The Fifth International Workshop on Ice Nucleation Phase 3 (FIN-03): Field Intercomparison of Ice Nucleation Measurements

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The third phase of the Fifth International Ice Nucleation Workshop (FIN-03) was conducted at Storm Peak Laboratory in Steamboat Springs, Colorado in September 2015 to facilitate the intercomparison of instruments measuring ice nucleating particles (INPs) in the field. Instruments included a subset of two online and four offline measurement systems for INPs, a subset of those utilized in the laboratory study that comprised the second phase of FIN (FIN-02). Composition of total aerosols were characterized by the Particle Ablation by Laser Mass Spectrometry (PALMS) and Wideband Integrated Bioaerosol Sensor (WIBS) instruments, and aerosol size distributions were measured by a Laser Aerosol Spectrometer (LAS). The dominant total particle compositions present during FIN-03 were composed of sulfates, organic compounds, and nitrates, as well as particles derived from biomass burning. Mineral dust containing particle types were ubiquitous throughout and represented 67% of supermicron particles. Total WIBS fluorescing particle concentrations for particles with diameter > 0.5 µm were 0.04±0.02 cm\(^{-3}\) (0.1 cm\(^{-3}\) highest, 0.02 cm\(^{-3}\) lowest), typical for the warm season in this region and representing ~9% of all particles in this size range as a campaign average.

The primary focus of FIN-03 was the measurement of INP concentration via immersion freezing at temperatures > –33 °C. Additionally, some measurements were made in the deposition nucleation regime at these same temperatures, representing one of the first efforts to include both mechanisms within a field campaign. INP concentrations via immersion freezing reported by all ice nucleation instruments generally agreed to within one order of magnitude for measurement and sampling times coordinated to within three hours. Sometimes, much better agreement was obtained. Outliers of up to two orders of magnitude occurred between –25 °C and –18 °C; better agreement was seen at higher and lower temperatures. INP activity in the
immersion freezing mode was generally found to be an order of magnitude or more efficient than
in the deposition regime at 95-99% water relative humidity, although this limited data set should
be augmented in future efforts.

To contextualize the study results an assessment was made of the composition of INPs
during the late Summer to early Fall period of this study, inferred through comparison to existing
ice nucleation parameterizations and through measurement of the influence of thermal and
organic carbon digestion treatments on immersion freezing ice nucleation activity. Consistent
with other studies in continental regions, biological INPs dominated at temperatures > –20 °C
and sometimes colder, while arable dust-like or other organic-influenced INPs were inferred to
dominate at most times below –20 °C.
1 Introduction

Atmospheric ice nucleation is one of the least certain aerosol-cloud interactions influencing climate (Kanji et al., 2017). Aerosols that physically catalyze freezing, known as ice-nucleating particles (INPs) (Vali et al., 2015), are found in the atmosphere in concentrations that span many orders of magnitude, ranging from $10^{-3}$ L$^{-1}$ or fewer at $-5$ °C to 1000 L$^{-1}$ or greater at $-35$ °C (Petters and Wright, 2015). INP number concentrations typically increase exponentially with degree of supercooling below 0 °C. However, chemical composition plays an important role in determining if, and at what temperature, individual particles may serve as INPs (Murray et al., 2012). INPs initiate the formation of ice in cold and mixed-phase clouds and in turn influence their physical and optical properties. An increase in INP concentration over a geographic area may increase the frequency of glaciated clouds at constant temperature, which in turn increases precipitation and decreases cloud lifetime (Lohmann and Feichter, 2005). Nevertheless, INP impacts on clouds simulated in global climate models are highly sensitive to how aerosol's ability to nucleate ice is parameterized (Boucher et al., 2013). Parameterizations can only be as accurate as the measurements on which they are based (e.g., Knopf et al., 2021).

Measurements of atmospheric INPs remain challenging due to the difficulty representing the physical processes involved in ice nucleation instruments. At temperatures below $\sim -38$ °C, micrometer-sized, dilute water droplets spontaneously freeze due to homogeneous freezing nucleation. Homogeneous freezing nucleation is well understood and included in most cloud formation models. However, at temperatures between 0 and $-38$ °C, freezing requires INPs to facilitate nucleation through a heterogeneous nucleation mechanism (Kanji et al., 2017; Murray et al., 2012; Vali, 1985). Nucleation is hypothesized to proceed through (1) immersion freezing, which occurs when an INP embedded within a water droplet enters a cooler environment, and nucleates an ice crystal, (2) condensation freezing, which occurs when freezing ensues as an
aqueous droplet condenses on the surface of an aerosol particle, (3) contact freezing, which occurs when an aerosol in contact with a water droplet surface initiates freezing (Durant and Shaw, 2005; Fornea et al., 2009), and (4) deposition nucleation, which is thought to occur through the direct deposition of water vapor on an INP surface. Of these mechanisms, immersion freezing nucleation is thought to be the most active heterogeneous nucleation process in the atmosphere, though there is considerable disagreement in the literature about the relative importance of other mechanisms (Kanji et al., 2017; Ullrich et al., 2017). When the ambient humidity is below water saturation, nucleation can occur via deposition of water from the vapor phase. In some cases, this behavior may be ascribable instead to water condensation in pores and cavities in aerosols facilitating freezing through a non-deposition mechanism (Marcolli, 2014; Wagner et al., 2016). However, this process is unlikely to be of importance at temperatures > -38 °C (David et al., 2020), which are the focus of this study. We will thus refer to ice nucleation at > –38 °C and below water saturation as happening within the “deposition regime”. Study of the efficiency of the deposition nucleation process in comparison to immersion freezing has been limited for natural INPs.

Ice nucleation measurements have been made with instruments designed and built by individual scientists, and more recently with commercial instruments. The ice nucleation community has a history of collaborating to address instrument performance and inconsistencies through participating in instrument intercomparisons, in which the custom-built instruments were operated side-by-side, to evaluate instrument response to the same aerosol populations (Coluzza et al., 2017; DeMott et al., 2011, DeMott et. al, 2018; Knopf et al, 2021; Lacher et al., 2024). To compare concentrations and compositions of INPs, a three-part workshop series, the Fifth International Ice Nucleation Workshop, or “FIN” was held in 2014-2015. The first two phases
were held at the Karlsruhe Institute of Technology’s Aerosol Interactions and Dynamics in the Atmosphere (AIDA) facility. FIN-01 focused on determination of composition of INPs by mass spectroscopy (Shen et al., 2024, in preparation), while FIN-02 entailed a laboratory ice nucleation instrument comparison (DeMott et al., 2018). FIN-03, the mountaintop field intercomparison of ice nucleation instruments is the focus of this manuscript.

Ice nucleation measurements have experienced a renaissance in the past decade, resulting in a proliferation in both the number of custom-built instruments and a diversification of measurement techniques employed (Zenker, 2017; DeMott, 2018; Möhler, 2021). Participation in FIN-02 was twice that of the previous formal international workshop intercomparison in 2007 (the International Workshop on Comparing Ice Nucleation Measuring Systems, or ICS-2007 held at the (AIDA) facility (Jones et al., 2011; Kanji et al., 2011). During FIN-02, online and offline instruments sampling the same population of aerosolized particles reported INP concentrations that generally agreed within one order of magnitude across a broad temperature range. Agreement was best in tests of immersion freezing on soils, dusts and bacteria but spanned up to 2 orders of magnitude (or 3 °C in temperature for the same active site density) for illite NX and K-feldspar (DeMott et al., 2018). While relatively good agreement in the laboratory between different measurement methods during FIN-02 represents significant progress for the atmospheric ice nucleation community, intercomparisons in ambient atmospheric settings are more difficult due to lower typical INP concentrations (Lacher et al., 2018) and variations in the chemistry and size of source aerosol and INPs (DeMott et al., 2017; Knopf, 2021; Lacher et al., 2024).

To evaluate how a suite of instruments operating collectively perform under the greater measurement challenges of the field setting, FIN-03 was conducted from September 12 to 28, 2015 at Storm Peak Laboratory (SPL) in Steamboat Springs, CO, USA (Elevation: 3220 m MSL).
Unlike the subsequent closure studies of Knopf et al. (2021) and the similar comparative sampling studies of Lacher et al. (2024), both of which occurred in regions surrounded by agricultural activities and possible nearby urban influences, this remote continental mountaintop site at an elevation of 3220 m provided the opportunity to sample both regional and long-range INP sources within both the boundary layer and free troposphere. The site is typically in the free troposphere during the nighttime and early morning, and in the boundary layer from the late morning to early evening, although topography and wind direction influence this timing (Collaud Coen et al., 2018).

When in the free troposphere, the site is more likely to reflect influences by regional or long-range transport of aerosols. For example, during FIN-03, the variety of air masses that were sampled and sensed by aerosol instruments included ones passing over phosphate mines in Idaho (on September 18 and 20) and mined deposits of rare earth metals at Mountain Pass, CA (on September 27) (Zawadowsicz et al., 2017). When the convective boundary layer height reaches the elevation of the laboratory, the site is likely more impacted by local/regional aerosol sources. Additionally, meteorological transitions can occur (e.g., frontal boundary passage, wind direction shifts), driving changes in aerosol sources that may indirectly occur in response to those changes (e.g., biological aerosols, carbonaceous particles from biomass burning, and mineral/soil dust). While the constantly fluctuating environmental conditions during FIN-03 added an additional challenge to the intercomparison, they also provided a realistic setting for atmospheric INP measurements. In addition to adding challenges, conducting the intercomparison in the presence of complex aerosols in the field provided the opportunity to survey instrument response to varied aerosol sources.

Participation in FIN-03 included online continuous flow diffusion chambers (CFDCs) and aerosol collections for offline INP measurements, representing a subset of the instruments that operated in FIN-02 (DeMott et al., 2018). Since aerosol physical and chemical properties strongly
influence their ability to activate as INPs (Hoose and Möhler, 2012; Kanji et al., 2017; Murray et al., 2012), measurements of aerosol sizes and composition (see Section 2) were included to lend context to the variable composition of aerosols and evaluate their potential role in ice nucleation activity. Rather than use these data for attempting closure, FIN-03 focused on using data to constrain existing parameterizations to diagnose INP compositions during the study period. Also, in contrast to other recent studies, special effort was made to characterize deposition nucleation activity in addition to immersion freezing.

2 Methods

2.1 Aerosol property measurements

Measurements of aerosol physical, chemical, and biological particle properties were made during FIN-03 to provide context to INP measurements. Sampling manifolds, which draw air into SPL from outdoors at high flow, are as follows: Inlets were located in each of the two wings of SPL that frame the living area, referred to as the “instrument” laboratory (facing north) and the “chemistry” laboratory (facing south). The “original” inlet system in the instrument laboratory (Haller et al. 2011; Petersen et al. 2019) feeds a nephelometer (see below) and a standard suite of aerosol instruments (not operational for FIN-03). This 15 cm diameter aluminum inlet rises 4 m above the roofline. At ~1 m inside the laboratory, it transitions to a 15 cm horizontal manifold. With a flow of ~500 L min⁻¹, aerosol transmission calculations have characterized the system to have a 50% upper particle size cut-off at an aerodynamic diameter of 5 µm (Haller et al., 2011). The “new” inlet system consists of two identical stainless steel, turbulent-flow, ground-based inlets described by Petersen et al. (2019), which are straight and enter the laboratory vertically. One is in the SPL instrument laboratory, and one is in the chemistry laboratory. These inlets that extend 10 m above the laboratory roof have been
demonstrated to have 50% upper particle size cut-offs at an aerodynamic diameter of approximately 13\(\mu\)m for a wind speed of 0.5 m s\(^{-1}\). Additional computational fluid dynamics simulations suggest that this size cut off remains above 5 \(\mu\)m even for exterior wind speeds up to 15 m s\(^{-1}\) (Petersen et al., 2019), higher than achieved at any time during FIN-03 sampling. Little bias was seen in ambient aerosol sampling between the original inlet system and the new, turbulent flow-based inlets based on the metric of total aerosol scattering (Petersen et al., 2019). Flow rates and transfer lines to individual instruments are described after the aerosol property measurements are introduced, at the conclusion of this section.

A Laser Aerosol Spectrometer (LAS, model 3340, TSI Inc., St. Paul, Minnesota, USA) was used to measure the aerosol size distribution over the diameter range 0.089-10 \(\mu\)m. Aerosols were assumed dry based on relative humidity always remaining below 30% when measured from its sample line. Sample was drawn at 0.1 L min\(^{-1}\) and sampling was done from the turbulent flow inlet system located in the SPL chemistry laboratory, as described further below. Size calibrations were performed using polystyrene latex spheres (PSL, Duke Scientific). PSL diameters were converted to ammonium sulfate equivalent diameters using Mie theory (Froyd et al., 2019). Particle concentrations are reported as a function of equivalent ammonium sulfate diameter. Volume and surface area distributions are derived assuming spherical particles. Number concentrations and surface areas, further informed by aerosol composition measurements, allows for connection to INP concentration predictions, and this information is used herein to diagnostically infer mineral and soil dust influences on INPs during the study. We will particularly reference the parameterizations of Niemand et al. (2012) that links mineral surface area to INP concentrations and DeMott et al. (2015) that links dust number concentrations at sizes larger than 0.5 \(\mu\)m to INP concentrations.
Measurements using a three-wavelength integrating nephelometer (TSI Model-3563, Shoreview, MN) also provided information on aerosol distributions via their optical properties. This nephelometer is part of the National Oceanic and Atmospheric Administration Federated Aerosol Network (Andrews et al., 2019). The nephelometer splits scattered light into red (700 nm), green (550 nm), and blue (450 nm) wavelengths. Impactors to cut aerosols at aerodynamic sizes below 1 and 10 μm are alternately used upstream of air flowing into the instrument. The nephelometer sampled within the original inlet in the SPL instrument laboratory. A blunt tap from this original SPL inlet manifold provided air samples to the nephelometer system via 1” i.d. conductive tubing.

The Particle Analysis by Laser Mass Spectrometry (PALMS) instrument performed measurements of the composition of 0.2 to 3.0 μm aerosol particles. The PALMS was designed and operated by the National Oceanic and Atmospheric Administration (NOAA) as described in Thomson et al. (2000). Particles are sampled, focused, and accelerated via an aerodynamic lens inlet (Schreiner et al., 2002) before passing into a vacuum chamber where they successively pass through two continuous-wave detection laser beams (532 nm Nd:YAG) and scatter light. Vacuum aerodynamic diameter is determined based on the transit time. The detection signal triggers an ArF excimer laser that emits a 193 nm pulse to simultaneously ablate and ionize single particles. The resulting ions are analyzed with a unipolar time-of-flight mass spectrometer, which allows polarity switching during the particle flight and thereby producing positive or negative mass spectra for individual particles. PALMS spectra are classified into compositional categories, and fractions are averaged over 5 min sample periods. Number, surface area, and mass concentration products for the different particle types are generated by combining PALMS size-dependent fractional composition data with absolute particle concentrations measured by the
LAS instrument (Froyd, et al. 2019; Froyd et al., 2022). When PALMS compositional concentrations are referenced in the results of FIN-03 aerosol compositions in Section 3.2, they have been determined by these methods.

The NOAA Wideband Integrated Bioaerosol Sensor, Model 4A (WIBS-4A; Droplet Measurement Technologies, Longmont, CO) was used to detect fluorescent properties of individual particles and assess the presence of biological particles. Measurements are presumed to characterize dry particles. The WIBS-4A is described in detail elsewhere (Gabey et al., 2010; Kaye et al., 2005; Perring et al., 2015) and is only briefly summarized here. As described in Zawadowicz et al. (2019), the gain for the WIBS-4A used at SPL was set to detect and classify particles between 0.4 and 10 μm. First, the optical diameter of particles entering the detection cavity is determined by light scattered during transit through a 635 nm laser beam. This signal triggers the sequential firing of two xenon flash lamps filtered to produce narrow excitation wavebands centered at 280 and 370 nm. The resulting fluorescence is detected by two wideband photomultiplier detectors observing 310-400 nm and 420-650 nm. Fluorescing particles were categorized according to the intensity of the signal in each of three channels (channel A excitation 280 nm/emission 310-400 nm, channel B excitation 280 nm/emission 420-650 nm, channel C excitation 370 nm/emission 420-650 nm). Particles for which the measured emission intensity in only one channel met the threshold (such that the signal intensity exceeded the value equal to three standard deviations above the mean) were assigned Type A, B, or C, and particles for which the measured emission intensity in two or more channels met the threshold were assigned Type AB, BC, BC, or ABC (Perring et al., 2015). The interpretation of particle composition according to the seven WIBS-4A channels is not straightforward, as many fluorophores are active in each channel, including non-biological components (Perring et al.,
Channel A fluorophores include biological components such as tryptophan, phenylalanine as well as nonbiological components which interfere with the determination of biological content, including polycyclic aromatic hydrocarbons (PAHs) (pyrene, naphthalene, phenanthrene). Biological fluorophores, which produce a signal in channel C, include the reduced form of nicotinamide adenine dinucleotide (NADH), nicotinamide adenine dinucleotide phosphate (NADPH), and riboflavin, and potential non-biological interference in channel C may result from the presence of humic acid in aerosol particles.

Channel B fluorophores are not generally considered to be biological in nature, though riboflavin and dry cellulose both produce signals in this channel.

We report WIBS-4A channel data herein under these noted caveats and further utilize these data to explore links to immersion freezing biological INP concentrations, as has been done in some previous efforts. Tobo (2013) previously reported relations of biological INPs acting in the immersion freezing mode (measured by the CSU CFDC) to fluorescent biological aerosol particles (FBAP) at sizes > 0.5 μm measured in the understory of a Ponderosa pine forest in Colorado. In that work, an ultraviolet aerodynamic particle sizer (UV-APS) with excitation wavelength at 355 nm and emission wavelengths 420-575 nm was used as a reference for FBAP concentrations. Due to differences between the excitation and emission wavelengths, UV-APS measurements correspond most closely with Type C particles detected by the WIBS-4A (Healy et al., 2014). Consequently, a conservative or “low” estimate of FBAP for use in the parameterization of Tobo et al. (2013) we employ herein uses the sum of C, AC, BC and ABC particles. A “high” FBAP for this parameterization has also been used by Twohy et al. (2016), considering all non-B-only particles (A, AB, ABC, AC, BC, C). We will use both definitions in our presented results and partly justify the higher estimate because the CSU CFDC assuredly
does not capture all biological INPs due to the use of the upstream impactor. A final class of particles defined by WIBS-4A data for relation to immersion freezing INPs are denoted as FP3 particles (Wright et al., 2014). FP3 particles are particles that show strong emission in the 310 to 400 nm spectral band when excited by 280 nm light (A type) but are only weakly represented as B and C types. A threshold of 1900 arbitrary fluorescence units in the 310 to 400 nm band is used to denote FP3 particles (Wright et al., 2014). FP3 particles have been connected to immersion freezing INP concentrations in multiple environments (Wright et al., 2014; Suski et al., 2018; Cornwell et al., 2023).

Flow rates and transfer lines to each instrument are summarized as follows. The PALMS, LAS, and WIBS-4A sampled from the SPL turbulent flow inlet stack at 0.75, 0.1, and 0.3 vlpm, respectively, via a common 1/4" o.d. aluminum tube. The total flow was held at 1.2 vlpm using a variable dump flow, and the line was split into multiple 1/8" o.d. stainless steel tubing sections connecting to each instrument. All tubing junctions employed Y-splitters, and all reducing fittings were internally beveled to prevent impaction losses. Sample lines were not actively dried, but relative humidity was < 30% in LAS and WIBS-4A. For the LAS instrument, the theoretical transmission of the inlet system was 98%, 84%, and 57% for 1, 3, and 5 μm aerodynamic diameter particles, respectively, with gravitational settling being the dominant loss process. Transmission to WIBS-4A for the same sizes was 99%, 90%, and 76%. Size distributions were not corrected for transmission losses. The nephelometer sampled from the original inlet in the SPL instrument laboratory via a blunt tap manifold and 1" i.d. conductive tubing.
Table 1 Descriptions of INP instruments.

<table>
<thead>
<tr>
<th>Instrument Type</th>
<th>Institute</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Online/direct Continuous flow diffusion chamber (CSU-CFDC)</td>
<td>Colorado State University</td>
<td>(Eidhammer et al., 2010; Rogers, 1988; Rogers et al., 2001)</td>
</tr>
<tr>
<td>Spectrometer for ice nuclei (MIT-SPIN)</td>
<td>Massachusetts Institute of Technology</td>
<td>(Garimella et al., 2016; Garimella et al., 2017; Kulkarni &amp; Kok, 2012)</td>
</tr>
<tr>
<td>Offline/post-processing Frankfurt Ice Nuclei Deposition Freezing Experiment deposition mode (FRIDGE-DEP)</td>
<td>Goethe University Frankfurt</td>
<td>(Schrod et al., 2016)</td>
</tr>
<tr>
<td>Frankfurt Ice Nuclei Deposition Freezing Experiment immersion freezing mode (FRIDGE-IMM)</td>
<td>Goethe University Frankfurt</td>
<td>(Schrod et al. 2020; DeMott et al. 2018)</td>
</tr>
<tr>
<td>Ice spectrometer (CSU-IS)</td>
<td>Colorado State University</td>
<td>(Hill et al., 2016; Hiranuma et al., 2015)</td>
</tr>
<tr>
<td>Cold stage (NC State-CS) Cold stage droplet freezing array (on hydrophobic glass slides)</td>
<td>North Carolina State University</td>
<td>(Wright &amp; Petters, 2013; Yadav et al., 2019)</td>
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2.2 INP measurement methods

A combination of direct-processing (online) and post-processing (offline) ice nucleation instruments were employed during the FIN-03 field campaign. All these instruments were also used in the FIN-02 laboratory campaign. Online instruments have the advantage in that the aerosol being evaluated as INPs remain free-floating and unaltered, never touching a substrate nor requiring shipment of samples to a laboratory. Online techniques can also monitor INP
concentration changes occurring over short time scales. Nevertheless, they are limited in the thermodynamic conditions that can be represented over a given time frame, and they are limited by volume sampling rates in assessing the low concentrations of INPs at modest supercooling. Offline techniques, i.e., those in which samples are collected in the field and subsequently processed in laboratory, provide the opportunity to capture large sample volumes (albeit over longer time scales) and consequently assess a wider temperature range of INP activation properties. A summary listing of all ice nucleation instruments is provided in Table 1. Detailed operating principles, siting of samplers (rooftop versus within SPL), and experimental methods for each instrument follow below. In this work, we will refer to the FIN-03 “intercomparison period” to define the times that all INP instruments co-sampled air with substantial temporal overlap for direct comparison. Other times of sampling by the different instrument groups were devoted to special science investigations that are not covered herein.

2.2.1 Online INP measurements

Two online instruments participated in intercomparison experiments in FIN-03. One, the Colorado State University CFDC (CSU-CFDC), has the most established history as an online technique for activating and counting INPs. The CSU-CFDC operating principles are described in several prior works (Rogers, 1988; Rogers et al., 2001; Eidhammer et al., 2010). Application and considerations for interpreting data have been described in detail in several publications, most recently by DeMott et al. (2018). The CSU-CFDC is composed of nested cylindrical copper walls that are chemically ebonized to be hydrophilic so they can be evenly coated with ice. The chamber is divided into two sections vertically. For FIN-03, the CSU-CFDC was operated to establish a temperature gradient between the colder (inner) and warmer (outer) ice walls in the
upper ~50 cm “growth” section to produce either water subsaturated or water supersaturated conditions at various temperatures within a central lamina. For the flow rates used (10 vlpm total flow, 1.5 vlpm sample flow) the residence time was ~5 s in the growth region. Aerosol particles were directed into that central lamina. Ice crystals forming on INPs in the growth section continued to grow for ~2 s in the lower ~35 cm “evaporation” section of the chamber where the outer wall temperature was adjusted to be at an equivalent temperature to the inner (cold) wall to promote evaporation of liquid drops. When operating in the water supersaturated regime, water relative humidity was controlled to be nominally at 105% during FIN-03 to stimulate droplet growth and subsequent freezing, for best comparison to offline immersion freezing methods. For probing ice nucleation in the deposition nucleation regime, relative humidity (RH) was controlled to ~95%.

The CSU-CFDC sampled from one of the turbulent aerosol inlet ports, located in the SPL instrument laboratory. Connection was via ¼” o.d. conductive tubing. Prior to entering the CFDC, aerosol was further dried using two inline diffusion driers and then size-limited using dual single-jet impactors that achieve a 50% upper particle size cut-off at an aerodynamic diameter of 2.5 μm. This limitation on aerosol sizes helps to remove ambiguity when distinguishing ice crystals at ~4 μm sizes from aerosol particles using an optical particle counter at the CSU-CFDC outlet. Temperature uncertainty is ± 0.5 °C at the reported CSU-CFDC lamina processing temperature and relative humidity uncertainty depends inversely on temperature, as discussed by DeMott et al., (2018), estimated for example as 2.4 % at −25 °C.

To correct for background counts that can occur due to ejection of frost emanating from interior surfaces of the CSU-CFDC over operational periods, and for defining measurement uncertainties, we follow Levin et al. (2019). Frost corrections are defined through intervals of
sampling ambient air through a HEPA filter. Sample data were background corrected by subtracting the interpolated filter period concentration before and after each sampling period. Background corrected data were then averaged to ~5-min sampling times to increase statistical confidence. Poisson counting errors during filtered and ambient sampling periods were added in quadrature, and INP concentrations were judged statistically significant at the 95% confidence level if they were greater than 1.64 times this combined INP error (one-tailed z test). Interior inlet tubing losses are not considered in the reported INP data because they have been estimated at 10% or less in the past. INP concentration correction underestimates inferred (by 3 time) to be due to aerosols spreading outside of the lamina during measurements specifically of mineral dust INPs (DeMott et al., 2015) are not generally applied to the data herein, though this is discussed regarding INP parameterizations in this paper.

An aerosol concentrator (MSP Model 4240) was used at selected times during FIN-03 to improve sampling statistics, in the same manner as in previously published studies (Tobo et al. 2013; Suski et al., 2018; Cornwell et al., 2019). The aerosol concentrator was positioned open to the air on the roof of the instrument laboratory room (covered and not used during rainfall), with a short ¼” o.d. copper line containing the concentrated aerosol entering the laboratory from about 3 m above the CFDC. Concentration factors were evaluated in the same manner as Tobo et al. (2013), leading to an average increase of INPs by 90 times during operation of the aerosol concentrator compared to ambient inlet periods during the study (not shown here because analysis repeats numerous past efforts).

A three-way manual stainless-steel valve was used to direct sample air to the CFDC from either the turbulent flow inlet or the aerosol concentrator. At times, a high temperature heating tube (Suski et al., 2018) or a single particle soot photometer (Schill et al., 2016) was placed in-line.
following the three-way valve for removing aerosol organics or black carbon (not reported here) prior to INP measurements. The use of a tube heater upstream of the CSU CFDC to expose single particles to 300°C, following the methods of Suski et al. (2018), is intended to isolate the action of total organic versus inorganic INPs via comparison of ambient versus heat-treated particle streams. Simultaneous measurements of both aerosol streams is of course not possible with a single CFDC, so sampling was conducted by alternating the inlet chosen during subsequent 10-minute periods, and ignoring any very short-term aerosol changes that might occur over such times. This was a special contribution by the CSU CFDC group, for comparison to bulk aerosol treatments discussed in the next subsection.

A second online instrument, the SPectrometer for Ice Nuclei operated at the time by the Massachusetts Institute of Technology (MIT-SPIN; Droplet Measurement Technologies, Boulder, CO), a commercially produced, parallel-plate continuous flow diffusion chamber style instrument, also sampled during FIN-03. Operating principles are described in Garimella et al. (2016) and Garimella et al. (2017). SPIN consists of two flat walls separated by 1.0 cm and coated in approximately 1.0 mm of ice. Aerosol particles are fed into the chamber in a lamina flow of about 1.0 liters per minute and are constrained to the centerline with a sheath flow of about 9.0 liters per minute. The temperature and relative humidity that the aerosol lamina experiences are controlled by varying the temperature gradient between the two iced walls (Kulkarni & Kok, 2012). After exiting the nucleation chamber, the particles enter SPIN’s optical particle counter, which sizes aerosol on a particle-by-particle basis for diameters between 0.2 and 15 µm. For FIN-03, the SPIN sampled from one of the turbulent flow inlet systems, located within the SPL aerosol chemistry laboratory. It was connected to the inlet system port with a short section of ¼” o.d. conductive tubing.
Data processing for SPIN was performed following a similar procedure as the CSU-CFDC instrument. Particle counts from the OPC were first filtered to only consider particles larger than 5 µm. A low-pass filter was applied next to remove all 1 Hz data that exceeded a total of three counts. Particle data was then converted from counts per second to number density (n L\(^{-1}\)) using the combined sheath and sample flow exiting the OPC. A SPIN specific particle concentration correction factor of 1.4 is applied to account for non-ideal instrument behavior resulting in underestimation of INP as described by Garimella et al. (2017). As the field measurements from this study predate the laboratory experiments performed to determine SPIN uncertainties, we select the minimum reported correction factor to remain conservative in our measurements. A depolarization filter was then applied to isolate particle data specific to ice using 1 Hz averaged backscattering data from the SPIN’s OPC, with instrument specific values of 3.5 and -0.25 for the \(\log_{10}(\text{Size})\) and \(\log_{10}(S1/P1)\) measurements respectively. Frost ejected from the plates of the SPIN chamber was characterized by particle-free sampling periods when the sample flow was diverted through a HEPA filter by an automated three-way valve. Linear interpolation was used to approximate background frost throughout the measurement period and smoothed using a five-minute moving average. Sample data was background frost corrected by subtracting this smoothed background frost density from total number density. Lastly, data points that exceeded water saturation were excluded from analysis.

Estimation of measurement error for the MIT-SPIN follows a similar procedure to the CSU-CFDC. Assuming the background corrected INP concentration follows a Poisson distribution, the Poisson error for both INP and background frost concentrations were defined as the square root of the sample mean. The significance test statistic was defined by the quadrature sum of counting errors multiplied by the z-score for a one-tailed z-test at the 95% confidence
interval. INP measurements were deemed statistically significant if the mean INP concentration was greater than this test statistic.

A third online instrument, the Texas A&M CFDC that shares the same design aspects of the CSU CFDC, was used for special studies conducted outside of intercomparison exercises (Zenker et al., 2017).

### 2.2.2 Offline INP measurements

Offline methods have undergone many improvements in recent years and have been successfully demonstrated for being used in a complementary manner to online methods in other recent intercomparisons (DeMott et al., 2017; DeMott et al., 2018; Hiranuma et al., 2015; Wex et al., 2015). In FIN-03 samples were collected with liquid impingers and filter samplers and analyzed for immersion freezing of distributed liquid particle suspensions using the North Carolina State University Cold Stage (Wright et al., 2013), the CSU Ice Spectrometer (Hiranuma et al., 2015; DeMott et al., 2018), and the FRankfurt Ice Nuclei Deposition FreezinG Experiment (FRIDGE) instrument (Schrod et al., 2016).

**The North Carolina State University Cold Stage (NC State-CS)**

The North Carolina State University cold stage (NC State-CS), previously described by Wright and Petters (2013) and Hader et al. (2014). Procedures used for collecting immersion freezing spectra are described below and by Yadav et al. (2019). During FIN-03, filter samples, impinger samples and precipitation samples were collected for analysis using the NC State-CS. For the intercomparison, the filter and impinger results are considered. Filter samples were collected from the roof of Storm Peak Lab for 3–4 hrs twice daily using 47 mm Nuclepore polycarbonate filters (0.2 μm pore size) housed in an open-faced stainless-steel filter holder operated at 14 L min⁻¹ (at altitude) or ~9 L min⁻¹ at standard temperature and pressure conditions.
(STP) of 1013 mb and 0 °C. Filter holders were directed downward and sheltered from precipitation by a large, inverted metal bowl. Images are shown in supplemental Section S1.

Impinger samples were collected directly into water using a glass bioaerosol impinger (SKC, Inc.) as described by Hader et al. (2014) and DeMott et al. (2018). The impinger jets air at 10.6 L min\(^{-1}\) (~7 L min\(^{-1}\) STP) into a 20 mL water reservoir, impacting 80% of particles ≥ 200 nm in diameter and ~100% of particles ≥ 1 μm (Willeke et al., 1998). Impinger samples were collected in the same manner as was done for all shared liquid samples for the FIN-02 intercomparison (DeMott et al., 2018) excepting that Teflon tape replaced stopcock grease to seal the impinger glass lid to prevent jamming. Water evaporating from the reservoir was replaced hourly; the impinger was in a rooftop shelter with its inlet extending through a hole in the shelter wall, into the open air at a height of ~6 feet below the position of filter sampling units that were mounted on an outside railing. Water used onsite was filtered (0.2 μm) Milli-Q water. All samples were stored at −20 °C onsite, shipped on dry ice, then stored at −80 °C until analysis.

Freezing statistics for each liquid sample were acquired by pipetting an array of approximately 256 droplets of 1 μL ± 0.88% volume on four hydrophobic glass slides under dry N\(_2\) gas. Temperature was ramped at a rate of −2 °C min\(^{-1}\) and freezing was detected optically by a microscope at a temperature resolution of 0.17 °C (every 5 s). Freezing temperature spectra are expected to be independent of cooling rate (Wright et al., 2013). The concentration of ice nuclei at temperature \(T\) per unit volume of liquid is given by Vali (1971):

\[
c_{\text{IN}}(T) = \frac{-\ln(f_{\text{unfrozen}}(T))}{V_{\text{drop}} T} \quad (1)
\]

where \(f_{\text{unfrozen}}\) is the fraction of unfrozen droplets at \(T\) and \(V_{\text{drop}}\) is the population-median droplet volume. The concentration of ice nucleating particles (INP) in the atmosphere is given by:
\[ c_{\text{INP}}(T) = \frac{c_{\text{IN}(T)}}{V_{\text{air}} \cdot \frac{V_{\text{liquid}}}{V_{\text{air}}} \cdot \frac{V_{\text{air}}}{V_{\text{air}}} = 0} \]  

(2)

where \( f \) accounts for sample dilution, \( V_{\text{liquid}} \) is sample volume, and \( V_{\text{air}} \) is the volume of air sampled (flow rate at STP \( \times \) duration). Freezing spectra were collected \( 3 \times \) per sample and binned into \( 1 \) °C intervals. Confidence intervals reported in archived data are \( \pm 2 \) standard deviations of the mean. We will refer to the processed filter samples as NC State-CS (F) and processed impinger samples as NC State-CS (I).

**CSU Ice Spectrometer (CSU-IS)**

The CSU-IS also post-processed particles sampled onto filters during FIN-03. This instrument has been described in Hiranuma et al. (2015) and Suski et al. (2018). Samples were collected for approximate periods of 4 hours for intercomparison periods (longer for overnight samples – not part of the intercomparison) using pre-cleaned 0.2 \( \mu \)m pore diameter, 47 mm polycarbonate Nuclepore filters (Suski et al., 2018) mounted in disposable, sterile open-faced and face-up holders (Nalgene), with a typical sample flow rate of 14.9 L min\(^{-1}\) (ambient) and 9.5 L min\(^{-1}\) (STP). Filters were collected on the same exterior laboratory roof railing as the NC State filters, approximately 2 m distance away. All filter samples were frozen following collection, and until processing at CSU. Pre-sterilization procedures and overall clean protocols for preparation and handling of filters are detailed in Suski et al. (2018). Particle re-suspension was done through 20 minutes of shaking filters in sterile 50 mL Falcon polypropylene tubes (Corning Life Sciences) with 6-10 mL of 0.02 \( \mu \)m pore diameter filtered deionized water. Further 20-fold dilutions using filtered water were made as needed to permit measurement of freezing spectra to the low temperature limit of operation of the CSU-IS.

Immersion freezing INP temperature spectra were obtained by distributing 24 - 32 aliquots of 50 \( \mu \)L particle suspensions into the sterile 96-well PCR trays that mount in the CSU-IS. The
cooling rate was –0.33 °C min⁻¹. Frozen wells were counted at 0.2 - 1 °C degree intervals to a limit of about -28 °C, and cumulative numbers of INP mL⁻¹ of suspension estimated following Vali (1971) and Eq. 1. Conversion to ambient air concentrations std L⁻¹ were made based on distributed suspension volume and the total air volumes collected (Eq. 2). Filter blanks were collected during FIN-03, one was tested and used to obtain background INP numbers per filter. Blank INPs were found to account for <5% of INPs at –20 and –25 °C. Binomial sampling confidence intervals (95%) were derived for INP concentrations following Agresti & Coull (1998). The temperature uncertainty of INP measurements is ±0.2 °C.

As a special contribution to FIN-03, portions of IS aerosol suspensions were set aside (e.g., suspensions of 6 to 10 ml can serve up to three or more IS aliquot fills) for treatments to proximally isolate total biological, other organic and inorganic contributions to measured immersion freezing INP concentrations. To assess removal of heat labile INP entities, a 2 mL aliquot of suspension was re-tested in the IS after heating to 95 °C for 20 min (McCluskey et al. 2018). To attempt to remove all organic INPs, 1 mL of 30% H₂O₂ was added to a 2 mL aliquot of suspension and the mixture heated to 95 °C for 20 min while illuminated with UVB fluorescent bulbs to generate hydroxyl radicals (residual H₂O₂ is then removed using catalase) (Suski et al. 2018), and the INPs were again assessed for freezing spectra in the IS. Herein we describe a subset of samples collected on September 15, September 23, and September 25 that were subjected as IS suspensions to the two treatments. These treatments are based on well-established methods which have been used to assess biological components in samples for more than 60 years (Alsante et al., 2023, and references therein). The interpretation of data from exposure of particle suspensions to 95 °C is that the reduction of INP concentrations under thermal treatment is a proxy for the concentration of biological (proteinaceous and microbial) INPs which have been eliminated or deactivated.
through treatment. A strong reduction in INP activity observed after peroxide treatment indicates dominant organic INP populations, whereas a lack of response to this treatment is assumed to indicate that inorganic INPs such as mineral dusts dominate non-heat labile INPs. This assessment for bulk suspensions of particles could be directly compared to measurements of 300°C heat treated single particles in the online CSU CFDC measurements on these same days, providing a more insightful investigation of INP compositions.

The use of such treatments and the insights they convey for atmospheric ice nucleation studies has been reported in published studies of INPs for a variety of locations (McCluskey et al., 2018; Suski et al., 2018; Barry et al., 2021a; Knopf et al., 2021; Testa et al., 2021). Taken together, such treatment studies show general utility for estimating biological contributions to INP, overall organic contributions and the importance of inorganic contributions. However, we note that not all biological materials may be completely denatured or removed by heat (Testa et al., 2021; Daily et al., 2022; Alsante et al., 2023) and not all organics may be removed by peroxide. For example, denaturation is the disruption of higher order (secondary, tertiary, and quaternary structure) in a protein which leads to a loss or lessening of function. It follows that simpler proteins or peptides, such as glutathione, have no higher order structure, and thus cannot be denatured (Alsante et al., 2023). Consequently, estimates of biological contributions to INP based on these treatments may be considered as lower limits.

FRIDGE Cold Stage and Deposition Nucleation Measurements

The FRIDGE instrument can be used to measure the concentration of INPs by two independent methods: a) a droplet freezing assay on a cold stage (hereafter: FRIDGE-CS Schrod et al., 2020; DeMott et al. 2018; Hiranuma et al. 2015) which addresses immersion freezing similarly to the NC State-CS and the CSU-IS and b) the diffusion chamber method (hereafter:
FRIDGE-DC), that addresses the deposition nucleation and condensation freezing modes introduced in Schrod et al. (2016) and referred to as the “standard” method in previous publications (e.g., DeMott et al, 2018). The ice nucleation analysis is performed inside the FRIDGE instrument for both methods, yet the sampling process, addressed nucleation modes and the specific analytical procedure differs as described below.

For the FRIDGE-CS method, aerosol particles were sampled via a short ¼” conductive tube from the shared turbulent flow aerosol inlet in the SPL instrument laboratory on Teflon membrane filters (Fluoropore PTFE, 47 mm, 0.2 μm, Merck Millipore Ltd.). The sampling duration ranged from 50 to 240 minutes, resulting in air volumes between 250 and 1000 std. L. The particles were extracted in 10 ml deionized water by shaking. Approximately 150, 0.5 μL droplets from that solution were pipetted onto a clean, silanized silicon wafer on the cold stage of the FRIDGE instrument and cooled by –1°C min⁻¹ at ambient pressure. A CCD camera detects freezing events and counts the number of frozen droplets as a function of temperature. This process is repeated with fresh droplets and fresh substrates until approx. 1000 droplets are attained. The INP number concentration is derived by Eqs. 1 and 2.

For the FRIDGE-DC measurements, an electrostatic aerosol collector EAC (Schrod et al., 2016) was connected to the same aerosol flow inlet via a short ¼” conductive tube. The EAC consists of a cylindrical sampler, whose inlet is concentrically surrounded by 12 gold wires that are at 12 kV against a grounded silicon wafer, which is used as the sample substrate, at the bottom of the sampler. Once the airflow is pumped inside, aerosol particles are charged by electrons emitted from the gold wires and are precipitated onto the silicon wafer. The analysis at certain pairs of T and RH follows in a separate step. For that, the wafer is placed on the cold stage inside a diffusion chamber. After the chamber is evacuated, the temperature is set to the
first analysis temperature. In a second, much larger volume, pure water vapor is regulated by pressure control to the desired supersaturation. Once the water vapor diffuses into the chamber, ice forms on the activated INPs and a CCD camera is used to record and count the emerging ice crystals, which appear as bright objects. It is assumed that one ice crystal represents one INP. The water vapor atmosphere and thus the growth of ice crystals is maintained for up to 100 seconds until the valve to the water vapor source is closed and the chamber is evacuated again. The process is repeated at increasing humidity first, and then at progressively lower temperatures. At SPL samples were taken from the electrostatic sampler for 50, 75 and 120 minutes, resulting in volumes of approximately 64-150 sL. The samples were analyzed by default at –20°C, –25°C and –30°C and 95%, 99% and 102% water saturation. In addition, a few samples were analyzed at –15°C. This was a special contribution by the FRIDGE group for comprehensive analysis of INP activation in the deposition regime, and for comparison to online data in this regime collected for some days.

2.3 INP processing and sampling strategies

As a campaign strategy, samples were collected over different time periods in the day to reflect both varied weather conditions and aerosol populations arriving at the mountain laboratory. For intercomparison, a select number of 4-hour sampling periods were allocated in which online instruments attempted to operate at a few predesignated temperature and relative humidity ranges, while samples were collected continuously for off-line analysis. While aerosol conditions can change within a 4-hour time frame, this was agreed upon as a minimal reasonable period for comparability. Similar sampling strategies have been employed in the past intercomparisons (DeMott et al., 2017; Knopf et al., 2021). Overall, measurements were conducted over a wide range of temperatures (–7 to –34 °C) in the heterogeneous ice nucleation
regime. Sampling such a broad data set allows for the consideration of instrument performance in response to the presence of highly active INPs (i.e., those facilitating freezing at temperatures $\geq -10 \, ^\circ C$) as well as more modestly effective INP ($-10 \, ^\circ C \geq T \geq -30 \, ^\circ C$). In addition, the range of INP concentrations includes lower concentrations which challenge instrumental limits of detection.

3 Results and discussion

3.1 Meteorological context

Weather conditions during FIN-03 were characterized using auxiliary measurements. Weather data (temperature, humidity, winds and pressure) were obtained for Storm Peak Laboratory through the MesoWest (https://mesowest.utah.edu/cgi-bin/droman/meso_base_dyn.cgi?stn=STORM) mesonet (STORM site), supplemented with measurements from instruments operated at SPL through the Western Regional Climate Center (WRCC) (https://wrcc.dri.edu/weather/strm.html) for the two days that were absent in the MesoWest record. Air temperature, relative humidity, and barometric pressure time series are shown in Figure 1(a), 1(b) and 1(c), respectively. Precipitation was measured via a rain gauge at Storm Peak Laboratory provided by NC State. Precipitation rate was calculated from the quotient of precipitation (in mm) and time collected (in hours), as shown in Figure 1(d). Back trajectories for all the sampling days in FIN-03 are reported by Zawadowicz et al. (2017), showing 72-hr air mass transits from regions that included Southern California, Washington State and Eastern Nebraska.
Figure 1. Weather conditions over the course of FIN-03, including (a) air temperature, (b) relative humidity, (c) barometric pressure, and (d) precipitation rate.

Relatively warm, dry conditions were observed initially at the Storm Peak Laboratory. Clear skies on September 11 and 12, 2015 gave way to clouds and haze on September 13. Cooler temperatures, lower barometric pressure, and higher relative humidity (generally above >
70%) accompanied rainfall on September 14. This was followed by continued rain on September 15, intermittent rain and short periods of hail on September 16, a mixture of rain, snow, and sleet on September 17, and snow on September 18. The next and longest period in the study, September 19 to 28, was marked by an increase in temperature, an increase in barometric pressure, lower relative humidity, and a lack of precipitation. More detailed weather records including daily photographs and a summary of human-produced daily observations are summarized in supplemental Section S1. Daily wind rose plots are provided in Figure S1.

3.2 Aerosol context

3.2.1 Aerosol size distribution and surface area

The time series of aerosol size distribution measured by the LAS (in three hour means) is shown in Figure 2a. The maximum and minimum total LAS concentrations were 706 cm$^{-3}$ and 74 cm$^{-3}$ respectively, and the mean and standard deviation of the total LAS concentration throughout FIN-03 were 410 cm$^{-3}$ and 138 cm$^{-3}$, respectively. The highest total LAS concentration recorded during FIN-03 (706 cm$^{-3}$) occurred in the early hours on September 25. Elevated aerosol concentration (at least one standard deviation above the mean) was also observed during midday on September 13, before and during midday on September 14, before midday on September 25, in the afternoon on September 26, and around midday on September 27.

The timeline of LAS aerosol surface area in Figure 2b emphasizes that surface area was predominately submicron throughout the study, with a mode at about 0.16 μm. This is important to note, in combination with chemical composition information discussed in the next section because it is relevant to understanding the likely sizes and surface areas of INPs. We will revisit the surface area of INPs for use in parameterizations in a later section. Quantitative timelines of
Figure 2. Time series of dry particle number concentration distribution (ambient conditions, not STP) measured by the laser aerosol spectrometer (LAS) in a), shown as three-hour means at ambient pressure. Time series of particle surface area distribution is in b). Timeline of nephelometer scattering (1-hr data) in the red channel for < 1 µm and 1 - 10 µm size ranges, 3-hr LAS number concentration > 0.5 µm, 3-hr LAS surface area at sizes below and above 0.5 µm, and Angström exponent (dashed, right axis).
LAS surface area above and below 0.5 μm are shown in Figure 2c. Surface area at above 0.5 μm is about a factor of 30 lower than at below this size over most of the study period. Also shown in Figure 2c is nephelometer scattering ($b_{sp}$) in the red channel (700 nm) showing a dominant contribution when the upstream impactor was set to 1 μm (aerodynamic) and a much lower level of 1 – 10 μm scattering. This scattering from coarse mode particles is consistent with and trends with the LAS surface area in the supermicron regime, while the Angström exponent (calculated using red and blue channels) being close to 2 (small particle dominance) throughout the study is consistent with the dominance of submicron contributions to total surface area. Figure 2 also emphasizes that the lowest aerosol concentrations and surface areas occurred during varied time in the wet period of the study from midday on the 14th through the 17th of September.

3.2.2 Aerosol composition

The number concentration of aerosol particles from 0.2 to 3 μm with characteristic spectra belonging to eight composition categories (sulfate/organic/nitrate, biomass burning, elemental carbon, sea salt, mineral dust, meteoric, alkali salt, and fuel oil combustion), and the number concentration of unclassified aerosol particles by the PALMS, were assessed for three-hour averages through the FIN-03 period. For simplicity, four of these categories (elemental carbon, meteoric, alkali salt, and fuel oil combustion) were combined into a category called “other” due to the low concentration of particles in each of these categories resulting in 6 total classifications (categories (SulfOrgNit = sulfates/organics/nitrates, BioBurn = products of biomass burning, Sea salt, Mineral dust, and Unclassified), as shown in Figure 3a. The three-hour averages of the number fractions of each particle type were also calculated as the fraction of the total aerosol number concentration measured by the PALMS in each of the six classifications, as shown in Figure 3b. The dominant categories throughout the FIN-03
Figure 3. Subplots (a) and (b) show the aerosol particle number (ambient conditions, not STP) and relative fractions (by cumulative count at all sizes) of each of the six PALMS compositional particle types for the three-hour periods during which the PALMS was used to sample ambient air. Subplots (c) and (d) show the aerosol particle number concentration and relative fractions (by count) of particles with diameter $>0.5 \, \mu m$ in each of the channels (A, B, AB, C, AC, BC, and ABC, which are described in Perring et al., 2015) over the course of the FIN-03 field campaign.
Figure 4. a) Total aerosol versus mineral/soil dust (ambient) number size distribution and dust fraction interpreted from PALMS and LAS data for all times that the PALMS was sampling during FIN-03. b) Surface area distribution differentiated for PALMS compositional types during the same sampling times. c) Expanded plot from b) for the coarse mode size range to emphasize progressive dominance of dust components at diameters > 0.5 µm.
campaign were BioBurn (mean 26 ± 43 cm$^{-3}$, maximum 177 cm$^{-3}$), SulfOrgNit (mean 22 ± 13 cm$^{-3}$, maximum 48 cm$^{-3}$), and mineral dust (mean 3 ± 11 cm$^{-3}$, maximum 55 cm$^{-3}$). The mineral dust type also includes soil particles (crustal species mixed with organic material) (Zawadowsicz et al., 2019). The highest total particle number concentration measured by the PALMS (218 cm$^{-3}$) occurred on September 14 (of which 177 cm$^{-3}$ consisted of biomass burning and 34 cm$^{-3}$ consisted of sulfates/organics/nitrates). This biomass burning plume impacted the site for several hours. Mineral/soil dust particles were ubiquitous throughout the study, with a concentration of 0.128 ± 0.446 cm$^{-3}$ (median and interquartile range). Anomalous concentrations >10 cm$^{-3}$ observed for a few 5-min sample periods are likely due to road dust emitted from site. Dust concentrations were <1 cm$^{-3}$ for 90% of the PALMS samples. Mineral/soil dust represented a median of 0.3% of particles in the >0.2 µm size range, increasing to 23% and 67% for >0.5 and >1.0 µm particles (Figure 4a). Similarly, mineral dust contributions to total surface area are inconsequential for total aerosol (Figure 4b) but dominate in the coarse mode regime for the study (Figure 4c). We will revisit this result in discussions of parameterization of INPs in Section 3.5.

The daily average number concentration of fluorescing aerosol particles corresponding with each of the seven WIBS-4A types with diameter > 0.5 µm is shown in Figure 3(c), and the daily average number fraction of each WIBS-4A type is shown in Figure 3(d). The dominant types of fluorescent aerosol particles throughout the FIN-03 field campaign were types B, AB, and A, which on average accounted for 63.2%±8.7%, 16.0%±6.3%, and 12.5%±3.9% of the particles detected by the WIBS respectively.

In contrast with the daily average number fraction in each PALMS category, the relative contributions of each of the seven WIBS-4A particle types did not vary much over the course of
the study when the WIBS-4A was operational, with perhaps the exception that Type AB decreased in prevalence from September 18 (42.9%) to September 21 (10.1%). A modest trend from lower total fluorescing particle concentrations (0.02 to 0.04 cm\(^3\) at STP) through September 21 to higher concentrations (0.07 to 0.15 cm\(^3\) at STP) from September 22 through 26\(^{th}\). WIBS-4A data was not collected on September 13-16, nor on September 27. The first period was somewhat critical to evaluating INP relations to bioaerosols, so we note here in advance this caveat. Time-resolved size distributions for each WIBS-4A channel, as well as the total particle concentration measured across these seven channels, are shown in supplemental Figure S2. FBAP assignments related to INP predictions will be discussed in Section 3.5.

### 3.3 Immersion freezing measurements

A summary of the number concentrations of immersion freezing INPs (\(N_{\text{INP}}\)) over the course of the field campaign, binned for one degree temperature intervals, is shown in Figure 5. The concentration of INP detected over this range ranged over five orders of magnitude (0.01 to 160 L\(^{-1}\)). At any one temperature, differences up to a little more than one order of magnitude are apparent in comparing average data from individual methods, mirroring results presented in previous laboratory and field studies (Hiranuma et al., 2015; DeMott et al., 2017, 2018; Knopf et al., 2021; Lacher et al., 2024).

As expected, a trend of increasing \(N_{\text{INP}}\) with decreasing temperature was observed for the FRIDGE-CS, CSU-IS, NC State-CS, and CSU-CFDC. The data represented by the single MIT-SPIN processing temperature condition also falls well within the concentration range reported for the other instruments. Incremental changes in \(N_{\text{INP}}\) with decreasing temperature was similar for all measurements that spanned a broad temperature range. This comparability of \(dN_{\text{INP}}/dT\) contrasts with an apparent increasing high bias of drop suspension freezing measurements versus...
Figure 5. Campaign average immersion freezing INP concentrations (sL$^{-1}$) in 1 °C bins for instruments participating in intercomparison studies. Error bars represent one standard deviation in the measurement means collected at the specified temperature and not measurement uncertainties. The star labels indicate when these exceeded the mean for the FRIDGE-CS data. The times over which the INP concentration has been averaged for each instrument is explained in the text.

CFDC measurements during comparable sampling at various surface sites (non-mountaintop or free troposphere) found in DeMott et al. (2017) but agrees with FIN-02 laboratory studies (DeMott et al., 2018) and recent atmospheric studies at Puy de Dome (Lacher et al., 2024). INP concentration variability at single temperatures, reflected in Figure 5 as a standard deviation of bin means, is likely due to variations in aerosol properties in response to production and scavenging processes upstream of the site. Nevertheless, generally higher $N_{\text{INP}}$ measurements...
were obtained with the FRIDGE-CS and the CSU-IS than the CSU-CFDC and NC State-CS (F) and NC State-CS (I) analyses. Such biases in other studies have been attributed to different efficiencies in sampling of largest particles (e.g., Lacher et al., 2024; Cornwell et al., 2023), but the collection methods for offline measurements in this study were substantially similar, as discussed further below. Hence, we cannot attribute measurement differences to a systematic source. Comparability of impinger versus filter sampling methods for immersion freezing measurements via the NC State-CS mirrors the findings in DeMott et al. (2017).

To compare the operation of these instruments over time, the mean and standard deviation (when applicable) of immersion freezing $N_{INP}$ were calculated over three-hour periods for each instrument (except for the MIT-SPIN) at 1 °C intervals (± 0.5 °C). Means are plotted as a time series in Figure 6. Although some differences appear in comparing instrument by instrument, as will be discussed, some general observations from the temporal data of Figure 6 are that INP concentrations at temperatures > –20 °C were at a maximum during the precipitation period, as might be expected for rainfall production of biological INPs (Huffman et al., 2013; Mignani et al., 2021; Testa et al., 2021; Cornwell et al., 2023), while the strongest differences between the concentrations of INPs active at higher and lower temperatures occurred during period of warming under high pressure later in the study. The latter observation might be expected for a strong contribution of dust-like INPs, with a steeper $dN_{INP}/dT$.

Periods of agreement and discrepancy are clearer in examining the ratios of time-matched and temperature-matched three-hour immersion $N_{INP}$ values that were calculated for each pair of instruments, as shown in Figure 7. As a positive note, the mean $N_{INP}$ reported by different instruments for all temperature conditions taken together generally fell within a span of one order of magnitude. Figure S3 shows the percent of immersion INP measurements in which all
Figure 6. Time series of immersion-freezing mode INP concentrations (sL$^{-1}$) measured during intercomparison periods by (a) the FRIDGE from Goethe University Frankfurt, (b) the CSU ice spectrometer, (c) the NC State cold stage (collected using an impinger), (d) the NC State cold stage (collected using the filter), and (e) the CSU continuous flow diffusion chamber. INP concentrations shown in this figure are averaged over three-hour periods.
Figure 7. Ratios of the immersion freezing INP concentrations measured by each instrument, to the immersion INP concentrations measured by each other instrument (three-hour averages). Each instrument (FRIDGE, CSU-IS, NCSU-CS (I), NCSU-CS (F), and CSU-CFDC) is represented by one of the five columns as well as one of the five rows.

Instrument pairs agreed within one order of magnitude. This level of agreement compares well with the findings from FIN-02, for which the immersion $N_{\text{INP}}$ measured by several online and offline instruments agreed within an order of magnitude. This is encouraging given that FIN-02 was a laboratory intercomparison on single composition aerosol samples consisting of particles with diameter < 2 µm whereas FIN-03 was a field campaign in which constant changes in the concentration, size distribution, and composition of the ambient aerosol population at Storm...
Peak Laboratory were all potential factors. This shows that field data can be collected with nearly the same level of accuracy as laboratory experiments. While also mimicking the results of DeMott et al. (2017) for a smaller instrument comparison exercise, agreement was slightly poorer than found in another recent intercomparison where INP concentrations were stated to match within a factor of 5 (Lacher et al., 2024).

Reiterating what is apparent from campaign-wide results in Figure 5, the best agreement for short-term periods throughout the study as shown in Figure 7 was observed between the FRIDGE-CS and the CSU-IS, in which only one out of seventy-two 3-hour, time- and temperature-matched \(N_{\text{INP}}\) (1.4%) did not agree within an order of magnitude. Nevertheless, discrepancies of a few to several times did occur from September 19\(^{th}\) onward, focused more often at \(>-22^\circ\text{C}\), with the CSU-IS measuring higher from the 19\(^{th}\) to the 22\(^{nd}\) and the FRIDGE-CS higher at some other times, notably the 23\(^{rd}\) and 26\(^{th}\) of September. None of these periods were distinguished in any discernible manner by weather or aerosol properties. For example, LAS and PALMS concentrations were no more than 20% different than the FIN-03 campaign means during any of these periods. Aerosol surface areas were about a factor of two lower overall during the 19\(^{th}\) to 22\(^{nd}\) period than for the period after the 23\(^{rd}\) (Figure 2), which does not imply a special sampling bias for larger particles for the IS filter that was open to the air, a point we will discuss further below.

Both the FRIDGE-CS and CSU-IS showed high bias from a few to more than 10 times versus NC State-CS(I) or CS(F), primarily at processing temperatures below \(-20^\circ\text{C}\), whereas agreement was generally very much better at \(>-20^\circ\text{C}\). The poorest agreement overall was observed for the CSU-IS compared to the NC State-CS(I), a combination for which 12 out of 44 (27%) immersion \(N_{\text{INP}}\) means did not agree within an order of magnitude. Agreement between
the FRIDGE-CS and the NC State-CS(I) was only slightly better, as 13 out of 52 (25%) time-
matched N_{INP} means did not agree within an order of magnitude. Higher than order of magnitude
such discrepancies at lower temperatures were markedly present on September 14, 23 and 26.
Based on PALMS data, the 14\textsuperscript{th} was richer in compounds from biomass burning, poorer in
sulfates, organics, and nitrates, and slightly poorer in mineral dust than average, as discussed in
Section 3.2. The concentration of > 0.5 µm particles measured by the LAS during this time was
also relatively high (2.5 cm\textsuperscript{3} compared to the campaign mean 0.45±0.62 cm\textsuperscript{3}). However, the
14\textsuperscript{th} is not markedly distinguished overall in the timeline of all INP measurements in Figure 6, so
perturbations to composition and concentrations of all particle sizes due to the biomass burning
event did not appear to specially perturb the INP populations. We have already noted that the
23\textsuperscript{rd} and 26\textsuperscript{th} of September had aerosol populations that were not much different than the project
mean on those days.

The CSU-CFDC INP measurements generally agreed with the other measurements within
an order of magnitude for data collected on the same day and temperature, and its measurements
of INP concentration were in best agreement with all methods for temperatures > −20 °C. CSU-
CFDC INP concentrations tended to be lower than those from the FRIDGE-CS and CSU-IS at
temperatures below −20 °C. A similar divergence in online versus offline N_{INP} measurements in
this temperature range was reported by DeMott et al (2017) for ground-based sampling, with
online measurements tending to measure progressively lower INPs than offline integrated filter
or impinger collections at below −20 °C, approaching one order of magnitude below −25 °C. At
the Puy de Dome mountain station (Lacher et al. 2024), only modest and insignificant
underestimates were made by the CSU-CFDC (again, with a 2.5 µm impactor) versus offline
INP concentrations when all were measured from a PM10 inlet. CSU-CFDC INP measurements
were comparable on average with measurements from the NC State-CS(I) and NC State-CS(F), consistent with the mean results shown in Figure 5. Comparing the timeline of ratios of NC State-CS(I) to NC State-CS(F), only 5 out of 130 (4%) of the INP concentrations obtained through analysis by the identical off-line apparatus differed by more than an order of magnitude.

A possible explanation for INP measurement discrepancies that has been tendered in other intercomparison campaigns sampling ambient air is that INPs are highly sensitive to the size range of collected aerosol, and systematic size-dependent differences in collection efficiencies vary for different collection types (DeMott et al., 2017; Knopf et al., 2021; Lacher et al., 2024). For example, Lacher et al. (2024) found significant underestimates of INPs by both online and offline methods measuring from the PM10 inlet versus offline measurements from filter collections made on the laboratory rooftop. In this study, as we have noted above, a similarly consistent difference between rooftop versus laboratory or between online and offline measurements is not found. FRIDGE-CS measurements from the turbulent-flow inlet and CSU-IS measurements from the rooftop filter agreed on average over the course of the study. CFDC INP measurements agreed reasonably well with the NC State (F) and (I) measurements. Larger particles do tend to have higher ice nucleation efficiency, so biases in their collection can lead to sometimes large differences in assessed INP concentrations (Mason et al., 2016). Disaggregation of the very largest collected particles when placed in water suspensions has also been implicated for discrepancies between different substrate collections (DeMott et al., 2017; Lacher et al., 2024). An obvious size-based collection bias existed for the online INP instruments, which had impactors upstream to limit particles >2.5 µm (50% cut-size) from entering. There may have been additional line losses for these instruments sampling from an inlet and using tubing to transfer particles, though these tend to be of minor influence at below the impactor size cut.
(Knopf et al., 2021). The impinger is known to be less efficient for small (<200 nm) and large (>10 µm) particle capture, but unless the relatively light to moderate wind conditions at the inlet during FIN-03 conferred some special bias, Hader et al. (2014) predict a 50% capture efficiency at near 10 µm. The filter samplers on the rooftop should have been equivalent, with the only difference in the orientation of filters for the NC State samples being mounted face-down. The size bias in this configuration is unknown. The FRIDGE filter should have captured particles with the same efficiency as the turbulent flow inlet, since only a very short line connected the filter to the interior inlet structure in the laboratory. Only if very large INPs > 13 µm were dominant by number amongst total INPs, which is unexpected, would the FRIDGE filter collection have been expected to differ from the rooftop filter collections.

In the end, it seems more likely that unquantifiable random and non-random sources of discrepancy, related to such things as sample size, instrument temperature sensor drift, varied instrument cooling rates and inconsistency in sample materials or handling and storage (e.g., Barry et al., 2021b; Beall et al., 2021), may also contribute to the fact that measurements of immersion freezing INP concentrations from ambient air are generally uncertain by up to one order of magnitude, as this study once again supports.

### 3.4 Relation of immersion freezing INPs to aerosol properties

While establishing correlations between INPs and aerosol properties were not a focus of the intercomparison, the ancillary aerosol data did allow for inspecting some simple linear correlation analysis. This provides insight into the size range of greatest relevance for the INP intercomparison period. Throughout the campaign, a positive and significant trend between total LAS particle concentration (i.e., > 0.1 µm) and $N_{\text{INP}}$ was observed for FRIDGE-CS ($R = 0.55-0.74$ and $p < 0.05$ for measurements at $-28 \, ^\circ C < T < -15 \, ^\circ C$), but no clear statistically significant
trend was observed between total LAS particle concentration and $N_{\text{INP}}$ for the other four instruments (Figure S4a). A greater number of significant positive trends were found between the concentration of particles with diameter $> 0.5 \, \mu\text{m}$ and $N_{\text{INP}}$. This was the case for the FRIDGE-CS (R = 0.54-0.94 and $p < 0.05$ for measurements at $-28 \, ^\circ\text{C} < T < -19 \, ^\circ\text{C}$), CSU IS (R = 0.46-0.72 and $p < 0.05$ for measurements at $-21$ to $-25 \, ^\circ\text{C}$), NC State CS(I) (R = 0.46-0.61 and $p < 0.05$ for measurements at $-29 \, ^\circ\text{C} < T < -24 \, ^\circ\text{C}$), and the NC State CS(F) (R = 0.51-0.64 and $p < 0.05$ for measurements at $-26 \, ^\circ\text{C} < T < -22 \, ^\circ\text{C}$).

No consistent, significant ($p < 0.05$) correlation was found between changes in composition (from the PALMS categories and WIBS-4A types) and immersion freezing $N_{\text{INP}}$ across the range of setpoint temperatures employed during FIN-03 (Figure S4b).

**3.5 Inferences to INP compositions during FIN-03**

To provide context for the discussed intercomparisons and because this study provides data needed for testing the relevance of existing parameterizations of ice nucleation in regional and global climate models (Andreae & Rosenfeld, 2008; Morris et al., 2011; Seifert et al., 2011), we utilize some previously-developed ice nucleation parameterizations for specific compositions to diagnose consistency or not with INP compositions in the high altitude environment of FIN-03. We examine parameterizations for mineral dust INPs that have different links to larger size particle concentrations (DeMott et al., 2015) versus mineral dust surface area (Niemand et al., 2012), and biological INPs as linked to fluorescent particle concentrations (Tobo et al., 2013; Twohy et al., 2016). Hereafter we will refer to these parameterizations as DeMott 2015, Niemand 2012, and Tobo 2013. We also utilize a more direct method of probing INP compositions using the IS sample treatments discussed in Section 2.2.2.
Each of the above-noted deterministic parameterizations was used to predict $N_{\text{INP}}$ at –30°C, –25°C, –20°C, and –15°C using the equations and inputs described in Table 2 and summarized below. We do not attempt an analysis using stochastic parameterizations.

1) DeMott 2015 is based on CSU-CFDC laboratory measurements of ice nucleation on mineral dust soil samples as well as field data from situations dominated by mineral dusts (i.e., dust plumes from major deserts), collected for CFDC operational conditions essentially the same as for this study (i.e., simulated immersion freezing conditions at 105% RH) (DeMott et al., 2015). For FIN-03, aerosol concentrations measured by the LAS (> 0.5 µm dry diameter) and converted to STP concentrations were used as the input for this parameterization for comparison to INP data that is also reported at STP concentrations. Predictions also depend on temperature (Table 2). Since PALMS data indicates that dust particles dominated the coarse mode only at sizes above 1 µm in diameter (Figure 4), we first adjust LAS data accordingly for the percentage of dust particles with diameters > 0.5 µm as input to this parameterization, which we have already stated is 23%. A correction factor (CF) of 3 was also applied (as indicated in Table 2) according to the results in DeMott et al. (2015) which showed that when applying the parameterization to represent immersion freezing INP concentrations in a model or in comparison to other immersion freezing methods, this CF is needed to account for CFDC underestimates of immersion freezing INPs (see Methods). The CF is applied in this case because calculations will be compared to the average $N_{\text{INP}}$ from all measurements.

2) The Niemand 2012 parameterization (Table 2) for mineral dust INPs is based entirely from laboratory measurements and incorporates measurements of temperature and
particle surface area as the basis for prediction of INPs. It is especially important to limit the size range of aerosols for which this parameterization is applied, because total surface area was dominated by small particles in FIN-03. Therefore, with reference to Figure 4, we will assume that all dust surface area occurs at sizes larger than 0.5 \( \mu m \) and represents 50\% of that surface area.

Table 2 Summary of INP parameterizations.

<table>
<thead>
<tr>
<th>Parameterization</th>
<th>Equation</th>
<th>Constants</th>
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| Mineral dust INPs: Niemand et al. (2012) | \( N_{INP}(T_C) \approx n_s(T_C)S_{tot} = (a \exp (b(T_C) + c))(S_{tot}) \) | \( a = 1 \times 10^{-9} \)  
\( b = -0.515 \)  
\( c = 8.821 \) |
| Mineral dust: DeMott et al. (2015) | \( N_{INP}(T_K) = (cf)(n_{a>0.5\mu m})\exp(\gamma T_F + \delta) \) | \( \alpha = -0.74 \)  
\( \beta = 3.8 \)  
\( \gamma = 0.414 \)  
\( \delta = -9.671 \) |
| Fluorescing biological aerosol particle INPs: Tobo et al. (2013) | \( N_{INP}(T_K) = (N_{FBAP>0.5\mu m})\exp(\gamma' T_F + \delta') \) | \( \alpha' = -0.108 \)  
\( \beta' = 3.8 \)  
\( \gamma' = 0 \)  
\( \delta' = 4.605 \) |
| Fluorescing biological aerosol particle INPs: Cornwell et al. (2023) | \( N_{INP}(T_C) = f(T_C)1000N_{FBAP>0.5\mu m} \) | N/A |

3) As discussed earlier, we use two definitions of FBAP at sizes larger than 0.5 \( \mu m \) to and temperature to predict biological INP concentrations based on Tobo 2013 as defined in Section 2.1, presuming to bracket low and high estimates of their links to INPs. We also
explore links of higher temperature freezing data (> −20 °C) to FP3 particles, using the same scalings of the relation between FP3 concentrations and INP concentrations as a function of temperature that were established by Cornwell et al. (2023) for a coastal California environment. While we have no reason to expect that these scaling factors listed in Table 2 are valid for the high altitude, continental environment of FIN-03, they are starting points to explore this additional link of certain FBAP particles to INPs.

To compare these parameterized values with observations, an overall mean observed immersion freezing \( N_{\text{INP}} \) was calculated for each three-hour period based on all the available data from all the instruments. This was considered as a reasonable approach since it factors in the inherent variability found between methods. Immersion freezing \( N_{\text{INP}} \) was predicted for each parameterization using mean WIBS-4A, and LAS data, both at STP concentrations, collected in the coincident 3-hour periods of time as the INP data. The observed and predicted immersion freezing \( N_{\text{INP}} \) are plotted against each other in Figure 8. Four temperatures of comparison (−15, −20, −25 and −30 °C) are presented in Figure 8 for DeMott 2015, Niemand 2012, and Tobo 2013, while two temperatures of comparison (−15, −20 °C) are used for links to FP3-based prediction of biological INPs. Temperatures are indicated via levels of shading of the data points.

Using the constraint on mineral particles from the combination of PALMS and LAS data for the campaign average, predicted INPs underestimate the mean INP concentrations at all temperatures (Figure 8a). The Niemand 2012 surface-area-based INP estimates come modestly closer to observations, averaging 25% of the total INP concentrations for all times and all temperatures, while the DeMott 2015 predictions average 4% of INP concentrations, with large variability apparent. These results can be expected to be highly sensitive to the assessed average
Figure 8. a) Comparison of mean observed N$_{\text{INP}}$ (all instrument average) and predicted N$_{\text{INP}}$ calculated from DeMott et al. (2015) (DeMott 2015) and Niemand et al. (2012) (Niemand 2012) mineral dust INP parameterizations at temperatures -30 °C, -25 °C, -20 °C, and -15 °C (gradations in shading from dark to light) for the PALMS estimated percentages of dust particle number and surface area at sizes above 0.5 µm. Mean N$_{\text{INP}}$ are averaged over three-hour periods and plotted uncertainties are standard deviations. Predicted N$_{\text{INP}}$ uncertainties are propagated based on 25 % uncertainty in aerosol number and surface area concentrations. b) Comparison of mean observed N$_{\text{INP}}$ and predicted N$_{\text{INP}}$ calculated from parameterizations linking to FBAP concentrations from Tobo et al. (2013) (T13_low and T13_high; see text for description) and from Cornwell et al. (2023) (C23_FP3) following the FP3 particle definition of Wright et al. (2014). Only -15 and -20 °C comparisons are shown for the FP3 prediction. The solid line in each plot is the 1:1 line and the dashed lines represent an order of magnitude in both directions.
mineral particle fraction at sizes above 0.5 µm (varied over the study) and on whether particles that have a source from regional soils will be represented only by those with mineral content. Therefore, for comparison, parameterization results using an assumption that all particles at diameters exceeding 0.5 µm were dust particles are presented in Figure S5. In this case, admittedly a somewhat unrealistic maximum assumption on mineral dust numbers and surface area, Niemand 2012 estimates a dust source for 50% and DeMott 2015 estimates 25% of observed INPs on average. Thus, the predictions of the two parameterizations become more closely aligned for assumption of more overall mineral dust particles in the size range larger than 0.5 µm. Discrepancy has been noted previously in applying these parameterizations to link to the aerosol model in an Earth System model for the Southern Ocean region (McCluskey et al., 2023). In that case, calculations were based on aerosol model derived dust distributions and occurred under very low dust loading scenarios where neither parameterization has been firmly tested in the laboratory or field. Under both assumptions on mineral particle number, since DeMott 2015 was developed based on CFDC measurements for particles < 2.5 µm in the field and laboratory, a low bias compared to Niemand 2012 might be expected in comparison to average immersion freezing data that includes larger particles.

The timeline of predicted $N_{\text{INP}}$ for the two dust parameterizations in comparison to mean observed $N_{\text{INP}}$ is shown in Figure 9 for the same temperatures used in Figure 8. These analyses emphasize that 1) INP observations do not show a special enhancement during the biomass burning event at the start of FIN-03, and hence closer agreement of the dust parameterizations with observations at that time is likely an artifact of attributing dust-like INP activation properties to the dominant biomass burning compositions at that time; 2) the predicted $N_{\text{INP}}$ trends better with observed $N_{\text{INP}}$ at temperatures $<-20$ °C, as expected for a dominance of dust-
like INPs; and 3) the predictions fare less well in describing the observed INP populations at > – 20 °C where biological INPs may be expected to have greater influence. Thus, these analyses overall suggest the presence of a dust-like immersion freezing INP type during FIN-03, but that the typical INP efficiency (INP as a function of dust concentration and temperature) attributed to mineral dust underestimates the freezing behavior of INPs overall during the period of study.

**Figure 9.** Time series of aerosol number concentration and surface area (3-h averages at STP) in a), and observed mean measured immersion freezing N\textsubscript{INP} (INP mean) plotted with predicted N\textsubscript{INP} from the mineral dust parameterizations of Niemand 2012 and DeMott 2015 as described in the main text (all three-hour averages at STP) at temperatures of -15, -20, -25, and -30 °C in b) to e), respectively. Dashed lines are intended only to connect data points and do not imply knowledge of intermediate values. Uncertainties mark one standard deviation above and below the mean values of all parameters.
Figure 10. a) Timelines of WIBS-based fluorescent particles assignments (all fluorescing in any channel, low and high FBAP, and FP3 particles), as defined in the text, during FIN-03. b) INP observed mean concentrations and biological INP parameterization predictions linked to high FBAP following Tobo et al. (2013) (T13-high) and FP3 particles following Cornwell et al. (2023) at -15 °C in b) and -20 °C in c).
For FIN-03, the Tobo parameterization of biological INPs consistently underpredicted $N_{\text{INP}}$, independent of the WIBS FBAP definition used, denoted as T13_low and T13_high in scatterplot comparison of measured versus predicted values at all times and temperatures in Figure 8b and the timeline comparisons at $-15$ and $-20 \, ^\circ\text{C}$ shown in Figure 10. Figure 10 also shows the timeline WIBS total fluorescent particle concentrations, the high and low FBAP concentrations, and FP3 concentrations. The higher FBAP prediction of INPs falls much closer to the observations than the low FBAP prediction in Figure 8b and shares some proximal equivalence to observations at $-15$ to $-20 \, ^\circ\text{C}$ at times. This result is like that found by Twohy et al. (2016) for air over the site where Tobo et al. (2013) collected their data, with the higher FBAP estimate bounding the upper end of measured immersion freezing INP concentrations at temperatures $> -20 \, ^\circ\text{C}$. Also notable in Figure 8b and Figure 10 is that the C13-FP3 INP concentration predictions filled a similar space as the T13_high estimates, coming closest together at $-20 \, ^\circ\text{C}$. While these results suggest that biological INP parameterizations can explain the higher temperature INP concentrations observed during FIN-03, with caveats on the large and likely not fully quantifiable uncertainty in such predictions, the temporal analysis (Figure S6) indicates that there is no consistent temporal agreement between predicted and measured INPs, even if different scaling factors were applied to the predictions. Predictions at $-20 \, ^\circ\text{C}$ again show the best overall agreement, while those at $-15 \, ^\circ\text{C}$ suggest that the Cornwell et al. (2023) scaling factor should be higher for the SPL site at the time of FIN-03 to better describe mean values of biological INP concentrations using the FP3 particle signal.
Figure 11. Summary of treated IS filter suspensions using heat and peroxide (a, b, c) and dry heat-treated CSU CFDC single particle data (d, e, f), for September 15, 23 and 25 (a-c, d-f, respectively). Error bars represent 95% confidence intervals for individual experimental spectra for the CSU-IS and for individual CSU CFDC measurements.

The results of CSU-IS and CSU-CFDC treatments on INP concentrations measured for three (of 21 overall) intercomparison time periods are shown in Figure 11, for examination of consistency with the results of the diagnostic parameterization analysis just discussed. In Figure 11a-c, it is seen that thermal treatments indicated the strong contribution of inferred biological INPs primarily at temperatures higher than about −20 °C, but that peroxide digestion of organic compounds lowered INP activity at all tested temperatures by an order of magnitude on average. Similar reductions of INPs measured for single particles by the CSU-CFDC following dry heating (Figure 11d-e) demonstrate strong consistency with the IS results for bulk immersion freezing on the dominance of organic INP compositions, even though CSU-CFDC measured unamended INP concentrations were always lower. The CSU-IS heat treatment results (Figure...
11a-c) suggest that biological INPs may have been ubiquitous during FIN-03 at temperatures above –20 °C, and extended to lower temperatures at times, as indicated by the results from September 25. This is broadly consistent with the parameterization results based on FBAP measurements, although the Tobo 2013 and FP3 parameterizations did not capture all the influence of apparent biological INPs during the study. Whether for size-limited (<2.5 µm) as in CSU-CFDC measurements, or bulk aerosol collected for CSU-IS immersion freezing measurements, the inferred INP compositions typically dominated by organics at temperatures < –20 °C could reflect origins from arable soil dusts (Testa et al., 2021) that surround the region of study. Biomass burning aerosols also have influence as organic INPs (Schill et al, 2020; Barry et al., 2021). However, while biomass burning type particles were noted as a prevalent composition in FIN-03, these types of potential INPs likely cannot explain INP concentrations in FIN-03 because Barry et al. (2021) showed that Western U.S. biomass burning INPs have active site densities about 3 orders of magnitude lower than those attributed to dust particles that also were ubiquitous at modest number concentrations during FIN-03. Furthermore, the strong biomass burning event noted on September 14 had only modest, if any, apparent impacts on INP concentrations despite greatly elevated aerosol concentrations and surface areas, as already mentioned above (Figure 9).

Finally, in Figure 12 we address whether the treatment results support the conclusion of the diagnostic parameterization analysis suggesting that inorganic INPs (mineral particles in particular) were of minor influence during FIN-03. We introduce the additional parameterization of Tobo et al. (2014) (Tobo 2014) for arable soil dust INPs as part of this discussion. Tobo et al. (2014) parameterized the ice nucleation behavior of soil dusts from Wyoming, regionally proximal to the FIN-03 site at SPL, specifically using the CSU CFDC and the dry heat method at
Figure 12. a) Comparison of all untreated CSU CFDC data (black circles), cases after passing through the upstream 300 °C tube heater (purple diamonds), and calculations from the DeMott 2015 dust parameterization in (orange squares) and with $CF = 1$ as appropriate for a direct comparison to CSU CFDC data (see text). b) The same exercise as in a) but using predictions of total soil organic INP concentrations and inorganic INP concentrations within soil INPs, both from Tobo et al. (2014). c) The same exercise but for all CSU-IS data and the cases with peroxide digestion. In this case, $CF = 3$ must be used in DeMott 2015 and the mineral dust INP prediction of Niemand 2012 is also shown.
300 °C to indicate organic versus inorganic INP contributions from such soil particles. This parameterization, like Niemand 2012, is based on the surface area of dust particles and so we apply the same assumptions as before to restrict to the proportion of dust larger than 0.5 µm. Since the CSU-CFDC is also restricted to measuring INPs at diameters below 2.5 µm, we apply a correction factor to the surface area to account for the fact that the surface area at below this size was 90% of the project average total surface area. No significant impact of the treatments is assumed on aerosol concentrations or surface area at sizes above 0.5 µm in Figure 12.

Figures 12a and 12b focus on specific comparisons to CSU-CFDC data. In Figure 12a, it is seen that INP concentrations predicted by the DeMott 2015 parameterization for sampling periods during the entire campaign show remarkable agreement with the 300 °C CSU-CFDC data on selected days when applying CF = 1 in the parameterization, as is appropriate for a direct comparison to instrument data that is uncorrected for the underestimates that led to selecting CF = 3 for modeling studies. In Figure 12b, it is shown that the Tobo 2014 parameterizations for untreated soil dusts and the inorganic remnants also give very good agreement with CFDC untreated and treated data, supporting the likely important influence of such arable soil dusts during FIN-03. Predictions for untreated soils do not quite reach the level of the observed INPs, but this could be explained by the additional contribution of biological INPs that has already been discussed.

In Figure 12c, direct comparisons of the Niemand 2012 and DeMott 2015 predictions for mineral dust INPs for the entire project are shown in comparison to the CSU-IS untreated and H₂O₂ treated data on selected days. The DeMott 2015 prediction of INP concentrations uses CF = 3 in this case, as appropriate. The same discrepancy between the DeMott 2015 and Niemand 2012 predictions as discussed already regarding Figure 8a appears in this comparison.
Nevertheless, it is seen that both parameterizations grossly underestimate untreated CSU-IS INP concentrations and the treated CSU-IS results fall between the predicted values, agreeing better with the Niemand 2012 parameterization. While one might wish to allude to the fact that the IS filters sample particle sizes, to 10 µm and possible larger that may have higher ice nucleation efficiencies, while the CSU-CFDC was restricted to sampling particles <2.5 µm as a source for the lower DeMott 2015 estimate in comparison to CSU-IS data, we have already addressed that there was no general consistency in INP concentrations for methods that sampled similar size particles overall. The best that can be stated is that the parameterization exercises and treatment data strongly support that inorganic INPs were of weak influence during FIN-03 and that arable soil dusts and biological INPs accounted for the strongest influences during sampling, akin to the findings of Testa et al. (2021).

3.6 Observations of INPs in the deposition nucleation regime

Measurements of deposition nucleation N\textsubscript{INP} are summarized in Figures 13 and 14. FRIDGE-DEP nucleation substrates were collected for 1 to 5 periods on many days during FIN-03 and processed at 5-degree interval temperatures from −15 to −30 °C, and for setpoint humidity of 95% and 99% RH (uncertainties to 2%). Data collected at 102% via the standard FRIDGE methods are not included herein. CSU-CFDC and MIT-SPIN deposition data were collected nominally at 95% RH with an uncertainty of about 2.5% RH, and at a range of temperatures on different days. Mean values and standard deviation error bars of the FRIDGE-DEP data are shown in Figure 13a and median values of FRIDGE-DEP N\textsubscript{INP} (with interquartile values as error bars) are shown in Figure 13b. Standard deviations were large over the course of the study for comprehensive FRIDGE-DEP data when binned at 5-degree interval temperatures. Nevertheless, average concentrations of deposition INPs measured by the FRIDGE-DEP
Figure 13. Summary of deposition-mode N$_{\text{INP}}$ (sL$^{-1}$) as a function of temperature. In a), mean FRIDGE-DEP data at 95% (open orange circles) and 99% (open orange squares) RH are shown along with mean immersion freezing data from the FRIDGE-CS (filled orange circles) and the mean for the few cases of statistically significant CSU-CFDC data (filled purple circle) at 95% RH. Error bars are one standard deviation of the means. In b), median FRIDGE-DEP data are shown and error bars for these are the 95% confidence intervals. The significant CSU-CFDC measurement points at 95% RH are also shown with their 95% confidence intervals. Data measured at 95% RH from the CSU-CFDC and MIT-SPIN that were positively valued but failed significance testing are shown without errors as open purple and open blue circles, respectively.

indicated a consistent 3-5 factor increase between 95 and 99% RH over the range of temperatures investigated. N$_{\text{INP}}$ concentration differences at the two RH values were slightly smaller for median values (Figure 13b), and the median values are slightly lower than the means. Finally, FRIDGE-CS values are plotted in each panel of Figure 13, indicating that FRIDGE-DEP N$_{\text{INP}}$ concentrations averaged for 99% RH are factors 10 to 30 lower than average immersion freezing N$_{\text{INP}}$ concentrations, depending on temperature.
Figure 14. Time series of FRIDGE-CS (immersion freezing) and FRIDGE-DEP (deposition) N\textsubscript{INP} measured at a) -20 °C, and b) -25 °C. Data are from individual filters or wafer collections and error bars are 95% confidence intervals.

One day of significant data was obtained for the CSU-CFDC deposition measurements while using the aerosol concentrator, on September 14, containing three different time periods. These are averaged to create the only online data point represented as a mean in Figure 13a. The individual period measurements from this day, with confidence intervals as errors, are shown for the CSU-CFDC in Figure 13b. Thereby it is seen that these measurements at close to –25 °C agree very well with the mean FRIDGE deposition N\textsubscript{INP} at –25 °C and 95% RH. No measurements of significance were achieved with the MIT-SPIN when operating in the deposition regime. In fact, the most common CSU-CFDC and MIT-SPIN deposition nucleation N\textsubscript{INP} results were below instrument detection limits, not meeting the test for significance, as shown for all periods from 6 common days of such observations represented in Figure 13b. Understanding that these data represent a failure to collect statistically-defensible data, the non-significant data generally scatter about the significant CSU-CFDC data and the FRIDGE-DEP
data at 95% RH, with a higher bias for the MIT-SPIN data. This indicates the difficulty for
online continuous flow instruments to capture low deposition \( N_{\text{INP}} \) concentration data that fall
below 1 sL\(^{-1}\) at most times, considering the FRIDGE-DEP data as the standard. Higher sample
volumes and limited background frost conditions are needed to sense these low atmospheric INP
concentrations.

Time series of the FRIDGE-DEP measurements at –20 °C and –25 °C are shown in
Figure 14. Deposition-mode \( N_{\text{INP}} \) has been averaged over three-hour periods for this analysis.
The FRIDGE immersion freezing data is included in this figure to allow for direct comparison
temporally. Immersion freezing \( N_{\text{INP}} \) generally exceeded deposition-mode \( N_{\text{INP}} \) when both types
of measurements were collected by the two FRIDGE operational methods within the same period
(or during adjacent time periods). This difference ranged from 0 to 2 orders of magnitude, with
the largest differences seen at –25 °C and a period of insignificant differences between the mode
results seen only from the 18\(^{\text{th}}\) to the 22\(^{\text{nd}}\) of September at –20 °C (Figure 14a).

Based on these FRIDGE-CS and FRIDGE-DEP results, immersion-mode ice nucleation
dominates at most times at mixed-phase cloud temperatures. Nevertheless, deposition-mode ice
nucleation contributes modestly to the pool of INP at mixed-phase cloud temperatures in the
atmosphere, and thus may bear consideration for parameterization in atmospheric models. The
ability of online ice nucleation instruments to measure \( N_{\text{INP}} \) in the deposition mode in
correspondence to offline measurements has not been confirmed due to the mentioned inability
of the online instruments used in FIN-03 to capture the low deposition nucleation \( N_{\text{INP}} \)
concentrations. More work should be carried out on measurements of INPs in the deposition
mode to understand variabilities in time and their relation to INP size and composition, as well as
to resolve if online measurements can be improved. For the time being, the substrate methods
appear to be recommended for ambient atmospheric measurements in the realm below water saturation at mixed-phase cloud temperatures.

4. Summary and conclusions

FIN-03 was an ice nucleation instrument intercomparison conducted in the challenging environment of the high-altitude mountaintop field setting. Two online systems (CSU-CFDC, MIT-SPIN) and three offline systems (FRIDGE, CSU-IS, NC State-CS) were represented in FIN-03. The immersion freezing INP concentrations measured in FIN-03 spanned a dynamic range of over five orders of magnitude ($10^{-3}$ to $\sim 10^2 \text{ L}^{-1}$) over the temperature range from $-35 \degree C$ to $-5 \degree C$. Agreement within one order of magnitude in immersion freezing $N_{\text{INP}}$ was generally observed between all ice nucleation instruments measuring immersion INP concentrations at any given temperature if measurement and sampling times were matched to within 3 hours. Better than one order of magnitude agreement was found at temperatures lower than $-25 \degree C$ and higher than $-18 \degree C$, with occasional deviations larger than an order of magnitude in the temperature range from $-25 \degree C$ to $-18 \degree C$. We do not have a full understanding of what controls better or worse agreement at different times or different temperatures, though some factors have been previously discussed in documenting FIN-02 laboratory studies (DeMott et al., 2018). In this study, there was some inference that the different filters and impinger used did not equally capture particles in all size ranges, which is something to improve on in future studies. Given the constant changes in the concentration, size distribution and composition of the ambient aerosol population, inevitable with any field campaign, this level of agreement represents state-of-the-art, at least as judged based on recent laboratory and other field comparisons using similar instrumentation (e.g., Knopf et al., 2021; Lacher et al., 2024).
Although FIN-03 was not conducted as a closure study per se, ancillary data on aerosol sizes and compositions as recommended in more recent discussions of needs for true closure exercises (Knopf et al., 2021; Burrows et al., 2022) were purposefully collected for integration into analyses. This included explicit measurements of the aerosol size distribution, and single particle measurements of aerosol chemical and biological composition. These measurements allowed inferences to be made about INP compositions that provide context for the period of study and establish an example for future intercomparison and long-term measurement efforts.

Through comparing INP data to some current parameterizations describing biological, mineral and soil dust INPs, and additional direct investigations of INP composition via certain pre-treatments to remove biological and organic immersion-freezing INPs, these investigations revealed ubiquitous biological and organic-influenced soil-dust-like INP influences that mimic those found over other continental regions (Knopf et al., 2021; Testa et al., 2021; Lacher et al., 2024). Biological INPs were indicated via selected immersion freezing heat treatments to be dominant at $\sim -20^\circ{\text{C}}$, although of potential influence at all mixed-phase temperatures.

Prediction of these based on parameterizations that utilize single particle fluorescence data (Tobo et al., 2013; Wright et al., 2014; Cornwell et al., 2023) suggest the average utility of such parameterizations but these were unable to predict the full temporal variation of biological INPs. This suggests that local variations of these INPs, which may in fact represent multiple biological particle types, is an area that requires more effort. Based on relatively good consistency between predicted and measured mineral influences on immersion-freezing $N_{\text{INP}}$ concentrations, strictly mineral or other inorganic components of INPs were suggested to have a modest contribution to total INP concentrations at most times and at the freezing temperatures probed during this study.

As in most prior studies, the mineral influence became stronger at the lowest temperatures.
assessed. In contrast, it was found by comparison to a parameterization based on proximally regional soil particles that arable soil INPs likely explained the second most important contribution of INPs during FIN-03, those emanating from other organic particle components that may have been internally mixed with minerals. Biomass burning influences were possible but appear to have not contributed greatly to the climatology of INPs during the study. It was critically important in arriving at these conclusions to have single particle aerosol composition data, from a mass spectrometer that could discern the sizes and fractional contribution of minerals and from a laser-based single particle fluorescence measurement to estimate the biological character of particles. Nevertheless, a great amount of work is still needed to generally parameterize the mixed INP populations that may occur temporally in the atmosphere at higher altitude sites like SPL, or anywhere for that matter.

Importantly, FIN-03 included an assessment of the separate relative contributions of deposition and immersion freezing INP concentrations, one of the few existing data sets of this kind. The offline FRIDGE-DEP method was used to acquire comprehensive deposition $N_{\text{INP}}$ measurements in dependence on RH (95 and 99%), while the CSU-CFDC and MIT-SPIN instruments attempted focused deposition nucleation measurements at (nominally) 95% RH on several days. As expected, FRIDGE-DEP measurements indicated factor of a few increases in deposition $N_{\text{INP}}$ concentrations between 95 to 99% RH. Also, deposition $N_{\text{INP}}$ concentrations were nearly always lower than immersion freezing $N_{\text{INP}}$ concentrations. Deposition INP concentrations at most times at 99% RH (always at 95% RH) were lower by an order of magnitude than immersion freezing INP concentrations at $-20\,^\circ C$ and by more than an order of magnitude at $-25\,^\circ C$. For the online instruments, only limited periods of deposition INP measurements on one day achieved statistical significance from the CSU-CFDC data. While
these data were in good agreement with FRIDGE-DEP data at –25 °C and 95% RH, the most striking result was that all other measurement periods for the CSU-CFDC and MIT-SPIN gave measurements that were not significant at the 95% confidence level. Thus, currently, offline methods for measuring deposition INPs appear to offer the best chance for success in measuring the lower concentrations of INPs that activate below water saturation in the mixed-phase temperature regime. It would be useful to make such assessments at a variety of sites to confirm measurements made during FIN-03 on the relative contributions and variability of INPs active in these conditions toward ice formation in clouds. Additional instrument developments for online measurements of these, and future intercomparisons, will be useful.

In summary, the relative agreements amongst instruments during FIN-03 that match those found in the FIN-02 laboratory studies are encouraging and represent steady improvement in the community’s collective ability to detect and quantify atmospheric ice nucleation. There was not a clear divide between the ability of online and offline systems to measure INP concentrations from the data collected in this study, although the need to carefully consider aerosol sampling efficiencies for different instruments was highlighted as a potential issue in this study, and one requiring close attention in future studies. In principle, both types of instruments show excellent promise for future field studies. For full closure studies of ice nucleation by atmospheric aerosols, methods for identifying INP composition as demonstrated herein and recommended by other recent discussions in Knopf et al. (2021) and Burrows et al. (2022) are critical for understanding and improving INP measurements overall.
Data availability

All data used for the figures in this paper can be accessed at https://radar.kit.edu/radar/en/dataset/eGhfvcOhsOyADZXN (persistent doi:10.35097/eGhfvcOhsOyADZXN)

Author contributions

Paul J. DeMott, Jessica A. Mirrielees and Sarah D. Brooks wrote the paper with assistance from all teams and authors contributing information on instrument descriptions and comments on all results and conclusions, with contributions from Jake Zenker on some data analysis. Paul J. DeMott, Ezra J.T. Levin, Thea Schiebel, Kaitlyn Suski, and Tom Hill provided data and analyses from the CSU-CFDC and IS instruments. Daniel J. Cziczo, Martin J. Wolfe, Sarvesh Garimella, and Maria Zawadowicz provided MIT-SPIN team measurements and analyses. Markus D. Petters and Sarah S. Petters provided data and analysis for the NC State-CS instrument. Heinz G. Bingemer, Jann Schrod, and Daniel Weber provided data and analyses for the FRIDGE instrument. Anne Perring provided data and analyses for the WIBS-4A. Karl Froyd provided data and analyses for the LAS and PALMS. Anna Gannet Hallar and Ian McCubbin oversaw field operations, coordinated with visiting teams at Storm Peak Laboratory, and provided nephelometer and meteorological measurements. Paul J. DeMott, Daniel J. Cziczo, Ottmar Möhler contributed to organize the campaign in connection with the other FIN activities.

Competing interests

The contact author has declared that none of the authors has any competing interests.

Acknowledgements
Partial financial support for this project was provided by the U.S. National Science Foundation, Grant No. AGS-1339264 and U.S. Department of Energy’s Atmospheric System Research, an Office of Science, Office of Biological and Environmental Research program, under grant no. DE-SC0014487. Paul J. DeMott, Ezra J.T. Levin, Thea Schiebel, Kaitlyn Suski, and Tom Hill acknowledge partial and in-kind research support during FIN-03 from NSF grant no. AGS-1358495. Markus Petters acknowledges partial and in-kind support during FIN-03 from NSF grant no. AGS-1450690. Jann Schrod acknowledges research support from the European Union’s Seventh Framework Programme (FP7/2007-2013) project BACCHUS under grant agreement no. 603445. Heinz G. Bingemer and Daniel Weber acknowledge research support under DFG grant BI 462/3-2. Thea Schiebel and Ottmar Möhler received support through the German Science Foundation Projects INUIT and INUIT-2 (MO 668/4-1 and MO 668/4-2). Anne Perring acknowledges support from the NOAA Health of the Atmosphere Program and the NOAA Atmospheric Composition and Climate Program. Special thanks to Romy Fösig (Ullrich) for assistance with data archival.
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https://doi.org/10.5194/egusphere-2024-1744
Preprint. Discussion started: 21 June 2024
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