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Analysis of isothermal and cooling rate dependent immersion freezing by a unifying stochastic ice nucleation model

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Abstract. Immersion freezing is an important ice nucleation pathway involved in the formation of cirrus and mixed-phase clouds. Laboratory immersion freezing experiments are necessary to determine the range in temperature (T) and relative humidity (RH) at which ice nucleation occurs and to quantify the associated nucleation kinetics. Typically, isothermal (applying a constant temperature)

- 5 and cooling rate dependent immersion freezing experiments are conducted. In these experiments it is usually assumed that the droplets containing ice nucleating particles (INPs) all have the same INP surface area (ISA), however the validity of this assumption or the impact it may have on analysis and interpretation of the experimental data is rarely questioned. Descriptions of ice active sites and variability of contact angles have been successfully formulated to describe ice nucleation experimental
- 10 data in previous research, however, we consider the ability of a stochastic freezing model founded on classical nucleation theory to reproduce previous results and to explain experimental uncertainties and data scatter. A stochastic immersion freezing model based on first principles of statistics is presented, which accounts for variable ISA per droplet and uses parameters including the total number of droplets (N_{tot}) and the heterogeneous ice nucleation rate coefficient, $J_{het}(T)$. This model is
- 15 applied to address if (i) a time and ISA dependent stochastic immersion freezing process can explain laboratory immersion freezing data for different experimental methods and (ii) the assumption that all droplets contain identical ISA is a valid conjecture with subsequent consequences for analysis and interpretation of immersion freezing.

The simple stochastic model can reproduce the observed time and surface area dependence in 20 immersion freezing experiments for a variety of methods such as: droplets on a cold-stage exposed to air or surrounded by an oil matrix, wind and acoustically levitated droplets, droplets in a continuous flow diffusion chamber (CFDC), the Leipzig aerosol cloud interaction simulator (LACIS), and the aerosol interaction and dynamics in the atmosphere (AIDA) cloud chamber. Observed time dependent isothermal frozen fractions exhibiting non-exponential behavior can be readily explained

- by this model considering varying ISA. An apparent cooling rate dependence of J_{het} is explained by 25 assuming identical ISA in each droplet. When accounting for ISA variability, the cooling rate dependence of ice nucleation kinetics vanishes as expected from classical nucleation theory. The model simulations allow for a quantitative experimental uncertainty analysis for parameters N_{tot} , T, RH, and the ISA variability. The implications of our results for experimental analysis and interpretation of the immersion freezing process are discussed.
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1 Introduction

Ice crystals in tropospheric clouds form at altitudes where temperatures fall below the ice melting point, also known as supercooled temperatures, and for conditions in which water partial pressure exceeds the saturation vapor pressure with respect to ice (Pruppacher and Klett, 1997; Hegg and Baker,

- 2009). Cirrus or mixed-phase clouds consist entirely of ice crystals or of ice crystals coexisting 35 with supercooled aqueous droplets, respectively. These clouds can significantly impact the global radiative budget and the hydrological cycle (Baker, 1997; Rossow and Schiffer, 1999; Chen et al., 2000; Liu et al., 2007; Lohmann and Hoose, 2009; Tao et al., 2012; Rosenfeld et al., 2014), however, their formation is not well understood or constrained in cloud and climate models (Boucher et al.,
- 40 2013). Ice nucleation precedes the formation of ice crystals. Homogeneous ice nucleation occurs from supercooled aqueous aerosol particles or cloud droplets. Ice formation can also occur at temperatures higher than the homogeneous freezing limit initiated by insoluble particles acting as ice nucleating particles (INPs). Heterogeneous ice nucleation can occur when INPs are immersed in supercooled aqueous droplets, termed immersion freezing, when INPs make physical contact with
- supercooled droplets, termed contact freezing, or when ice nucleates on INPs directly from the su-45 persaturated vapor phase, termed deposition ice nucleation. It is impossible to observe in situ ice nucleation in the atmosphere and very difficult to infer the ice nucleation pathway (Haag et al., 2003; Hegg and Baker, 2009). Despite the established importance of the impact of heterogeneous ice nucleation on cirrus and mixed-phase cloud formation, it is not included in global radiative forcing
- 50 estimates (Myhre et al., 2013).

Laboratory studies are necessary to investigate at which thermodynamic conditions, i.e. temperature, T, and relative humidity, RH, and by which mode ice nucleation occurs for predictive use in cloud and climate models. This study presents a newly developed model simulation applied for analyses of previously published laboratory immersion freezing data obtained by different experimental

55 methodologies. It allows prediction of atmospheric ice particle production under relevant scales of time and INP surface area (ISA).

Classical nucleation theory (CNT) is currently the only available physical theory to describe ice nucleation. Simply stated, CNT quantifies a maximum Gibbs free energy barrier corresponding to the minimum number of water molecules in a cluster that has to be overcome to initiate ice nucleation

60 (Pruppacher and Klett, 1997). Cluster formation and thus, ice nucleation, occurs stochastically and is dependent on time, *t*, and in the case of homogeneous ice nucleation, the supercooled liquid volume, *V*. Koop et al. (2000) parameterized the theoretical homogeneous ice nucleation rate coefficient, *J*_{hom}, as a function of *T* and water activity, *a*_w (*a*_w = 1.0 for pure water and *a*_w < 1.0 for aqueous solution). This approach yields *J*_{hom} to be independent of the nature of the solute and avoids the
65 weakness of the capillary approximation in CNT (Pruppacher and Klett, 1997).

Immersion freezing can be described by CNT by reducing the free energy barrier due to the presence of a solid surface. Ice nucleation remains a stochastic process, but is dependent on the available ice nucleating surface area, A, instead of V (Pruppacher and Klett, 1997; Zobrist et al., 2007). The heterogeneous ice nucleation rate coefficient, J_{het} , is a physically and experimentally defined param-

- eter which gives the rate of nucleation events for given surface area and unit time. Knopf and Alpert (2013) parameterized J_{het} as a function of T and a_w following Koop et al. (2000) using direct measurements of J_{het} and J_{het} derived from previous studies (Archuleta et al., 2005; Alpert et al., 2011a, b; Knopf and Forrester, 2011; Murray et al., 2011; Broadley et al., 2012; Iannone et al., 2011; Pinti et al., 2012; Rigg et al., 2013). Known as the a_w based immersion freezing model (ABIFM) (Knopf and Alpert,
- 75 2013), J_{het} can be derived for different types of INPs such as mineral dusts, organic, surfactant and biogenic, applicable for $a_{\text{w}} \leq 1.0$, and independent of the nature of the solute. The ABIMF is a holistic and computationally efficient description of the immersion freezing process for prediction of ice nucleation for atmospherically relevant conditions and applicable for a variety of experimental methods, including the droplet-on-substrate approach (Zobrist et al., 2007; Knopf and Forrester, 2011;
- Alpert et al., 2011a, b; Iannone et al., 2011; Murray et al., 2011; Broadley et al., 2012; Rigg et al., 2013), oil-encased droplets (Murray et al., 2011; Broadley et al., 2012; Wright and Petters, 2013), differential scanning calorimetry (Marcolli et al., 2007; Pinti et al., 2012), and continuous flow diffusion (Rogers et al., 2001; Archuleta et al., 2005; Hartmann et al., 2011; Kulkarni et al., 2012; Wex et al., 2014). These previous studies represent a subset of a much broader selection of experimental meth-

85 ods and designs.

Different parameterizations of J_{het} exist. Zobrist et al. (2007) investigated droplet freezing initiated by organic surfactant monolayers and parameterized experimentally derived J_{het} values using parameterizations of the Gibbs free energy and diffusion activation energy. Reduction in the Gibbs free energy barrier is described by the parameter known as the contact angle, α , which is defined as

90 the angle of contact between an ice embryo and substrate surrounded by the liquid parent phase and derived by the balance between interfacial surface tension between the three (Pruppacher and Klett, 1997). It was found that a single value of α could not reproduce the experimental freezing data for organic monolayers, but when allowing α to be a linear function of T the data could be represented (Zobrist et al., 2007), corroborated by others and for different INPs (Knopf and Forrester,

- 95 2011; Alpert et al., 2011a, b; Welti et al., 2012; Rigg et al., 2013). It is important to note that a self assembled organic monolayer is a completely uniform surface down to the molecular level, i.e. the notion of different ice active sites present does not apply (Gavish et al., 1990; Popovitz-Biro et al., 1994; Majewski et al., 1995). For particles that have an uneven morphology, cracks, pits, or ridges such as mineral dust, a single α value has been shown also to not reproduce experimental data (e.g.
- 100 Marcolli et al., 2007; Lüönd et al., 2010; Niedermeier et al., 2011b; Welti et al., 2012; Rigg et al., 2013; Niedermeier et al., 2014; Wheeler et al., 2015). Unlike Zobrist et al. (2007), some of these studies do not consider that α can vary with T, but instead randomly distribute α values on particles immersed in droplets while using all the other same parameterizations (i.e. the Gibbs free energy and diffusion activation energy) to calculate J_{het} and not experimentally derive J_{het} . This procedure is
- 105 similar to distributing J_{het} values over different droplets containing INPs as done by Broadley et al. (2012) and Herbert et al. (2014). These are successful approaches to describe the freezing data leading one to think that allowing α to be a linear function of T or distributing α across particles are in principle the same, both resulting in the necessity to change the contact angles to represent the freezing data. However, application of these frameworks are conceptually and mathematically very
- 110 different and in fact result in very different interpretations of the data and underlying ice nucleation processes. The method of Zobrist et al. (2007) derives a single continuous function of $J_{het}(T)$ for a specific particle type, while an active site approach randomly distributes multiple $J_{het}(T)$ functions across particles and their surfaces. In other words, one can either use a single $J_{het}(T)$ function (i.e. not applying a single α value) or use multiple $J_{het}(T)$ functions (different only by utilizing different
- 115 values of α), constituting completely different pictures of ice nucleation. One major advantage of the ABIFM approach chosen here is that it uses a single function of $J_{het}(T)$ and avoids any use or calculation of α and instead uses water activity as a parameter. It inherently allows α to be both T dependent (Zobrist et al., 2007) or distributed in a particle population (an active site approach). A caveat to our approach is that it cannot make a statement about the active site distribution. An-
- 120 other advantage of our approach is that ABIFM can be used simultaneously for immersion freezing from pure water and aqueous solution droplets. When using the ABIFM, a uniform ice nucleating surface is not assumed, however, a single function of $J_{het}(T)$ for a single particle type is assumed to describe the experimental data without invoking the presence of different (rare) and non-detectable ice nucleating sites or components present in some but not all droplets.
- The major difficulty with a variety of experimental techniques is how accuracy and uncertainty of *T*, RH, *t*, and *A* are assessed and how these uncertainties affect extrapolation of laboratory derived ice nucleation parameterizations to atmospherically relevant conditions. Previous investigations have developed state of the art instrumentation and methods to constrain uncertainties (Connolly et al., 2009; Lüönd et al., 2010; Niedermeier et al., 2010; DeMott et al., 2010; Niedermeier et al., 2011b;
- 130 Hoose and Möhler, 2012; Niemand et al., 2012; Rigg et al., 2013; Hiranuma et al., 2015; Vali and Snider,

2015). Additionally, progress in understanding ice nucleation behavior has been made by validating empirical parameterizations or models based on the concept of ice active sites, i.e. that surface sites on a particle have variable ice nucleating efficiencies, can be used to reproduce experimental data. However, there is no physical basis for these interpretations (Niedermeier et al., 2010) and

- 135 they may be inherently constrained to the investigated range of *T*, RH, *t*, *A* and concentration of INPs from which they are derived (Rigg et al., 2013; Knopf and Alpert, 2013). These include the multi-component model (Murray et al., 2011), the time-dependent freezing rate parcel model (Vali and Snider, 2015), parameterizations of INPs per liter of air (DeMott et al., 2010), the α -PDF model (Marcolli et al., 2007; Lüönd et al., 2010), the active site model (Marcolli et al., 2007;
- 140 Lüönd et al., 2010), the singular description (Vali, 1971; Connolly et al., 2009; Alpert et al., 2011a, b; Vali, 2008; Murray et al., 2011; Hiranuma et al., 2015) and the soccer ball model (Niedermeier et al., 2011b). According to the singular hypothesis, the number of active sites, $n_s(T)$, is dependent on Tonly and neglects ice nucleation kinetics. We suggest that further analytical efforts regarding ice nucleation kinetics can improve our understanding on the governing parameters of immersion freezing.
- The immersed ISA per droplet is important for experimental derivation of J_{het} and for deriving empirical quantities such as $n_s(T)$ or fitting functions and their parameters. In previous experimental studies, droplets for ice nucleation experiments were dispensed from a bulk solution containing INPs (Broadley et al., 2012; Rigg et al., 2013; Wright and Petters, 2013; Herbert et al., 2014; Diehl et al., 2014). In other investigations, solid particles were size selected by their electrical mobility and then
- 150 injected into, or continuously flown through, an ice nucleation chamber where water condensation precedes ice nucleation (Archuleta et al., 2005; Niedermeier et al., 2010; Kulkarni et al., 2012; Welti et al., 2012; Wex et al., 2014). In these studies and those that used polydisperse aerosol (e.g. Niemand et al., 2012), surface area calculations assumed that particles with the same mobility diameter are spherical with identical surface area. Despite this assumption, advancement in accounting
- 155 for particle size variability considering multiple charged particles in ice nucleation experiments has been made (Lüönd et al., 2010; Augustin-Bauditz. et al., 2014). However, extensive theoretical and experimental literature exists on aerosol sizing instrumentation and morphology characterization, which consider particle density, void fraction, shape and electrical charge effects implying their non-sphericity (DeCarlo et al., 2004; Slowik et al., 2004; Zelenyuk et al., 2006; Schmid et al., 2007;
- 160 Park et al., 2008). In general, neglecting these effects may influence surface area estimates. Surfaces of particles may differ depending on different generation techniques, and previous studies have made progress in understanding why or why not these result in differences in ice nucleation efficiency for particle types such as hematite (Hiranuma et al., 2014), Fluka kaolinite (Lüönd et al., 2010), KGa-2 and KGa-1b type kaolinites (Pinti et al., 2012), NX illite (Pinti et al., 2012; Hiranuma et al., 2012)
- 165 2015), or Arizona Test Dust (ATD) (Marcolli et al., 2007; Niedermeier et al., 2011a). In the present study, we consider laboratory generated particles of ATD (Wright and Petters, 2013), NX illite (Broadley et al., 2012; Diehl et al., 2014), KGa-1b kaolinite and K-feldspar (Herbert et al., 2014),

Fluka type kaolinite (Wex et al., 2014) and natural dust (Niemand et al., 2012) each with their own respective generation method and argue that accurate quantification of ISA is crucial to discriminate

- 170 how surface properties affect ice nucleation efficiency. Frequently, distributions of immersed ISA per droplet are typically assumed to be monodispersed, or in other words, each droplet is assumed to contain identical ISA. Furthermore, the number of droplets applied in an ice nucleation experiment may also affect the significance of the freezing data and thus interpretation of the experiment. It is necessary to question if a potential variability in ISA and/or the assumption of monodisperse ISA
- 175 and a limited number of observed freezing events become important for interpretation of immersion freezing experiments with subsequent ramifications for the analytical ice nucleation description.

We introduce a newly developed model simulation in which ice nucleation is treated explicitly as a stochastic process applicable for isothermal and cooling rate experiments. Previous experimental results using different experimental methods are simulated and compared for a wide range of

- atmospherically relevant conditions. However, this analysis is applied to laboratory generated particles only and may not be applicable to field or natural samples because of the difficulty to separate INPs from others. Sensitivity studies on frozen fraction data and experimentally derived J_{het} are performed as a function of ISA assumptions, the number of droplets employed in the experiment, T, and RH. The validity of typical assumptions of ISA variability and uncertainty are tested. Then,
- 185 a detailed analysis of the ability of the model simulation to reproduce experimental results with strict uncertainty estimation is presented for 7 independent immersion freezing studies utilizing 8 different instrumentation: (i) droplets on a cold-stage exposed to air, (ii) droplets on a cold-stage covered in oil, (iii) oil-droplet emulsions, (iv) droplet acoustic levitation, (v) droplet wind tunnel levitation, (vi) the Leipzig aerosol cloud interaction simulator (LACIS), (vii) a continuous-flow diffusion chamber
- 190 (CFDC) and (viii) the aerosol interaction and dynamics in the atmosphere (AIDA) cloud chamber. A rigorous uncertainty analysis of the ice nucleation kinetics for typical ranges in experimental conditions is presented and discussed for laboratory application.

2 Immersion freezing model based on classical nucleation theory

2.1 Simulation of isothermal freezing experiments

- 195 Stochastic immersion freezing simulations (IFSs) are performed to evaluate the effect of variable ISA on droplet immersion freezing experiments conducted in the laboratory. As discussed above, different droplets in a laboratory experiment will possess different ISA. To account for this fact, ISA in each simulated droplet is sampled from a distribution to mimic this variability. Surface area can be any real positive value and can change by orders of magnitude. For this reason, a lognormal
- 200 distribution can be assumed with the most probable ISA being A_g or a mean distribution parameter $\mu = \ln(A_g)$. The distribution width parameter is $\sigma = \ln(\sigma_g)$, where σ_g represents the factor by which ISA can vary. A different distribution can also be assumed with knowledge of experimental meth-

ods used in particle generation, e.g. assuming a bipolar charge distribution for electrical mobility diameter selected particles. Knowledge of ISA for each droplet can be directly used as an alterna-

205 tive without a need for random sampling. Droplet freezing for isothermal experiments can then be described by

$$\delta N_{\rm ufz} = -J_{\rm het} A_{\rm tot} \delta t \,, \tag{1}$$

where $\delta N_{\rm ufz}$ represents the change in the number of unfrozen droplets after a certain interval of time, δt , and J_{het} is the heterogeneous ice nucleation rate coefficient. The total available ISA is $A_{\text{tot}} =$ 210 $\sum A_j$, where A_j is the ISA in the *j*th droplet. An assumption typically made is that all droplets contain the same ISA, or $A_{\text{tot}} = A_{\text{g}} N_{\text{ufz}}$, where A_{g} is the ISA for all droplets (e.g. Marcolli et al., 2007; Lüönd et al., 2010; Niedermeier et al., 2010; Murray et al., 2011; Rigg et al., 2013). Using this assumption and assuming a continuous differential in Eq. (1) leads to,

$$\frac{\mathrm{d}N_{\mathrm{ufz}}}{N_{\mathrm{ufz}}} = -J_{\mathrm{het}}A_{\mathrm{g}}\mathrm{d}t.$$
(2)

Integrating Eq. (2) further results in the commonly used expression for the fraction of frozen droplets, 215

$$f_{\rm frz} = \frac{N_{\rm frz}}{N_{\rm tot}} = 1 - e^{-J_{\rm het}A_{\rm g}t}.$$
(3)

The form of the expression given in Eq. (3) is used in many studies although modified slightly when considering multiple components or contact angle distributions (e.g. Niedermeier et al., 2010; Murray et al., 2011; Broadley et al., 2012; Rigg et al., 2013), and when particle or droplet sizes are discretized or binned (e.g. Marcolli et al., 2007; Lüönd et al., 2010). The major weakness of this 220 exponential form to describe $f_{\rm frz}$ lies entirely in the assumption it is based on, i.e. it is only valid if the ISA is exactly the same for all droplets considered. When taking into account individual droplet ISA for all droplets, this formulation is not valid. Thus, application of this formula to interpret ice nucleation studies, or use in mathematical frameworks, strictly speaking, is also invalid when ISA on a droplet per droplet basis is different.

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The ISA in a single droplet is a measurable quantity with a corresponding measurement uncertainty. It is unlikely that every droplet prepared in an immersion freezing experiment has identical ISA. For the same particle type, there will exist a systematic ISA uncertainty with respect to a particular droplet preparation technique. This systematic uncertainty is σ_g and can be determined by

directly measuring ISA in a population of independently prepared droplets. Since the ISA variabil-230 ity may not be typically resolved in previous experiments, a droplet freezing simulation must be employed to model ice nucleation for interpretation purposes. To accomplish this, freezing of each single droplet is assumed to be stochastic, or in other words, there exists a probability of the *j*th droplet to freeze, $P_{j,\text{frz}}$, within δt . The probability for a single droplet not to freeze, $P_{j,\text{ufz}}$, is realized 235 as an exponential decay law (Pruppacher and Klett, 1997; Koop et al., 1997) and therefore,

$$P_{i,\text{frz}} = 1 - P_{i,\text{ufz}} = 1 - e^{-J_{\text{het}}A_j\delta t}.$$
(4)

A time and surface area dependent immersion freezing process which follows CNT is assumed and as a result, all simulations employ J_{het} having units of cm⁻² s⁻¹. However, J_{het} does not explicitly depend on time and ISA, but on T and a_w . A droplet can either remain in an unfrozen state or freeze and therefore, is described exactly by a binomial distribution, B($k; n, P_{j,frz}$), with parameters $P_{j,frz}$ given by Eq. (4) and n = 1 meaning that only one trial is given for an individual droplet to freeze in

 δt . A randomly sampled number, k = 0 or 1, is obtained from the distribution

$$B(k; n = 1, P_{j, frz}) = P_{j, frz}^{\kappa} (1 - P_{j, frz})^{1 - \kappa}$$
(5)

for each droplet with a normalization prefactor, n!/(k!(n-k)!) = 1. When k = 1, freezing occurs
for the *j*th droplet and if k = 0, the droplet does not freeze and another k is sampled in the next time interval. For a collection of multiple droplets, the number of freezing events that occur in a given time interval is n_{frz} and the cumulative sum as a function of time is N_{frz}(t). For a single IFS starting with N_{tot} liquid droplets, the fraction of unfrozen droplets is f_{ufz}(t) = 1 - N_{frz}(t)/N_{tot}.

- A record of $n_{\rm frz}$ and corresponding droplet ISA, i.e. A_j , is kept for a single IFS. This record can be thought of as a simulated experimental immersion freezing data set, i.e. it gives a record of droplet freezing time while tracking A_j . Due to the stochastic nature of nucleation, repetition of isothermal IFSs will not result in identical values of $f_{\rm ufz}$ over t. Likewise, repetition of a laboratory experiment will not result in exactly the same $f_{\rm ufz}(t)$ curve. Therefore it is necessary to repeat the simulations in order to reveal a range of $f_{\rm ufz}(t)$ values of which the mean unfrozen fraction, $\overline{f}_{\rm ufz}(t)$, can be derived
- from all simulations. We choose an ensemble of 10^5 IFSs to accurately determine $\overline{f}_{ufz}(t)$. This procedure is a basic form of a Monte Carlo method and yields upper and lower percentile bounds at 5 and 95% serving as a stochastic uncertainty of the immersion freezing process. We define stochastic uncertainty as the scatter in the data due to the occurrence of random (i.e. stochastic) freezing events upon repeat experiments as a result of a set number of observed freezing events.
- An ensemble of IFSs, referred to as a model simulation, requires the selection of parameters, e.g. N_{tot} , A_{g} , σ_{g} , and J_{het} . For demonstration purposes, the parameter choice is arbitrary. However, when reproducing a laboratory derived data set, a parameter selection process is applied. Parameters which can be directly accessed from previous laboratory studies are first selected to mimic experimental conditions. For example, if a study reports that 100 droplets were examined in an immersion freez-
- 265 ing experiment, then $N_{\text{tot}} = 100$. Some previous studies report only average ISA per droplet, A_{avg} , and neglect information for estimating σ_{g} . If A_{avg} is reported as $7.1 \times 10^{-6} \text{ cm}^2$, then for simplicity we set $A_{\text{g}} = 7.1 \times 10^{-6} \text{ cm}^2$. For all studies in which a parameter or multiple parameters are not available or readily calculated, the model derived f_{ufz} or f_{frz} are fitted simultaneously to experimentally derived f_{ufz} or f_{frz} , and either critically assessed whether or not the parameter best reproduces
- 270 experimental conditions or the fitted parameter value is compared with independently derived values from other published literature. We define 'model derived' to refer to calculated frozen fraction, unfrozen fractions or J_{het} values which are not model input parameters. Details about the selection of parameters or whether or not parameters are or are not fitted for each simulation will be discussed

in results section 3. In many isothermal immersion freezing laboratory studies, droplet freezing con-

tinues over time when all other conditions remain constant, i.e. at constant T (Wright and Petters, 2013; Murray et al., 2011; Broadley et al., 2012; Herbert et al., 2014). Therefore, the J_{het} parameter is selected to be constant for isothermal IFSs.

2.2 Simulation of cooling rate dependent immersion freezing experiments

2.2.1 Experimentally derived J_{het} for model input

- 280 When a cooling rate is applied in model simulations, droplet freezing is simulated in discrete temperature intervals and therefore J_{het} at every step is required for deriving $P_{j,frz}$. In this study, only water droplets are considered and therefore, it is assumed that $a_w = 1.0$ and J_{het} becomes a function of Tonly. Ideally, experimentally derived $J_{het}(T)$ should be used for prediction of immersion freezing. However, these data sets are usually limited in T range and are discrete in nature. Knopf and Alpert
- 285 (2013) compiled experimental data which was parameterized as a continuous function over T following the ABIFM expressed as,

$$\log_{10}(J_{\text{het}}) = m\Delta a_{\text{w}} + c \,, \tag{6}$$

where m and c are slope and intercept parameters, respectively, and Δa_w is the independent variable following the formulation of Koop et al. (2000). The Δa_w at which a droplet freezes is calculated by subtracting the a_w of the droplet (= 1.0 for pure water) from the water activity point that falls on the ice melting curve, $a_{w, ice}(T)$, at the same temperature or

$$\Delta a_{\rm w} = a_{\rm w}(T) - a_{\rm w,\,ice}(T) \,, \tag{7}$$

where

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$$a_{\rm w,\,ice}(T) = p_{\rm ice}(T)/p^{\circ}_{\rm H_2O}(T) , \qquad (8)$$

and $p_{ice}(T)$ and $p^{\circ}_{H_2O}$ are the vapour pressure with respect to planar ice and water, respectively (Murphy and Koop, 2005).

Resulting calculations from Eqs. (6) to (8) are not computationally demanding and conveniently derive $J_{het}(T)$ for model input. Details about parameter selection, i.e. m and c, the treatment of ISA variability or whether or not parameters are or are not fitted will be discussed in the results section 3.

2.2.2 Simulated droplet freezing

Cooling rate dependent IFSs are performed to evaluate the effect of stochastic freezing and variable ISA in laboratory immersion freezing experiments. Again, the ISA for a single droplet is sampled, however, Eqs. (1) and (4) are modified to

$$305 \quad \delta N_{\rm ufz} = -\frac{A_{\rm tot}}{r} J_{\rm het}(T) \delta T , \qquad (9)$$

and

$$P_{j,\text{frz}} = 1 - P_{j,\text{ufz}} = 1 - e^{-\frac{A_j}{r} J_{\text{het}}(T)\delta T},$$
(10)

respectively, where δT is a temperature interval and $r = \delta T / \delta t$ is the cooling rate. $J_{\text{het}}(T)$ is calculated from Eq. (6) and used in Eq. (10). Once the probability for the jth droplet to freeze is calculated

- for all droplets, freezing is determined by sampling from $B(k; n, P_{j,frz})$ (Eq. 5). The number of freez-310 ing events that occur in a given δT is $n_{\rm frz}$, and the cumulative sum as a function of T is $N_{\rm frz}(T)$ and used to calculate frozen fractions of droplets, $f_{\rm frz}(T) = N_{\rm frz}(T)/N_{\rm tot}$. Similar to isothermal freezing, a single r dependent IFS yields a droplet immersion freezing record analogous to an experimental data set. In this case, the record of droplet freezing and corresponding A_i is a function of T. The average frozen fraction for 10^5 simulations, $\overline{f}_{frz}(T)$, is calculated along with percentiles at 5 and
- 95% used as a stochastic uncertainty.

It is important to note that application of r dependent IFSs presented here do not require the AB-IFM, as it is only used as a parameterization of previously published immersion freezing data sets to calculate $J_{het}(T)$. Any other published $J_{het}(T)$ will work equally as well. The ABIFM parameteri-

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zation is INP type dependent and suitable for saturated and subsaturated conditions, i.e. $a_{\rm w} \leq 1$, or $RH \le 100\%$, if the droplet is in equilibrium with the water vapor phase. Therefore, the ABIFM is a useful and convenient tool for model input $J_{het}(T)$.

3 **Results and discussion**

3.1 Isothermal model simulations of individual droplet freezing experiments

- 325 Figure 1a shows 5 and 95 % bounds of $f_{\rm ufz}$ from 4 model simulations for different $N_{\rm tot}$ applying either uniformly equal ($\sigma_g = 1$) or lognormally distributed ($\sigma_g = 10$) ISA per droplet as given in Table 1. Two of these test cases, Iso1 and Iso2, have uniform ISA both resulting in f_{ufz} (on a logarithmic scale) linear with t. However, the spread of the 5 and 95% bounds is much wider for Iso2 having $N_{\text{tot}} = 30$ than for Iso1 having $N_{\text{tot}} = 1000$. It is clear that a larger spread in simulated f_{ufz} is entirely due to
- applied smaller N_{tot} . This implies that a laboratory experiment using a small N_{tot} , is statistically 330 less significant compared to an experiment with greater $N_{\rm tot}$. A single experimentally derived $f_{\rm ufz}$ curve under the same conditions as Iso2 will fall anywhere between the upper and lower bounds, and thus may even appear to deviate from a log-linear relationship over time. Therefore, interpretation about the nature of the heterogeneous ice nucleation process from the slope of $f_{\rm ufz}$ over time for an experiment using small N_{tot} should be conducted with care.

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Model simulations Iso3 and Iso4 are shown in Fig. 1a where $N_{\text{tot}} = 1000$ and 30, respectively, and the ISA per droplet is sampled from lognormal distribution with $\sigma_{\rm g} = 10$. In Iso3, $f_{\rm ufz}$ significantly deviates from a log-linear relationship with t. In Iso4, the same curvature exists, however the percentile bounds are much wider due to applied smaller N_{tot} . It is important to note that J_{het} is the

- same and constant for all simulations shown in Fig. 1a. The nucleation rate of each *j*th droplet can be calculated as, $\omega_{\text{het},j} = J_{\text{het}}A_j$ with units of s⁻¹. The droplets having a larger or smaller ISA will result in larger or smaller $\omega_{\text{het},j}$, respectively. The fact that f_{ufz} is linear for $\sigma_g = 1$, the curvature effect in f_{ufz} seen for $\sigma_g = 10$ must be entirely due to ISA variability. This is because droplets with greater values of $\omega_{\text{het},j}$ will tend to nucleate more rapidly than those having smaller $\omega_{\text{het},j}$ values. In
- 345 other words, the curvature of $f_{ufz}(t)$ is entirely due to those droplets having larger and smaller ISA that freeze within shorter and longer time scales, respectively. In addition, the spread in the 5 and 95 percentiles is very similar for Iso1 and Iso3, and for Iso2 and Iso4. This is seen most clearly at the intersection of the blue and green shaded regions ($t \simeq 1.3 \text{ min}$). In isothermal freezing experiments, variability in ISA will not significantly affect stochastic uncertainty estimates, but will cause $f_{ufz}(t)$
- 350 to deviate from a log-linear relationship.

In some previous experimental isothermal immersion freezing studies, the number of liquid droplets and an estimate of the average ISA per droplet are provided or can be derived. However, the validity of the assumption that all droplets possess the same ISA is rarely investigated or quantified. Similarly, J_{het} is not often reported. However, laboratory data do provide an opportunity to test our model

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for robustness while using parameters similar to those reported in the experimental studies. In fact, our model can also provide estimates for parameters typically unreported or unavailable, such as J_{het} and σ_{g} .

Experimental data by Wright and Petters (2013) for isothermal immersion freezing by ATD is very well reproduced by model simulation IsoWR as demonstrated in Fig. 1b. Parameters for IsoWR are given in Table 1 and chosen to mimic experimental conditions in which droplets contained 1 yrt ⁹⁷

- 360 given in Table 1 and chosen to mimic experimental conditions in which droplets contained 1 wt %ATD held at 251 K. Bounds at 5 and 95 % of simulated f_{ufz} are shown in Fig. 1b and envelop the laboratory data. A repeat experiment by Wright and Petters (2013) should result in a f_{ufz} curve falling within the percentile bounds 95 % of the time when considering only stochastic uncertainty.
- To further evaluate the validity of the simulations, the parameters used are compared with exper-365 imental conditions given in Wright and Petters (2013). IsoWR uses $N_{tot} = 1000$ which agrees with the reported range of 300–1500. Parameters A_g , σ_g and J_{het} used in IsoWR were fitted simultaneously so that the average f_{ufz} from 10⁵ simulations (or \overline{f}_{ufz}) best reproduced observed f_{ufz} . The first parameter in question is $\sigma_g = 9.5$, which can be interpreted as a systematic standard error in ISA due to the experimental methods of generating or dispensing droplets containing ATD acting as INPs. We
- 370 note this is different from an absolute ISA measurement error. Wright and Petters (2013) emulsified a mixture of oil and a bulk solution of water and ATD particles to form droplets with diameters of 50–250 µm. The variability in ISA should scale directly with the variability in droplet volumes (i.e. a factor of 125 or just over 2 orders of magnitude). Additional uncertainty will certainly arise from the variability in ATD particle numbers and the variability in ATD particle size. The surface area
- 375 distribution and a random sampling is given in Fig. S2 for the IsoWR model simulation. A factor of 125 in range in ISA is a lower estimate of uncertainty, but already accounts for about 75% of the

total sampled droplets. While not directly defined by Wright and Petters (2013), we are confident that the overall range in ISA should be well over 2 orders of magnitude and therefore, $\sigma_g = 9.5$ is a reasonable value for the lognormal distribution width parameter employed in the simulations in

- Fig. 1b to reproduce the experimental data. The third parameter in question is $A_g = 6.4 \times 10^{-3} \text{ cm}^2$. Unfortunately, an average ISA was not reported by Wright and Petters (2013), but can be estimated using literature values of specific surface area (SSA) applying the Brunauer, Emmett and Teller gas adsorption method (Brunauer et al., 1938). It is important to note that surface area measurements are not unambiguous due to the fact that heterogeneous ice nucleation may involve layers of wa-
- 385 ter molecules interacting with surface molecules (Cox et al., 2013). The BET technique is one of many in which particle surface area is measured, and can be used to represent molecular available surface area. Bedjanian et al. (2013) report SSA for ATD used in Wright and Petters (2013) as 85±10 m² g⁻¹. The ISA per drop can then be estimated from the drop volume, V_{drop}, and the density of water, ρ_w, using the equation V_{drop} · ρ_w · wt % · SSA. Considering only the variability in
- 390 V_{drop}, average ISA per drop should range between 5.5 × 10⁻⁴ and 7.0 × 10⁻² cm². The A_g parameter in model simulation IsoWR falls within this range. Finally, J_{het} for ATD in water droplets was investigated by Pinti et al. (2012) who reanalyzed ATD immersion freezing data by Marcolli et al. (2007) but did not report J_{het} values. However, estimates can be made following Knopf and Alpert (2013) accounting for f_{frz} = 0.01 and a nucleation time assumed to be 1 s, which yields J_{het} ranging
 395 from 5 × 10⁶ to 1 × 10² cm⁻² s⁻¹ between T = 247.4 and 252.8 K, in reasonable agreement with

 $J_{\rm het} = 2.6 \times 10^3 \, {\rm cm}^{-2} \, {\rm s}^{-1}$ used in IsoWR at $251 \, {\rm K}$.

The new model simulation presented here based entirely on CNT can describe freezing experiments by Wright and Petters (2013) accounting for long nucleation time scales and a large number of droplets considering variability in ISA. In addition, all crucial parameters applied are experimen-

- 400 tally supported, in particular J_{het} which is in agreement with independent studies (Marcolli et al., 2007; Pinti et al., 2012). Therefore, the isothermal immersion freezing data set of Wright and Petters (2013) may be explained by a time and ISA dependent stochastic freezing process, in which each droplet contains variable ISA. More experimental investigation and model analysis should be conducted to verify their agreement at different temperatures, time scales and surface areas. Droplet to
- 405 droplet variability in ice nucleation efficiency is typically parameterized with a variable efficiency of sites to nucleate ice, as done successfully in Wright and Petters (2013) or different contact angles (e.g. Niedermeier et al., 2011b; Broadley et al., 2012). Droplet to droplet variability parameterized in these ways and employing identical ISA can result in a deviation of f_{ufz} from a log-linear relationship, similar to what is seen if Fig. 1. However, using a known ISA variability (Broadley et al.,
- 410 2012; Wright and Petters, 2013), we reveal that the observed deviation from a log-linear relationship can be accounted for entirely by the ISA distribution. This may imply that the droplet to droplet variability in ice nucleation efficiency is entirely due to variable ISA.

Figure 2 shows results of isothermal freezing experiments by Broadley et al. (2012) for illite compared to model simulation IsoBr and experimental results by Herbert et al. (2014) for the INP types

- 415 kaolinite and feldspar compared to model simulations IsoHE1 and IsoHE2, respectively (see Table 1). The experimental data and \overline{f}_{ufz} for all model simulations are in agreement and fall within the percentile bounds. Notice that the scatter in the isothermal immersion freezing data points is much larger than for Wright and Petters (2013) shown in Fig. 1b. As previously discussed, this is entirely due to a smaller number of droplets used in the laboratory experiments by Broadley et al. (2012)
- 420 $(N_{tot} = 63)$ and Herbert et al. (2014) $(N_{tot} = 40)$ and thus, may be entirely attributed to the stochastic nature of immersion freezing as expected by CNT. The model simulations capture this effect by producing a wide range in f_{ufz} . Only one experiment was performed for each of the laboratory data sets presented in Fig. 2 and if these experiments were repeated, f_{ufz} values would very likely not be the same and may even exhibit a more linear or curved behavior with time. Repetition of experiments
- should provide better estimates of \overline{f}_{ufz} and σ_g , but for any single experiment, f_{ufz} may still fall within the given percentile bounds. In other words, additional experiments would better define the mean of f_{ufz} and the uncertainty in the mean of f_{ufz} , but will not decrease the uncertainty bounds. Only by using more droplets, e.g. Wright and Petters (2013), would a single experiment be more statistically significant.
- The selection of parameters and ISA distribution used in IsoBR are discussed. $N_{\text{tot}} = 63$ applied in IsoBR is used here (Broadley et al., 2012). In the analysis of the experiment "run 20" simulated by IsoBR, the authors sub-selected droplets $10 - 20 \,\mu\text{m}$ in diameter from a droplet population. The droplet volume and the ISA variability should scale to the 3rd power of the droplet diameter, i.e. by a factor of about $(20/10)^3 = 8$. Therefore, the simulated ISA is assumed to follow a uniform
- 435 probability density function between 9.4×10^{-8} and 7.5×10^{-7} cm². This ISA range spans a factor of ~ 8 with a geometric average of 2.65×10^{-7} cm² as reported in Broadley et al. (2012). We note that a factor of 8 is a lower limit of variability as any additional uncertainty in illite particle size distribution or the numbers of illite particles per droplet is not considered. The parameter $J_{\text{het}} =$ 2.82×10^3 cm⁻² s⁻¹ was not fitted, but instead selected in such a way that this single value resulted in
- 440 $f_{\rm ufz}$ data falling entirely within the stochastic uncertainty limits. This value is in agreement with the previous ABIFM parameterization, $J_{\rm het} = 1.25 \times 10^3 \,{\rm cm}^{-2} \,{\rm s}^{-1}$, at the same temperature and water activity (Knopf and Alpert, 2013). We note that the decay of simulated $f_{\rm ufz}$ over time appears linear in Fig. 2a although the experimental data appears curved. This curvature led Broadley et al. (2012) to assume an active site model to describe the data. However, we find that when using too small
- 445 numbers of droplets, experiments may be too uncertain to make any solid claims about the nature of the ice nucleation process. The stochastic uncertainty bounds in Fig. 2a are sufficiently large that the data is still in agreement with the model simulations presented here. The IsoBR simulation demonstrates that freezing due to illite can also be described by a stochastic freezing approach with

one value of J_{het} . Thus, laboratory derived isothermal immersion freezing of illite may be explained by CNT accounting for the stochastic nature of immersion freezing and variability in ISA.

In the model simulation IsoHE1, parameters A_g , σ_g and J_{het} are are fitted to the experimental data. The parameter $A_g = 1.2 \text{ cm}^2$, is in good agreement with experimentally derived $A_g = 2.4 \text{ cm}^2$ reported in Herbert et al. (2014), for kaolinite using SSA = $11.8 \text{ m}^2 \text{ g}^{-1}$ (Murray et al., 2011), 1.0 wt % concentration and $V_{drop} = 1 \mu \text{L}$. Herbert et al. (2014) did not report sufficient in-

- formation to estimate an overall variability in ISA, therefore, comparison of σ_g to experimental conditions is difficult. As previously discussed, a repeat experiment may result in f_{ufz} exhibiting more linear or non-linear behavior with t within the calculated percentile bounds, i.e. within the stochastic uncertainty. Figure 1a shows that a more linear or non-linear relationship of f_{ufz} with t implies a smaller or larger value of σ_g . We note that due to the lack of quantitative information
- 460 about the variability in ISA, the assumed lognormal surface area distribution may be over or underestimated. Performing more experiments or employing a larger number of droplets will decrease the stochastic uncertainty and better constrain the curvature of \overline{f}_{ufz} over time. The ABIFM yields $J_{het} = 1.75 \times 10^{-2} \text{ cm}^{-2} \text{ s}^{-1}$ at T = 255.15 K and $a_w = 1.0$ which is within an order of magnitude of J_{het} used in IsoHE1. The agreement between simulated and experimental parameters implies
- that CNT may be able to explain observed immersion freezing of kaolinite when variable ISA and stochastic uncertainty is considered. Herbert et al. (2014) and Murray et al. (2011) came to the conclusion that this particular type of kaolinite, KGa-1b, is a "single-component system", which means that a single J_{het} function of T can reproduce the experimental data. Our model simulations lead to the same conclusion, and the derived J_{het} value is in agreement with the independently formulated

470 ABIFM parameterization by Knopf and Alpert (2013).

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Immersion freezing data of Herbert et al. (2014) for feldspar is reproduced by the model simulation IsoHE2. The parameters for IsoHE2 are given in Table 1. Those that are fitted are A_g , σ_g and J_{het} . Average ISA for the data in Fig. 2 is $1.85 \times 10^{-2} \text{ cm}^2$, similar to $A_g = 2.0 \times 10^{-2} \text{ cm}^2$ used in IsoHE2. Droplets used in Herbert et al. (2014) were dispensed with a digital micropipet with high

- 475 accuracy, thus it can be expected that the contribution of droplet volume variability to the σ_g parameter is low. However, it is impossible to make any estimate of σ_g for comparison with the fitted σ_g due to the lack of quantitative experimental information about ISA variability. As in simulation IsoHE1, a more linear or non-linear f_{ufz} curve may imply that our assumed lognormal distribution width is over or underestimated, respectively. However, it will be demonstrated that the same σ_g can
- 480 be used to reproduce the cooling rate dependent experiments of feldspar, which gives confidence that this σ_g value may be appropriate. To better constrain σ_g , more stochastic certainty is required by application of more droplets or conducting multiple experiments. Values of J_{het} for feldspar independent from Herbert et al. (2014) in the same temperature range to our knowledge do not exist making comparison difficult.

- Model simulations IsoDI1-3 of isothermal immersion freezing experiments by Diehl et al. (2014) for illite in wind tunnel levitation experiments are shown in Fig. 3. Simulation parameters are given in Table 1. Only one droplet was observed in each experiment, and approximately 45 experiments were conducted for each of the 3 data sets shown in Fig. 3. This is equivalent to 1 experiment with $N_{\text{tot}} = 45$ droplets, since droplet freezing is independent of the freezing of other droplets. Ex-
- 490 cellent agreement is observed between simulated and experimental f_{ufz} . The parameter A_g for the three simulations match the average surface area per drop reported in Diehl et al. (2014). Parameters σ_g , $J_{het}(-18 \,^\circ\text{C})$ and $J_{het}(-21 \,^\circ\text{C})$ are simultaneously fitted to experimental data. It can be expected that σ_g is the same for all three simulations, due to the fact that Diehl et al. (2014) likely used identical bulk water-illite stock solutions. Therefore a single fitted value of $\sigma_g = 3.2$ is used
- for all simulations. IsoDI2 and IsoDI3 simulate data taken at $T = -21^{\circ}$ C and use the fitted parameter $J_{het}(-21^{\circ}$ C) = 1.0×10^{0} cm⁻² s⁻¹. IsoDI1 simulated data at $T = -18^{\circ}$ C uses the parameter $J_{het}(-18^{\circ}$ C) = 1.8×10^{-2} cm⁻² s⁻¹. At T = -18 and -21° C, the ABIFM yields $J_{het} = 1.8 \times 10^{-2}$ and 2.6×10^{-1} cm⁻² s⁻¹, respectively, and is in excellent agreement with derived values in IsoDI1-3. An adequate constraint of σ_g could not be established due to a lack of information about the ISA
- 500 variability. However, it is evident that the fitted σ_g value may be justified due to the fact that the same value reproduced all 3 isothermal data sets. We find that a time dependent and stochastic immersion freezing process may reconcile observations when variable ISA is considered.

Depending on ISA variability, trajectories of model derived f_{ufz} over time are significantly altered and thus assuming identical ISA may not be valid. It is well known that immersion freezing depends on surface area, i.e. an increase in ISA translates to an increase in nucleation rate. However, we note that variability in both t and ISA equally affect calculations of droplet freezing probabilities (Eq. 4) used in model simulations, and therefore neglecting time dependence will cause erroneous interpretation of immersion freezing data to the same degree as if the surface area dependence is ne-

510 the isothermal ice nucleation data of various experiments without invoking empirical parameterizations, assumptions of particle surface composition, and/or other modifications in parameters and interpretations.

glected. This simple stochastic immersion freezing model accounting for ISA variability can explain

3.2 Cooling rate model simulations of individual droplet freezing experiments

Cooling rate IFSs were performed to investigate the effects of variable ISA and N_{tot} on experimentally derived J_{het} and f_{frz} as a function of T. For a single cooling rate IFS, variable ISA per droplet is applied and used to calculate $P_{j,\text{frz}}$ from from Eq. (10) in discrete temperature steps, δT , and then Eq. (5) simulates freezing. The IFS stops after some T or when all droplets freeze, and the simulated freezing record is kept detailing which droplets froze or remained liquid at each T and their corresponding ISA. This is analogous to running an immersion freezing experiment in a laboratory

setting and recording the observed number of frozen droplets or ice crystals as a function of T.

The simulated freezing record is treated as a freezing data set from which the assumption of identical ISA can be tested. This is accomplished by re-calculating J_{het} from the simulated data. These (re-)calculations use n_{frz} , the length of the time interval, $\delta t = \delta T/r$, and either of two different approaches in determining A_{tot} . For the first approach, A_{g} is assumed to be identical for all droplets,

525 i.e. without the knowledge that immersion freezing was simulated for droplets with variable ISA in the first place. This is equal to assuming a monodisperse INP population in laboratory immersion freezing experiments resulting in an "apparent" heterogeneous ice nucleation rate coefficient, $J_{het}^{apparent}(T)$, calculated by

$$J_{\text{het}}^{\text{apparent}}(T) = \frac{n_{\text{frz}}(T)}{n_{\text{ufz}}(T)A_g \frac{\delta T}{r}},\tag{11}$$

530 where $n_{ufz}(T)$ is the number of unfrozen droplets at T and $A_{tot} = n_{ufz}A_g$. The second approach accounts for the variable ISA present in droplets resulting in the "actual" heterogeneous ice nucleation rate coefficient, $J_{het}^{actual}(T)$, calculated by

$$J_{\rm het}^{\rm actual}(T) = \frac{n_{\rm frz}(T)}{\sum A_j \frac{\delta T}{r}},\tag{12}$$

and $A_{tot} = \sum A_j$ is the total surface area contribution from droplets that remain liquid. Comparing results from Eqs. (11) and (12) allows evaluation of the assumption that all droplets have the same ISA, when they actually do not. In this way a null hypothesis is considered, that is if $J_{het}^{apparent}(T)$ and $J_{het}^{actual}(T)$ are the same, then the assumption of identical ISA is valid.

Poisson statistics are used to derive upper and lower fiducial limits of J^{apparent}_{het}(T) and J^{actual}_{het}(T) at x = 0.999 confidence for n_{frz} following Koop et al. (1997). The upper fiducial limit of the heterogeneous ice nucleation rate coefficient, J^{up}_{het}, accounts for additional freezing events occurring with a probability of x, than observed n_{frz}. Likewise, a lower fiducial limit of the heterogeneous ice nucleation rate coefficient, J^{low}_{het}, accounts for less than the observed n_{frz} occurring with a probability of x. We refer to the upper and lower limits of n_{frz} as n^{up}_{frz} and n^{low}_{frz}, respectively (Koop et al., 1997). The fiducial limits of J^{apparent}_{het} and J^{actual}_{het} for a single simulation can be calculated using Eqs. (11)
and (12), but replacing n_{frz} with n^{up}_{frz} or n^{low}_{frz}, respectively. Each simulation results in different J^{apparent}_{het} and J^{actual}_{het} values and different fiducial limits at the same T due to random sampling, therefore, averages are reported.

Figure 4 shows the results of two model simulations, Cr1 and Cr2, having r = 0.5 and 5.0 K min⁻¹, respectively. For all 10⁵ IFSs, J^{apparent} and J^{actual} are shown in Fig. 4a and b as dashed lines, respectively, along with corresponding f_{frz} curves displayed in Fig. 4c and d. The parameterization of J_{het}(T) for illite dust (Knopf and Alpert, 2013) with m = 54.5 and c = -10.7 used in Eq. (10) for each simulation is shown as the red line in Fig. 4a and b and referred to as the model input J_{het}. Simulation parameters for Cr1 and Cr2 are given in Table 2.

According to CNT, two immersion freezing cooling rate experiments conducted at different rshould result in identical J_{het} values due to the fact that J_{het} is independent of r. CNT is violated if significantly different J_{het} values are derived at different r. Figure 4a shows that values of $\overline{J}_{het}^{apparent}$ are significantly different between model simulations Cr1 and Cr2. Also $\overline{J}_{het}^{apparent}$ is overestimated at higher freezing temperatures and underestimated at lower freezing temperatures compared with model input $J_{het}(T)$. These significant differences do not support the null hypothesis and imply

- that when experimentally deriving J_{het} , the assumption that ISA per droplet is identical is invalid. Figure 4b shows that accounting for variable ISA, $\overline{J}_{het}^{actual}$ for Cr1 and Cr2 is consistent. It is important to note that $\overline{J}_{het}^{apparent}$ shows a cooling rate dependence, i.e. when the cooling rate varies by an order of magnitude, $\overline{J}_{het}^{apparent}$ evaluated at one temperature is also about an order of magnitude different. Values of $\overline{J}_{het}^{actual}$ on the other hand, are not dependent on cooling rate. We reiterate that $\overline{J}_{het}^{apparent}$
- and $\overline{J}_{het}^{actual}$ are calculated from the same simulated freezing record having lognormally distributed ISA. The only difference between "actual" and "apparent" J_{het} is the surface area assumptions used in their respective calculation (Eqs. 10 and 11), i.e. $J_{het}^{apparent}$ intentionally assumes identical ISA as done commonly in experimental analysis and J_{het}^{actual} accounts for the variable ISA. Thus, the apparent cooling rate dependence in simulations Cr1 and Cr2 is a direct result of assuming identical ISA. This
- 570 is the case when a broad ISA distribution is simulated, i.e. $\sigma_g = 10$. When the ISA distribution is very narrow, uniform or σ_g is about 1.0, then J_{het}^{actual} will equal $J_{het}^{apparent}$. This is demonstrated in Fig. S7 by model simulations Cr3 and Cr4 using $\sigma_g = 1$. If an experimental study succeeds to create a narrow enough distribution, then assuming identical ISA may be applicable.
- Towards warmer (T > 248 K) and colder temperatures (T < 238 K), the difference in upper and lower fiducial limits derived in Cr1 and Cr2 are much greater than for the mid temperature range (238 < T < 248 K). In fact the smallest difference occurs at $\overline{f}_{frz} \simeq 0.5$. This is because calculations are statistically more significant at the median freezing where n_{frz} is largest. Fewer droplets freeze at the beginning and end of a cooling process resulting in a wide fiducial limit range reaching up to 4 orders of magnitude (Fig. 4a and b) in spite of a high number of dropets used ($N_{tot} = 1000$).
- 580 The corresponding percentile bounds of $f_{\rm frz}$ shown in Fig. 4c and d do not reflect a considerable uncertainty compared to the upper and lower fiducial limits (Fig. 4a and b). It is important to note that $f_{\rm frz}$ are identical in Fig. 4c and d, because surface area is not used to derive $f_{\rm frz}$. This analysis suggest that values and uncertainties of $f_{\rm frz}$ are not suited to derive $J_{\rm het}$ and any corresponding error. Previous immersion freezing experiments by Herbert et al. (2014) are modeled in CrHE1 and
- 585 CrHE2 where r = 0.2 and $2.0 \,\mathrm{K\,min^{-1}}$, respectively, for the case of feldspar acting as INPs. Herbert et al. (2014) used the same weight fraction of feldspar per droplet in isothermal and cooling rate dependent experiments. Therefore, it is reasonable to suspect that the parameters should be the same or very close. The parameters σ_g and A_g are not fitted in model simulations CrHE1 and CrHE2 and instead are taken from the values fitted in IsoHE2. The parameters m and c used to calculate J_{het} as
- 590 a function of T following the ABIFM are fitted so that experimentally derived f_{frz} is best reproduced by model derived $\overline{f}_{\text{frz}}$. Since Herbert et al. (2014) assumed identical ISA, experimentally derived J_{het} can be directly compared with $J_{\text{het}}^{\text{apparent}}$ from cooling rate model simulations.

Figure 5 shows experimentally derived f_{frz} and $J_{\text{het}}^{\text{apparent}}$ from Herbert et al. (2014) compared to results of model simulations CrHE1 and CrHE2. Parameters m = 122.83 and c = -12.98 are used in

- Eq. (6) to reproduce frozen fraction data (Fig. 5a) within 5 and 95% bounds. The laboratory data falls within percentiles and fiducial limits of $f_{\rm frz}$ and $J_{\rm het}^{\rm apparent}$, respectively. Previous studies have been successful in interpreting immersion freezing studies to follow an active site approach by considering both cooling rate and isothermal experiments (Vali, 2014; Herbert et al., 2014). Alternatively, we have found that our isothermal and cooling rate simulations based on a single function of $J_{\rm het}(T)$ for
- 605 data. Note that simulated and experimentally derived $J_{het}^{apparent}$ for r = 0.2 and $2.0 \,\mathrm{K \,min^{-1}}$ (5b) are different by about 1 order of magnitude at the same T.

Figure 5c shows $\overline{J}_{het}^{actual}$ with upper and lower fiducial limits derived from CrHE1 and CrHE2. For comparison, J_{het} derived in model simulation IsoHE2 for feldspar is also shown and in agreement with J_{het}^{actual} within our stochastic uncertainty estimates. When accounting for variable ISA, $\overline{J}_{het}^{actual}$

- 610 are in excellent agreement with the ABIFM parameterization derived in this study for feldspar INPs. Furthermore, $\overline{J}_{het}^{actual}$ calculated for different r are identical as predicted by CNT, a similar finding as in the model simulations Cr1 and Cr2 (Fig. 4b). Therefore, $J_{het}(T)$ used here can be considered a new $J_{het}(\Delta a_w)$ parameterization for feldspar valid for $0.078 < \Delta a_w < 0.120$.
- The differences between $J_{het}^{apparent}$ and J_{het}^{actual} shown in Figs. 5b and c can be attributed to two 615 reasons. i) A potential misrepresentation of the slope of J_{het} versus T and ii) a potential misrepresentation of a dependence of J_{het} on r. Regarding the slope of J_{het} vs. T, note that droplets with ISA less than A_g likely freeze at colder T compared to droplets with ISA greater than A_g , that likely freeze at warmer T. However, assuming identical ISA equal to A_g for all droplets either overestimates or underestimates the actual ISA present in droplets that freeze at colder and
- 620 warmer temperatures, respectively. Due to the inverse relationship between A_g and J_{het} , calculations of $J_{het}^{apparent}$ from Eq. (11) may then be underestimated and overestimated at colder and warmer temperatures, respectively. When comparing $J_{het}^{apparent}$ against J_{het}^{actual} in model simulations (calculated from the same freezing record generated using lognormally distributed ISA), $J_{het}^{apparent}$ is underestimated and overestimated at colder and warmer temperatures, respectively. Therefore, this analysis
- 625 clearly shows that assuming identical ISA in each droplet may potentially lead to misrepresentation of the slope J_{het} vs. T. Regarding a dependence of J_{het} on r, we find that $J_{het}^{apparent}$ for simulations CrHE1 ($r = 0.2 \text{ K min}^{-1}$) and CrHE2 ($r = 2.0 \text{ K min}^{-1}$) are different by about 1 order of magnitude at the same T. In Herbert et al. (2014), experimentally derived J_{het} applying r = 0.2 and 2.0 K min^{-1} differ by about 1 order of magnitude at the same T. In separate model simulations not

shown here, applying r different by 2 orders of magnitude yields J^{apparent} values that differ by 2 orders of magnitude. We note that this is the case for a wide distribution width of σ_g = 8.5 and that the stochastic uncertainty in J_{het} is large, not considering additional uncertainties, e.g. in temperature or determining the surface area measurement and variability. Nevertheless, experimentally derived J_{het} data from Herbert et al. (2014) is in good agreement with model simulated J^{apparent}_{het} indicating a potentially erroneous cooling rate dependence which may be caused by the assumption of identical

ISA.

Model simulations CrDI1 and CrDI2 of immersion freezing experiments by Diehl et al. (2014) for illite acting as INPs probed in acoustic levitation experiments are shown in Fig. 6. A non-linear r was used in Diehl et al. (2014) and was the same for both experiments and model simulations, but the ISA

- 640 per droplet was varied. Diehl et al. (2014) reported an ISA per drop of 7.1×10^{-1} and 7.1×10^{-3} cm² in the 2 different sets of experiments. These surface areas are not the same as for isothermal experiments, and so we suspect that different stock solutions were prepared for cooling rate experiments by Diehl et al. (2014). Therefore, parameters from IsoDI1 to 3 are not used in cooling rate model simulations and instead newly fitted values of simulation parameters σ_g and A_g for CrDI1 and CrDI2
- 645 were derived and are given in Table 2. A continuous function of J_{het} was not fitted, but calculated using the ABIFM for illite (Knopf and Alpert, 2013). When using the A_g values reported in Diehl et al. (2014) in conjunction with the other parameters, model simulations cannot reproduce experimental f_{frz} . This is in spite of the excellent performance of IsoBr for reproducing droplet freezing initiated by illite from Broadley et al. (2012). In attempt to reconcile results from Diehl et al. (2014) with
- 650 previous literature data (Broadley et al., 2012; Knopf and Alpert, 2013), model derived $f_{\rm frz}$ are fit to experimental $f_{\rm frz}$ yielding two different parameter values of $A_{\rm g} = 2.94$ and $2.91 \times 10^{-2} \,{\rm cm}^2$ used in CrDI1 and CrDI2, respectively. We note that fitted $A_{\rm g}$ values differ only by a factor of 4 from values reported by (Diehl et al., 2014) and therefore, are in reasonable agreement. However, calculated $J_{\rm het}^{\rm apparent}$ values shown in Fig. 6b still use ISA of 7.1×10^{-1} and $7.1 \times 10^{-3} \,{\rm cm}^2$ as reported by
- 655 Diehl et al. (2014).

Figure 6a shows that simulated and experimental $f_{\rm frz}$ are in agreement when accounting for ISA variability ($\sigma_{\rm g} = 5.7$). Experimental values of $J_{\rm het}^{\rm apparent}$ displayed in Fig. 6b are in agreement with model derived $\overline{J}_{\rm het}^{\rm apparent}$. This result is robust since experimental $J_{\rm het}^{\rm apparent}$ data was not used in fitting $f_{\rm frz}$ and the same value of the fitted parameter $\sigma_{\rm g}$ is used in the two surface area dependent

- 660 cooling rate experiments, which gives confidence that the ABIFM parameterization and the ISA distribution width are appropriate. Accounting for the actual variability in ISA used to simulate freezing, J_{het}^{actual} shown in Fig. 6c is in perfect agreement with the ABIFM parameterization for illite (Knopf and Alpert, 2013). For comparison, J_{het} derived in model simulation IsoDI1 to 3 for illite are also shown and in agreement with J_{het}^{actual} within our stochastic uncertainty estimates. Again, the
- 665 data and model supports a stochastic, time dependent immersion freezing process and may describe laboratory data considering variable ISA.

A major inconsistency between experiment and simulation is shown in Fig. 6b, evident from the agreement of $J_{het}^{apparent}$ for the case when different ISA but identical cooling rates are applied. According to CNT, J_{het} is independent of surface area. This means that if two experiments are

- 670 performed with different ISA but use the same r, J_{het} should be the same as a function of T. However, simulated and experimentally derived $\overline{J}_{het}^{apparent}(T)$ deviate by more than 1 order of magnitude as the surface area varies by two orders of magnitude. This would indicate that $J_{het}^{apparent}$ values violate CNT, but this is the cause of assuming identical ISA. In fact, this freezing behavior also contradicts all surface-based empirical parameterization of immersion freezing, such as determining $n_s(T)$, or the
- 675 number of actives sites per particle surface area (Murray et al., 2012; Hoose and Möhler, 2012). This result impacts immersion freezing experiments conducted as a function of ISA that assume identical ISA in each experiment, thereby implicitly imposing a surface area dependence on $J_{het}^{apparent}$ or $n_s(T)$. Accounting for the experimental uncertainty and variability in ISA may reconcile experimental data.

3.3 Continuous flow and cloud chamber immersion freezing experiments

- 680 Model simulations IsoCFDC and IsoLACIS (see Table 1) reproduce experimental results of Wex et al. (2014) who used 2 ice nucleation instrumentation, (i) a continuous flow diffusion chamber (CFDC) (Rogers et al., 2001; DeMott et al., 2010) and (ii) the Leipzig aerosol cloud interaction simulator (LACIS) (Hartmann et al., 2011), respectively, to observe immersion freezing of 300 nm mobility diameter selected kaolinite particles as a function of *T* and RH > 100%. It is important to note that
- for both instruments, droplet freezing is not observed and instead is optically detected and at the LACIS outlet, a self-built optical particle spectrometer, TOPS-Ice (Clauss et al., 2013), determines if the arriving hydrometeors are liquid droplets or frozen ice crystals, resulting in the determination of a frozen fraction. Thus, $f_{\rm frz}$ is calculated from the ratio between observed ice crystal and aerosol numbers per volume of air. The model simulation parameter $N_{\rm tot}$ is derived from known experimen-
- 690 tal parameters, including residence time of the CFDC, $t_r = 5 \text{ s}$, aerosol flow rate, $Q = 1.0 \text{ L min}^{-1}$, and kaolinite particle concentrations, $N_p = 10 \text{ cm}^{-3}$ (Wex et al., 2014). By defining a single IFS over an interval of time equal to t_r , $N_{tot} = N_pQt_r = 833$ particles per IFS. Similarly for LACIS, $Q = 0.08 \text{ L min}^{-1}$, $t_r = 1.6 \text{ s}$, and $N_{tot} = 21$ particles per IFS. Note that minimum f_{frz} values for CFDC and LACIS presented in Wex et al. (2014) are approximately equal to $1/N_{tot}$. We run 1440
- and 6000 isothermal IFSs for IsoCFDC and IsoLACIS, respectively, equivalent to 2 h averages as done in Wex et al. (2014). Simulation parameters for IsoCFDC and IsoLACIS are given in Table 1. Figure 7 shows that simulated $f_{\rm frz}$ for IsoCFDC and IsoLACIS agree very well with CFDC

and LACIS data by Wex et al. (2014). However, some data points fall outside of the 5 and 95 percentiles (Fig. 7a), which may imply that a greater uncertainty exists that cannot be explained

700 by a stochastic freezing process. This may be due, in part, to uncertainty in ice crystal detection which is not accounted for in model simulations. The surface area for spherical 300 nm particles is $A_{300\text{nm}} = 2.8 \times 10^{-9} \text{ cm}^2$. However, the assumption that a kaolinite particle with an electrical mobility diameter of 300 nm is equal to a 300 nm diameter sphere is likely not true, due to shape irregularities, variable density, void fractions, multiple charges, and other geometries (DeCarlo et al., 2004;

- Slowik et al., 2004; Zelenyuk et al., 2006; Schmid et al., 2007; Park et al., 2008) with a tendency for greater surface area than assumed. Additionally, particles of larger diameter, and thus larger surface area, may have the same electrical mobility due to the presence of multiple charges. Therefore, a distribution of particle surface area can be expected. Following Wiedensohler and Fissan (1988), the probability for particles having multiple charges as a function of particle diameter, $P(\ln D_p)$, at a
- constant electrical mobility diameter of 300 nm is shown in Fig. S8. The distribution P(ln D_p) is a probability density function from which particle diameters are sampled in simulations IsoCFDC and IsoLACIS. Individual sampled particle surface area is calculated assuming spherical particles. We note that a lognormal distribution is not used in IsoCFDC and IsoLACIS. Instead, it is assumed that the ISA distribution varies only due to the theoretical multiple charge distribution. Parameters *m* and *c* used to calculate J_{het} for Fluka kaolinite are fitted.

Calculations of $J_{het}^{apparent}$ and J_{het}^{actual} assuming constant ISA equal to A_{300nm} or accounting for variable ISA, respectively, in IsoCFDC and IsoLACIS are shown in Fig. 7b. We find agreement between $J_{het}^{apparent}$, data by Wex et al. (2014) and J_{het}^{actual} which accounts for multiple particle charges predicted by Wiedensohler and Fissan (1988). Within the uncertainties presented here, assuming that the elec-

- 720 trical mobility diameter corresponds to the physical particle diameter and calculating surface area from a spherical geometry may be a valid assumption. Hence, ice nucleation experiments in which particles are mobility selected may be good examples of cases where ISA variability plays a minor role. Studies which use pre-impactors to remove larger sized particles, i.e. the selected size is larger than the median size of the total size distribution (Wex et al., 2014; Augustin-Bauditz. et al., 2014),
- 725 may even have a more narrow size distribution than used here. On the other hand, a recent study by Hartmann et al. (2016) derived the numbers of multiple charges on 300 nm mobility diameter size selected particles using simultaneous measurements of cloud condensation nuclei activation curves and total particle counts with a condensation particle counter. The authors found that when utilizing a pre-impactor, the multiple charge distribution of mobility diameter selected particles was larger than
- theoretical predictions (Hartmann et al., 2016). For comparison to the charge distribution used in Iso-LACIS and IsoCFDC shown in Fig. S9, we also plot the distribution measured by Hartmann et al. (2016) for the same particle type (Fluka kaolinite) used in (Wex et al., 2014). Hartmann et al. (2016) claimed that when correcting for their measured multiple charge distribution in experiments, values of $n_s(T)$ are shifted by 2K. We note that $J_{het}^{apparent}$ shown in Fig. 8b is about +1K shifted from
- 735 J_{het}^{actual} . This shift is smaller than observed in Hartmann et al. (2016) due to the fact that we applied the narrower theoretical distribution. Despite these issues, the model input J_{het} represents a new parameterization for Fluka kaolinite where m = 53.32 and c = -8.61 following the ABIFM applicable for $0.220 < \Delta a_w < 0.305$.

Wex et al. (2014) presented a detailed immersion freezing analysis of various kaolinite particle

- sizes and types of coatings and found that both stochastic and active site approaches can be applied to describe the data. Simulating all these cases using our model simulations is beyond the scope of this paper, however, we are certain that model simulations which use the same $J_{het}(T, a_w)$ will hold for all systems at all T and RH due to the prediction of immersion freezing kinetics (i.e. using J_{het}) being independent of experimentally applied ISA, particle size, and particle coating type (assuming
- 745 the coating dissolves when water is taken up and does not react with the INP surface). These findings demonstrate that our new model simulations and the ABIFM are applicable for ice nucleation studies using a CFDC as previously shown by Knopf and Alpert (2013) and additionally LACIS.

IFSs are used to describe AIDA chamber immersion freezing experiments applying natural dust by Niemand et al. (2012) in model simulations CrNI1 and CrNI2. Among the different types of nat-

- valuation ural dust investigated, we choose 2 Asian dust experiments at -20.1 < T < -28.1 °C and -14.3 < T < -22.4 °C (see ACI04_19 and ACI04_16 in Tables 2 and 3 in Niemand et al., 2012). A continuous non-linear cooling rate with time due to adiabatic expansion is fitted to experimental trajectories using a 4th order polynomial function. In AIDA experiments water saturation is typically reached after cooling begins. To mimic this process, ice particle production in model simulations is allowed
- after 80 s of cooling (see Fig. 2 in Niemand et al., 2012). Ice crystal concentration in an aerosol sampling flow of 5 Lmin^{-1} , from the chamber is observed every 5s using an optical particle counter (Benz et al., 2005), thus a volume of 0.42 L of air is simulated. Total particle numbers in the simulated volume are on the order of 10^5 which agree well with minimum reported f_{frz} of about 10^{-5} . Niemand et al. (2012) reported lognormal surface-size distributions with parameters, $d_{\text{S,median}}$ and
- 760 $\sigma_{\rm g}$ of polydisperse aerosol population. In CrNI1 and CrNI2, A_j is derived by sampling particle diameters from the corresponding number-size distributions and assuming spherical particles. We note that a fitted lognormal distribution is not used in CrNI1 and CrNI2, due to the fact that reported size distributions are well defined. Sampling stops when $A_{\rm tot}$ equals total surface area reported by Niemand et al. (2012). Experimentally derived $J_{\rm het}$ is not available and so the ABIFM parameters
- 765 m and c are fitted to experimentally derived f_{frz} data. Model simulation parameters for CrNI1 and CrNI2 are given in Table 2.

Figure 8 shows simulated $f_{\rm frz}$ and $J_{\rm het}^{\rm actual}$ from CrNI1 and CrNI2 and the time evolution of simulated ice crystal concentration in CrNI1 observed during the experiments. Simulated $f_{\rm frz}$ (Fig. 8a) fall within the experimental uncertainty reported by Niemand et al. (2012) and the scatter in the data

- for all dust types. Narrow 5 and 95% bounds are attributable to large N_{tot} on the order of 10^5 droplets per cooling simulation. Ice particle concentrations over time in CrNI1 are shown (insert in Figure 8b) and capture the overall observed trend in observations. This is in spite of the fact that the observed time evolution of ice crystal numbers was not used for fitting parameters m and c. Figure 8b shows $\overline{J}_{het}^{actual}$ and upper and lower fiducial limits. As frozen fraction decreases the fiducial limits become
- broader ranging from 0.8 to 2.5 orders of magnitude. We conclude that our model simulations are

suitable for describing laboratory immersion freezing in AIDA cloud chamber and further support the necessity of quantification of ISA variability in the derivation of ice nucleation kinetics.

Notice that in Fig. 8a, the vertical scatter in the experimental data increases at warmer T and for low f_{frz}, which implies that uncertainty likely increases as f_{frz} decreases. Since aerosol numbers and surface area in the experiments by Niemand et al. (2012) are relatively the same for the two exper-

- iments, decreasing f_{frz} implies fewer detected ice crystals or decreasing numbers of ice nucleation events resulting in an increase in experimental uncertainty. Immersion freezing due to natural dust was parameterized using a deterministic (singular) approach, i.e. using $n_s(T)$, which captured the trend in experimental results (Niemand et al., 2012). However, a deterministic approach for inter-
- pretation and analysis of ice crystal production, which inherently ignores stochastic freezing, cannot explain the increase in the data scatter for smaller $f_{\rm frz}$ values at warmer T. These observations can be explained by a stochastic and time-dependent immersion freezing process. We note that other measurement uncertainties may exist which may not be captured either by a deterministic approach or by our model. However, we conclude that stochastic uncertainty is important to consider for future
- 790 ice nucleation studies. The fiducial limits of J_{het}^{actual} shown in Fig. 8b, in fact, capture this effect of larger scatter as T increases implying the uncertainty in observed ice nucleation kinetics increases. Since the freezing efficiency of Asian dust was shown to be similar for Saharan, Canary Island, and Israeli dust (Niemand et al., 2012), the new ABIFM parameterization of $J_{het}(T, a_w)$ derived here is applicable for natural dust.

795 4 Simulation findings and uncertainty analysis

Our results strongly suggest that laboratory immersion freezing studies should provide accurate estimates of ISA variability in droplets. We find that simplified assumptions about ISA can result in misinterpretation and miscalculation of J_{het} values. This includes assuming identical surface area, which implicitly imposes a dependence of J_{het} on both ISA and r. Future laboratory immersion

- 800 freezing studies should also consider the stochastic nature of ice nucleation following CNT and resulting uncertainties. When only a single ice nucleation experiment is performed or too few droplets are used, stochastic uncertainty can potentially be very large and may limit data interpretation. Once again, stochastic uncertainty refers to large or small expected data scatter from observing small or large numbers of freezing events, respectively. The surface area based deterministic approach deriv-
- 805 ing $n_s(T)$ is an alternative to calculating J_{het} , but does not consider stochastic effects or effect of time in analysis of immersion freezing. By design, $n_s(T)$ should therefore, not have any dependence on r. However, this is not supported as $n_s(T)$ has been observed to be dependent on r for feldspar and kaolinite (Herbert et al., 2014).

The model simulation and laboratory data sets investigated here were performed for INPs im-810 mersed in pure water droplets. However, aqueous solution droplets having $a_w < 1.0$ are frequently present in the atmosphere at supercooled temperatures and subsaturated conditions (i.e. RH < 100%). The ABIFM (Eqs. 6–8) inherently and accurately accounts for these conditions and thus, provides a complete description of immersion freezing for laboratory experiments, as well as cloud models under atmospherically relevant *T* and RH. We suggest that future isothermal and cooling rate depen-

815 dent immersion freezing studies investigate aqueous solution droplets in addition to water droplets (e.g. Archuleta et al., 2005; Alpert et al., 2011b; Wex et al., 2014), providing additional data sets to constrain ice nucleation kinetics and to validate and expand ABIFM and other parameterizations.

Uncertainty analysis is crucial for the interpretation of laboratory immersion freezing results. Here we present a quantitative uncertainty analysis of J_{het} , by defining ΔJ_{het} as the total uncer-

tainty derived from individual contributions of statistical uncertainty due to N_{tot}, temperature accuracy referred to as ΔT, a_w or RH accuracy referred to as ΔRH, ISA variability expressed as σ_g, and accuracy of measuring absolute surface area referred to as ΔA_g. This uncertainty analysis is applicable to both isothermal and cooling rate dependent immersion freezing experiments. It is convenient to quantify ΔJ_{het} in the form of a [×]/_± error instead of a typical ± error due to J_{het} varying
exponentially over a linear range in T. If J_{het} = 100 cm⁻² s⁻¹ with a factor of ±3 error for example, then ΔJ_{het} =[×]/_± 3 equivalent to J_{het} = 100^{×3}/_{±3} = 100⁺²⁰⁰/₋₆₇ cm⁻² s⁻¹. In the following analysis, ΔJ_{het}

is quantified as $\stackrel{\times}{\underline{}}$, representing a factor error.

The uncertainty due to stochastic freezing is derived by running 10^5 IFSs with different values of $N_{\rm tot}$ and calculating $\Delta J_{\rm het}$ where the widths of the fiducial limits are smallest, i.e. at $f_{\rm frz} \simeq 0.5$. Thus,

- 830 ΔJ_{het} derived from N_{tot} yields the smallest error estimate possible or the limit of greatest experimental accuracy. Figure 9a illustrates that smaller N_{tot} results in larger ΔJ_{het} . When $N_{\text{tot}} = 30$ for example, $\Delta J_{\text{het}} = \stackrel{\times 15}{\div 5}$, and when $N_{\text{tot}} = 1000$, $\Delta J_{\text{het}} = \stackrel{\times 1.3}{\div 1.3}$. The uncertainty contribution due to ΔT is calculated using the slope of J_{het} vs. T following a similar procedure as in Riechers et al. (2013). Using the ABIFM at various temperature ranges and for different INP types (Knopf and Alpert, 2013),
- 835 J_{het} varies by a factor of 7.5 ± 5.5 per degree K. This means that if $\Delta T = \pm 1.0$ K, $\Delta J_{\text{het}} = \stackrel{\times}{\pm} 7.5$ on average, but can be $\stackrel{\times}{\pm} 2$ or $\stackrel{\times}{\pm} 13$ depending on the INP type and the range in T and RH. For example, $\Delta T = \pm 0.5$ K translates to $\Delta J_{\text{het}} = \stackrel{\times}{\pm} 3.75$ as displayed in Fig. 9a. Considering the uncertainty in RH, Eq. (6) is used to derive $\Delta J_{\text{het}} = J_{\text{het}}(\Delta a_{\text{w}})/J_{\text{het}}(\Delta a_{\text{w}} \pm \Delta \text{RH}) = 10^{m\Delta\text{RH}}$. Values of m in Eq. (6) are taken from this study and from Knopf and Alpert (2013) ranging from 15–123 and re-
- sults in 69 on average. The mean and range of ΔJ_{het} due to ΔRH are shown in Fig. 9b. For example, if $\Delta RH = \pm 3\%$, then $\Delta J_{het} = \stackrel{\times}{\pm} 117$ on average. If ISA per droplet varies in an experiment, but is assumed to be uniform, J_{het} is overestimated for $f_{frz} < 0.5$ and underestimated for $f_{frz} > 0.5$. This effect is quantified by allowing σ_g to vary and calculating the ratio $\Delta J_{het} = \overline{J}_{het}^{apparent} / \overline{J}_{het}^{actual}$ evaluated at $f_{frz} = 0.1$ and 0.9. The resulting ΔJ_{het} is displayed in Fig. 9c as a function of σ_g . If $\sigma_g = 10$,
- for example, then $\Delta J_{\text{het}} = \stackrel{\times 4}{\div}_{20}$ at $f_{\text{frz}} = 0.1$ and 0.9. Finally, ΔJ_{het} is directly proportional to ΔA_{g} shown in Fig. 9c, e.g. if $\Delta A_{\text{g}} = \stackrel{\times}{\div}_{2}^{\times}$ 5, then $\Delta J_{\text{het}} = \stackrel{\times}{\div}_{2}^{\times}$ 5.

Figure 9 demonstrates that each experimental parameter contributes to the uncertainty in J_{het} . The total uncertainty in J_{het} can then be estimated by summing the error contributions due to N_{tot} , T, RH, σ_g , and A_g , respectively. Figure 9 shows dotted lines serving as example values of exper-

- 850 imental uncertainties and corresponding ΔJ_{het} . Applying $N_{\text{tot}} = 30$, $\Delta T = \pm 0.5 \text{ K}$, $\Delta \text{RH} = \pm 3\%$, $\sigma_{\text{g}} = 10$, and $\Delta A_{\text{g}} = \stackrel{\times}{\pm} 5$, results in $\Delta J_{\text{het}} = \stackrel{\times}{\pm} \stackrel{148}{\pm} 154$. If laboratory immersion freezing studies were to be conducted under these conditions, then the range in experimentally derived J_{het} should be over 4 orders of magnitude. Notice that the uncertainty due to RH alone can potentially dominate the total uncertainty. We hope that Fig. 9 provides guidance in conducting future immersion freezing studies.
- We test our analysis to reproduce experimentally derived uncertainty. In Knopf and Alpert (2013), all experimentally derived J_{het} fell within ± 2 orders of magnitude as a function of the a_w criterion (Eq. 7) and as a result, this range was adapted as a conservative uncertainty estimate for the ABIFM model. The root mean square error of over 18 000 droplet freezing events for 6 different INP types was experimentally derived independent from model simulations, as an alternative uncertainty esti-
- mate exhibiting values as high as ±1.3 orders of magnitude. Experimental parameters of studies used in the formulation of the ABIFM for pure water and aqueous solution droplets (Alpert et al., 2011a, b; Knopf and Forrester, 2011; Rigg et al., 2013; Knopf and Alpert, 2013) were about N_{tot} = 300, ΔT = ±0.3 K, ΔRH = ±1%, σ_g = 5, and ΔA_g = [×]/₂ 5. Applying the analysis displayed in Fig. 9 results in an uncertainty of ΔJ_{het} = ^{×16}/₂₁₈ (spanning about 2.5 orders of magnitude) for the ABIFM
 model. This estimate is in excellent agreement with independently derived root mean square errors
- of J_{het} (Knopf and Alpert, 2013) and demonstrates the accuracy of our uncertainty analysis.

Model simulations reproduced observations of immersion freezing due to illite by Diehl et al. (2014) and Broadley et al. (2012). These experimental data were included in a recent intercomparison study of illite immersion freezing by Hiranuma et al. (2015). Using 17 different instruments,

- 870 experimentally derived $n_s(T)$ values were observed to increase from 10^{-3} to 10^8 cm^{-2} when T decreased from 263 to 236 K, equivalent to a slope of 0.5 orders of magnitude per 1 K. The instruments used are grouped by common methods and include, (i) cold stage (Broadley et al., 2012; Bingemer et al., 2012; Schill and Tolbert, 2013; Wright and Petters, 2013; O'Sullivan et al., 2014; Budke and Koop, 2015), (ii) liquid aliquots (Hill et al., 2014), (iii) droplet levitation (Szakáll et al.,
- 2009; Diehl et al., 2014; Hoffmann et al., 2013), (iv) cloud chamber (Möhler et al., 2003; Niemand et al., 2012; Tajiri et al., 2013) and (v) continuous flow (Bundke et al., 2008; Stetzer et al., 2008; Welti et al., 2009; Lüönd et al., 2010; Chou et al., 2011; Friedman et al., 2011; Hartmann et al., 2011; Kanji et al., 2013; Tobo et al., 2013; Wex et al., 2014). The scatter in the n_s is roughly 3 orders of magnitude, but depending on T, a n_s range of 2 and 4 orders of magnitude can envelop the data. However,
- the authors provided no quantitative uncertainty analysis to explain this scatter. Since experimental methods and data reproduced by presented model simulations are included in Hiranuma et al. (2015) for illite, we apply the quantitative uncertainty analysis presented in Fig. 9 to provide a potential explanation of the data scatter. We note that the abscissa in Fig. 9 extends to a value of ΔJ_{het} equal to

a factor of 300, to encompass typical uncertainties of about ± 2 orders of magnitude. Although, J_{het}

and $n_{\rm s}(T)$ are different quantities, the contribution to their uncertainties is the same for ΔT , ΔRH , $\sigma_{\rm g}$, $\Delta A_{\rm g}$.

Experimental T uncertainty for all methods typically ranged from ± 0.2 to ± 1.0 K, and hence $\Delta T = \pm 0.5$ is chosen as a representative value. Considering the slope n_s vs. T, $\Delta T = \pm 0.5$ contributes a factor of ~ 2 uncertainty to $n_s(T)$, or $\Delta n_s = \stackrel{\times}{\pm} 2$. The ISA distribution width parameter of

- 890 simulated experiments (Tables 1 and 2) is averaged to represent $n_s(T)$ data, yielding a reasonable value of $\sigma_g = 7$, resulting in $\Delta n_s = \stackrel{\times 3}{\div}_{\pm 12}$. The ISA measurement error is considered to be $\Delta A_g = \stackrel{\times}{\pm} 5$, thus $\Delta n_s = \stackrel{\times}{\pm} 5$. Calculation of $n_s(T)$ is not stochastic by design, and thus any uncertainty contribution due to N_{tot} on $n_s(T)$ was previously not considered (Hiranuma et al., 2015). Additionally, the intercomparison analysis ignores differences in experimental time scales in $n_s(T)$ derivation.
- However, this study demonstrates that the stochastic uncertainty may be able to explain immersion freezing data and may contribute to the range of data scatter in $n_s(T)$. Typically, N_{tot} is about 50 which serves as a reasonable representation yielding $\Delta n_s = \stackrel{\times 8}{\div 4}$, although N_{tot} can vary between 10 and more than 1000 depending on the experiment. Previous immersion freezing experiments for illite have shown that when r or residence time differ by 1 order of magnitude, freezing temperatures
- shift by about 0.75 K on average (Broadley et al., 2012; Welti et al., 2012; Knopf and Alpert, 2013). As discussed in Hiranuma et al. (2015), cooling rates and residence times in the different instruments varied over ± 2 orders of magnitude, or $\Delta t = \stackrel{\times}{\pm} 100$, corresponding to $\Delta T = \pm 1.5$ K, and thus contributing to an error of ± 0.75 orders of magnitude or $\Delta n_{\rm s} = \stackrel{\times}{\pm} 6$. Accounting for all uncertainties and making use of Fig. 9 results in $\Delta n_{\rm s} = \stackrel{\times(2+3+5+8+6)}{\div(2+12+5+4+6)}$ for a total uncertainty of $\Delta n_{\rm s} = \stackrel{\times 24}{\div}$, or an
- 905 uncertainty range of 2.8 orders of magnitude. The vast majority of data in Hiranuma et al. (2015) fall within this uncertainty and implies that variability in $n_s(T)$ may be attributed to experimental, timedependent, and stochastic uncertainties. It is important to note that the uncertainty due to neglecting time, ISA variability and stochastic effect contributes more to Δn_s , than T and ISA measurement error. Hiranuma et al. (2015) hypothesized that experimental procedures of droplet or particle prepa-
- 910 ration, including particle generation, size selection, ice crystal detection, particle loss at instrument sampling inlets, contamination, inhomogeneous temperature, and differences in surface cation concentration between wet dispersed or dry dispersed particles may be the cause in measured scatter in $n_s(T)$ data. These effects are not considered in the uncertainty analysis presented here, but may also contribute.

915 5 Summary and conclusions

Immersion freezing simulations based on a droplet resolved stochastic ice nucleation process applicable for various types of INPs and experiments are presented here for both isothermal conditions and applying a cooling rate, r. The parameters in the IFSs are all physically defined and measurable, including the heterogeneous ice nucleation rate coefficient, J_{het} , the number of droplets at the start of

- an experiment, N_{tot}, and the immersed surface area (ISA) per droplet. When knowledge of ISA per droplet is not known, it may be assumed to be lognormally distributed. For IFSs in which a cooling rate, r, is applied, J_{het} as a function of T and aqueous solution water activity, a_w, can be calculated following the water activity based immersion freezing model (ABIFM) applicable for both pure water (a_w = 1.0) and aqueous solution (a_w < 1.0) droplets. These IFSs generate frozen and unfrozen
 droplet fraction data, f_{ufz} and f_{frz}, respectively, and using a Monte Carlo method in which 10⁵ IFSs
 - are performed under the same conditions, 5 and 95% bounds are derived as uncertainty estimates.

The sensitivity of f_{ufz} on σ_g and N_{tot} was tested using sets of isothermal IFSs, where a single set is referred to as a model simulation. Uniform ISA (i.e. $\sigma_g = 1$) resulted in f_{ufz} (on a logarithmic scale) being linear with t. When ISA varied lognormally with parameters $\mu = \ln(A_g)$ and $\sigma = \ln(\sigma_g)$,

- 930 where $\sigma_g > 1$, $\ln(f_{ufz})$ vs. t exhibit non-linear behavior. When larger or smaller N_{tot} was used, f_{ufz} had a smaller and larger uncertainty, respectively, due to the statistical significance of observing more freezing events. These results demonstrate that in laboratory immersion freezing experiments, variable ISA imposes changes in trajectories of f_{ufz} and f_{frz} over time, and that the number of investigated droplets significantly impacts experimental uncertainty.
- Cooling rate model simulations were used to test the validity of assuming uniform ISA. This was accomplished by recalculating J_{het} after simulation of immersion freezing in two ways, either (i) assuming uniform ISA referred to as the "apparent" ice nucleation rate coefficient, $J_{het}^{apparent}$, or (ii) accounting for variable ISA referred to as the "actual" ice nucleation rate coefficient, J_{het}^{actual} . When different r were applied in simulations, values of $J_{het}^{apparent}$ were significantly different from
- 940 each other. When comparing experiments with different ISA but identical r, $J_{het}^{apparent}(T)$ was again significantly different. For $f_{frz} < 0.5$ and $f_{frz} > 0.5$, $J_{het}^{apparent}$ was over and underestimated, respectively, compared to J_{het}^{actual} , yielding an erroneous slope of $J_{het}^{apparent}(T)$. These results demonstrate that the assumption of identical ISA implicitly imposes a cooling rate and surface area dependence on experimentally derived $J_{het}(T)$. However, derivation of J_{het}^{actual} from model simulations accounting
- for variable ISA were consistent for different r and ISA, supporting a stochastic immersion freezing description as predicted by CNT.

Model simulations in which variable ISA was considered reproduced laboratory experiments using Arizona test dust (ATD) (Wright and Petters, 2013), illite (Broadley et al., 2012; Diehl et al., 2014), kaolinite (Wex et al., 2014; Herbert et al., 2014), feldspar (Herbert et al., 2014), and natu-

- acting as INPs. Despite whether isothermal or linear and nonlinear cooling rates were applied, modeled and experimental f_{frz} and f_{ufz} were in agreement within the stochastic uncertainty. More importantly, experimentally derived $J_{\text{het}}(T)$ and simulated $J_{\text{het}}^{\text{apparent}}$ were in agreement for ATD, illite, kaolinite and feldspar possible indicating an imposed bias solely due to the assumption of uniform ISA
- and not to physical processes governing ice nucleation. On the other hand, variability of ISA in

experimental studies using size selected particles from a differential mobility analyzer (Wex et al., 2014), were modeled based on a bipolar charge distribution. There was a 1K difference between J_{het} assuming uniform ISA and accounting for variable ISA following the bipolar charge distribution, but this was within stochastic uncertainty limits. Thus, the assumption of identical ISA may be

960 valid when size selecting particles. However, the actual ISA distribution for each study should be measured to verify this assumption about ISA in droplets in order to correctly interpret immersion freezing (Hartmann et al., 2016). In general, model simulations can correct for any introduced bias or calculate "actual" values, or J_{het}^{actual} , when not provided. J_{het}^{actual} resulted in consistent agreement between different studies and additionally new a_w based parameterizations of $J_{het}(\Delta a_w)$ for feldspar

965 and natural dusts.

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A quantitative uncertainty analysis of J_{het} was presented applicable for experimental studies in which the contribution due to (i) N_{tot} , (ii) temperature accuracy referred to as ΔT , (iii) a_w or RH accuracy referred to as ΔRH , (iv) σ_g , and (v) the accuracy of A_g referred to as ΔA_g , were individually quantified. The following points summarize these error sources and give recommendations for future experimental studies:

- Applying too few N_{tot} or performing only a single ice nucleation experiment in laboratory studies results in highly uncertain freezing results. Therefore, repetition of immersion freezing experiments or a statistically significant number of droplets must be applied. We recommend using at least 100 droplets and three independent freezing cycles in order to better quantify data scatter and average J_{het} , f_{frz} , and f_{ufz} values. This contributes to a range of 0.75 orders of magnitude in the uncertainty of experimentally derived J_{het} .
- For different INP types, the slope of J_{het} vs. T is not the same and thus, the uncertainty due to ΔT is INP type dependent, but can be as high as 1 order of magnitude per 1 K. We recommended that ΔT remain $< \pm 0.5$ K to achieve an acceptable uncertainty contribution, i.e. half an order of magnitude.
- The greatest source of error stems from RH, or Δ RH. Immersion freezing experiments for RH < 100% should aim for Δ RH to be as small as possible. Current and future immersion freezing experiments should be designed to carefully control RH and quantify its uncertainty.
- Droplets in laboratory immersion freezing experiments will not have identical ISA, but will
 vary from droplet to droplet (σ_g) around some ISA value (A_g). Variability in ISA and corresponding uncertainty should be quantified and accounted for when analyzing ice nucleation experiments.
 - Surface area and nucleation time scales clearly affect immersion freezing data. Common assumptions of ISA and neglecting the impact of variable experimental time scales will lead to

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an incomplete experimental accuracy and uncertainty. Consideration of these effects is recommended to narrow the uncertainty in predicting ice crystal formation.

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Considering that ice nucleating particles have variable ISA may impact atmospheric ice crystal numbers. For a broad surface area distribution of INPs, ice nucleation should occur over a broader range of time and temperature, when compared with a narrow INP surface area distribution. This

- 995 results in greater ice particle production at warmer temperatures, important for mixed phase cloud formation and their evolution. We suggest that field measurements determine and consider the entire aerosol size distribution as a source of INPs for implementation of a stochastic, time-dependent ice nucleation process characterized by J_{het} , which is easily parameterized following the ABIFM in addition to current methodologies.
- 1000 Our findings concerning laboratory immersion freezing experiments emphasize the importance of setting constraints on the minimum number of droplets and experimental trials that need to be employed for improved characterization of ISA per droplet. The results presented here resolves commonly used assumptions that contribute to additional uncertainty in predicting immersion freezing data for model implementation. The simulations use ABIFM, shown to be valid for various INP
- 1005 types. We demonstrate that the ABIFM can reproduce immersion freezing by mineral dust for many vastly different experimental designs and measurement methods. Laboratory derived J_{het} values can aid in testing existing ABIFM parameterizations and formulating new ones. Their application to a very simple stochastic freezing model based on a binomial distribution in accordance with classical nucleation theory, can reconcile immersion freezing data for various INP types and measure-
- 1010 ment techniques when the applied INP surface areas are treated more realistically. These findings hopefully stimulate further discussion on the analytical procedure and interpretation of immersion freezing and its implementation in atmospheric cloud and climate models.

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Name	N_{tot}	$\sigma_{ m g}$	$A_{\rm g}/{ m cm}^2$	T / K	$J_{\rm het}$ cm ⁻² s ⁻¹	INP Type	Figure	Color
Iso1	1000	1	1.0×10^{-5}	_	1.0×10^3	_	1a	dark green
Iso2	30	1	1.0×10^{-5}	_	1.0×10^3	_	1a	light green
Iso3	1000	10	1.0×10^{-5}	-	1.0×10^3	_	1a	dark blue
Iso4	30	10	1.0×10^{-5}	-	1.0×10^3	_	1a	light blue
IsoWR	1000	9.5	6.4×10^{-3}	251.15	6.0×10^{-4}	ATD^{a}	1b	orange
IsoBR	63	$\mathrm{U} ext{-}\mathrm{pdf}^\mathrm{b}$	2.6×10^{-7}	243.3	2.8×10^3	illite	2a	orange
IsoHE1	40	2.2	1.2×10^0	255.15	4.1×10^{-3}	kaolinite	2b	orange
IsoHE2	40	8.5	2.0×10^{-2}	262.15	2.0×10^{-2}	feldspar	2c	orange
IsoDI1	45	3.2	5.1×10^{-1}	255.15	1.8×10^{-2}	illite	3	green
IsoDI2	45	3.2	5.1×10^{-2}	252.15	1.0×10^0	illite	3	orange
IsoDI3	45	3.2	5.1×10^{-1}	252.15	1.0×10^0	illite	3	blue
IsoCFDC	833	$\mathrm{MCD}^{\mathrm{c}}$	MCD	238.65 -	$\operatorname{ABIFM}^{\operatorname{e}}$	kaolinite	7	orange, black
				$247.65^{\rm d}$				
IsoLACIS	21	MCD	MCD	235.65 -	ABIFM	kaolinite	7	blue, green
				$238.65^{\rm d}$				

Table 1. Summary of parameters used in isothermal model simulations.

^a Arizona Test Dust.

^b A uniform probability density function (U-pdf) was used to define the surface area distribution centered on 2.6×10^{-7} cm², with distribution

endpoints at 9.4×10^{-8} and 7.5×10^{-7} cm². See text and Fig. S3 for further details.

^c A multiple charge distribution (MCD) was used to define the surface area distribution. See text and Fig. S9 for further details.

 $^{\rm d}$ Isothermal simulations were performed at $0.15\,\rm K$ increments within the stated temperature range.

^e Values of J_{het} are calculated from the water activity, a_w , based immersion freezing model (ABIFM) where m = 53.32 and c = -8.61 (Knopf and Alpert, 2013).

Name	$N_{ m tot}$	$\sigma_{ m g}$	$A_{\rm g}/{\rm cm}^2$	m	с	$r /\mathrm{Kmin^{-1}}$	INP Type	Figure	Color
Cr1	1000	10	1.0×10^{-5}	54.48	-10.67	0.5	illite	4	orange
Cr2	1000	10	1.0×10^{-5}	54.48	-10.67	5.0	illite	4	blue
Cr3	1000	1	1.0×10^{-5}	54.48	-10.67	0.5	illite	4	black
Cr4	1000	1	1.0×10^{-5}	54.48	-10.67	5.0	illite	4	green
CrHE1	40	8.5	2.1×10^{-2}	122.83	-12.98	0.2	feldspar	5	orange
CrHE2	40	8.5	2.1×10^{-2}	122.83	-12.98	2.0	feldspar	5	blue
CrDI1	45	5.7	2.9×10^{0}	54.48	-10.67	non-linear ^a	illite	6	orange
CrDI2	45	5.7	2.9×10^{-2}	54.48	-10.67	non-linear ^a	illite	6	blue
	$A_{\rm tot}/{ m cm}^2$		$D_{\rm p,g} / \mu{ m m}$						
CrNI1	6.5×10^{-4}	1.72	0.42	22.62	-1.35	non-linear $^{\rm b}$	ND ^c	8	blue
CrNI2	5.4×10^{-4}	1.69	0.40	22.62	-1.35	non-linear $^{\rm b}$	ND^{c}	8	orange

Table 2. Summary of parameters used in cooling rate model simulations.

 $^{\rm a}$ A continuous non-linear cooling rate with time is given in Diehl et al. (2014).

^b A continuous non-linear cooling rate with time due to adiabatic expansion is fitted to experimental trajectories (Niemand et al., 2012) using a 4th order polynomial.

 $^{\rm c}$ Natural dusts from Niemand et al. (2012): Asian, Saharan, Israeli and Canary Island dust.



Figure 1. Sensitivity calculations of the unfrozen droplet fraction, f_{ufz} , as a function of time, t, derived from model simulations for a total number of droplets, N_{tot} , and variability of ice nuclei surface area, σ_g . (a) Model simulated 5 and 95% bounds of f_{ufz} are shown as dark green (Iso1), light green (Iso2), dark blue (Iso3), and light blue (Iso4) shading. Parameter values are given in the legend. (b) Simulated 5 and 95% bounds of f_{ufz} derived from IsoWR are shown as the orange shading along with experimental data of isothermal immersion freezing by Arizona test dust (Wright and Petters, 2013) shown as black circles. Parameter values for all model simulations in (a) and (b) are given in Table 1.



Figure 2. Simulated and experimentally (Broadley et al., 2012; Herbert et al., 2014) derived unfrozen droplet fractions, f_{ufz} , as a function time, t. Model simulations and INP types used are: (a) IsoBR and illite, (b) IsoHE1 and kaolinite, and (c) IsoHE2 and feldspar, respectively. Orange lines and shading represent \overline{f}_{ufz} and corresponding 5 and 95% bounds, respectively. Parameter values for model simulations are given in Table 1.



Figure 3. Simulated and experimentally (Diehl et al., 2014) derived unfrozen droplet fractions, f_{ufz} , as a function time, t, using illite. Model simulated 5 and 95% bounds of f_{ufz} are shown as green, orange and blue shading for IsoDI1, IsoDI2 and IsoDI3, respectively. Temperature and average surface area per droplet reported by Diehl et al. (2014) are given in the legend. Parameter values for model simulations are given in Table 1.



Figure 4. Sensitivity calculations of heterogeneous ice nucleation rate coefficients, J_{het} , and frozen droplet fractions, f_{frz} , on cooling rate, r, derived from model simulations Cr1 (orange) and Cr2 (blue) where r = 0.5 and 5.0 K min^{-1} , respectively. J_{het} as a function of temperature, T, are shown in (**a**) assuming uniform ice nuclei surface area (ISA) per droplet yielding $J_{het}^{apparent}$, and (B) accounting for different ISA yielding J_{het}^{actual} . The dashed lines in (**a**) and (**b**) are $\overline{J}_{het}^{apparent}$ and $\overline{J}_{het}^{actual}$, respectively. Shadings in (**a**) and (**b**) correspond to upper and lower fiducial limits with x = 0.999 confidence and the solid red line is calculated from Eq. (6) for illite (Knopf and Alpert, 2013). Frozen droplet fractions, f_{frz} , are shown in (**c**) and (**d**) where dashed lines and shadings represent \overline{f}_{frz} and 5 and 95% bounds, respectively. Parameter values for Cr1 and Cr2 are given in Table 2.



Figure 5.

Figure 5. Frozen droplet fractions, $f_{\rm frz}$, and heterogeneous ice nucleation rate coefficients, $J_{\rm het}$, from immersion freezing cooling rate, r, dependent model simulations CrHE1 and CrHE2 where r = 0.2 (orange) and 2.0 K min⁻¹ (blue), respectively, and experimental data of feldspar acting as immersion INPs (Herbert et al., 2014). Dashed lines and shadings in (**a**) are $\overline{f}_{\rm frz}$ and 5 and 95% bounds, respectively. $J_{\rm het}$ as a function of temperature, T, are shown in (**b**) assuming uniform ice nuclei surface area (ISA) per droplet yielding $J_{\rm het}^{\rm apparent}$ and (**c**) accounting for variable ISA yielding $J_{\rm het}^{\rm actual}$. The dashed lines in (**b**) and (**c**) are $\overline{J}_{\rm het}^{\rm apparent}$ and $\overline{J}_{\rm het}^{\rm actual}$, respectively. Shadings in (**b**) and (**c**) correspond to upper and lower fiducial limits with x = 0.999 confidence. Experimentally derived $f_{\rm frz}$ and $J_{\rm het}$ are shown as circles in (**a**) and (**b**), respectively (Herbert et al., 2014). The red line in (**b**) and (**c**) is calculated from Eq. (6) (Knopf and Alpert, 2013) using new parameters derived for feldspar. Parameter values for CrHE1 and CrHE2 are given in Table 2. The fitted $J_{\rm het}$ value from model simulation IsoHE2 is shown in (**c**) and its corresponding error derived from Fig. 9 considering $N_{\rm tot} = 40$, $\sigma_{\rm g} = 8.5$ (see Table 1) and a temperature error, $\Delta T = \pm 0.4$ K.



Figure 6.

Figure 6. Frozen droplet fractions, $f_{\rm frz}$, and heterogeneous ice nucleation rate coefficients, $J_{\rm het}$, from immersion freezing model simulations CrDI1 (orange) and CrDI2 (blue), and experimental data of illite acting as immersion INPS are shown (Diehl et al., 2014). Dashed lines and shadings in (**a**) are $\overline{f}_{\rm frz}$ and 5 and 95% bounds, respectively. $J_{\rm het}$ as a function of temperature, T, are shown in (**b**) assuming uniform ice nucleating particle surface area (ISA) per droplet yielding $J_{\rm het}^{\rm apparent}$ and (**c**) accounting for variable ISA yielding $J_{\rm het}^{\rm actual}$. The dashed lines in (**b**) and (**c**) are $\overline{J}_{\rm het}^{\rm apparent}$ and $\overline{J}_{\rm het}^{\rm actual}$, respectively. Shadings in (**b**) and (**c**) correspond to upper and lower fiducial limits with x = 0.999 confidence. Experimentally derived $f_{\rm frz}$ and $J_{\rm het}$ are shown as circles in (**a**) and (**b**), respectively (Diehl et al., 2014). The red line in (**b**) and (**c**) is calculated from Eq. (6) for illite (Knopf and Alpert, 2013). Parameter values for CrDI1 and CrDI2 are given in Table 2. Fitted $J_{\rm het}$ values from model simulations IsoDI1 to 3 are shown in (**c**) and their corresponding error derived from Fig. 9 considering $N_{\rm tot} = 45$, $\sigma_{\rm g} = 3.2$ and a temperature error, $\Delta T = \pm 0.7$ K (see Table 1).



Figure 7. Frozen droplet fractions, $f_{\rm frz}$, and heterogeneous ice nucleation rate coefficients, $J_{\rm het}$, from isothermal model simulations IsoCFDC (orange and black) and IsoLACIS (blue and green), and experimental data of immersion freezing due to kaolinite by Wex et al. (2014) are shown. Dashed lines and shadings in (**a**) are $\overline{f}_{\rm frz}$ and 5 and 95% bounds, respectively. $J_{\rm het}$ as a function of temperature, T, are shown in (**b**) assuming uniform ice nucleating particle surface area (ISA) per droplet yielding $J_{\rm het}^{\rm apparent}$, and accounting for variable ISA yielding $J_{\rm het}^{\rm actual}$. The dashed lines in (**b**) are $\overline{J}_{\rm het}^{\rm apparent}$ and $\overline{J}_{\rm het}^{\rm actual}$ as indicated in the legend. Shadings in (**b**) correspond to upper and lower fiducial limits with x = 0.999 confidence and the red line is calculated from Eq. (6) for kaolinite (Knopf and Alpert, 2013). Parameter values for IsoCFDC and IsoLACIS are given in Table 1.



Figure 8. Frozen droplet fractions, $f_{\rm frz}$, and heterogeneous ice nucleation rate coefficients, $J_{\rm het}$, derived from adiabatic cooling immersion freezing model simulations CrNI1 (blue) and CrNI2 (orange). Simulated and experimentally observed ice crystal concentrations are shown in the insert of panel (**a**). Dashed lines and shadings in (**a**) are $\overline{f}_{\rm frz}$ and 5 and 95% bounds, respectively. Experimentally derived $f_{\rm frz}$ and uncertainties by Niemand et al. (2012) are shown as symbols and error bars. $J_{\rm het}$ as a function of temperature, T, is shown in (**b**) and accounting for variable ISA yielding $J_{\rm het}^{\rm actual}$, where dashed lines and shading are $\overline{J}_{\rm het}^{\rm actual}$ and fiducial limits with x = 0.999 confidence, respectively. The red line in (**b**) is calculated from Eq. (6) using new parameters derived for natural dust. Parameter values for CrNI1 and CrNI2 are given in Table 2.



Figure 9. Uncertainty analysis derived from immersion freezing model simulations. The relative error in the experimentally derived heterogeneous ice nucleation rate coefficient, J_{het} , is referred to as ΔJ_{het} . The y axis indicates ΔJ_{het} as a factor error, e.g. $\Delta J_{het} = 10$ indicates an error in J_{het} by a factor of 10 in the positive and negative direction. stochastic error due to the applied number of droplets, N_{tot} , is shown in (a) where red and blue represent the upper and lower fiducial limits of J_{het} , respectively. The error due to temperature accuracy, ΔT , for a variety of INP types is shown in (a) in orange color where the solid line is average ΔJ_{het} as a function of ΔT and the shading is for a range of INP types. The error due to the absolute uncertainty in water activity or equivalently relative humidity, ΔRH , is shown in (b) where the blue line is average ΔJ_{het} , and the shading represents the range of values for a variety of INP types. The uncertainty due to variability in INP surface area, σ_g , is shown in (c) as black and green lines evaluated at $f_{frz} = 0.1$ and 0.9, respectively. The uncertainty values given as dotted lines are described in the text.