et al., 2014	×	×		-22 °C < T < -10 °C	MCC: -22 °C < 7 < -18 °C	MCC (poly): 0.71- 4.59
									FC: -22 °C < T < -18 °C	FC (poly): 0.81-4.95
4	*EDB	Electrodynamic balance levitator	o N	Hoffmann et al., 2013a; 2013b	×			-40 °C < 7 < -1 °C	MCC: -32 °C < 7 < -29 °C	MCC (320 nm): 7.4 MCC (800 nm): 1.3
2	*,¹FRIDGE_default	Substrate-supported diffusion cell	Yes	Schrod et al., 2016	×			-30 °C < 7 < 0 °C	MCC: -30 °C < 7 < -16 °C	MCC (poly): 1.82
9	*INKA	Cylindrical plates CFDC	No	Schiebel, 2017	×			-60 °C < T < -10 °C	MCC: -32 °C < T < -25 °C	MCC (poly): 3.35
7а	*, ² LACIS_dry	Laminar flow tube	o N	Hartmann et al., 2011; Wex et al., 2014	×			-40 °C < 7 < -5 °C	MCC: -36 °C < T < -27 °C	MCC (poly): 7.00
7b	*,³LACIS_wet	Laminar flow tube	No	Grawe et al., 2016	×			-40 °C < <i>T</i> < -5 °C	MCC: -35 °C < 7 < -30 °C	MCC (700 nm): 5.70
∞	MRI-DCECC	Dynamic CECC	o N	<i>Tajiri et al.</i> , 2013; Hiranuma et al., 2015a	×			-100 °C < 7 < ~0 °C	MCC: -28 °C < 7 < -16 °C	MCC (poly): 1.36
6	PNNL-CIC	Parallel plates CFDC	Yes	Friedman et al., 2011	×			-55 °C < <i>T</i> < -15 °C	MCC: -28 °C < 7 < -20 °C	MCC (600 nm): 6.67
10	*BINARY	Cold stage-supported droplet	No	Budke and Koop., 2015	×	×	×	-30 °C < T < ~0 °C	MCC: -27 °C < 7 < -22 °C	MCC (bulk): 0.068
		assay							FC: -29 °C < <i>T</i> < -22 °C NCC: -25 °C < <i>T</i> < -20 °C	FC (bulk): 0.087 NCC (bulk): 1.24
11	_S CMU-CS	Cold stage-supported droplet assay	N _O	Polen et al., 2016; Beydoun et al., 2017	×	×	×	-30 °C < T < ~0 °C	MCC: -30 °C < <i>T</i> < -20 °C FC: -29 °C < <i>T</i> < -22 °C	MCC (bulk): 0.068 FC (bulk): 0.087
									NCC: -25 °C < T < -19 °C	NCC (bulk): 1.24

9	1		Colinion		Celli	Cellulose type	be	Investigable <i>T</i> -	Investigated T-range for	() v J J
2	mstrument	Describation	Mobiles	Reference	MCC	FC.	NCC	range	this study	(B -III) PCC
12	*FRIDGE-CS	Cold stage-supported droplet assay	Yes	Hiranuma et al., 2015b	×		×	-29 °C < T < 0 °C	MCC: -28 °C < <i>T</i> < -19 °C NCC: -23 °C < <i>T</i> < -13 °C ⁶	MCC (poly): 1.717 NCC (bulk): 1.24
13	Leeds-µL-NIPI	Nucleation by immersed particles instrument	Yes	Whale et al., 2015	×	×		-36 °C < <i>T</i> < ~0 °C	MCC: -21 °C < <i>T</i> < -17 °C FC: -20 °C < <i>T</i> < -16 °C	MCC (bulk): 0.068 FC (bulk): 0.087
14	⁸ LINDA	Immersion mode ice spectroemeter	Yes	Stopelli et al., 2014	×	×	*	-18 °C < 7 < ~0 °C	MCC: -18 °C < 7 < -12 °C FC: -18 °C < 7 < -11 °C NCC: -11 °C < 7 < -4 °C	MCC (bulk): 0.068 FC (bulk): 0.087 NCC (bulk): 1.24
15	*M-AL	Acoustic droplet levitator	ON.	Diehl et al., 2014	*	×	*	-30 °C < 7 < ~0 °C	MCC: -21 °C < 7 < -14 °C FC: -22 °C < 7 < -14 °C NCC: -24 °C < 7 < -13 °C	MCC (bulk): 0.068 FC (bulk): 0.087 NCC (bulk): 1.24
16	*M-WT	Vertical wind tunnel	o N	Szakáll et al., 2010; Diehl et al., 2011	*	×		-30 °C < 7 < ~0 °C	MCC: -23 °C < T < -12 °C FC: -24 °C < T < -22 °C	MCC (bulk): 0.068 FC (bulk): 0.087
17	NC State-CS	Cold stage-supported droplet assay	0	Wright and Petters, 2013	×	×	*	-40 °C < 7 < ~0 °C	MCC: -24 °C < 7 < -16 °C FC: -24 °C < 7 < -16 °C NCC: -24 °C < 7 < -17 °C	MCC (bulk): 0.068 FC (bulk): 0.087 NCC (bulk): 1.24
18a	NIPR-CRAFT	Cold stage-supported droplet assay	ON.	<i>Tobo,</i> 2016	*	×	*	-34 °C < <i>T</i> < ~0 °C	MCC: -28 °C < 7 < -20 °C FC: -28 °C < 7 < -21 °C NCC: -31 °C < 7 < -17 °C	MCC (bulk): 0.068 FC (bulk): 0.087 NCC (bulk): 1.24
18b	°NIPR-CRAFT_<10 µm	Cold stage-supported droplet assay	0	<i>Tobo</i> , 2016	×	×		-34 °C < 7 < ~0 °C	MCC: -28 °C < 7 < -17 °C FC: -28 °C < 7 < -17 °C	MCC (<10 µm): 3.35 ¹⁰ FC(<10 µm): 3.35 ¹⁰
19	WISDOM	Microfluidic device-supported droplet assay	0 N	Reicher et al., 2018	*		*	-40 °C < 7 < ~0 °C	MCC: -33 °C < T < -24 °C NCC: -35 °C < T < -20 °C	MCC (bulk): 0.068 NCC (bulk): 1.24
20	WT-CRAFT	Cold stage-supported droplet assay	O _N	<i>Tobo,</i> 2016	×	×		-34 °C < T < ~0 °C	MCC: -26 °C < T < -17 °C FC: -26 °C < T < -18 °C	MCC (bulk): 0.068 FC (bulk): 0.087

*Instruments of INUIT project partners, **Specific surface area; poly = polydispersed particles, homo = homogenised particles, bulk = bulk material, 1. Default deposition nucleation mode operation, 2. Experiments with a dispersed-aerosol injection, 3. Experiments with atomized-aerosol injection, 4. Homogenized-sample data, 5. 0.001-1 wt% aqueous suspensions employed, 6. Experiments with 1.2 wt% non-diluted suspension, 7. TSI-OPS basis, 8. Experiments with both dissolved mass in solution and dry powder mass, 9. Experiments with size-selected (<10 µm) particles, 10. The AIDA-derived geometric SSA value (3.35 m² g¹) was used since it accounts for only <10 µm particles.

Table 3. List of the Gumbel cumulative distribution fit parameters to the $n_{s,geo}(T)$ for T-binned ensemble datasets of MCC, FC and NCC (All). The datasets are fitted in the log space. Besides All, fit parameters for ensemble maximum values (All_{max}), ensemble minimum values (All_{min}), suspension subset (AS), and dry dispersed particle subset (DD) are also included in this table. The correlation coefficient, r, for each fit is also shown. All $n_{s,geo}(T)$ values are in m^{-2} . T is in °C.

Fitted dataset	Fitted <i>T</i> range	$[n_{s,ge}]$	Fit F $_{10}(T) = \exp(a$	arameters ·exp(-exp(+ <i>d</i>)]	
		а	b (°C ⁻¹)	c (°C)	d	r	$\Delta \log(n_{s,geo})/\Delta T$
All (MCC)	-36 °C < T < -12 °C	24.47	0.12	15.99	3.24	0.96	0.32
All _{max} (MCC)	-36 °C < <i>T</i> < -12 °C	23.19	0.19	14.36	3.28	0.83	0.33
All _{min} (MCC)	-36 °C < <i>T</i> < -12 °C	27.95	0.08	18.67	3.03	0.95	0.30
DD (MCC)	-36 °C < <i>T</i> < -16 °C	24.12	0.08	12.56	4.69	0.91	0.20
AS (MCC)	-33 °C < <i>T</i> < -12 °C	28.03	0.10	18.22	3.48	0.97	0.37
All (FC)	-29 °C < T < -11 °C	22.25	0.11	15.95	3.62	0.88	0.33
All _{max} (FC)	-29 °C < <i>T</i> < -11 °C	23.78	0.13	16.85	4.79	0.94	0.40
All _{min} (FC)	-29 °C < <i>T</i> < -11 °C	21.88	0.08	16.85	3.15	0.58	0.26
DD (FC)	-29 °C < <i>T</i> < -18 °C	26.97	0.07	18.12	6.85	0.89	0.28
AS (FC)	-29 °C < <i>T</i> < -11 °C	22.57	0.09	16.05	3.46	0.92	0.29
All (NCC)	-35 °C < <i>T</i> < -13 °C	19.30	0.14	19.48	6.59	0.90	0.31
All _{max} (NCC)	-35 °C < T < -13 °C	17.22	0.18	17.36	7.30	0.93	0.29
All _{min} (NCC)	-35 °C < <i>T</i> < -13 °C	17.39	0.21	19.88	6.30	0.89	0.32
DD (NCC)	-33 °C < <i>T</i> < -15 °C	16.40	0.18	17.33	7.45	0.97	0.29
AS (NCC)	-35 °C < <i>T</i> < -13 °C	15.35	0.28	20.83	8.53	0.98	0.30

reference immersion freezing $n_{s,geo}(T)$ values for MCC from H15a. The numbers in parentheses are maxima and minima of $n_{s,geo}$ at T. The numbers in brackets represent the number of instruments used to calculate the log average $n_{s,geo}(T)$ values derived from a single instrument are not included in this table. Table 4. T-binned ns,geo values (in m-2) of three different cellulose samples based on the log average of all available results at T [i.e., Fig. 1 (iv)]. The first MCC column represents

(5)													
	MCC (H15a)		MCC (Max-Min)	Min)			FC (Max-Min)	Min)			NCC (Max-Min)	Min)	
-34		5.94E+10	(1.26E+11 -	2.80E+10)	[2]								
-33		5.85E+10	(1.35E+11 -	2.62E+10)	[3]					1.09E+10	(1.37E+10 -	8.75E+09)	[2]
-32		3.33E+10	(3.41E+11 -	6.42E+09)	[9]					8.28E+09	(1.29E+10 -	5.30E+09)	[2]
-31		2.88E+10	(1.74E+11 -	4.77E+09)	[7]					8.10E+09	(1.20E+10 -	5.46E+09)	[2]
-30		1.07E+10	(8.20E+10 -	9.31E+08)	[11]					8.19E+09	(1.01E+10 -	6.64E+09)	[2]
-29		5.89E+09	(4.20E+10-	7.61E+08)	[10]	2.82E+09	(3.61E+10 -	3.55E+08)	[3]	5.36E+09	(8.02E+09 -	3.58E+09)	[2]
-28	1.18E+10	3.25E+09	(2.74E+10 -	2.51E+08)	[12]	6.55E+08	(1.05E+10-	6.47E+07)	[4]	2.79E+09	(6.40E+09 -	1.29E+09)	[3]
-27	8.55E+09	2.06E+09	(2.19E+10 -	2.21E+08)	[13]	4.22E+08	(4.08E+09 -	4.89E+07)	[4]	2.08E+09	(5.13E+09 -	1.02E+09)	[3]
-26	6.81E+09	1.23E+09	(2.03E+10 -	1.35E+08)	[12]	1.49E+08	(3.65E+08 -	2.84E+07)	[4]	1.27E+09	(2.90E+09 -	7.73E+08)	[3]
-25	5.03E+09	7.97E+08	(6.06E+09 -	4.55E+07)	[12]	1.13E+08	(4.68E+08 -	1.43E+07)	[2]	4.52E+08	(1.05E+09 -	2.62E+08)	[2]
-24	2.97E+09	4.46E+08	(4.34E+09 -	3.93E+07)	[11]	6.50E+07	(3.30E+08 -	7.12E+06)	[9]	1.12E+08	(2.67E+08 -	3.87E+07)	[9]
-23	1.87E+09	1.32E+08	(3.39E+09 -	4.15E+06)	[10]	1.51E+07	(2.19E+08-	1.19E+06)	[7]	1.89E+07	(9.37E+07 -	3.01E+06)	8
-22	1.23E+09	5.33E+07	(2.48E+09 -	8.72E+05)	[11]	7.43E+06	(1.42E+08-	3.79E+05)	[8]	3.66E+06	(5.52E+07 -	5.46E+05)	8
-21	6.30E+08	2.32E+07	(1.61E+09 -	2.54E+05)	[11]	5.45E+06	(1.05E+08-	1.29E+05)	[9]	1.07E+06	(2.83E+07 -	2.09E+05)	[_
-20	5.14E+08	1.21E+07	(7.99E+08 -	1.23E+05)	[11]	6.58E+06	(1.03E+08-	2.94E+05)	[9]	3.21E+05	(1.41E+07 -	3.21E+04)	[2]
-19	2.94E+08	6.63E+06	(2.95E+08 -	2.42E+05)	[6]	3.11E+06	(1.03E+08 -	1.34E+05)	[9]	2.31E+05	(1.20E+07 -	4.27E+04)	[2]
-18	1.60E+08	1.42E+06	(1.60E+08 -	9.33E+04)	[8]	5.46E+05	(1.70E+07 -	7.43E+04)	[9]	1.21E+05	(1.16E+07 -	1.77E+04)	[2]
-17	1.15E+08	6.31E+05	(1.15E+08 -	3.83E+04)	[7]	1.08E+05	(2.48E+05 -	4.27E+04)	[4]	6.42E+04	(1.13E+07 -	4.44E+03)	[2]
-16	9.69E+07	5.89E+05	(9.69E+07 -	3.76E+04)	[2]	4.36E+04	(1.12E+05 -	1.86E+04)	[4]	1.59E+05	(1.12E+07 -	5.05E+03)	[3]
-15		2.79E+04	(3.64E+04 -	2.14E+04)	[5]	4.40E+04	(5.85E+04 -	3.31E+04)	[2]	1.25E+05	(1.11E+07 -	3.30E+03)	[3]
-14		1.48E+04	(1.82E+04 -	1.21E+04)	[5]	2.88E+04	(3.79E+04 -	2.19E+04)	[2]	9.01E+03	(3.57E+04 -	2.27E+03)	[2]
-13										6.67E+03	(1.68E+04 -	2.65E+03)	[2]

Table 5. Summary of the geometric *SSA* of MCC and FC particles assessed by DFPC-ISAC. In general, high *SSA* values indicate the presence of small grains because the relative dominance of the mass to the surface becomes small.

Exp_ID	Avg. SSA (m ² g ⁻¹)	Stdev. SSA (m ² g ⁻¹)
MCC_Dry_7um_cut-size	0.8	0.09
MCC_Wet_no_cyclone	3.12	0.1
MCC_Wet_0.5um_ cut-size	3.48	0.13
MCC_Dry_1um_ cut-size	4.37	0.24
FC_Dry_7um_ cut-size	0.9	0.1
FC_Wet_no_cyclone	3.11	0.11
FC_Wet_0.5um_ cut-size	3.57	N/A
FC_Dry_1um_ cut-size	4.91	0.35