

An aerosol chamber investigation of the heterogeneous ice nucleating potential of refractory nanoparticles

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Abstract. Nanoparticles of iron oxide (crystalline and amorphous), silicon oxide and magnesium oxide were investigated for their propensity to nucleate ice over the temperature range 180–250 K, using the AIDA chamber in Karlsruhe, Germany.

All samples were observed to initiate ice formation via the deposition mode at threshold ice super-saturations (RHithresh) ranging from 105% to 140% for temperatures below 220 K. Approximately 10% of amorphous Fe₂O₃ particles (modal diameter = 30 nm) generated in situ from a photochemical aerosol reactor, led to ice nucleation at $RHi_{thresh} = 140\%$ at an initial chamber temperature of 182 K. Quantitative analysis using a singular hypothesis treatment provided a fitted function $[n_s(190K) = 10^{(3.33 \times s_{ice}) + 8.16}]$ for the variation in ice-active surface site density $(n_s:m^{-2})$ with ice saturation (sice) for Fe2O3 nanoparticles. This was implemented in an aerosol-cloud model to determine a predicted deposition (mass accommodation) coefficient for water vapour on ice of 0.1 at temperatures appropriate for the upper atmosphere. Classical nucleation theory was used to determine representative contact angles (θ) for the different particle compositions. For the in situ generated Fe₂O₃ particles, a slight inverse temperature dependence was observed with $\theta = 10.5^{\circ}$ at 182 K, decreasing to 9.0° at 200 K (compared with 10.2° and 11.4° respectively for the SiO₂ and MgO particle samples at the higher temperature).

These observations indicate that such refractory nanoparticles are relatively efficient materials for the nucleation of ice under the conditions studied in the chamber which correspond to cirrus cloud formation in the upper troposphere. The results also show that Fe_2O_3 particles do not act as



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ice nuclei under conditions pertinent for tropospheric mixed phase clouds, which necessarily form above ~ 233 K. At the lower temperatures (<150 K) where noctilucent clouds form during summer months in the high latitude mesosphere, higher contact angles would be expected, which may reduce the effectiveness of these particles as ice nuclei in this part of the atmosphere.

1 Introduction

Insoluble refractory material, in the form of primary and secondary aerosol, is thought to play a crucial catalytic role in the activation of ice crystal formation in cirrus clouds in the troposphere and in polar stratospheric clouds (PSCs) in the stratosphere (Pruppacher and Klett, 1997; Zondlo et al., 2000). In addition, secondary aerosol, commonly referred to as *meteoric smoke*, which forms from recondensed material derived from ablating meteoroids in the upper mesosphere, is the favoured candidate for the production of high altitude ice clouds (83 - 85 km) which occur at high latitudes during summer, and are known as noctilucent clouds (NLCs) (also termed polar mesospheric clouds or PMCs) (Gadsden and Schroder, 1989; Reid, 1997; Rapp and Thomas, 2006).

Whilst the presence of meteoritic aerosol at lower altitudes has been substantiated through direct sampling from aircraft in the 1960's (Mossop, 1963) and, more recently, via sampling and in situ measurements (Murphy et al., 1998; Cziczo et al., 2001, 2004; Curtius et al., 2005), the existence, although long predicted from theoretical and modelling work (Rosinski and Snow, 1961; Hunten et al., 1980), let alone the participation in ice cloud nucleation, of meteoric smoke in the mesosphere awaits definitive evidence. A series of rocket-borne sampling devices launched in the 1960's did recover a variety of particulate samples from the upper atmosphere (Hemenway et al., 1963; Farlow et al., 1970) which showed a range of sizes, morphologies and compositions, although whether any of these samples could be classed as meteoric smoke remains unanswered. However, two groups of collected particles, labelled "type 2" and "type 5" in the paper of Farlow et al. (1970), did show an aggregated nature and extended morphology, possibly indicative of meteoric smoke. More recently, in situ measurements of positively charged particles at 80-90 km (Rapp et al., 2005) and satellite solar occultation measurements (Hervig et al., 2009) have lent further support to the presence of nano-sized meteoric smoke in the upper atmosphere, whilst line features observed in radar spectra have been shown to be consistent with the effect of charged particles present in the upper atmosphere (Strelnikova et al., 2007).

The classical heterogeneous nucleation theory of Fletcher (1958, 1959, 1969), based upon thermodynamic considerations of ice crystal activation and the role of particle size and surface inhomogeneity i.e. preferential activation sites, is applied primarily to spherical particles of crystalline materials e.g. silver iodide, for which a close "lattice fit" with hexagonal ice I_h was originally proposed as the primary reason for the high activation efficiency of this material. However, this scenario is somewhat idealised and it seems more likely that meteoric smoke particles are porous or "fluffy" aggregates of spherical, 'primary' particles of amorphous material such as iron oxide or silicate (Saunders and Plane, 2006). In addition, the formation of the metastable cubic phase of ice I_c in the Earth's atmosphere (Whalley, 1983) has recently been shown to be likely to occur more readily than previously thought in the lower atmosphere via homogeneous freezing (Murray et al., 2005). It therefore seems likely that in the mesosphere, where temperatures well below 150K have been regularly measured at high latitudes in the summer mesopause region (von Zahn and Meyer, 1989; Lübken, 1999), where NLC formation is typically observed, and where significant water supersaturations $(S \gg 1)$ have also been identified (Seele and Hartogh, 1999), ice I_c formation through heterogeneous nucleation could occur. On this very point, the study of Roddy (1984) specifically addressed the favourable lattice fits of possible meteoric materials of cubic structure (i.e. FeO, MgO). However, such a consideration alone is insufficient to account for the ice nucleating potential of a material (Evans, 1965), particularly if it is of a non-crystalline nature. Indeed, another possibility is that of amorphous ice formation, which to date has not been observed in the Earth's upper atmosphere but has been shown to be relatively common in other environments such as outer planetary satellites and comets (Baragiola, 2003). In light of these considerations, theoretical predictions of ice particle nucleation and activation based upon the assumption of hexagonal ice formation on crystalline materials are, in the realm of the upper atmosphere, likely to be somewhat erroneous. It is therefore preferable, where possible, to directly measure the ice nucleation potential of candidate materials of realistic composition and morphology under conditions approaching those which exist in the atmosphere.

In terms of the heterogeneous ice nucleating potential of meteoric smoke, nothing exists in the current literature regarding the properties of the most likely aerosol species (metal and silicon oxides and metal silicates i.e. olivines) under conditions relevant to the upper atmosphere and at nanoparticle dimensions (< 10 nm), expected to be present at such high altitudes. Whilst deposition nucleation studies, with regard to the lower atmosphere, for materials of terrestrial origin such as mineral dusts (Archuleta et al., 2005; Möhler et al., 2006) and soot (Dymarska et al, 2006; Kärcher et al., 2007) are of great current interest, only the experimental study of Bigg and Giutronich (1967) seems to have considered the activating role of recondensed meteoric material in the atmosphere, and only then with respect to the troposphere. Other attempts to investigate the freezing capacity of meteoritic material, such as that of Biermann et al. (1996) with regard to acid-water solutions in the stratosphere, have reported a largely negative (inactive) potential, but the samples used (whole or ground micrometeorites) are unlikely to be representative of the composition and morphology of recondensed (smoke) particles, and consequently exhibit the appropriate particle surface and nucleation properties. This point was argued by Bigg and Giutronich (1967) with respect to earlier cited nucleation studies.

Although the study of Bigg and Giutronich indicated that low density particle aggregates, formed via the boiling of a chondritic meteorite, required relatively low threshold ice supersaturations at ~ 263 K, the study of heterogeneous nucleation under the extreme conditions encountered in the mesosphere, obviously presents a much greater experimental challenge.

This was the motivation that led us to conduct a series of low temperature nucleation studies on a variety of commercially available crystalline and amorphous nanopowders of potential meteoric smoke compositions, as well as a laboratory-generated amorphous iron oxide powder, and nanoparticles of the same iron oxide formed in situ using a photochemical aerosol system. The work was conducted at the AIDA (Aerosol Interactions and Dynamics in the Atmosphere) chamber in Karlsruhe, Germany. This is an 84.3 m³ aluminium chamber in which temperatures down to ~ 183 K and pressures less than 1 Pa can be attained for the investigation of the deposition mode ice nucleation of aerosol particles injected into the chamber volume (Möhler et al., 2006; Field et al., 2006; Cotton et al., 2007). The facility at the Karlsruhe Institute of Technology (KIT) includes a suite of analytical techniques for the in situ or on line monitoring of particle size distribution, ice crystal growth and light scattering along with continual monitoring of the ambient pressure, temperature and supersaturation levels with respect to both water and ice in the chamber, throughout the course of an experimental run.

Our primary aim in these experiments was to identify the critical threshold conditions of temperature and ice supersatuation under which heterogeneous ice nucleation is activated by the particle samples described and, if possible, to extend the analysis, via modelling, to include a derivation of ice formation rates in order to establish the likely role of such materials as ice nuclei (IN) in the lower, middle and upper atmosphere, with regard to their proposed participation in cirrus, polar stratospheric and noctilucent cloud formation.

2 Meteoric Smoke Particles (MSPs) and their experimental analogues

The compositions of the various particle samples were chosen to most closely reflect those thought to comprise meteoric smoke. These nanoparticles are formed in the upper mesosphere by the condensation of predominantly Fe, Mg and Si oxides (+ mixed silicates i.e. olivine minerals). The metal and silicon atoms are deposited in the atmosphere as a result of the ablation or frictional heating of meteors as they pass through the Earth's upper atmosphere (Vondrak et al., 2008). The particles are formed by homogeneous nucleation, grow through coagulation, are transported to lower altitudes by sedimentation and winds and can ultimately be deposited out at the surface. Iron oxide (FeO, Fe₂O₃ and Fe₃O₄) particles of extraterrestrial origin have been detected in air samples (Kopcewicz & Kopcewicz, 1996) and ice cores (Lanci et al., 2007) whilst silica (SiO₂) particles of both terrestrial and extraterrestrial origin have been implicated in the heterogeneous nucleation of stratospheric aerosol (Bogdan and Kulmala, 1999; Bogdan, 2000). As for magnesium oxide, no direct detection of this composition has to date been made in MSPs but laboratory work on oxidation pathways of Mg atoms (Plane & Helmer, 1995) and consideration of similar chemical processes in circumstellar environments (Köhler et al., 1996) indicate a possible role for MgO nucleation.

Two separate experimental sessions were performed at the AIDA chamber. The first took place over the period of a week in December 2007 (all data sets for this session are referred to in the following discussion by the prefix IN11). These experiments focused on an inter-comparison of the ice nucleation potential of 3 different forms of iron oxide (Fe₂O₃) powders: (i) an amorphous, laboratory-generated powder collected from the photo-oxidation of iron pentacarbonyl $\{Fe(CO)_5\}$ vapour in the presence of ozone in an aerosol reactor system at the University of Leeds (Saunders and Plane, 2006); (ii) a high purity (99.999%, Sigma-Aldrich 529311) crystalline sample and (iii) a sample of 'pseudocubic' Fe₂O₃ synthesised by a gel-sol process (Sugimoto and Sakata, 1992) in Karlsruhe. Figure 1 (top panel) shows an image of photo-chemically generated particles collected on a holey carbon TEM grid. The pseudocubic sample consisted of monodisperse crystalline particles with an edge length of \sim 0.6 microns (Fig. 1, bottom panel) – the interested reader is

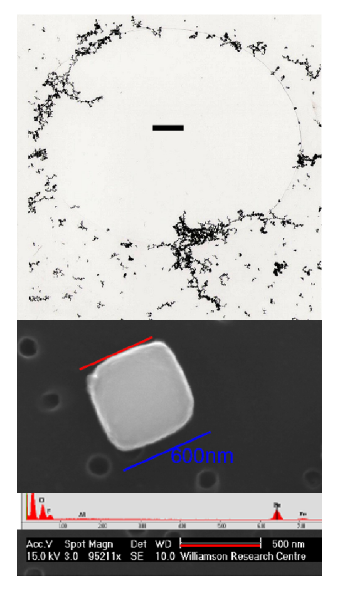


Fig. 1. (Top) Electron micrograph of amorphous iron oxide particles generated photo-chemically (scale bar in centre represents 2 microns). Deposited powder (referred to in the text as "IronOxide1") and in-situ generated particles were both used in ice nucleation experiments. (Bottom) Scanning electron micrograph and xray spectrum of a single crystal of the pseudocubic hematite sample described in Sect. 2.

directed to Park et al. (1996) for a detailed discussion of the morphology of such particles.

The second session at the chamber took place over two weeks in June 2008. All data sets for this session are referred to in the following discussion by the prefix IN12. For these experiments, commercial powders of MgO and SiO₂ (Sigma Aldrich 549649 & S5130 respectively), and nanoparticles of Fe₂O₃ composition, generated in situ adjacent to the chamber using the same aerosol system as mentioned previously, were used for nucleation tests.

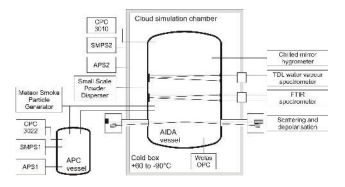


Fig. 2. Block schematic of the chamber and instrument set-up used to study the ice nucleating properties of the refractory nanoparticle samples – see Sect. 3 for definitions of all acronyms shown in the figure.

3 Experimental procedure and instrumentation

Figure 2 is a schematic diagram of the AIDA chamber and diagnostic instruments used for the experiments discussed here. Ice crystals are formed and observed in the central cylindrical chamber (4 m diameter, 84 m³ volume), which is encased in a large thermally insulated box. Prior to each series of ice nucleation experiments, the chamber was evacuated to a pressure below 0.1 hPa, flushed several times between 1 and 10 hPa, and filled with particle-free dry synthetic air. This procedure kept the aerosol background concentration typically below 0.1 cm^{-3} . The chamber system was humidified by adding a certain amount of water vapour to the evacuated chamber to achieve partial ice coverage on the inner chamber walls. At constant temperature and pressure conditions, the ice coverage maintains a high relative humidity between 90 and 95% in the chamber volume, which is stirred with a mixing fan operated in the bottom part of the chamber.

Aerosols were either added directly to the AIDA chamber or first added to the 3.7 m³ large stainless steel Aerosol Preparation and Characterisation (APC) chamber and then partially transferred into the larger AIDA chamber for the ice nucleation experiments. The APC chamber was also used to test the aerosol generators and to comprehensively investigate and characterise the resulting aerosols. During IN11, dry powder samples were dispersed with a Small Scale Powder Disperser (SSPD, TSI Inc.) and added directly to the AIDA chamber. During IN12, small agglomerates of nanometer-sized oxide particles were generated in situ as surrogates of meteor smoke particles and added to both the APC and AIDA chambers for aerosol characterisation and ice nucleation experiments, respectively. Aerosol number and size distribution measurements were performed with a modified CPC (condensation particle counter), SMPS (scanning mobility particle spectrometer) and APS (aerodynamic particle spectrometer) from TSI Inc. Comparison between the SMPS and APS data was made through conversion of the aerodynamic particle size (APS) to an equivalent mobility size (SMPS) using an appropriate particle density (ρ) and dynamic shape factor (χ) – see Möhler et al. (2006). Values of ρ (g cm⁻³) used for the samples described in Sect. 2 were: 5.0 (Fe₂O₃ – crystalline), 2.0 (Fe₂O₃ – amorphous), 2.2 (SiO₂) and 3.6 (MgO), whilst χ values of 1.5–1.6 were chosen (see DeCarlo et al. (2004) for more detailed discussion of this parameter).

The efficiencies of the various aerosols to act as ice nuclei in the deposition mode of heterogeneous ice nucleation were investigated in a series of expansion runs, using techniques and protocols established for AIDA cloud simulation studies over the past few years (see Möhler et al., 2003, 2005, 2006 and 2008 for details of chamber operation). The water vapour partial pressure is measured with a fast and sampling-free open-path tunable diode laser (TDL) absorption spectrometer (Ebert et al., 2005). In addition, a chilled mirror frost point hygrometer (MBW Calibration Ltd) measures the total water concentration in the chamber system.

During the course of an expansion experiment, the ice formation by aerosol particles is observed as a function of the changing temperature and ice saturation ratio (s_{ice}) /relative humidity with respect to ice (RHi). Ice crystals are detected and measured with an in situ light scattering and depolarization setup for the sensitive detection of droplet and ice particle growth, an infrared extinction spectrometer (FTIR) (Wagner et al., 2006; 2007), and two optical particle counters (Welas, Palas GmbH) (Benz et al., 2005). The aerosol number concentration is continuously measured with a condensation particle counter (CPC3010, TSI Inc.) modified for operation at low pressure and temperature (Seifert et al., 2004). Aerosol size distributions were only obtained at atmospheric pressure before starting the expansion run. Ice active aerosol particle fractions (f_{ice}) were calculated as the ratio of the ice particle to aerosol particle number concentration and were used as an initial measure of efficiency of the various aerosols to act as deposition mode ice nuclei.

4 Results

4.1 *IN11*(Fe₂O₃ powders)

Figure 3 shows a direct comparison of ice nucleation data sets obtained for the powder produced photo-chemically in the laboratory (left panel – labelled "IronOxide-1") and the commercial Fe₂O₃ powder (right panel – labelled "IronOxide-2") for the same initial chamber temperature of \sim 214 K.

The top panel in each case shows the variation in chamber air temperature (blue line) and pressure (black line) throughout an experimental run while the dot-dash red line shows the stability of the temperature of the chamber walls. The next panel down shows the temporal variation in relative p (hPa)

RH (%)

Scatter ratio

C, (cm⁻³)

(mu) b

10

100

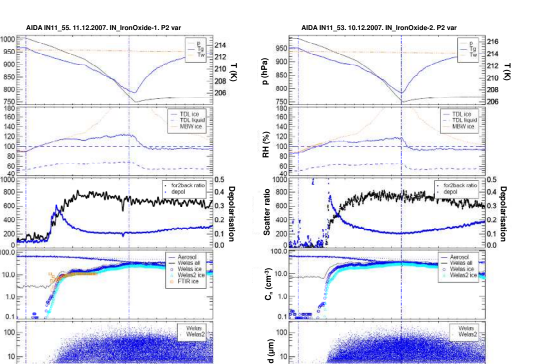


Fig. 3. AIDA chamber ice nucleation data (IN11, #55 – left and #53 – right) for Fe₂O₃ powders: left-hand side is for laboratory generated aerosol and right-hand side for the commercial sample (see Fig. 1 and text). The initial chamber air temperature (T_i) was \sim 214 K as indicated by the blue line at t = 0 s in the top panel of each data set. See text for further explanation of panel data and details of initial particle size distributions.

2000

humidity (RH) with respect to ice (RHi, solid blue line) and liquid water (RHw, dashed blue line below RH = 100%), both measured with the TDL system; "MBW" refers to frost point measurements taken with the chilled-mirror hygrometer. This instrument samples air from the cloud chamber through a heated stainless steel tube and therefore measures the total (gas-+condensed-phase) water concentration. The third panel shows the in-situ Ar⁺ laser data with the black line indicating the ratio of forward (2°) to backward (178°) scattering at 488 nm, while the blue line shows the change in depolarisation determined from backscattering perpendicular and parallel to the direction of linear polarisation of the laser, with increasing values corresponding to increasing non-spherical morphology resulting from ice crystal growth. The fourth panel gives the change in total aerosol particle concentration (blue line) and total ice particle concentration (black line). Also shown are the ice particle concentrations as measured by the WELAS#1 instrument (blue circles), from the second WELAS#2 detector, indicated by light blue triangles, and from the in-situ FT-IR spectrometer (orange squares). Finally, the bottom panel in each data set shows the single particle data from the two WELAS detectors as indicated in the panel caption. Each dot (grey - WELAS1, blue - "WELAS 2") in this plot results from detection of a

500

1000

Time (s)

1500

single ice particle at a specific size as indicated on the vertical axis and consequently the density of data points gives an indication of the concentration of detected ice particles at a particular size and time. All data is plotted with respect to the time (t) for which an experiment was conducted in the chamber.

1500

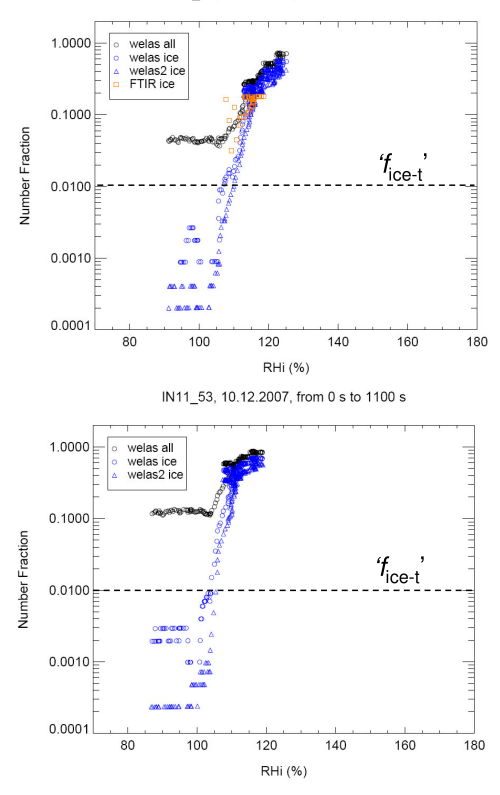
2000

500

1000 Time (s)

Measured initial smoke particle size distributions in the chamber were fitted with the following log-normal parameters ($\sum N$ – total particle concentration, D_m – modal size and σ – geometric standard deviation); laboratory Fe₂O₃ – $\sum N = 60 \text{ cm}^{-3}$, $D_m = 162 \text{ nm}$ and $\sigma = 2.1$ and commercial $\text{Fe}_2\text{O}_3 - \sum N = 58 \text{ cm}^{-3}$, $D_m = 260 \text{ nm}$ and $\sigma = 2.1$. In both experiments, RHw was maintained well below saturation.

Figure 4 summarises the variation in the ice number fraction, f_{ice} (ratio of the number of detected ice particles to the total initial concentration of aerosol particles injected into the chamber) as a function of RHi, taking into account the aerosol dilution effect during pumping (pressure change). Whilst some particle losses to the chamber walls (small particles) and settling to the bottom of the chamber (largest particles) are likely to occur as a result of such pumping, on the timescale of a typical expansion run with ice cloud formation these losses will be sufficiently small (<5% of total) to have any significant impact on the analysis and conclusions



IN11_55, 11.12.2007, from 0 s to 1100 s

Fig. 4. Variation of ice number fraction with relative humidity (with respect to ice) calculated from data in Fig. 3 (f_{ice-t} refers to a defined threshold level of 1% of the initial aerosol number).

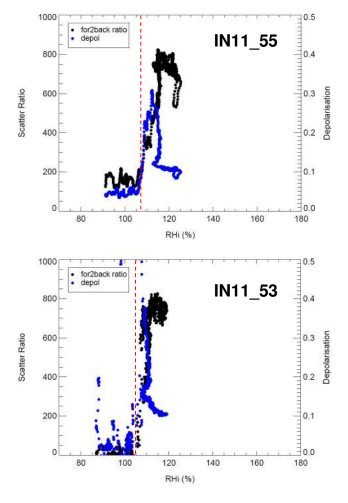


Fig. 5. Variation of light scatter (forward: back) ratio with relative humidity (ice) for nucleation experiments ($T_i = 214$ K) with laboratory-generated (top panel) and commercial (bottom panel) iron oxide powders. The dashed red lines indicate the RH_i values at which f_{ice} exceeds 0.01 in the respective experiment.

made. The nucleation data for the respective powder samples is plotted for a time period indicated at the top of each panel (in this case, the same time span), corresponding to the vertical dot-dash blue lines in Fig. 3.

In the case of the laboratory produced Fe₂O₃ sample (top panel), a defined threshold ice number fraction (f_{ice-t} : $f_{ice} = 0.01$) occurs ~300 s after the expansion cycle was started, corresponding to RHi ~107% and increases with increasing RHi to a maximum value ($f_{ice-max}$) of ~0.8. For the commercial Fe₂O₃ aerosol, f_{ice-t} occurs at around the same time as the laboratory produced sample at ~105% RHi with $f_{ice-max} = 0.85$. These "critical" RHi levels (also see Table 1) are further verified by the in situ laser scattering data shown in Fig. 5 where the scattering ratio increases significantly as does the depolarisation at the stated RHi values as indicated by the dashed red lines in this figure.

Figure 6 shows the ice fraction and laser scattering data for the same two Fe₂O₃ samples but this time from experiments conducted at a lower initial chamber temperature of 192–193 K. Measured initial smoke particle size distribution parameters were $\sum N = 150 \text{ cm}^{-3}$, $D_m = 68 \text{ nm}$ and $\sigma = 2.0$ (laboratory: left-hand panels) and $\sum N = 100 \text{ cm}^{-3}$, $D_m = 233 \text{ nm}$ and $\sigma = 1.8$ (commercial).

At the lower temperature, for both samples, f_{ice-t} takes longer to occur after the expansion cycle is initiated and significantly lower $f_{ice-max}$ values (0.12 for the laboratory sample and 0.5 for the commercial sample) are evident than at the warmer temperature. The commercial powder shows a relatively fast increase in f_{ice} , scatter ratio and depolarisation above RHi ~125% (panels b & d) whilst the data for the laboratory-generated powder shows significantly greater scatter, although it is still evident from the increased density of data points that a critical RHi occurs at ~140%.

Finally, Fig. 7 shows the variation in f_{ice} with RHi for the pseudocubic (monodisperse) Fe₂O₃ particles at an initial chamber temperature of 192 K. In this case, f_{ice-t} occurs after ~600 s, at RHi ~130% with $f_{ice-max}$ ~0.8.

4.2 IN12(Fe₂O₃ nanoparticles and MgO/SiO₂ powders)

Figure 8 gives data sets for the Fe₂O₃ nanoparticles generated in-situ at the chamber. Both experiments were conducted at an initial temperature of ~246 K but in the case of the right-hand set, the initial air pressure was significantly lower (~760 hPa) than that normally used (~1000 hPa), as was the minimum pressure reached in the expansion cycle. Measured initial size distributions parameters were $\sum N = 150 \text{ cm}^{-3}$, $D_m = 23 \text{ nm}$ and $\sigma = 1.5$ (left) and $\sum N = 40 \text{ cm}^{-3}$, $D_m = 21 \text{ nm}$ and $\sigma = 1.3$ (right).

At this warmer temperature, two nucleation modes (liquid droplets and ice particles) are evident in the bottom 2 panels of each data set, overlapping in the higher pressure run but resolved under the lower pressure conditions. In these panels, the symbols represent the ice particle measurements with the two WELAS instruments, whilst the grey line shows the total (droplet + ice particle) number. It appears that water droplets nucleate first, followed by ice particles at lower temperatures.

Figure 9 summarises the variation in the calculated nucleated number fraction f (droplet and ice) as a function of RHi (panel a) and RHw (panel b) from the experiment at higher pressure (Fig. 8, left-hand data set) for t = 0-750 s. Droplet formation exceeds the defined threshold limit ($f_{ice-t} = 0.01$) at below 100% RHw/130% RHi (dotted red lines), and ice formation at ~100% RHw/140% RHi (dashed red lines) with a maximum total f of ~0.3.

From panel (c), f_t occurs at ~241 K (droplet) and 237.5 K (ice), whilst panel (d) illustrates the cycle of RHi values attained in the experiment (orange line). Also shown for comparison in this last panel, are the data for liquid water saturation (solid blue line – calculated from Murphy and Koop,

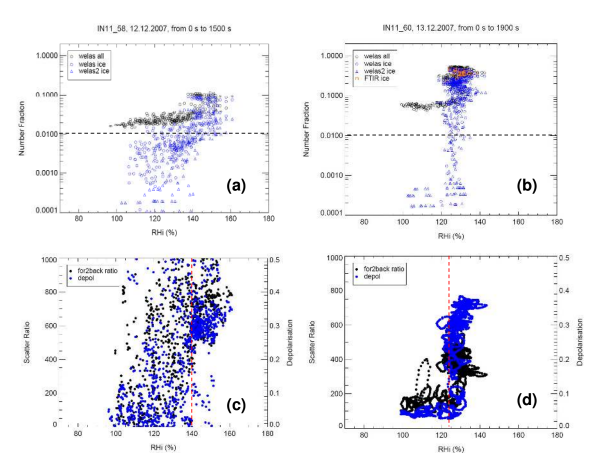


Fig. 6. Ice number fraction and laser scatter plots from nucleation experiments (IN11, #58 – left and #60 – right) carried out on laboratorygenerated (panels (a) and (c)) and commercial (panels (b) and (d)) iron oxide powders at $T_i = 193$ K.

2005) and for solution droplet homogeneous freezing at two different water activity offset values (Δa_w) of 0.3 (lower dashed blue line) and 0.32 (upper dashed blue line) – see Koop et al. (2000) and Möhler et al. (2008).

These results indicate that Fe_2O_3 particles do not nucleate ice below water saturation, but do serve as condensation nuclei (CN) at 241 K. These droplets then activated to ice at below 238 K. This is close to the temperature at which ice nucleates homogeneously within water droplets. This is a surprising result since it might be expected that water droplets contaminated with solid iron oxide particles would freeze heterogeneously.

In order to investigate the apparent inactivity of Fe₂O₃ particles when immersed in water droplets, we performed a series of separate experiments using a microscope equipped with a specialised cold stage. In this system, which is described elsewhere (Murray et al., 2010), droplets containing a known concentration of contaminating particles were placed on a hydrophobic surface and cooled down while monitoring for ice nucleation using the optical microscope. Tests with pure water droplets free of any solid impurities showed that this substrate did not nucleate ice heterogeneously and the droplets (\sim 10–30 μ m diameter) froze at the homogeneous

limit between 236 \pm 0.5 and 234 \pm 0.5 K. Droplets contaminated with up to 0.05 wt% of the same laboratory generated Fe₂O₃ particles and up to 1 wt% of the commercial Fe₂O₃ nanopowder used in the AIDA studies, also only nucleated at the homogeneous limit. Contamination of droplets with similar quantities of clay minerals such as illite causes ice to nucleate some 10 K higher (Murray et al., 2010). These results confirm that Fe₂O₃ particles are inactive in the immersion mode in pure water droplets.

Figure 10 shows nucleation data for Fe₂O₃ nanoparticles generated in-situ at the lower temperatures of 200 K (LHS) and 182 K (RHS) – please refer to Table 1 for a summary of threshold values. Measured initial size distributions parameters were $\sum N = 120 \text{ cm}^{-3}$, $D_m = 27 \text{ nm}$ and $\sigma = 1.5$ (LHS) and $\sum N = 110 \text{ cm}^{-3}$, $D_m = 27 \text{ nm}$ and $\sigma = 1.5$ (RHS).

Finally, Fig. 11 compares the number fraction data (top panels: against RHi, bottom panels: against *T*) for the commercial MgO (left-hand side) and SiO₂ (right-hand side) powders dispersed into the chamber at an initial temperature of 199–200 K. Measured initial size distributions parameters were $\sum N = 700 \text{ cm}^{-3}$, $D_m = 140 \text{ nm}$ and $\sigma = 2.2$ (MgO) and $\sum N = 55 \text{ cm}^{-3}$, $D_m = 111 \text{ nm}$ and $\sigma = 2.1$ (SiO₂). Again, threshold values are given in Table 1.

	Exp. # (Figure #)	(a) <i>T_i</i> (K)	(b) <i>D_m</i> (nm)	(c) t (thresh) (s)	(d) RH _i (thresh) (%)	(e) f _{ice-max} (% RH _{i-max})
(A)	IN11_60	192	233	500	125	0.50
Fe ₂ O ₃	(6)	215	260	300	105	(135)
(commercial)	IN11_53					0.85
	(3–5)					(115)
MgO	IN12_03	200	140	450	120	0.08
	(11)					(125)
SiO ₂	IN12_21	200	111	150	118	0.50
	(11)					(160)
(B)	IN11_55	213	162	300	107	0.80
Fe ₂ O ₃	(3–5)	193	68	1000	140	(120)
(powder)	IN11_58					0.12
	(6)					(160)
Fe ₂ O ₃	IN12_17	200	27	500	125	0.02
(nanoparticles)	(10)	182	27	800	140	(180)
	IN12_09					0.09
	(10)					(170)
(C)	IN11_58	193	68	1000	140	0.12
Fe ₂ O ₃	(6)					(160)
(amorphous)						
Fe ₂ O ₃	IN11_60	192	233	500	125	0.50
(crystalline	(6)					(135)
– hexagonal)						
Fe ₂ O ₃	IN11_62	192	600	600	130	0.80
Pseudocubic	(7)		(edge)			(150)

Table 1. Comparison of Nucleation Data for Different Forms ((A) composition, (B) size and (C) morphology) of Refractory Particles.

(a) $-T_i$: initial temperature of air in AIDA chamber prior to expansion cycle

(b) $-D_m$: modal diameter of measured aerosol size distribution

(c) – t(thresh): "threshold" nucleation time for fraction of initial aerosol to be detected as ice particles (f_{ice})>0.01

(d) – RH_i (thresh): relative humidity with respect to ice in the chamber at t_{thresh}

(e) $-f_{ice-max}$: maximum value of f_{ice} in experimental run, RH_{i-max}: maximum relative humidity with respect to ice reached in the chamber

5 Discussion

For an initial analysis of the nucleation data for the different aerosol samples described in Sect. 2, we will focus on the three primary properties which are likely to affect threshold conditions of relative humidity with respect to ice (RH_i) and the calculated fraction (f_{ice}) of initial aerosol particles which initiate ice crystal formation in each case. These critical particle properties are (a) composition, (b) size and (c) structure/morphology.

5.1 Particle composition

Table 1 – section A summarises the data for the first of the stated aerosol properties by comparing the commer-

cially available iron, magnesium and silicon oxide powders that were investigated. Two data sets are shown for Fe₂O₃ ("IronOxide-2"), because the initial air temperatures in the chamber (T_i) for those experiments straddle the 200 K temperature used in the experiments on the other oxides.

It can be seen that the SiO₂ particles seem to have similar nucleation properties to the Fe₂O₃ sample at 200 K, in terms of the experimental observables of threshold RH_i (for f > 0.01) and $f_{ice-max}$. Whilst there are many different structural forms of silica, which are likely to have very different water-adsorption and ice nucleating propensities, the amorphous form used in these studies is the most likely one to result via condensation from the gas-phase in the upper atmosphere. The sample used has a high quoted surface area of ~390 m² g⁻¹ and consists of aggregated chains of ~7 nm

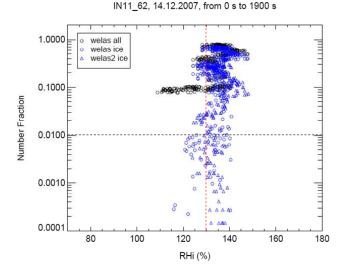


Fig. 7. Ice number fraction data for the pseudo-cubic Fe₂O₃ sample at $T_i = 192$ K

primary particles which are of the order of a few tenths of a micron in dimension. It is likely that the dispersal method used for admitting aerosol in powder form into the AIDA chamber will result in some degree of break-up of such aggregates, reflected in the measured modal size of 111 nm. The Fe₂O₃ particles from the dispersed commercial sample have a modal size almost twice that of the SiO₂ indicating that particle size alone does not explain the relatively high efficiency of the silica aerosol, but composition and/or structural differences are more important factors.

The MgO sample appears to be a weaker candidate for ice nucleation in comparison with iron and silicon oxides at these temperatures. The sample used is quoted as consisting of nanoparticles of <50 nm size and a surface area of $130 \text{ m}^2 \text{ g}^{-1}$ although the measured modal size of 140 nm seems to indicate some degree of aggregation. A significantly lower $f_{\text{ice}-\text{max}}$ of 0.08 was observed, although it should be noted that the maximum RHi value of 125% for this chamber run was lower than those attained for the Fe₂O₃ and SiO₂ samples at 200 K (see Table 1).

5.2 Particle size

Table 1 – section B compares data for the laboratorygenerated amorphous Fe₂O₃ samples ("Iron Oxide-1") in both powder form (mechanically dispersed into the chamber volume) and particle form (generated in situ adjacent to the chamber and passed directly into the chamber). Two T_i runs are shown for each sample. These are not identical for the two particle forms but are close enough for reasonable inferences to be made.

It has been shown in previous work (Saunders and Plane, 2006) that the photo-oxidation of $Fe(CO)_5$ in O_3 leads to the formation of amorphous Fe_2O_3 particles which form chain-

like aggregates of micron dimensions, consisting of primary particles of 6–7 nm diameter. Both of these aerosol samples were produced by this same method, although in order to sediment out enough powder for the subsequent nucleation experiments (*IN11*), the aerosol generator was run for much longer times and at much higher precursor concentrations and photolysis light intensities. The only difference expected therefore is that of particle/aggregate size as corroborated by the measured modal sizes. As mentioned before, the dispersal method used to admit the powder sample is likely to result in the mechanical break-up of the micron-size aggregates (see Fig. 1 – top panel). Indeed, the modal sizes of <200 nm suggest that such break-up is significantly more pronounced than in the case of the SiO₂ powder as discussed in the previous section.

For the powder sample, there is a marked decrease in $f_{ice-max}$ below 200 K, although at 193 K, still ~12% of the initial aerosol population was observed to lead to ice crystal formation with RHi-max = 160%. In contrast, the nanoparticles (D_m ~30 nm) formed in situ showed lower activation levels although the lower temperature run did result in a higher activated fraction. This was probably a consequence of the more rapid increase in air temperature after ~1500 s of the $T_i = 200$ K run which resulted in a faster drop in RH_i, hence most likely limiting any further activation of the aerosol population.

The lower values for the nanoparticles are consistent with the classical heterogeneous nucleation theory of Fletcher (1958) which predicts a pronounced size effect to ice nucleation directly from the vapour-phase for spherical particle sizes below 100 nm. On this basis, for the case of a polydisperse initial aerosol distribution, it is reasonable to suppose that the larger particles within the distribution will nucleate ice formation sooner (at lower RHi) than the smaller particles. Whereas the particle size range for the powder sample extended up to a few hundreds of nanometers, for the nanoparticles formed in situ, the upper limit of the measured size range was \sim 80 nm and so the observed maximum ice number fractions at a particular temperature/ice supersaturation were lower. In addition, the maximum RHi value reached in a particular nucleation run is a complex function of a number of factors including the rates of ice crystal formation and growth and the cooling rate within the chamber (Möhler et al., 2006). As such, the RHi increase within the chamber, after ice crystal formation has been initiated by the sample, is somewhat self-limiting and consequently, higher ice super-saturation (>200%) conditions required for smaller particles within a distribution to nucleate ice are prevented. Therefore, the maximum reported ice number fractions for each experiment do not represent the "true" activation efficiency of a particular particle material. For this, it is more instructive to use the observed threshold RHi values for determination of the contact angle with respect to ice embryo formation at the particle surface, or the variation of f_{ice} with RHi to evaluate ice formation rates (see Sect. 6). Clearly,

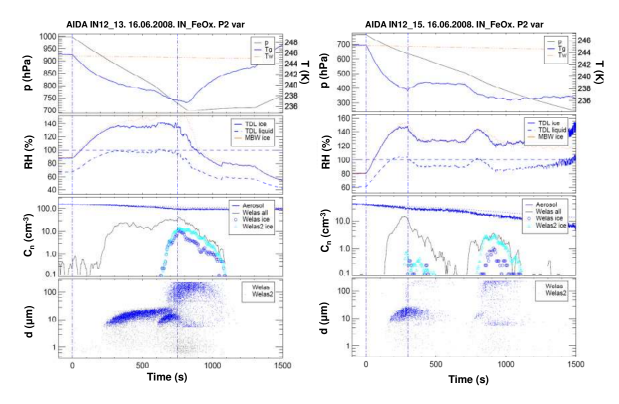


Fig. 8. Nucleation data sets (*IN12*, #13 – left and #15 – right) for the Fe₂O₃ nanoparticles generated in-situ at the chamber with $T_i \sim 246$ K. The left-hand side set was obtained with the initial air pressure at the "normal" level whereas the right-hand side data was from a lower initial air pressure run in the chamber.

even after discounting these effects, the 9% activation level of a sub-80 nm particle size distribution at 182 K suggests that aerosols of such dimensions and compositions have the potential to nucleate ice at sufficient supersaturation levels in the atmosphere.

5.3 Particle structure

Finally, Table 1 – section C illustrates differences between the three structural forms of Fe₂O₃ aerosol used i.e. the laboratory-generated amorphous powder, the commercial crystalline powder and the synthesised pseudocubic form, for the same value of T_i . At 192–193 K, the preferred form of ice is predicted to be cubic (I_c) . Data comparison shows that the pseudocubic form of Fe₂O₃ has a significantly higher $f_{ice-max}$ value than either the amorphous or non-cubic crystalline forms. However, this observation must be qualified by the fact that the pseudocubic particles were much larger in size and so it may be that both size and structural factors are responsible for the observed differences in nucleating ability. The non-cubic aerosol particles showed similar nucleating efficiencies although the larger modal size of the crystalline sample could mean that, once again, size effects were the primary reason for differences in the nucleation data, rather than structural considerations.

6 Data analysis

For quantitative analysis of the chamber ice nucleation data, we have adopted two common approaches (see Pruppacher and Klett, 1997): (i) the stochastic (classical) determination of the contact angle (θ) between the smoke particle and growing ice embryo, which inherently assumes that θ is the same for any particle of the same size and composition, and (ii) a more recently developed approach based upon the "singular hypothesis" which determines a particle-specific parameter known as the ice-active surface site density (n_s).

Whilst the classical approach provides a comparison with θ values assumed in previous theoretical studies of heterogeneous ice nucleation by MSPs (e.g. Keesee, 1989; Rapp and Thomas, 2006), it should be noted that the values derived from the observed threshold nucleation data and measured aerosol distributions can only be taken as representative of the efficiency of the most active sites on the particles. In practice, there is likely to be a distribution of contact angles in an aerosol population as a result of variations of the surface sites available for ice nucleation. Consequently, the second approach, which specifically treats such inherent particle site variability, is more rigorous, but due to its relatively recent development, has a smaller literature database for comparison of evaluated n_s values.

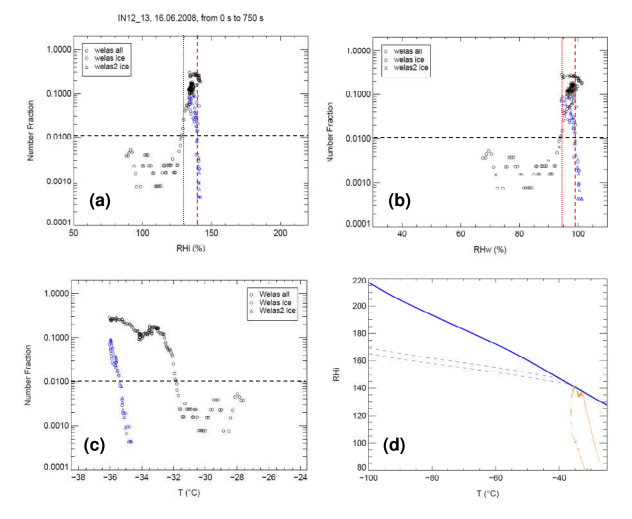


Fig. 9. Summary plots for the ice nucleation experiment (*IN12*, #13) on in-situ-generated Fe₂O₃ nanoparticles at $T_i \sim 246$ K (calculated from left-hand data set in Fig. 8). Panel (a) shows the number fraction (f) of detected ice particles (blue symbols) and total fraction (black symbols) i.e. droplets plus ice, as a function of RHi, panel (b), f versus RHw, panel (c), f versus T, and panel (d) RHi versus T. Red lines mark positions of critical RH values (dot: droplet detection, dash: ice detection) and the dashed black line indicates the defined threshold value (f_{ice-t}).

6.1 Classical theory: contact angle determination

From classical heterogeneous nucleation theory, the critical parameter which defines the efficiency of a particular material as an ice-nucleating substance is the contact angle (θ), which is determined from the contact parameter ($m = \cos i \theta \theta$). For a flat or a curved surface (i.e. particle), m can be estimated from experimental measurements of the minimum saturation ratio (s_{crit}) at which ice nucleation is observed for a given temperature. Nucleation efficiency increases as $m \to 1$ or $\theta \to 0^{\circ}$.

The recent paper by Trainer et al. (2009) reported such data for a silicon wafer at temperatures down to 150 K and determined a temperature-dependent contact parameter using the classical equations from Pruppacher and Klett (1997). Following this method, we have used the AIDA data for the

particle samples described earlier to ascertain the values of m (and hence θ) at the temperatures used in the respective nucleation runs as indicated in Table 1.

The surface nucleation rate $(J: \text{cm}^{-2} \text{s}^{-1})$ in each chamber run, required for determination of the matching function f(m,x) with respect to curved (particle) surfaces (see equation 9 of Trainer et al., 2009), and hence θ , were calculated following the procedure outlined in Sect. 3.2 (Eq. 3–5) of the paper by Möhler et al. (2006). For the specific particle type/chamber temperature, the *J* value is obtained from the ratio of the rate of ice crystal formation $(dn_{ice}/dt: \text{cm}^{-3} \text{s}^{-1})$, calculated from the observed threshold RHi value at a given temperature, to the particle surface area $(\text{cm}^{-2} \text{ cm}^{-3})$, determined from the measured initial particle distribution.

Figure 12 shows the temperature-dependent contact angle values determined as described for the three Fe₂O₃ particle

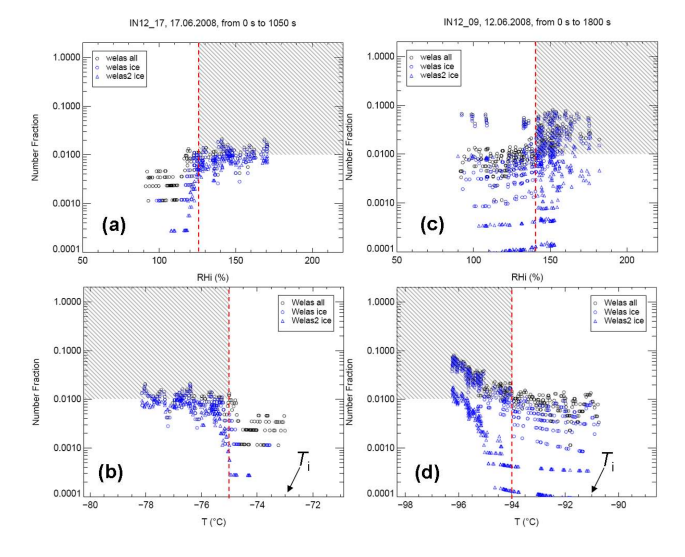


Fig. 10. Calculated ice number fraction data (*IN12*, #17 – left and #9 – right) for nucleation experiments with Fe₂O₃ nanoparticles with $T_i = 200$ K (left) and 182 K (right). Top panels show f_{ice} as a function of RHi; bottom panels as a function of chamber air temperature (*T*). Red dashed lines indicate critical RHi and *T* values for the respective experiments for which $f_{ice} > 0.01$ (f_{ice-t}). Data points within the hatched areas therefore correspond to conditions for which f_{ice-t} is exceeded.

samples as indicated. The data shows clearly that the contact angle (as defined schematically in the figure) increases with decreasing temperature for all samples. Analogous behaviour was observed by Trainer et al. (2009) for a planar silicon surface. They reported a much sharper rise in θ , particularly for T < 180 K, which of course we are not able to verify with our samples as these low temperatures are not achievable in the AIDA chamber. However, the angles for the iron oxide particles are significantly lower than those for the silicon surface. For example, at 182 K, Trainer et al. (2009) report m = 0.896 ($\theta = 26.4^{\circ}$) compared with that of $\sim 10.5^{\circ}$ for the nanoparticles generated in situ at the chamber. This most likely reflects the more hydrophilic nature of the metal oxide compared with pure silicon. It is noticeable that at T > 210 K, the calculated contact angles are very similar (5–7°) for all three samples, whilst at lower temperatures the particles generated in situ display a smaller temperature dependence and so seem to be the most efficient nucleating species at T < 200 K. The larger contact angles found for the powder samples in Fig. 12 are most probably an indication of a greater hindrance to the formation of stable ice embryos at the surfaces of the more aggregated particle samples. In addition, all values lie well below the horizontal line at $\theta = 18.2^{\circ}$ (m = 0.95), the "nominal" value usually assumed in modelling of NLC formation (Rapp and Thomas, 2006), although at summer mesopause temperatures below 150 K it is probable that the contact angles for iron oxide aerosol will approach or possibly exceed this value, due to the likelihood of non-hexagonal or metastable (cubic or amorphous) ice formation directly from the vapour phase being increasingly

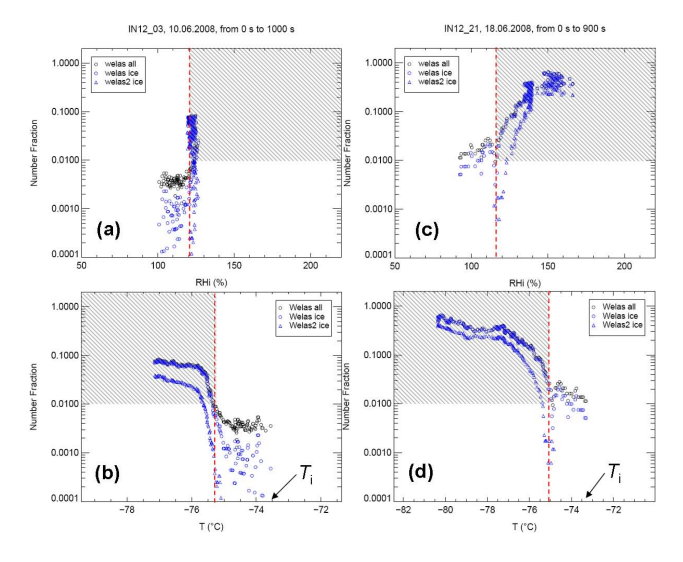


Fig. 11. Calculated ice number fraction data (*IN12*, #03 – left and #21 – right) for nucleation experiments with MgO powder (left) and SiO₂ powder (right) at $T_i \sim 200$ K. Top panels show f_{ice} as a function of RHi; bottom panels as a function of chamber air temperature (*T*). Red dashed lines indicate critical RHi and *T* values for the respective experiments for which $f_{ice} > 0.01$ (f_{ice-t}). Data points within the hatched areas therefore correspond to conditions for which f_{ice-t} is exceeded.

favoured under such conditions (Murphy and Koop, 2005; Shilling et al., 2006).

Of course, particles larger than the modal size will most probably tend to nucleate ice first at somewhat lower s_{crit} values than for the modal size particle, but as these are at much lower number concentrations, they probably lead to the formation of ice crystal numbers below the threshold levels. For example, for the IN12_#9 run (in situ generated particles, $T_i = 182$ K and $s_{crit} = 1.4$), at the same contact angle of 10.5° ($m \sim 0.983$) determined for the modal particle size (from the surface area distribution) of ~ 27 nm, an s_{crit} value of 1.33 is predicted at 40 nm.

For the SiO₂ and MgO samples at 200 K, the contact angles are 10.2° and 11.4° respectively.

6.2 Singular hypothesis method

6.2.1 Analysis

For a given aerosol type/chamber temperature experiment, a log normal fit to the measured size distribution of initial aerosol (prior to the start of the nucleation experiment) was used to define the available surface area available for nucleation. Then, the measured ice crystal number concentration from the WELAS OPC, and the measured RH from the TDL and temperature probes were used to determine the number of ice active surface sites that become active at a set RH. This is equivalent to the "singular hypothesis" i.e. the surface of each particle is assumed to have a characteristic active site density as a function of the driving nucleation variable. Such

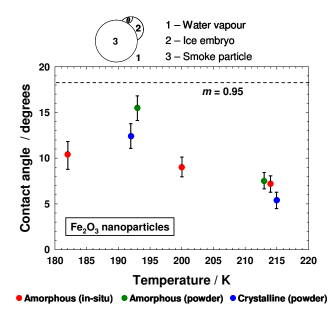


Fig. 12. Variation in estimated contact angle (θ) values with temperature for the three iron oxide particle samples indicated. The schematic at the top of the figure is used to illustrate the definition of θ with respect to heterogeneous ice nucleation on a particle.

a treatment is analogous to the theory discussed in Connolly et al. (2009) with respect to the heterogeneous freezing mode (for which temperature is the critical parameter), but adapted for the scenario of heterogeneous deposition nucleation (see Sect. 3.3.2 of Connolly et al., 2009), for which RH (or saturation ratio) is the parameter of interest. To date, this approach has only been applied to heterogeneous ice nucleation at warm cloud temperatures for micron-sized terrestrial dust samples.

Figure 13 provides a schematic explanation of the analytical approach taken using a representative particle nucleation data set. It is important to note that in this analysis we assume that the area provided by the smoke particles is given by the equivalent spherical surface area.

Analysis was performed for experiments that used the commercial (crystalline) sample, the prepared (amorphous) laboratory sample and the in situ generated amorphous particles of iron oxide composition. Figure 14 shows the derived values of ice-active surface site density (n_s) for the iron oxide particles at two chamber temperatures (~190 and 215 K) for the experiments as indicated in the left-hand plot.

These experiments clearly show that heterogeneous deposition nucleation is more efficient at the higher temperature, indicated by the higher values of n_s for lower values of RH_i. This can be understood by classical nucleation theory which predicts that the energy of germ formation is inversely proportional to temperature (as predicted by Boltzmann statistics) and also inversely proportional to the natural logarithm of the supersaturation (a result of the surface energy of germ

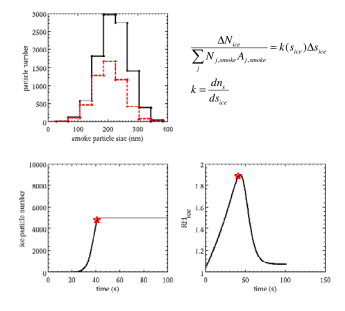


Fig. 13. Illustration of how the ice nucleation data were analysed. Top left: black shows the initial smoke particle size distribution, red shows the smoke distribution after nucleation has occured. Note that the largest particles are depleted first as these have the larger surface area for nucleation to occur. Bottom left: shows the temporal variation of measured ice particle number concentration with the star indicating the ice concentration that corresponds to the point in time where the smoke distribution is as in the red histogram. Bottom right: shows the variation in RHi with chamber expansion time. Note that no nucleation occurs once the peak RH has occurred. This data is used to define the ice active surface site density per saturation interval (k(sice)) by dividing the change in ice particle number by the total surface area available for nucleation $(\Sigma N_{j,\text{smoke}}A_{j,\text{smoke}})$ where N and A are the number and surface area of smoke particles in size bin *j*). This provi This provides a function that describes the number of ice active sites per unit area of aerosol at saturation ratio sice.

formation). Normally at warmer temperatures the supersaturation effect is dominant over the temperature fluctuation effect and we observe that nucleation is more efficient at the ice supersaturation. However, at very cold temperatures the molecular fluctuations associated with the heat of the substance become important and, in principle, the opposite behaviour may be observed. It is interesting to note that there is no significant difference between the IN activity of the crystalline samples vs the amorphous samples when the surface area is taken into account (e.g. comparing IN experiments 59 with 61).

The function fitted to the data at 190 K, $n_s 10^{(3.33 \cdot s_{ice})+8.16}$ (green line – bottom panel), was used to model ice formation under different conditions i.e. updraft, initial temperature, different mode radius of particles.

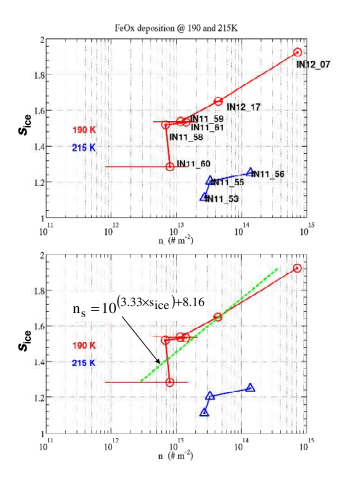


Fig. 14. Top: shows the calculated number of active sites per unit area of material that become active at the given s_{ice} for the iron oxide aerosol experiments as indicated. Errorbars denote 5 and 95 percentiles assuming that the Poisson distribution describes the uncertainty. Bottom: for the colder experiments, a curve (green line) was fitted through the data using the least squares method. The equation for this fit was used in the modelling studies.

6.2.2 Modelling

The model used is a cloud parcel bin model which includes treatments of both ice and liquid microphysics, and the role of aerosol surfaces (see appendix of Connolly et al., 2009 for details). The fitted RHi vs n_s data derived in the previous section with respect to iron oxide nanoparticles were used along with log-normally distributed particle size distributions to predict the activated fraction of the particles and the corresponding peak saturation ratio (s_{ice}) under conditions appropriate for the upper atmosphere. Model simulations were started at 180 K at $s_{ice}=0.95$ with a smoke particle concentration of 100 cm⁻³ (in line with the recent 2-D & 3-D modelling studies of Megner et al., 2008a & b and Bardeen et al., 2008).

For vertical motions we adopted a gravity wave scheme that results in a peak cooling rate of 0.01 to $0.1 \,\mathrm{K \, s^{-1}}$ (or equivalently 0.1 to $1 \,\mathrm{m \, s^{-1}}$ assuming adiabatic expansion).

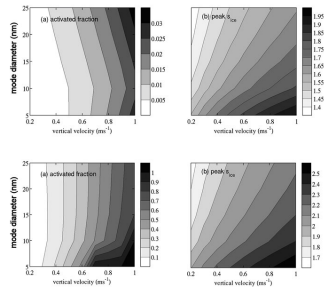


Fig. 15. Simulations with a cloud parcel model to infer the activated fraction of particles as IN (chamber data for iron oxide nanoparticles were used in the model runs). Top panels: left shows the activated fraction for different updraught velocities and initial mode diameter of the smoke particles and right shows the peak supersaturation achieved in the simulation (simulations assume an ice deposition coefficient of 1.0). Bottom panels: As in top panels but simulations assumed an ice deposition coefficient of 0.1. Note that in these simulations the activated fraction is significantly higher.

Gravity waves of this order are fairly typical in the upper mesosphere (e.g. Hoppe and Fritts, 1995). Although the diurnal and semidiurnal tides result in weaker vertical motions, the range of gravity wave cooling rates was chosen for an upper boundary on the cooling rate. Results from the model are shown in Fig. 15.

Note that if we use a deposition (mass accommodation) coefficient (α_D) of 1.0 for water vapour uptake on ice we are unable to activate sufficiently high fractions of smoke particles as IN since the water vapour is depleted quickly by the growing ice crystals (Fig. 15, top panels). In order to nucleate sufficiently high numbers of smoke particles as IN, the super-saturation must build up to high enough values. For the range of values in our simulations this can be achieved by reducing the deposition coefficient for ice to around 0.1 (Fig. 15, bottom panels). It is interesting to note that higher activated fractions occur for the smaller mode diameter. This is because in this regime the deposition coefficient has the largest sensitivity for small particles.

In order to explain growth rates of small ice crystals using the standard depositional growth model, values of α_D much less than unity were inferred from the laboratory measurements of Magee et al. (2006), while a value of 0.1 is consistent with literature values tabulated in Choularton and Latham (1977) for the appropriate temperature range.

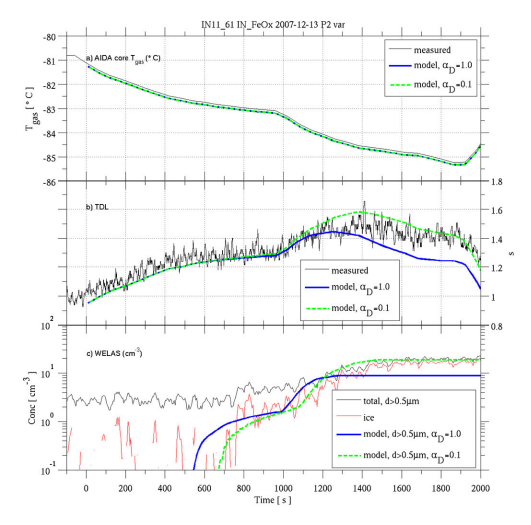


Fig. 16. Observations and model simulation of experiment *IN11*, #61 (commercial Fe₂O₃). Note that temperature and pressure were constrained in the model to the observations as was the total water (condensed plus vapour) in the simulation (as measured by the chilled mirror hygrometer). The free variables then are the ice water concentration and mass (and thus the water vapour supersaturation) calculated by the model. We note that for a deposition coefficient of 1.0 the ice crystals grow too fast and deplete the available water vapour so that ice concentrations in the model are too low. Better agreement is found using a deposition coefficient of 0.1 where the simulated water vapour supersaturation and the ice crystal number concentration give good agreement to observations. Initial smoke particle size distribution parameters were: $N_L = 55 \text{ g}^{-1}$ (of air), $\ln(\sigma) = 0.4762$ and $D_m = 0.26\mu\text{m}$.

Figure 16 shows the results of model simulations of the temporal evolution of ice crystal number (bottom panel) from a representative experiment (IN11, #61; crystalline Fe₂O₃ particles; $T_i \sim 196$ K), assuming α_D values of 1.0 and 0.1. In this experiment and others (not shown) the best agreement between observations and modelling is found when using a deposition coefficient of 0.1.

7 Atmospheric implications

The reported chamber ice nucleation results at temperatures below 230 K for amorphous iron oxide nanoparticles/aggregates and silicon- and magnesium oxide powders with particle dimensions ranging from $\sim 10-300$ nm suggest that these refractory aerosol species are likely to play an important role in ice cloud/PSC formation in the upper troposphere/stratosphere. The inherently alkaline nature of metal oxide particles will certainly favour the uptake and condensation of sulphuric acid onto particle surfaces within the Junge layer, and thus lead to a role in the formation of PSCs (Turco et al., 1981).

In terms of ice cloud formation in the mesosphere, studying the heterogeneous nucleating abilities of meteoric smoke particles smaller than 10 nm diameter (typically 0.5–2 nm) and at temperatures below 150 K currently remains beyond the experimental capabilities of large volume chambers and particle detection instruments. Consequently, any detailed extrapolation of our nucleation studies to the mesosphere is not possible, but some general comments can be made. For example, the classic 1-D model prediction (Hunten et al., 1980) of smoke particle concentrations of a few thousand per cm³ at NLC-forming altitudes (80-90 km) would require ice activation levels of greater than $\sim 5\%$ to explain typical ice crystal numbers of around $100 \,\mathrm{cm}^{-3}$ in these clouds as indicated by lidar studies (e.g. von Cossart et al., 1999). However, more recent 2-D and 3-D modelling treatments of the growth and transport of smoke particles (Megner et al., 2008a; Bardeen et al., 2008) suggest that the concentrations of smoke particles are much lower ($<200 \,\mathrm{cm}^{-3}$) at the summer mesopause because of rapid meridional transport to lower latitudes. Clearly at these levels, it becomes increasingly difficult to envisage a major role for MSPs in heterogeneous ice nucleation, even if activation levels approached 100%. However, there are two ways in which MSPs could still act as efficient IN. First, if a significant fraction of the MSP population attains a steady-state charge within the plasma which exists in the upper atmosphere (see Rapp, 2009), then certain critical constraints (primarily the minimum particle size) with respect to ice nucleation will be removed (Megner et al., 2008b, Gumbel and Megner, 2009), thus increasing the likely activation levels. Indeed, higher levels would be needed to account for the numbers $(>1000 \text{ cm}^{-3})$ of smaller charged ice particles (inferred from radar detection) which give rise to the phenomenon of polar mesosphere summer echoes (PMSE) in this altitude region (Rapp and Lübken, 2004). Second, individual molecules containing meteoric metals (e.g. NaHCO₃, FeSiO₃) have very high electric dipole moments (>8 D), which result in significantly lower binding energies for polar H₂O molecules (Plane, 2000). With these considerations in mind, it seems most likely that MSPs nucleate ice particles in the upper mesosphere alongside other mechanisms, including ion-induced (Keesee, 1989; Gumbel and Witt, 2002) and homogeneous nucleation (Murray and Jensen, 2010; Zasetsky et al., 2009).

8 Conclusions

All refractory particles (Fe₂O₃, SiO₂ and MgO) studied at the AIDA nucleation chamber were observed to nucleate ice to a significant degree under conditions of super-saturation (with respect to ice), between 182 K and 215 K. The powder samples in all cases produced ice number fractions (f_{ice}) of ≥ 0.08 for temperatures of 200 K or below, with the crystalline (commercial) iron oxide and amorphous silica samples showing the highest (maximum) values of 0.5 at 192 K and 0.5 at 200 K respectively, and the lowest threshold RHi values (defined as the level at which $f_{ice} > 1\%$) of 125% and 118% respectively. MgO particles (with cubic structure) showed a significantly lower maximum f_{ice} of 0.08 (RHi_{thresh} = 120%) at 200 K, whilst the pseudo-cubic form of Fe₂O₃ produced a much higher ice fraction of 0.8 (RHi_{thresh} = 130%), although this observation must be coun-

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tered with the fact that these particles were significantly larger (edge-length of 500–600 nm) than the particles from the powder samples (poly-disperse with modal sizes of 100 – 250 nm).

The Fe₂O₃ nanoparticles generated in situ in a photochemical reactor adjacent to the nucleation chamber were much smaller (typical modal size \sim 30 nm) and were still observed to initiate ice crystal formation (RHi_{thresh} = 140% at 182 K) with $f_{ice-max}$ = 0.09. Contact angle values determined for these particles (e.g. 10.5° at 182 K) indicate that amorphous iron oxide nanoparticles are relatively efficient nucleating surfaces when compared with recently reported data for pure silicon. Direct comparison of determined contact angles of the three oxide nanoparticle types at 200 K indicate a nucleation efficiency order of Fe₂O₃>SiO₂>MgO.

Modelling of the low temperature (<200 K) nucleation data for iron oxide nanoparticles using a fitted function obtained from evaluated ice-active surface site density values, indicates a deposition coefficient of 0.1 for water vapour onto nucleated ice crystals for conditions relevant to the upper atmosphere i.e. significantly smaller than the value of 1 typically assumed in NLC model studies to date (see Rapp and Thomas, 2006).

The nucleation runs conducted at 246 K and with RHw \rightarrow 100% (see Figs. 8 and 9) clearly indicate a different mechanism characterised by initial droplet formation and subsequent freezing to form ice crystals, as observed previously at the AIDA chamber in the case of Asian desert dust (Cotton et al., 2007). This indicates that under such conditions, the particles are acting as CN rather than IN and the droplets only freeze at or close to the homogeneous limit. The conclusion is that Fe₂O₃ particles in either crystalline or amorphous forms will not initiate ice nucleation in tropospheric mixed phase clouds.

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