Dear Ryan,

Thanks again for your input. We have added the Boydoun et al. (2017) and Polen et al. (2018) references to sections 3.2 and 2.2, respectively. Please see the tracked changes below.

Kind regards,

Jessie

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Using freezing spectra characteristics to identify ice nucleating particle populations during the winter in the Alps

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Abstract. One of the least understood cloud processes is modulation of their microphysics by aerosols, specifically of cloud ice by ice nucleating particles (INPs). To investigate INP impacts on cloud ice and subsequent precipitation formation, measurements in cloud environments are necessary but difficult given the logistical challenges associated with airborne measurements and separating interstitial aerosol from cloud residues. Additionally, determining the sources of INPs is important given the dependency of glaciation temperatures on the mineral or biological components and diversity of such INP populations. Here, we present results from a comparison of INP spectral characteristics in air, cloud rime, and fresh fallen snow at the High Altitude Research Station, Jungfraujoch. The goal of the study was two-fold: (1) to assess variability in wintertime INP populations found in-cloud based on wind and air mass direction during snowfall and (2) to evaluate possible INP sources between different sample types using a combination of cumulative INP ($K(T)$) and differential INP ($k(T)$) spectra. INP freezing temperatures and concentrations were consistently higher on average from the southeast as compared to the northwest for rime, snow and especially aerosol samples which is likely a result of air mass influence from predominantly boundary layer terrestrial and marine sources in Southern Europe, the Mediterranean, and North Africa. For all three sample types combined, average onset freezing temperatures were −8.0 and −11.3 °C for southeasterly and northwesterly days, respectively, while $K(T)$ were 3 to 20 times higher when winds arrived from the southeast. Southeasterly aerosol samples typically had a clear mode in the warm temperature regime (i.e., ≥−15 °C) in the $k(T)$ spectra—indicating a putative influence from biological sources—while the presence of a warm mode in the rime and snow varied. Evaluating $K(T)$ concert with $k(T)$ spectra exhibited variable modality and shape—depending on the types of INPs present—and may serve as a useful method for comparing different sampled substances and assessing the possible relative contributions of mixed mineral and biological versus only biological INP sample populations.

1 Introduction

Aerosols are key players in the atmospheric radiation budget, cloud microphysics, and precipitation development. Aerosol-induced ice microphysical modifications influence cloud lifetime and albedo (Albrecht, 1989; Twomey, 1977; Storelvmo et al., 2011), as
well as the production of precipitation (DeMott et al., 2010). Mixed-phase clouds (MPCs) are ubiquitous in the troposphere over the entire annual cycle yet are difficult to quantify globally in part due to an inadequate understanding of aerosol-cloud interactions in mixed-phase environments (Korolev et al., 2017). Thus, a close evaluation of aerosol-cloud processes is crucial to evaluating weather and climate processes. However, one of the most significant challenges with regard to aerosols is quantifying their impacts on cloud ice formation through serving as ice nucleating particles (INPs) (Boucher et al., 2013). Constraining aerosol-cloud impacts in models, specifically when parameterizing INPs in MPC systems, remains a significant challenge due to limited observations (Cziczo et al., 2017; Coluzza et al., 2017; DeMott et al., 2010; Kanji et al., 2017; Korolev et al., 2017). Observations directly in-cloud are even more scarce—given the logistical costs and resources required by airborne platforms, caveats associated with aircraft probes and instrumentation, and instrumental artefacts caused by flying through clouds at high speeds (Cziczo et al., 2017)—but are useful for assessing the impacts of INPs on MPC microphysics as compared to most surface measurements which are geared towards evaluation of INP sources.

In the absence of conditions with $-38 ^\circ C$ and relative humidity with respect to ice above 140%, INPs are required for initiation of tropospheric cloud ice formation (Kanji et al., 2017). Aerosols such as dust and primary biological aerosol particles (PBAPs) are some of the most abundant and efficient INPs found in the atmosphere, respectively (Murray et al., 2012; Hoose and Möhler, 2012; DeMott et al., 1999; Conen et al., 2011; Creamean et al., 2013). PBAPs originating from certain bacteria, pollens, and vegetative detritus are the most efficient INPs known, capable of initiating freezing near $-1 ^\circ C$, while most PBAPs (e.g., fungal spores, algae, and diatoms) tend to nucleate ice at temperatures similar to those of mineral dust (Despres et al., 2012; Murray et al., 2012; Tobo et al., 2014; Hader et al., 2014a; O'Sullivan et al., 2014; Hill et al., 2016; Tesson et al., 2016; Alpert et al., 2011; Knopf et al., 2010; Fröhlich-Nowoisky et al., 2015). In general, previous works collectively indicate that PBAP INPs that nucleate ice at temperatures greater than approximately $-10 ^\circ C$ are bacterial (Murray et al., 2012; Hu et al., 2018; Hoose and Möhler, 2012; Despres et al., 2012; Fröhlich-Nowoisky et al., 2016), but could also be pollen or certain fungal spores (von Blohn et al., 2005; Hoose and Möhler, 2012; O'Sullivan et al., 2016), although the latter two are less likely. Plant bacteria such as *Pseudomonas syringae* are deemed omnipresent in the atmosphere and precipitation (Despres et al., 2012; Stopelli et al., 2017; Morris et al., 2014), and facilitate cloud ice formation up to $-1 ^\circ C$ (Despres et al., 2012). Only a few laboratory-based studies have reported known inorganic or mineral materials with ice nucleation activity at such temperatures (Ganguly et al., 2018; Atkinson et al., 2013). Mineral and soil dust serving as atmospheric shuttles for organic microbial fragments can be transported thousands of kilometres and serve as effective INPs, even from highly arid regions such as the Sahara (Kellogg and Griffin, 2006). The origin of the ice nucleation germ forming at the warmest temperatures is thought to be due to the ice binding proteins or macromolecules of the biological components in mixed mineral-biological INPs (O'Sullivan et al., 2014; O'Sullivan et al., 2016; Conen and Yakutin, 2018). In general, the previous studies on the climate relevance of PBAPs demonstrate the importance of such INPs at MPC temperatures and precipitation enhancement (Morris et al., 2004; Bergeron, 1935; Christner et al., 2008; Morris et al., 2014; Morris et al., 2017; Stopelli et al., 2014; Fröhlich-Nowoisky et al., 2016).

Although biological constituents, from cellular material to intact bacteria and spores, are thought to be omnipresent in the atmosphere (Burrows et al., 2009b; Burrows et al., 2009a; Jaenicke, 2005; Jaenicke et al., 2007), modeling studies constraining global emission estimates of biological INPs and PBAPs are very limited, subject to significant hurdles, and often yield conflicting results due to not having a sufficient set of observations and complexity of atmospheric PBAPs (Hummel et al., 2015; Burrows et al., 2013; Twohy et al., 2016; Fröhlich-Nowoisky et al., 2012; Despres et al., 2012; Hoose and Möhler, 2012; Morris et al., 2011). Yet, biological aerosols such as bacteria have been shown to cause significant perturbations in cloud ice in numerical weather prediction models, affording modulations in cloud radiative forcing and precipitation formation (Sahyoun et al., 2017). In addition,
measuring and quantifying PBAPs is non-trivial—methodologies for counting, culturing, and nucleic acid sequencing of PBAPs and especially for those which fall in the warm temperature INP regime (i.e., INPs that nucleate ice > −15 °C) are: (1) time and labor intensive, (2) require specific expertise or at times substantial resources, (3) require substantial sample volumes, or (4) are species- or genera-specific or limited to viable microorganisms (Despres et al., 2012). Although such techniques are required to adequately assess the atmospheric microbiome and PBAP sources, a simpler approach could be applied to evaluate and even quantify warm temperature biological INP populations as compared to colder temperature PBAPs or mineral dust.

The objectives of the study presented here are: (1) to conduct an comparison of INP measurements of aerosol, cloud rime, and snow directly in-cloud and (2) evaluate different types of INP spectra in a manner such that we can estimate the relative contribution from biological INPs in the warm temperature regime relevant to MPCs. A secondary goal is to assess the variability in INP spectra and possible sources from different dominant air mass transport directions observed during sample collection under the current study. Sampling was conducted at the High Altitude Research Station Jungfraujoch (JFJ), a unique location for evaluating populations of INPs that affect winter storms in the European Alps, and where MPCs are particularly common (Lohmann et al., 2016). Recent studies at JFJ have provided valuable insight into INP concentrations, sources, and removal processes under a variety of conditions and during various times of the year. Conen et al. (2015) measured INPs at −8 °C over the course of a year at JFJ, and found a strong seasonality in such INPs, with two order of magnitude higher concentrations observed during the summer. They also suggested INPs measured at this temperature may be limited most of the year by microphysical processing during transit. Stopelli et al. (2015) verified this removal mechanistic process through INP measurements and isotopic composition of fresh fallen snow at JFJ, concluding that warm temperature INPs are rapidly depleted by precipitating clouds at lower elevations. Stopelli et al. (2016) expanded their INP analyses to 2-years of data at JFJ, concluding that a high abundance of INPs at −8 °C is to be expected whenever high wind speed coincides with air masses having experienced little or no precipitation prior to sampling, yet a separate study by Stopelli et al. (2017) found that only a small fraction of the INPs were cultivable cells of Pseudomonas syringae. In contrast, Lacher et al. (2018a; 2018b) conducted an interannual synopsis of INP measurements at JFJ and found anthropogenic influence on INP concentrations but only during boundary layer intrusion (BLI) and at relatively cold temperatures (i.e., approximately −30 °C), and higher INP concentrations during Saharan dust events (SDEs) and marine boundary layer air arriving at JFJ. Eriksen Hammer et al. (2018) characterized ice particle residuals and concluded that silica and aluminosilicates were the most important ice particle residuals at JFJ during the mixed-phase cloud events during Jan – Feb 2017, while carbon-rich particles of possible biological origin were a minor contribution.

Here, we demonstrate how variable sources influence INP populations depending on air mass transport and direction, and spectral modality between the rime, snow, and aerosols can help explain the exchange of INPs from air into cloud then into precipitation. Our results expand upon previous studies by evaluating INPs via a combination of aerosol, rime, and snow, and at a temperature range that comprises common biological and mineral INPs.

2 Methods

2.1 Aerosol, cloud rime and snow collection at Jungfraujoch

Collocated collection of snow, cloud rime, and aerosol samples for the Ice Nucleation Characterization in the Alps of Switzerland (INCAS) study took place 15 Feb – 11 Mar 2018 in the Sphinx observatory at JFJ (46.55 °N, 7.98 °E; 3580 m above sea level (m a.s.l.); https://www.hfsjg.ch/en/home/). Snow was collected as described by Stopelli et al. (2015) using a Teflon-coated tin (0.1 m², 8 cm deep) for a duration of 1 – 18 hours, but typically for 1 – 4 hours. Collection quantities and time of collection were...
dependent upon snowfall rates but additionally on winds blowing snow out of the collection pans. Because of this possibility, we cannot determine actual snowfall rates with certainty. Cloud rime was collected using a slotted plexiglass plate placed vertically during snow sample collection (Lacher et al., 2017; Mignani et al., 2018). Sample collection times were at times longer than the duration of in-cloud conditions (see section 2.3) and were dependent on when manually changing the sampling tin and plate was possible. Daily size-resolved aerosol samples were collected using a Davis Rotating-drum Universal-size-cut Monitoring (DRUM) single-jet impactor (DA400, DRUMAir, LLC.) (Cahill et al., 1987; Bukowiecki et al., 2009; Creamean et al., 2018a) from a 1-m long inlet constructed of 6.4-mm inner diameter static-dissipative polyurethane tubing (McMaster-Carr®) leading to outside of the Sphinx and connected to a funnel covered with a loose, perforated plastic bag to prevent rime ice build-up or blowing snow from clogging the inlet. The DRUM collected aerosol particles at four size ranges (0.15 – 0.34, 0.34 – 1.20, 1.20 – 2.96, and 2.96 – >12 µm in diameter) and sampled at 27.7 L min⁻¹ (volumetric flow), equalling 39888 total L of air per sample. Such size ranges cover a wide array of aerosols—particularly those that serve as INPs (DeMott et al., 2010; Fridlind et al., 2012; Mason et al., 2016)—while the large volume of air collected promotes collection of rarer, warm temperature biological INPs, but may represent a lower fraction of overall INP concentrations (Mossop and Thorndike, 1966). Samples were deposited onto 20 x 190 mm strips of petrolatum-coated (100%, Vaseline®) perfluoroalkoxy plastic (PFA, 0.05 mm thick) substrate secured onto the rotating drums (20 mm thick, 60 mm in diameter) in each of the four stages at the rate of 7 mm per day (5 mm of sample streaked onto the PFA followed by 2 mm of blank). It is possible local sources of aerosol, such as tobacco smoke or emissions from touristic infrastructure, were collected by the DRUM (Bukowiecki et al., 2016), but did not likely affect the 2.96 – >12 µm particles which we focus on herein. Intervals in which snow, rime, and aerosol were sampled did not fully overlap during a day because conditions were changing often unpredictably between out-of-cloud and in-cloud conditions, the latter with or without precipitation. At the same time we intended to collect enough material from either component for a robust analysis of warm temperature INPs. Consequently, the combined data of a day integrate over a larger air mass, including clouds and cloud-free regions. Comparing data from snow, rime, and aerosol samples still makes sense as long as wind direction and the influence of planetary boundary layer did not change substantially during a day.

2.2 Ice nucleation measurements

All samples were analysed immediately after collection for INPs using a drop freezing cold plate system described by Creamean et al. (2018b; 2018a). Briefly, snow and cloud rime samples were melted into covered 50-mL glass beakers for analysis, resulting in approximately 10 mL of liquid per sample. Samples were manually shaken prior to analysis. Aerosols deposited onto the PFA were prepared for drop freezing by cutting out each daily sample and placing in a 50-mL glass beaker with 2 mL of molecular biology reagent grade water (Sigma-Aldrich®). Beakers were covered and shaken at 500 rpm for 2 hours (Bowers et al., 2009). In between sampling, beakers were cleaned with isopropanol (99.5%), sonicated with double-distilled water for 30 minutes, and then heated at 150 °C for 30 minutes. Copper discs (76 mm in diameter, 3.2 mm thick) were prepared by sonicating in double-distilled water for 30 minutes, cleaning with isopropanol, then coated with a thin layer of petrolatum (Tobo, 2016; Bowers et al., 2009). Following sample preparation, a sterile, single-use syringe was used to draw 0.25 mL of the suspension and 100 drops were pipetted onto the petrolatum-coated copper disc, creating an array of ~2.5-µL aliquots. Drops were visually inspected for size; however, it is possible not all drops were the same exact volume, which could lead to a small level of uncertainty (Hader et al., 2014b; Bigg, 1953; Langham and Mason, 1958; Creamean et al., 2018b). The copper disc was then placed on a thermoelectric cold plate (Aldrich®) and covered with a transparent plastic dome. Small holes in the side of the dome and copper disc permitted placement of up to four temperature probes using an Omega™ thermometer/data logger (RDXL4SD; 0.1 °C resolution and accuracy of ± (0.4% + 1 °C) for the K sensor types
During the test, the cold plate was cooled at 1 – 10 °C min⁻¹ from room temperature until around –35 °C. Control experiments at various cooling rates within this range show very little discernible dependency of drop freezing on cooling rate (Creamean et al., 2018b), akin to previous works (Wright and Petters, 2013; Vali and Stansbury, 1966).

A +0.33 °C correction factor was added to any temperature herein and an uncertainty of 0.15 °C was added to the probe accuracy uncertainty based on DFCP characterization testing presented in Creamean et al. (2018b), to account for the temperature difference between the measurement (i.e., in the plate centre) and actual drop temperature. Frozen drops were detected visually but recorded through custom software. The software records the time, probe temperature, and cooling rate for every second of the test. When a drop is identified as frozen, a button is clicked on the software graphical user interface so that it records that exact time, probe temperature, and cooling rate of that drop in a separate file. The test continued until all 100 drops were frozen. Each sample was tested three times with 100 new drops for each test. The fraction frozen was calculated from all detected drops frozen combined from the three tests (typically, > 90% of the drops were detected). The results from the triplicate tests were then binned every 0.5 °C to produce one spectrum per sample. Cumulative INP concentrations were calculated using the equation from Vali (1971):

\[ K(T) = -\frac{1}{V_{\text{drop}}} \times \ln[1 - f(T)] \]

Where \( V_{\text{drop}} \) is the volume of each drop and \( f(T) \) is the fraction of drops frozen at temperature \( T \). Aerosol cumulative INP concentrations were corrected for the total volume of air per sample \( (K(T) \times V_{\text{suspension}} / V_{\text{air}}) \) while melted rime/snow residual cumulative INPs were adjusted to the total used during analysis \( (K(T) \times V_{\text{suspension}}) \), where \( V_{\text{suspension}} \) and \( V_{\text{air}} \) represent the total liquid volume analyzed per sample (0.75 mL for the three tests) and total volume of air drawn per sample (39888 L), respectively.

Differential INP spectra—which as the name indicates, correspond to the differential of the cumulative spectra (Vali et al., 2015)—were used early in earlier studies (Vali, 1971; Vali and Stansbury, 1966). Spectra from these previous studies only reached a minimum of –20 °C due to the limitations of background artifacts in the water used at that time. Recent work by Vali (2018) and Polen et al. (2018) revisits the use of differential spectra, expanding to lower temperatures. We employ the calculation for differential INP concentrations from Vali (2018):

\[ k(T) = -\frac{1}{V_{\text{drop}} \times \Delta T} \times \ln \left( 1 - \frac{\Delta N}{N(T)} \right) \]

where \( N \) is the number of unfrozen drops and \( \Delta N \) is the number of freezing events observed between \( T \) and \( (T - \Delta T) \). Differential concentrations were divided by the maximum concentration per sample (i.e., to normalize) then smoothed using a moving average.

### 2.3 Supporting meteorological and source analysis data

Auxiliary surface meteorological observations, including but not limited to hourly mean air temperature measured 2 m above ground level (a.g.l.) (°C), relative humidity measured 2 m a.g.l. (%), scalar wind speed (m s⁻¹) and direction (degrees), and incoming longwave radiation (W m⁻²) were acquired from MeteoSwiss (https://gate.meteoswiss.ch/idaweb/). From the longwave measurements, in-cloud conditions were determined by calculating the sky temperature and comparing to air temperature measured at the station, per the methodology of Herrmann et al. (2015) from a 6-year analysis of JFJ observations. There were no in situ
measurements of cloud presence or extent. For the current work, each hourly measurement was categorized as out-of-cloud or in-cloud based on such calculations and averaged to obtain daily cloud coverage percentage.

Radon \(^{222}\text{Rn}\) concentrations have been continuously measured at JFJ since 2009. Details on the detectors themselves and the measurements can be found in Griffiths et al. (2014). Briefly, 30-minute radon concentrations were measured using a dual-flow-loop two-filter radon detector as described by Chambers et al. (2016). Calibrated radon concentrations were converted from activity concentration at ambient conditions to a quantity which is conserved during an air parcel’s ascent: activity concentration at standard temperature and pressure (0 °C, 1013 hPa), written as Bq m\(^{-3}\) STP (Griffiths et al., 2014). Time periods with BLI were classified as radon concentrations > 2 Bq m\(^{-3}\) (Griffiths et al., 2014). Particle concentrations from approximately 0.3 to > 20 μm in diameter were measured with a 15-channel optical particle sizer (OPS 3300; TSI, Inc.) at a 1-minute time resolution (Bukowiecki et al., 2016). Due to operational complications, OPS data were not collected prior to 23 Feb during INCAS. Air was drawn through a heated total aerosol inlet (25 °C) which, besides aerosol particles, enables hydrometeors with diameters < 40 μm to enter and to evaporate, at wind speeds of 20 m s\(^{-1}\) (Weingartner et al., 1999). SDEs were determined from existing methodology using various aerosol optical properties, but specifically, the Ångström exponent of the single scattering albedo (\(\alpha_{\text{SSA}}\)), which decreases with wavelength during SDEs (Collaud Coen et al., 2004; Bukowiecki et al., 2016). SDEs are automatically detected by the occurrence of negative \(\alpha_{\text{SSA}}\) that last more than four hours. Based on previous work, most of the SDEs do not lead to a detectable increase of the 48-h total suspended particulate matter concentrations at JFJ (Collaud Coen et al., 2004). Additionally, we consider these events probable SDEs, but may have influences from other sources in addition.

Air mass transport analyses were conducted using the HYbrid Single Particle Lagrangian Integrated Trajectory model with the SplitR package for RStudio (https://github.com/rich-iannone/SplitR) (Draxler, 1999; Draxler and Rolph, 2011; Stein et al., 2015). Reanalysis data from the National Centers for Environmental Prediction (NCEP) National Center for Atmospheric Research (NCAR) (2.5° latitude-longitude; 6-hourly; https://www.ready.noaa.gov/gbl_reanalysis.php) were used as the meteorological fields in HYSPLIT simulations. Air mass transport directionality and frontal passages were verified by NCEP/NCAR reanalyses of wind vectors and geopotential height at 600 mb (i.e., approximate pressure at the altitude of JFJ; https://www.esrl.noaa.gov/psd/data/composites/day/). Trajectories were initiated at 10, 500, and 1000 m a.g.l. every 3 hours daily, but only the 500-m trajectories are shown. Trajectories were only simulated for each northwesterly, southeasterly, SDE, and BLI case study day (i.e., Table 1). It is important to note that “northwesterly” is a contribution of north, west, and northwest winds, while “southeasterly” includes south, east, and southeast winds. SDE and BLI days were predominantly (not entirely) southeasterly.

### 3 Results and discussion

#### 3.1 Directional dichotomy of air masses arriving at JFJ during INCAS

Local surface meteorology was variable at JFJ during INCAS, with air temperatures ranging from –27.5 to –4.8 °C (average of –13.7 °C)—temperatures relevant to heterogeneous nucleation of cloud ice—and relative humidity ranging from 18 to 100% (Figure 1a). All days contained some fraction of in-cloud conditions that varied between 12% and 100%, on average. Due to the topography surrounding JFJ, predominant wind directions were northwest followed by southeast, with the fastest winds recorded originating from the southeast (Figure 2). Such conditions are typical for JFJ during the winter (Stopelli et al., 2015; Collaud Coen et al., 2011). Out of the entire study, several days were classified as northwesterly (5 days) or southeasterly (2 days) conditions when a combination of aerosol, cloud rime, and snow samples were collected (i.e., a full 24 hours of northwesterly or a full 24 hours of southeasterly winds during snowfall; Table 1), which are herein focused on as the case study days (indicated by the blue and red
in Figure 1b, respectively). These days were also deemed days with “storm” conditions since clouds and snow were both present at JFJ. There were 4 days that maintained predominantly southerly wind directions as indicated in green in Figure 1b and Table 1 and were characterized as days influenced by SDEs or BLI as discussed herein. Rime and snow were only collected on one of these days, while remaining SDE or BLI cases had only aerosol collected. Aside from 22 Feb (missing data), the remaining days in the study were characterized as FT and did not exhibit influences from warm temperature INPs (see section 3.2 and 3.3).

Most southeasterly case days (and 06 Mar) apart from the SDE days experienced longer residence times in what was likely the boundary layer (i.e., 1000 m or less) compared to northwesterly cases, which is supported by radon data (Figure 1c). Griffiths et al. (2014) determined that radon concentrations > 2 Bq m⁻³ signify BLI, which in the current work was clearly observed on 23 Feb, 27 Feb, 28 Feb, 06 Mar, and 11 Mar case days, indicating samples collected on these days were likely influenced by continental boundary layer sources. Relatively low radon concentrations were observed the remaining case study days, indicating these samples were predominantly affected by free tropospheric (FT) air and thus, lower aerosol concentrations and/or more distant, including marine, sources. Although OPS data were missing until 23 Feb, source information can be gleaned from the available data. For example, 23 Feb had episodic high concentrations of particles (maximum of 9.6 cm⁻³) towards the beginning of the day coincident with the largest spike in radon, with a steady decrease as time transpired, indicating the boundary layer was an ample source of > 0.3 μm particles. A similar episode with the OPS and radon concentrations was observed 27 – 28 Feb, where the highest concentrations of each were observed during the entire study period. Selected days were subject to diurnal winds (not shown), such as 06 Mar, where boundary layer air reached JFJ and a midday maximum in OPS particle concentrations was observed, indicating lower elevations were the dominant source of aerosol. Although, diurnal variations in aerosol from local sources have been shown to not be common in the winter at JFJ (Baltensperger et al., 1997). In contrast, 11 Mar was exposed to boundary layer air based on radon observations, but particle concentrations were low (average of 0.2 cm⁻³ compared to a study average of 3.0 cm⁻³), signifying that although BLI occurred at JFJ, it was not a substantial source of aerosol. These relationships corroborate the ice nucleation observations, as discussed in detail below.

Extending past local conditions, air mass transport 10 days back in time prior to reaching JFJ on case study days was, as expected, dissimilar between northwesterly (Figure 3) and southeasterly/SDE/BLI (Figure 4) conditions. The main distinctions between northwesterly and southeasterly/SDE/BLI days are: (1) northwesterly days originated from farther west, with some days reaching back to North America, while air masses on southeasterly/SDE/BLI days predominantly hovered over land and oceanic sources closer to Europe, (2) southeasterly/SDE/BLI air masses travelled closer to the surface relative to northwesterly days, while northwesterly air masses were typically transported from higher altitudes (i.e., more FT exposure), and (3) aside from 6 Mar (which is discussed in more detail in the following section), northwesterly air masses did not travel over the Mediterranean and northern Africa, whereas the southeasterly/SDE/BLI air masses above JFJ arrived from over such regions within less than 2 days before arriving to JFJ. One obvious inconsistency is that the air mass trajectories on 24 Feb do not indicate transport occurred from Northern Africa even though this day was characterized as an SDE. Collaud Coen et al. (2004) reported that in 71% of all cases they evaluated at JFJ, 10-day back-trajectories were able to reveal the source of Saharan dust and that back trajectories cannot always explain SDEs. Boose et al. (2016) reported similar transport pathways for JFJ during multiple consecutive winters and concluded that marine and Saharan dust served as dominant sources of INPs at −33 °C. Reche et al. (2018) also reported similar pathways and sources for bacteria and viruses, but during the summer in southern Spain. These disparate sources and transport pathways of air support the variability in the ice nucleation observations as discussed in more detail in the following section.
3.2 Variability in INP spectra based on air mass source

Out of the 25 aerosol, 30 rime, and 39 snow samples collected, 7 aerosol, 19 rime, and 23 snow were collected northwesterly or southeasterly storm case study days, while 4 aerosol, 1 rime, and 2 snow were collected on SDE or BLI days (Table 1). Most mixed wind direction days were excluded, as sources from both directions would contribute to the daily aerosol sample. Figure 5 shows the cumulative \( K(T) \) and differential \( k(T) \) INP concentrations from aerosol, snow, and rime samples on the case days. The use of \( k(T) \) demonstrates the presence of 1 – 2 INP populations by having a mode in the warm regime (i.e., warm mode or likely biological) and enables us to intercompare between the different types of samples from different air mass sources at JFJ.

In addition to containing higher concentrations of warm temperature INPs, most southeasterly and SDE/BLI samples contained a clear mode in the warm temperature regime compared to northwesterly samples which typically did not contain such a mode in the \( k(T) \) spectra. This warm mode, or “bump” at temperatures above approximately \(-15^\circ C\) has been observed in a wide range of previous immersion mode ice nucleation studies including but not limited to some of the earliest studies of total aerosol (Vali, 1971), residuals found in hail (Vali and Stansbury, 1966), sea spray aerosol (McCluskey et al., 2017; DeMott et al., 2016), soil samples (Hill et al., 2016), agricultural harvesting emissions (Suski et al., 2018), and in recent reviews of aerosol (Kanji et al., 2017; DeMott et al., 2018) and precipitation (Petters and Wright, 2015) samples. Most previous studies that show spectra with the warm mode typically: (1) report a wide range of freezing temperatures such that it can be observed relative to the steady increase of INPs at colder temperatures or (2) are of samples that include a mixture of biological and mineral or other less efficient INP sources. For example, several previous studies report INP concentrations down to only \(-15^\circ C\) (e.g., Conen and Yakutin, 2018; Hara et al., 2016; Schnell and Vali, 1976; Vali et al., 2014; Wex et al., 2015), namely because the goal was to target efficient, warm-temperature biological INPs. However, the warm mode may not be evident in such studies, given it cannot be visualised next to the drastic increase in INPs with temperatures below \(-15^\circ C\). In contrast, studies conducting INP measurements on known mineral dust samples also are not able to observe the warm mode (e.g., Price et al., 2018; Atkinson et al., 2013; Murray et al., 2012). Recently, Beydoun et al. (2017) demonstrate that mixtures of biological and mineral particles can be resolved through careful analysis based on their controlled experimentation using known Snowmax and illite mineral mixtures. Together, it is apparent that a mixed biological and mineral (or less efficient biological INPs) sample is needed to assess the modal behaviour in the INP spectra. However, we note the tentative nature of characterizing these modes based on the previous body of literature and that more controlled, quantitative experiments of mixed biological-mineral INPs is needed in the future.

Only the largest size range of the aerosol is shown because the remaining size ranges (i.e., < 2.96 μm) were not distinct with respect to wind direction. The fact that size, alone, exhibited directionally-dependent results and that such dependencies were only observed in the coarse mode aerosol indicate: (1) the sources were indeed different between northwesterly, southeasterly, and SDE/BLI transport—supporting the air mass source analyses—and (2) the coarse mode aerosols were likely from a regional source as opposed to long-range transported thousands of kilometres. This is because gravitational settling typically renders transport of coarse particles inefficient especially within the boundary layer (Creamean et al., 2018a; Jaenicke, 1980). Previous work by Collaud Coen et al. (2018) concludes that the local boundary layer infrequently influences JFJ in the winter, supporting the current work (i.e., more FT days (17 of 25 days); Table 1).

Generally, INPs from southeasterly and SDE/BLI days were higher in concentration and more efficient (i.e., were warm temperature INPs that facilitated ice formation > \(-15^\circ C\)) than northwesterly samples. Our results are parallel to those by Stopelli et al. (2016), who also observed higher \( K(T) \) in snow samples collected during southerly conditions at JFJ from Dec 2012 to Oct 2014 (Figure 6a). However, \( K(T) \) reported here were generally higher than those reported by Stopelli et al. (2016), especially for...
the northwesterly samples. Unlike Stopelli et al. (2016), there was no clear correlation between $K(T)$ with air temperature and wind speed in the current work (not shown).

Onset freezing temperatures (i.e., the highest temperature in which the first drop in each sample froze) were typically higher for southeast/SDE/BLI samples as compared to the northwest (Figure 6b), indicating influences from sources that produce warm temperature INPs on these days. The temperatures in which 10% ($T_{10}$) and 50% ($T_{50}$) of the samples froze were also typically higher for the southeast/SDE/BLI as compared to the northwest samples, especially for the aerosol samples, indicating higher concentrations of more efficient warm temperature INPs.

Regardless of the snow, it is possible that surface processes generate airborne ice particles, which contribute to a snow sample collected at a mountain station (Beck et al., 2018). However, snow that is re-suspended during a snowfall event largely consists of the most recently fallen snow crystals covering wind-exposed surfaces. These particles are unlikely to be different from concurrently falling snow. Hence, their contribution will not change INP abundance or spectral properties of the collected sample. Another matter are hoar frost crystals, which can be very abundant in terms of number, but because of their small size (i.e., < 100 µm (Lloyd et al., 2015)) can only make a minor contribution to the mass of solid precipitation depositing in a tin placed horizontally on a mountain crest. The majority of small crystals will follow the streamlines of air passing over the crest. All that an increased influence of hoar frost particles would do to our observations is to decrease measured differences between snow and rime samples, because additions of hoar frost, a form of rime, would render the collected snow sample a bit more similar to rime.

### 3.3 Potential components of INP populations at JFJ

Taking the spectral characteristics in the context of air mass direction and transport can help elucidate the possible sources of INPs at JFJ during INCAS. Quantitative evaluation of the warm mode, or likely, the relative contribution of warm temperature biological INPs, are calculated and shown in Figure 6c – g. Additionally, normalizing such spectra affords a qualitative comparison of spectra signatures between aerosols and residuals found in cloud rime and snow. We offer some possible explanations for the observed variability between the samples. Naturally, the boundary layer more frequently than not contains higher concentrations of warm temperature INPs—and INPs in general—as compared to the free troposphere given the proximity to the sources (e.g., forests, agriculture, vegetation, and the oceans) (Burrows et al., 2013; Despres et al., 2012; Frohlich-Nowoisky et al., 2016; Wilson et al., 2015; Burrows et al., 2009a; Burrows et al., 2009b; Frohlich-Nowoisky et al., 2012; Suski et al., 2018). Although, microorganisms and nanoscale biological fragments are episodically lofted into the free troposphere with mineral dust and transported thousands of kilometres (Creamean et al., 2013; Kellogg and Griffin, 2006).

Air arriving at JFJ on 15 and 16 Feb originated from the farthest away and were not heavily exposed to boundary layer air, as evidenced by the air mass trajectory analysis (Figure 3) and radon (Figure 1c), indicating long-range transport in the free troposphere. This could explain why the warm mode (and higher $T_{10}$ and $T_{50}$) was observed for the rime and snow, but not the aerosol—the aerosol had sufficient time to nucleate ice during free tropospheric transport and especially the warm temperature INPs that would likely become depleted in-cloud first (Stopelli et al., 2015), assuming the clouds formed along the air mass transport pathways. Cloud fraction was relatively low (12.5 to 25%), but air temperatures were relatively high (−8.4 to −7.1 °C), suggesting conditions were amenable for long-range transported warm temperature INPs to nucleate cloud ice. However, from the available data, we cannot determine with certainty if the local conditions were the same as those when nucleation initially occurred. For 19 and 20 Feb, air temperature was cold (−16.4 and −19.6 °C, respectively) cloud fraction was high (92 and 54%, respectively), and not all samples contained a warm mode. One possible explanation is that any warm temperature INPs that were present in the...
clouds had already snowed out prior to reaching the sampling location, as observed by Stopelli et al. (2015) at JFJ. Although, given the low radon concentrations and erratic transport pathways, it is possible such air masses did not contain a relatively large concentrations of warm temperature INPs due to deficient exchange with the boundary layer. It was not until the southeasterly cases that the aerosol samples exhibited a warm mode. Specifically, on 23 Feb local winds shifted to southeasterly (147 degrees on average) and air masses arrived from over the eastern Alps, Eastern Europe, Scandinavia, and earlier on in time, the Atlantic Ocean. Thus, these samples were predominantly influenced by the continental (mostly over remote regions) and marine boundary layers, where sources of warm temperature INPs are more abundant (Frohlich-Nowoisky et al., 2016).

The northwesterly case of 06 Mar is somewhat interesting in that the local wind direction was clearly from the northwest, but air mass source analyses show brief transport in the boundary layer (radon) from the south, when looking farther back in time, traveling over the Mediterranean and North Africa. The aerosol sample had a high onset temperature for INPs relative to other northwest samples (Figure 6b) and snow samples exhibited a warm mode (Figure 6g). It is the only one of the northwesterly case samples that encountered boundary layer exposure according to the radon observations. Combined, these results suggest a somewhat mixed-source sample, and that 06 Mar may not be directly parallel to the other northwesterly cases. Transitioning back to a southeasterly case on 11 Mar, only the rime and snow unveiled a warm mode from air transported from similar regions as the 06 Mar sample. Additionally, OPS concentrations were very low (Figure 1c). These results suggest the aerosols already nucleated cloud ice prior to reaching JFJ on 11 Mar (i.e., low ambient aerosol), where the aerosol did not contain a warm mode, but rime and snow did.

When evaluating the SDE and BLI days, there is a bit of variability. On 24 Feb, clouds were present at JFJ (a cloud fraction of 37.5%), but riming was insufficient to collect enough quantity for INP analysis and no snowfall occurred. Interestingly, the warm mode was the second highest for the aerosol sample, indicating a relatively large contribution of warm temperature INPs as compared to the other samples from the study. Air mass transport was very similar to 23 Feb signifying similar INP sources even though this day was characterized as an SDE, but it is probable that a slightly warmer (–6.0 as compared to –9.8 °C air temperature), drier (79 versus 89% relative humidity), and higher pressure (649 versus 645 mb) postfrontal system moved over JFJ on 24 Feb, inhibiting removal of warm temperature INPs during transport relative to the day prior (corroborated by reanalysis from NCEP/NCAR of geopotential height at 600 mb). The BLI case of 28 Feb was very similar to 24 Feb in that: (1) only an aerosol sample was collected and (2) the warm mode was much higher in concentration ($k(T)$) than the remaining samples. As compared to 27 Feb where a warm mode was not observed, 28 Feb was warmer (–20.0 as compared to –26.2 °C), drier (52 versus 62%), higher pressure (635 versus 630 mb), and had a warmer onset temperature (–6.8 versus –14.8 °C). Wind direction was slightly different: southeasterly (153 degrees) on 27 Feb as compared to southwesterly on 28 Feb (221 degrees). However, conditions were cloudier than the 23 – 24 Feb coupling and completely cloudy on 27 Feb (100 and 66.7% cloud fraction on 27 Feb and 28 Feb, respectively). Additionally, radon and OPS concentrations were the highest on 27 – 28 Feb as compared to the rest of the days during INCAS (Figure 1c). Combined, these results suggest a very local, boundary layer source of INPs started on 27 Feb, but were quickly depleted in the very cloudy conditions. Once clouds started to clear and a shift in frontal characteristics occurred, a similar source of very efficient warm temperature INPs affected JFJ but were able to be observed in the aerosol.

4 Conclusions and broader implications

Aerosol, cloud rime, and snow samples were collected at the High Altitude Research Station Jungfraujoch during the INCAS field campaign in Feb and Mar 2018. The objectives of the study were to assess variability in wintertime INP sources found in cloudy environments and evaluate relationships between INPs found in the different sample materials. To directly compare air to liquid
samples, characteristics of differential fraction frozen and INP spectra were compared in the context of cumulative INP spectra statistics, air mass transport and exposure to boundary layer or free tropospheric conditions, and local meteorology. Distinction between northwesterly and southeasterly conditions yielded differing results regarding INP efficiency and concentrations, biological versus non-biological sources, and meteorological conditions at the sampling location. In general, cumulative INP concentrations were 3 to 20 times higher for all sample types when sources from the southeast infiltrated JFJ, while the INP spectra of the aerosol contained a warm mode but the presence of a warm mode was variable for the rime and snow depending on meteorological context.

In general, comprehensive measurements of INPs from aerosol, and rime and snow when possible, affords useful information to compare and explain exchange between aerosols, clouds, and precipitation in the context of local and regional scale meteorology and transport conditions. Assessment of different INP spectral types, modality, and spectra statistics adds another dimension for qualitative and semi-quantitative intercomparison of sampling days and evaluation of associations between aerosol, cloud, and precipitation sampling. Extending INP analyses past reporting cumulative concentrations affords more detailed information on the population of INPs and enables comparison between samples from aerosols, clouds, precipitation, and beyond (e.g., seawater, soil, etc.). Using auxiliary measurements and air mass simulations, in addition to context provided by previous work at JFJ, we have addressed possible sources for INCAS. However, more detailed source apportionment work should be imminent to comprehensively characterize INP sources based on spectral features. Future studies should ideally use such statistical analyses in tandem with focused chemical and biological characterization assessments to provide direct linkages between INP spectral properties and sources. Such investigations could yield valuable information on INP sources, and aerosol-cloud-precipitation interactions, which could then be used to improve process-level model parameterizations of such interactions by rendering quantitative information on INP source, efficiency, and abundance. Improving understanding of aerosol impacts on clouds and precipitation will ultimately significantly enhance understanding of the earth system with respect to cloud effects on the surface energy and water budgets to address future concerns of climate change and water availability.

Author contributions

JMC collected the samples, conducted the DFA sample analysis, conducted data analysis, and wrote the manuscript. CM and FC also contributed to collecting rime and snow samples. JMC, CM, and FC designed the experiments. NB provided quality controlled OPS data. CM, NB, and FC helped with manuscript feedback and revision prior to submission.

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do not necessarily reflect the official views of the Swiss Government. We are grateful to the International Foundation High Altitude Research Stations Jungfraujoch and Gornergrat (HFSJG), 3012 Bern, Switzerland, for providing through its infrastructure comfortable access to mixed-phase clouds. Special thanks for go to Joan and Martin Fischer, and Christine and Ruedi Käser, the custodians of the station. The authors gratefully acknowledge the NOAA Air Resources Laboratory (ARL) for the provision of the HYSPLIT transport and dispersion model and/or READY website (http://www.ready.noaa.gov) used in this publication.

References


Draxler, R. R.: HYSPLIT user’s guide, NOAA Air Resources Laboratory, Silver Spring, MD, 1999.


Table 1. Dates and times for cloud rime, snow, and aerosol samples collected during the 2018 winter INCAS study at Jungfraujoch. Also shown are the dominant air mass sources (free troposphere or FT or boundary layer intrusion of BLI) for each day based on radon data. Samples highlighted in blue and red are the northwest and southeast case studies, respectively. Samples highlighted in green represent predominantly southerly days that were classified as Saharan dust events (SDEs) or BLI days.

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Figure 1. Daily averaged meteorological data at JFJ from INCAS, including a) percentage of cloudiness in the vertical profile over JFJ per the estimation by Herrmann et al. (2015), station relative humidity (%), and station air temperature (°C) and b) station wind direction. The background of b) is shaded horizontally by north (light blue) or south (light red) directions. Additionally, days with combined aerosol, cloud rime, and snow collections are vertically shaded grey. Blue and red markers for wind direction represent case study storm days that were entirely northwesterly or southeasterly, respectively. Green markers represent predominantly southerly days that were classified as Saharan dust events (SDEs) or heavy boundary layer influence days. c) Time series of radon (²²²Rn) corrected for standard temperature and pressure and OPS particle concentrations. The black dashed line indicates a threshold of 2 Bq m⁻³ whereby boundary layer intrusion likely occurred at JFJ. OPS data were missing prior to 23 Feb. Error bars represent standard deviation.
Figure 2. Rose plot for wind data during INCAS. Values correspond to wind direction binned by 45 degrees and wind speeds binned by 5 m s\(^{-1}\). The probability for wind speed to fall within these bins is plotted.
Figure 3. 10-day air mass backward trajectories initiated every 3 hours during sample collection for northwesterly case days ending at 500 m above ground level (a.g.l.). Trajectories are plotted by latitude-longitude (left) and altitude-time (right) profiles for 15 Feb (a, b), 16 Feb (c, d), 19 Feb (e, f), 20 Feb (g, h), and 06 Mar (i, j). Darker shades of blue represent trajectory points farther back in time.
Figure 4. Same as Figure 3, but for southeasterly (23 Feb and 11 Mar), SDE (24 Feb and 10 Mar), and BLI (27 and 28 Feb) case days.
Figure 5. Cumulative INP spectra ($K(T)$, on left) and differential INP spectra ($k(T)$, on right) for the same samples of a) cloud rime, b) snow, and c) aerosol for the size range 2.96 – >12 µm in diameter. Spectra shown are for samples from the northwest (blue) and southeast (red) case study dates, in addition to SDE and boundary layer intrusion (BLI) case days (green). Multiple cloud rime and snow samples were collected while one aerosol sample from each size range was collected on northwest and southeast case study days (see Table 1). The warm mode region is indicated by the dark grey shading in the $k(T)$ spectra. Note the axes are not all the same ranges.
Figure 6. a) Comparison of INCAS cumulative snow INP concentrations for northwest (blue) and southeast (red) within the same range of those reported by Stopelli et al. (2016) for measurements at JFJ during the 2012/13 winter season. Summary of INCAS INP concentrations from aerosol (squares), cloud rime (circles), and snow samples (diamonds), including b) freezing onset temperatures, c) the temperature in which 10% of drops froze, and d) the temperature in which 50% of the drops froze calculated from fraction frozen. From the $k(T)$ spectra, warm mode temperatures are shown for e) aerosol, f) rime, and g) snow samples. Blue, red, and green markers represent northwesterly cases, southeasterly cases, and SDE and BL I case days, respectively. Some data points overlap and thus plots may appear to not have the same number of points per sample.