A slow cooling-rate in situ cell for long-duration studies of mineral precipitation in cold aqueous environments on Earth and other planetary bodies

Stephen P. Thompson,a* Hilary Kennedy,b Sarah J. Day,a Annabelle R. Baker,a Benjamin M. Butler,c Emmal Safi,a,d Jon Kelly,a Andrew Male,a Jonathan Potter,a Tom Cobb,a Claire A. Murray,a Chiu C. Tang,a Aneurin Evansd and Ronaldo Mercadoa

aDiamond Light Source, Harwell Science and Innovation Campus, Didcot, Oxfordshire OX11 0DE, UK, bSchool of Ocean Sciences, Bangor University, Menai Bridge, Anglesey LL59 5AB, UK, cEnvironmental and Biochemical Sciences, The James Hutton Institute, Craigiebuckler, Aberdeen AB15 8QH, UK, and dAstrophysics Group, Lennard-Jones Laboratories, Keele University, Keele, Staffordshire, ST5 5BG, UK. E-mail: stephen.thompson@diamond.ac.uk

Abstract

Long considered defining features of the Earth, space missions and observations have shown that liquid oceans and ice caps, along with ice crusts, are in fact common features among many of the Solar System’s outer planets and their satellites. Interactions with rock forming materials have produced saline oceans not dissimilar in many respects to those on Earth, where mineral precipitation within frozen seawater plays a significant role in both determining global properties and regulating the environment in which a complex ecosystem of extremophiles exists. Since water is considered
an essential ingredient for life, the presence of oceans and ice on other Solar System bodies is of great astrobiological interest. However the details surrounding mineral precipitation in freezing environments are still poorly constrained, due to difficulties of sampling and ex situ preservation for laboratory analysis, meaning that predictive models have limited empirical underpinning. To address this we present the design and performance characterisation of a transmission geometry sample cell for use in long-duration synchrotron X-ray powder diffraction studies of in situ mineral precipitation from aqueous ice-brine systems. Capable of very slow cooling rates (e.g. 0.3 °C/day or less), we demonstrate the cell’s performance with the results from a year-long study of the precipitation of the hydrated magnesium sulphate phase meridianiite (MgSO$_4$ · 11H$_2$O) from the MgSO$_4$-H$_2$O system. Evidence from the Mars Rover mission suggests this hydrated phase is widespread on the present day surface of Mars. However, as well as the predicted hexagonal ice and meridianiite phases, we observe an additional hydrated sulphate phase and a disordered phase.

1. Introduction

1.1. Low-temperature aquatic mineral precipitation

Cold aqueous environments play a critical role in defining surface geochemistry both on Earth and on certain other planetary bodies in our solar system (e.g. Mars and the icy satellites of Jupiter and Saturn, see Table 1) where water is found in close association with ice (Marion & Farren, 1999; Hussmann et al., 2006; Massé et al., 2014; Nimmo & Pappalardo, 2016). For example, the freezing of aqueous solutions containing dissolved salts such as NaCl, KCl, Na$_2$SO$_4$, MgCl$_2$ and CaCl$_2$ – the main dissolved salts in terrestrial oceans – results in the formation of highly concentrated electrolyte solutions (brines) caused by the separation of pure water into solid ice via eutectic freezing (Cox & Weeks, 1983). The precipitation of minerals within such
ice-brine environments means that they tend to be retained rather than lost to the underlying water column (Light et al., 2003) and are therefore retained close to the ice–atmosphere (or ice–space) interface. The presence of mineral precipitates affects the structural and optical properties of the ice due to their size distribution and regulatory effect on brine volume (Assur, 1960; Maykut & Light, 1995; Light et al., 2003, 2009) which in turn may influence global properties such as albedo and the effect that varying insolation has on climate change (e.g. Cheng et al., 2009).

Understanding the chemical and physical properties of mineral precipitation from highly concentrated electrolyte solutions is thus fundamental to understanding the precise role played by planetary aquatic processes. In particular, their contribution to climate change on Earth (both present and historical, since the planet cycles between greenhouse, icehouse and possibly snowball phases), present and past hydrological activity on Mars and core-ocean-ice crust interactions on bodies such as Europa and Enceladus where saline subsurface oceans exist beneath a surface ice crust (e.g. Hussmann et al., 2006). These latter ocean worlds are also considered to be highly plausible habitats for extra-terrestrial life and, post-Cassini, are the targets of future planned space missions (e.g. NASA Ocean Worlds program, ESA JUICE mission). Indeed, on Earth, sea ice brine channels host a wide range of sympagic organisms (bacteria, microalgae, viruses, fungi, protozoans and metazoans) where osmotic conditions (salinity) are defined by temperature dependent salt precipitation in the Na-K-Mg-Ca-Cl-SO\(_4\)-H\(_2\)O sea water system (e.g. the precipitation of mirabilite Na\(_2\)SO\(_4\)-10H\(_2\)O below \(-6.4^\circ\text{C}\); Butler et al., 2016). Mineral precipitates can also provide solid surfaces for biogenic interactions (Cleaves et al., 2012), and in the context of the precursor RNA world hypothesis for the abiogenic origin of life, cold ice-brine environments may have been an essential low-temperature step in the early replication of nucleic acids (Trinks et al., 2005; Price, 2007; Vincent et al., 2004; Feller, 2017). It would seem plausible
to assume that cryogenic brines and their low-temperature precipitates could play (or could have played) a similar role for low-temperature biogenic reactions in the development of putative life in the low-temperature environments, for example, on Mars, Europa or Enceladus (Price, 2007, 2010).

1.2. Limitations of current models and ex situ measurements

Mathematical models, such as FREZCHEM (Marion & Kargel, 2008), use basic physical-chemical thermodynamic properties of multi-component solutions to predict the chemical composition of aqueous solutions at temperatures down to \(-113^\circ\text{C}\) and pressures up to 0.1 MPa, suitable for investigation of environmental settings on Earth, Mars and the icy satellites (Marion & Farren, 1999, Marion et al., 2003, Kargel et al., 2000). Although theoretical models have been used to predict the changing chemistry of sea ice brines on Earth and other planets during freezing, there is very little in situ field data to support the predictions. A further limitation to modelling is that predictions of the chemical composition of electrolyte solutions at sub-zero temperatures are rarely made at sub-zero temperatures. Our understanding of chemical equilibrium in cryogenic brines can thus be limited by large uncertainties, due to the estimation through extrapolation of the existing salinity-temperature functions down to the physical conditions for temperatures below 0 °C and salinities above 50 g/kg (Papadimitriou et al., 2017). In addition, thermodynamic modelling may not predict variations in solution chemistry, or mineralogy, in natural settings at sub-zero temperatures, since metastable states can both exist and, in some cases, persist; while environmental changes can often occur rapidly over timescales that are shorter than those required to reach thermodynamic equilibrium (Ewert & Deming, 2013).

Even on Earth there are problems with obtaining reliable field data. The inhospitable conditions that occur during austral winter limit the in situ sampling and
analysis of sea ice. Although retrieval of cores may be possible, post-collection changes in storage temperature and the methodology used to separate minerals from ice may result in changes, such that the sample under investigation no longer represents the environment from which it was extracted. Exploration of other planetary bodies necessitates the use of landers, with limited sampling and analysis capabilities, or by spectroscopic remote sensing via spacecraft fly-by missions, which have limited spatial resolution and surface penetration.

Laboratory simulations on the other hand offer the prospect of investigating low-temperature phases in a controlled environment. However compromises are often made due to managerial constraints such as time-limited experimental manipulations, access to facilities, or operational constraints such as minimum cooling rates achievable by the experimental apparatus. At worst, this can mean experimental outcomes are driven by kinetics rather than thermodynamics, such that the end point phases obtained are not those that would arise from thermodynamically driven systems, or that crucial intermediary phases are not observed (Butler et al., 2017).

1.3. Investigating aqueous precipitation at the Diamond Long-Duration Experiments facility

The recently constructed Long-Duration Experiments Facility (LDE) on Beamline I11 at the Diamond Light Source has been designed specifically to address the issues of long-term access to structure-probing facilities for slowly evolving systems, and is an ideal platform on which to conduct experiments and measurements of mineral precipitation in cold aquatic environments. The I11 LDE has been described in detail by Murray et al. (2017), however, in brief, the facility houses multiple long-running experiments (e.g. 6 months to 2 years), each mounted on individual motorised xy-stages, which are mounted on dedicated linear stages that run across a large granite sample table (Fig. 1). This enables each experiment to be brought separately into
the beam, such that X-ray transmission powder diffraction data can be collected via a large area detector (currently a Pixium RF4343, Thales), which is mounted on a linear drive that runs along the length of the sample table, allowing a predefined sample–detector distance to be set. Data collections are made weekly during each synchrotron machine run (~6-8 wks) and the wavelength and sample–detector distance each week is obtained by refinement of a standard CeO$_2$ calibrant that is either built-in to each sample cell, or permanently mounted on the sample stage. During machine shutdown periods (typically ~ 4 weeks), although X-ray data cannot be collected, parametric data (e.g. cell temperature) and ongoing control and status monitoring are continuously recorded and archived.

To study slow mineral precipitation in cold aqueous environments using the I11 LDE facility, a bespoke, transmission geometry cold cell has been designed that is capable of achieving very slow cooling ramp-rates. The cell can house five separate aqueous samples, of varying ionic composition, allowing the dynamic changes between ice, brine and mineral phases to be investigated simultaneously over the same controlled temperature range (room temperature to $-30^\circ$C). In this paper we describe the cell design and report results from a year-long study of the low-temperature precipitation of the mineral phase meridianiite (MgSO$_4$·11H$_2$O) from a MgSO$_4$-rich solution. Meridianiite is believed to be widespread on the present day surface of Mars and is a diagnostic for the loss of liquid water within a freezing, rather than evaporative, environment (Kargel, 1991; Hogenboom et al., 1995).

2. Cell design

2.1. Mechanical design

Computer Aided Design (CAD) renderings and a photograph of the installed cell are shown in Fig. 2. At the heart of the cell, the sample chamber consists of two bolt-
together solid copper body components with two 6 mm diameter circular apertures for beam transmission. A 60:40 glycol anti-freeze and demineralised water refrigerant flows in closed circuit through the blocks via an external Lauda ECO RE1050 chiller unit (GOLD control head). The small volume sample compartment is formed by two 0.05 mm thick diamond windows and two 0.5 mm thick Silicone O-ring with an inner diameter of 7 mm, such that the sample fluids do not come into direct contact with the copper body (see inset Fig. 2 middle). The sample block is enclosed within an insulating body made from 9 horizontally stacked layers of 25 mm thick Palight PVC foam board, each with a 0.025 mm Kapton window to give an insulating multi-glazing effect while still allowing the transmission of the incident and diffracted beams. The insulating layers are held together by brackets and through-bolts top and bottom. The overall outside dimensions of the cell are 300 mm \((w)\) × 250 mm \((h)\) × 225 mm \((l)\). The cell was mounted on one of the “medium” sized sample stages (see Murray et al., 2017 for details) on the large sample table in the I11 LDE hutch. Insulated services provide connection to the chiller unit located under the sample table, while a thermocouple embedded in one of the copper blocks connects to patch panel connections built into the sample stage itself. Since it is undesirable to thermally cycle the wavelength calibration standard, this is mounted to one side on the front Palight board with its own dedicated beam path through the upstream boards. There is a fixed 88.46 mm offset along the beam between the sample and standard positions.

The eutectic freezing of concentrated aqueous solutions results in pure-phase water ice, with any salts that were dissolved in the original liquid water being confined to saturated liquid brines, which form inclusions or channels than run through the mass of the frozen ice (Kutschan et al., 2010). Thus during the course of the experiment the 7 mm diameter sample compartment size means that the cell can be scanned both vertically and horizontally through the beam in order to provide representative aver-
age sampling, or to identify the brine-rich regions in which low-temperature mineral precipitation occurs.

2.2. Temperature control

The cell temperature is regulated via the Lauda chiller unit which is controlled externally – in open loop – via a computer script running on a dedicated EPICS IOC within the beamline controls system. This provides a programmable smooth decrease in temperature over a 24 hour period. Given a starting temperature, ramp rate and the total number of days over which the cell is to be cooled, the script calculates and updates the required chiller setpoint temperature every 5 s. No closed feedback exists between cell temperature and chiller temperature (i.e. no cell temperature dial-up capability) since the intended aim is to provide a selectable linear ramp rate irrespective of the actual temperature at any one time. The cooling rate of 0.3°C/day used in the work presented here was selected as it is broadly representative of the seasonal rate of temperature change during Arctic sea ice formation (Cox & Weeks, 1988) and also allows a reasonable difference in temperature to be achieved within the weekly measurement regime of the I11 LDE facility.

Repeated test runs showed a near perfect linear ramp for both chiller coolant temperature and the cell temperature (Fig. 3), with a very gradually increasing offset difference between the two of ~5 °C over the entire operating range. However this coolant-cell difference and the achieved final base temperature was found to be sensitive to the cooling rate and a rate of ~0.5 °C/day or less, allowed base temperatures below −29°C to be obtained. This matched the original design specification of −30°C, intended to cover the region of formation of hydohalite (NaCl·2H₂O), the most abundant mineral formed in terrestrial sea ice below −22°C (Butler et al., 2017). Faster ramp rates (>~1 °C/day) however resulted in raising the achievable base temperature
by several degrees.

Since the chiller is driven in open loop, in its current form the script applies a specific ramp rate for a set number of days, after which the achieved temperature can be maintained, or the sample thermally cycled by ramping up and down. Furthermore, as it is unlikely that an experiment running in a central facility environment continuously for one or more years will escape experiencing unforeseen outages, the script can also be stopped, edited and restarted, during which time the chiller holds at its last received set point temperature. Similarly, if, for whatever reason, network communications are lost between the IOC running the script and the chiller (or indeed loss/failure of the IOC itself), the chiller again maintains its last setpoint temperature. The open loop script approach thus provides a high degree of flexibility of control and ongoing management of the experiment. Backup power to the LDE facility itself similarly ensures continued electrical supply to the chiller in the face of power outages, planned or otherwise.

3. Experimental details

3.1. Commissioning experiment: precipitation in the MgSO$_4$--H$_2$O aqueous system

Evidence for the presence of magnesium sulphate in Martian soils and rocks first emerged from the Viking landings in 1977 (Clark & Van Hart, 1981; Clark, 1993) and was subsequently confirmed by later missions: the Pathfinder landers in 1997 (Rieder et al., 1997; Wänke et al., 2001) and the Mars Exploration Rovers Opportunity and Spirit in 2004 (Gellert et al., 2004; Rieder et al., 2004). In situ chemical analyses during these missions showed a strong correlation between sulphur and magnesium, pointing to magnesium sulphate being present in significant quantities. Later orbital observations showed hydrated sulphate outcrops, several km thick in the walls of Valles Marineris (Bibring et al., 2006). Although kieserite (MgSO$_4$.H$_2$O) was the first phase
to be considered likely (Baird et al., 1976), data from the Mars Express spacecraft (Gendrin et al., 2005) and the Mars Reconnaissance Orbiter (Roach et al., 2008) pointed to more highly hydrated sulphates. Indeed, observations show salts (sulphate, chloride and perchlorate) to be widespread across the planet’s surface (e.g. Gendrin et al., 2005; Squyres et al., 2006; Wang et al., 2006a; Hecht et al., 2009; Massé et al., 2010; McClennan, 2012). The presence of MgSO$_4$ on Mars is believed to have arisen from the reaction of basaltic material with sulphuric acid of volcanic origin and subsequent evaporation (Tosca et al., 2005).

Data from the Mars Odyssey Orbiter (Feldman et al., 2004a,b; 2011) allowed the global distribution of water-equivalent-hydrogen (WEH) down to $<1$ m depth to be measured. In two large regions near the Martian equator and in mid-latitude deposits buried in a large region of Arcadia Planitia, WEH concentrations of $\sim$50 wt% were found, and are much higher than could be accommodated by open pore-space volumes within the surface regolith. Three possible WEH carriers are thus likely (Feldman et al., 2004a,b; 2011): (i) ice, (ii) water adsorbed by mineral grains and (iii) hydrous minerals. At mid- to low-latitudes ground ice is unstable and while phyllosilicates and zeolites with 5-21 wt% structural H$_2$O/OH are common, their observed abundances cannot account for the observed WEH. However, with up to 62 wt% structural H$_2$O/OH, sulphates require less surface material to be present in order to account for the observed WEH. Although rare on Earth and limited to a few – mostly glacial and sea ice – occurrences, the undecahydrate sulphate meridianiite (MgSO$_4$·11H$_2$O) has the highest hydration state of all the magnesium sulphates and, under present day conditions, chemical models predict meridianiite to be the most abundant and widespread hydrous sulphate phase on the Martian surface (Toner et al., 2014). Indeed meridianiite derives its mineralogical name from the Opportunity rover landing site in Meridiani Planum (a plain located 2 degrees south of the Martian equator) where
its presence was deduced in situ from rover observations (Peterson & Wang, 2006; Squyres et al., 2007). Meridianiite forms at low temperatures as a precipitate from freezing staturated MgSO$_4$–H$_2$O solutions (Kargel, 1991; Hogenboom et al., 1995).

Laboratory studies and thermodynamic modelling show liquid water could theoretically exist – at least transiently – on the present day Martian surface (Brass, 1980; Haberle et al., 2001; Knauth & Burt, 2002; Chevrier & Altheide, 2008; Chevrier et al., 2009a,b; Möhlmann, 2011; Gough et al., 2011), but because surface conditions are close to the triple point of water, pure liquid water is likely to be unstable (Haberle et al., 2001). However, the presence of salts should lower both the freezing point and the evaporation rate of water. As a consequence, brines should be more stable and therefore more likely to be present on the Martian surface (e.g. Brass, 1980; Altheide et al., 2009; Zorzano et al., 2009; Möhlmann & Thomsen, 2010). These could have become systematically enhanced in Ca, Mg or Na depending on whether they are in contact with the CO$_2$-rich Martian atmosphere, or isolated in the sub-surface region (Knauth & Burt, 2002; Burt & Knauth, 2003). Experiments have shown halotolerant and sulphate-reducing bacteria can survive in highly concentrated sulphate-rich brines (Marnocha et al., 2011; Crisler et al. 2012). Under freezing conditions, the precipitation of hydrated mineral phases such as meridianiite from these enriched brines will occur. Consequently, the melting of meridianiite may thus be the most likely source of transient liquid water on the present day surface, especially at low latitudes where ice is less common in the regolith (Chou & Seal, 2007) and may contribute to the formation of the chaotic outflow terrains observed on the Martian surface (Peterson & Wang, 2006).

Understanding the precipitation of hydrated MgSO$_4$·$n$H$_2$O phases such as meridianiite thus impinges on understanding both the present-day hydrological cycle on Mars as well as the formation and distribution of relic hydrated phases associated with the
historic loss of Mars’ ancient oceans. Therefore, as a commissioning experiment for the LDE cold cell, we monitored over a period of one year the behaviour of a slowly cooling MgSO₄·H₂O solution formulated to precipitate MgSO₄·11H₂O. During this time we observed the precipitation of two hydrated MgSO₄ phases and a disordered ice/sulphate phase and monitored their long-term dynamic behaviour.

3.2. Sample preparation, data collection and reduction

In what follows we will, for ease of comparison, refer to the various MgSO₄·nH₂O phases as MSn, such that meridianniite, with 11 waters of hydration will be MS11, epsomite (MgSO₄·7H₂O) will be MS7 etc. When necessary, use of mineral names will be restricted to the scientific discussion and their MSn given in parenthesis.

For the commissioning experiment an aqueous solution of 8 wt% concentration of MgSO₄ was made by mixing 5 g of reagent grade MS7 with 24 g of 18 MΩ.cm deionised water. With part of the copper block assembly on its back and a diamond window and O-ring in place, the solution was dropped into the O-ring centre using a pipette. The second window and O-rings were inserted, followed by the second half of the copper body and the cell assembled and mounted on the beamline (Fig. 2).

Prior to the start of the experiment, the detector was set to a nominal 350 mm from the sample, sufficient to give a reasonably wide 2θ range of 42°, and the centre position of each cell was determined by scanning the cell chamber both horizontally and vertically through the beam. This position was entered into a data acquisition script such that the chamber returns to the same sampling position each week. Diffraction data were then collected each week (see Murray et al., 2017 for details) at this optimal in-beam position and for the present work at a number of plus-or-minus vertical positions offset from the centre by 1-2 mm. The beam size was 400 μm × 400 μm, achieved by slitting down the incident 25 keV X-ray beam. Each week, using data
from the CeO$_2$ reference standard, the wavelength and detector distances were refined by the beamline data acquisition and reduction pipeline that incorporates the DAWN software suite (Filik et al., 2017), which was also used to integrate and convert the 2-D images collected from each cell position to conventional 2$\theta$ versus Intensity ASCII files.

In this type of aqueous brine system and environment, ice formation can be problematic for the collection of powder diffraction data. In particular the formation of large ice crystals can result in very intense diffraction spots occupying a significant area of the detector. While it is, in principle, possible to mask these out prior to integration, the volume of data accumulated and the variability in their occurrence from week to week (such that a new mask would need to be drawn for each individual image) meant that this was not a feasible option in the first instance (see discussion section for future developments); while simple thresholding produced variable results such as halo artefacts, missed spots and loss of legitimate signals. Instead all images were integrated and the 1-D data plotted together on the same axes. Those datasets collected at different positions showing large, intense and wide non-background features were rejected, while the remaining data were averaged, thus producing a single data file for each week.

4. Results

4.1. Long-term behaviour of the MgSO$_4$-H$_2$O system

In order to start from an ice-brine system, the cold cell was supercooled to $-8^\circ$C to promote initial freezing, after which the temperature was ramped at $-0.3^\circ$C per day under computer control. Taking this as day 0, Fig. 4 plots the measured cell temperatures on the days X-ray data were collected, while Fig. 5 shows a typical image collected from the area detector. In this figure can be seen (a) relatively good powder
rings due to the cell’s diamond windows, (b) very spotty rings due to a combination of crystalline ice and sulphate precipitate and (c) diffuse scatter coming from an amorphous component. In Fig. 4 three key points, A, B and C are indicated, corresponding to 30, 70 and 184 days. The corresponding 1-D integrated and averaged diffraction patterns for these are shown in Fig. 6. In this figure the large Bragg peak just below 14°2θ is diamond (as are smaller features at ~22.5, 26.5, 32.2 and 35.2 °2θ), with other Bragg peaks being due to hexagonal ice and sulphate precipitate. There is also a very clear difference in the background signal. Since the pattern at C was collected at a much lower temperature than the pattern at A, and after some considerable time, this suggests a time/temperature evolution. Fig. 7 plots the intensity of the background signal at 12.5 °2θ as a function of time. This angle was selected because it represents a high point in the background and does not coincide with any Bragg features in the data. To account for the variation in beam intensity from week to week due to variations in the synchrotron beam fill and small errors in the repeatability and alignment of the beamline optical components, the measured background intensity was normalised to the total scattered intensity of the CeO$_2$ standard attached to the cell. This reveals an interesting behaviour. Firstly, the background only starts to increase once point B in Fig. 4 is reached after ~70 days. Secondly, the background increases to a peak value at point C and then decreases, reaching a minimum somewhere between 290 and 320 days (the large gaps between points being due to machine shutdowns), after which it increases in intensity again. Fig. 4 identifies point B as the point when the cell temperature drops below −22°C (~251 K). Below this temperature the number of Bragg peaks in the integrated patterns also increased and could be fitted by a mixture of hexagonal ice, MS7 and MS11 (see section 4.3). The rise in the strength of the background is caused by the precipitation of the sulphate phases from the brine. The associated removal of Mg$^{2+}$ and SO$_4^{2−}$ promotes instantaneous ice formation to
re-establish ice–brine equilibrium at these low temperatures, which concentrates the brine and results in further MgSO$_4 \cdot n$H$_2$O precipitation. This cycle propagates in tandem until ice–brine–MgSO$_4$ equilibrium is attained. Because the temperature is so low the precipitation of ice occurs rapidly, resulting in a disordered ice phase and an increase in the diffuse scatter.

4.2. Sample temperature

Fortes et al. (2008) analysed the thermal expansion behaviour of MS11 from 250 to 4 K in terms of an Einstein oscillator model for the specific heat and we use this here to show that the temperature achieved by the sample is accurately represented by the temperature recorded by the thermocouple embedded in the cell body. The Einstein model assumes all atoms in the solid vibrate at the same characteristic angular frequency, $\omega_E$, expressed in terms of an Einstein temperature, $\theta_E = h\omega_E/k_B$. Thus the values of each of the MS11 lattice parameters, $X$, at a given temperature, $T$, can be expressed relative to their values at absolute zero, $X_0$ by,

$$X(T) = X_0 + \frac{E}{\exp(\theta_E/T) - 1}, \quad (1)$$

where $E$ is a constant given by

$$E = 3R\gamma\theta_E/K_T, \quad (2)$$

for which $\gamma$ is the Grüneisen ratio relating the volume thermal expansion coefficient, $\alpha_V$, specific heat capacity, $C_V$, molar volume, $V$, and the isothermal bulk modulus, $K_T$. Assuming $\gamma$ and $K_T$ are independent of temperature, the thermal expansion behaviour is then a function of the internal energy of the crystal via its heat capacity. Although not as rigorous as Debye formalism, the Einstein model has (as noted by Fortes et al.) been successfully used to model various related inorganic solids such as CaSO$_4$·2D$_2$O (Knight et al., 1999) and is mathematically simple. Fortes et al. parameterised $E$ by
a polynomial function,

\[ E = e_0 + e_1 T + e_2 T^2 + e_3 T^3, \]  

obtaining values for the \( e_n \) coefficients from fits to their measured data. Since the temperature range of the cold cell overlaps the temperature range covered by the Fortes et al. neutron study, we used their values of \( e_n \) and \( \theta_E \) to calculate expected values for the meridianiite lattice parameters near the base temperature recorded by the cold cell thermocouple (−23 to −28°C) and compared them (Table 3) to values obtained from fits (see next section for details) to the cold cell diffraction data collected at these temperatures. Given the limitations of the Einstein model, the agreement between calculated and fit parameters is to within \( \sim 1\% \) and shows that the temperatures being achieved by the sample within the cell are well represented by the thermocouple measurement, confirming the cell design and its appropriateness for this type of medium-to low-temperature study.

4.3. Data analysis: ice formation and sulphate precipitation

A major problem in the MgSO₄·H₂O system is the nature of the precipitate. Many of the possible MgSO₄·nH₂O phases \((n = 1, 2, 3, 4, 5, 6, 7, 11)\) share the same unit cell structures (see Table 2) and during profile refinement, due to the large numbers of closely spaced reflections, can almost all fit the same medium resolution data out to high angle. Thus, to distinguish between precipitated phases, we restricted our analysis to low angles where differences in cell dimension offer the prospect of distinct diffraction peak positions (see also discussion in section 5.1 for improvements to data collection). Conveniently the cubic unit cell of the diamond windows yields a strong and narrow Bragg peak at \( \sim 14^\circ 2\theta \) and provided both a useful cut-off and additional intensity calibrant (see below). We also used this feature to refine the \( 2\theta \) zero point, allowing the diamond lattice parameter to float to account for the change in temper-
ature. The similarity of the pattern of diffuse background scattering recorded in the area detector images to patterns of hexagonal ice with static disorder (e.g. compare Fig. 8 with Fig. 1 in Wehinger et al., 2014) meant that our first step in obtaining Pawley fits to the low-angle diffraction data was to include two hexagonal ice phases: one to account for the narrow Bragg features of crystalline ice and the other to account for the broad diffuse contributions. Following this, profiles for MS7 and MS11 were first refined separately and then together to obtain the relative contributions of each phase. However, in doing this we found that the fits could be improved by including an amorphous sulphate phase (sometimes two) to better model the background and suggesting that the rise in diffuse scatter may also contain contributions from amorphous or nanosized sulphate. Although it is unrealistic to suggest that an amorphous sulphate component can be identified in this way as, in the fit, it is really only accounting for deficiencies in the background function, we found certain phases gave better improvements than others. For example amorphous MS2, MS4 and MS5 tended to provide improvements for the earlier data, while MS4, MS5 and MS6 gave improvements for data collected later in the run. Including amorphous MS1 and MS3 rarely gave improvements and adding in additional MS7 or MS11 phases never improved the fit. It is however interesting to note that the formation of amorphous MS2 and MS3 from MS7 under desiccating conditions at \(-8^\circ\text{C}\) has previously been observed (Wang et al., 2006b), which although amorphous by X-ray diffraction, were readily identifiable by the position of the strong Raman feature near 1000 cm\(^{-1}\).

To gain a picture of how the precipitate system evolved with time, the area under the fit for each component was normalised to the area of the 14 °2\(\theta\) diamond feature, and are shown in Fig. 9. Interestingly, a peak in both MS11 and MS7 is observed at \(\sim340\) days that immediately follows the minimum seen in the background signal (Fig. 7).
Over the course of the experiment we have thus observed the following sequence: formation of a first ice phase as the system initially cools, followed at $\sim 70$ days, once the cell temperature has dropped below $\sim 23$ °C, by the simultaneous precipitation of MS7 and MS11 and the formation of a disordered phase. The disordered phase continues forming, while the MS7 and MS11 levels increase little, if at all. However, after 100 days the levels of MS7 and MS11 start to increase, until at $\sim 280$ days MS7 increases more steeply, followed at $\sim 320$ days by MS11. Meanwhile, at $\sim 200$ days the disordered phase starts to decrease, reaching a minimum somewhere around $300$ days, after which it starts to increase again. At the end of the first MS7/11 growth period the ratio of MS11:MS7 is approximately 4:1, while after the second it is approximately 2:1, suggestive of some ongoing processing relationship between the sulphate, ice and/or the disordered phase.

5. Discussion

5.1. Sulphate precipitation sequence

The 8 wt% MgSO$_4$ concentration and base temperature (−28.2°C) are marked on the MgSO$_4$-H$_2$O phase diagram shown in Fig. 10. However, the MgSO$_4$-H$_2$O phase diagram derives from those presented by Kargel (1991) and Hogenboom et al. (1995) and are based on much earlier studies (citations in Kargel and Hogenbloom et al.) combined with results obtained by Hogenboom et al. (note that until Peterson & Wang (2006) solved the structure by single crystal X-ray diffraction and confirmed the asymmetric unit contains only 11 waters, MS11 was misidentified as MS12 in earlier work).

Tracing down the 8 wt% line superimposed on the MgSO$_4$-H$_2$O phase diagram suggests the sequence of

$$\text{liquid} \rightarrow \text{ice + liquid} \rightarrow \text{ice + MS11}$$

(4)
should be observed. At 0.1 MPa (atmospheric pressure) the initial ice phase should be hexagonal ice, which we observe. However the MgSO$_4$-H$_2$O equilibrium phase diagram does not indicate the formation of a disordered phase or MS7. Our observed sequence is

$$\text{liquid} \rightarrow \text{Ih + liquid} \rightarrow \text{Ih + D + MS7 + MS11},$$  

(5)

where D represents the phase responsible for the diffuse scattering. Although Hogenboom et al. did not explore the behaviour of a 8 wt% solution at atmospheric pressure, they did observe the precipitation of metastable MS7 in the MS11 stability field for higher concentrations (e.g. 22 wt% solution) and at high pressures. They also noted that MS11 forms more slowly than MS7 (also noted by Kargel, 1991 at atmospheric pressure) and may be related to the difficulty in achieving equilibrium crystal nucleation during cooling of the saturated MgSO$_4$ solution. This phenomenon is common among binary and multicomponent systems and especially so in MgSO$_4$ solutions (Kargel, 1991), occurring when the cooling rate is too fast and the system supercools. This suggests that even at a cooling rate of 0.3°C/day the MgSO$_4$-H$_2$O system within the cold cell is still out of equilibrium. However, we do note that in order to promote the first ice freeze and to form liquid brines within the ice the cell was fast ramped to $-8^\circ$C over the course of $\sim$1 hour. The first ice freezing will increase the wt% concentration of MgSO$_4$, moving the red dashed line in Fig. 10 to the right. When the temperature drops to $-23^\circ$C and MS7 and MS11 start precipitating out, the formation of the disordered phase (which we assume is a mixture of disordered ice and amorphous MS) concentrates the brine further. Thus even though MS11 is thermodynamically favoured, and initially forms in greater quantities (i.e. 4:1) the faster growth rate for MS7 means that by the end of the observed growth period the ratio of MS11:MS7 has reduced to approximately 2:1. The fact that we observe two different growth phases for both MS7 and MS11 points to a complex interplay between the
precipitated phases and the trapped brine.

Towards the end of the experiment, at around \( \sim 340 \) days, MS11 and M7 reach a peak and start to decline, while the background signal, which had previously reduced in intensity, had been increasing since \( \sim 310 \) days. Although we have no definitive explanation at this point, it may be possible that the sulphate phases reach a point where they are unstable with respect to the remaining brine and begin re-dissolving, releasing their waters of hydration, which due to the low temperature rapidly freezes as part of the disordered phase. Whether this process results in nano or amorphous MS4/MS5/MS6 components (as suggested in section 4.3) derived from the excess MS11/7 being incorporated into the disordered phase remains to be seen. If this explanation holds true, then we might predict that this growth and decline behaviour is part of a longer-term behaviour, akin to damped oscillations, as the system eventually moves towards equilibrium. Operational constraints unfortunately meant that we were unable to run the experiment beyond a year.

5.2. Future upgrades and improvements

As discussed above, the similarity in structure of many of the precipitating salt systems means that resolving closely located or overlapped diffraction features is important. In the I11 LDE setup there are several limiting factors that may be possible to improve or optimize.

5.2.1. Resolution  Small beam sizes are currently achieved by slitting down the beam to improve resolution and is obviously at the expense of incident intensity, requiring longer collection times to compensate. To overcome this, an F-switch ‘random access’ compound refractive lenses manipulator (Duller et al., 2016) is currently being commissioned. Based on the transfocator concept (Vaughn et al., 2011), the F-switch
houses 120 selectable compound refractive lenses which can be chosen to provide selectable spot sizes at a given position, typically from 100 μm down to 5 μm diameter, while still proving a gain in intensity and a reduction in data collection times. This in turn will provide either for increasing the number of spatially resolved points across the sample at which data is collected, or for the direct sampling of individual brine inclusions and channels. A further advantage of introducing a focusing element will be to increase the divergence of the incident beam over that of the undulator beam, which should improve powder averaging around the diffracted Debye-Scherrer rings.

Neglecting the sample thickness contribution, for a regular shaped beam of dimension $d$, in transmission from a sample to a detector with pixel size $p$ at distance $D$ away, the $2\theta$ resolution (towards the centre of a flat detector) is approximately given by

$$\Delta 2\theta \approx 2\arctan \left( \frac{d + p}{2D} \right)$$

(6)

Fig. 11 shows $\Delta 2\theta$ as a function of detector distance, calculated for the Pixium 4343 area detector currently deployed on the I11 LDE and which has a pixel size of 148 μm. For $400 \times 400$ μm beam size, distances of 500 and 950 mm should give $\Delta 2\theta$ of $\sim 0.063^\circ$ and $\sim 0.033^\circ$ respectively, while a beam size of $300 \times 300$ μm should yield $\sim 0.051^\circ$ and $\sim 0.027^\circ$ respectively. Also shown in Fig. 11 is $\Delta 2\theta$ calculated for a Perkin Elmer 4343 XRPAD area detector, which has a pixel size of 100 μm. Not available when the LDE facility was constructed, a Perkin Elmer detector is currently in commissioning and at the long detector distance and smaller beam size should in principle provide further improvement in resolution. Shown in Fig. 12 are the (Pixium) measured 1 1 1 Bragg peaks of the cold cell’s CeO$_2$ reference standard for the same detector distances and beam sizes, obtained by integrating around the ring. At 500 mm from the sample the FWHM for $400 \times 400$ μm and $300 \times 300$ μm are 0.1 and 0.08 and are clearly
quite large, while at 950 mm the FWHM are 0.086 and 0.036 respectively, with the latter lying closer to the calculated $\Delta 2\theta$. The discrepancy at shorter detector distance likely originates from the use of mechanical slits to define the beam, sample thickness, increased path length to higher detector angles at shorter distances, and the associated increase in spread of the scattered beam as the incident angle increases. Fig. 12 also highlights the decrease in the diffracted signal strength incurred by slitting down the beam to increase resolution.

A detector distance of 950 mm means that the powder ring from the cold cell’s diamond window is just contained within the active area of the detector, allowing maximum separation of the lower angle precipitate peaks. However there is a practical limitation to the maximum usable detector distance imposed by the need to simultaneously calibrate both the wavelength and detector distance. As the detector distance increases, the number of CeO$_2$ rings captured by the detector also decreases, such that the uncertainty in the wavelength and $2\theta$ zero point also increase. For 25 keV X-rays at this distance seven CeO$_2$ rings are captured, aided by the fact that the CeO$_2$ standard is offset closer to the detector than the sample by 88.46 mm. Increasing this offset could allow for more rings to be captured; but to be useful the standard mount would need to extend outside the sample stage footprint and risk fouling adjacent experiments.

5.2.2. Data reduction and pipelining The current data collection and reduction pipeline per weekly beamtime collection consists of the following steps: (i) using area detector collect $N$ patterns at different positions of the sample; (ii) integrate the 2-D detector images to $N$ 1-D data files; (iii) rejecting those 1-D patterns that contain significant contributions from large ice spots; (iv) sum or average the remaining 1-D files and normalise to the integrated intensity of the CeO$_2$ reference standard. Steps (ii) to
(iv) are currently done entirely manually and are dependent on the operators sense of what constitutes a “bad” pattern and is wasteful of data in that a single large spot in a detector image means that the useful information contained in the rest of the image is being discarded. To improve on this, the process ideally needs the individual area detector images to have a mask applied prior to integration. In normal circumstances an operator would define the mask position interactively using the reduction software, and this would be applied to all subsequent images. However in the case of the cold cell data, not only are the large spots in different places on the image from position to position in the sample, but also vary from week to week due to the dynamic nature of the ice-brine system. Automatic detection and masking will solve this problem, but will need to be sufficiently robust that (a) it does not inadvertently remove good data and (b) does not introduce artefacts into the signal.

5.3. Current deployment and future experiments

Following on from these commissioning experiments a number of low-temperature aqueous precipitation experiments are in progress with, future experiments planned. Although the outcome of these will be reported in separate publications (e.g. Butler et al., 2018) we highlight some of them here.

5.3.1. Sea ice in Earth’s polar oceans  As mentioned in the introduction, the formation of minerals in sea ice has an important role to play from both an ecological and climatological perspective. However, in situ measurement of the composition, distribution and dynamics (solubility, precipitation and dissolution) of authigenic mineral assemblages in freezing ocean waters have only recently begun. This and the dependence on thermodynamic modelling, reliant on interpolation and extrapolation, clearly limit our ability to understand the evolutionary sequence of mineral dynamics in cold environ-
ments, where the kinetics may mean that thermodynamic equilibrium is not attained or maintained before environmental conditions change. Such environments in Earth’s polar oceans are the least understood in terms of their input to climate change. Compositional changes take place within thickening sea ice, where ice temperature varies both with depth and with season. The complexity of the slowly evolving sea ice and changes in brine composition and consequent mineral formation has not yet been measured using conventional laboratory techniques. The 0.3 °C/day cooling rate used in the commissioning work reported here also represents a good proxy for the change in temperature observed in the natural setting at the ice-air interface in the Arctic ocean during a seasonal cycle of sea ice formation.

Seawater has a typical dissolved salt content (salinity) of ~35 g/kg and currently three conservative seawater brines are under investigation using the cold cell, with salinities of 100, 75 and 35 g/kg. Of particular interest will be the verification of the time minerals such as hydrohalite take to reach equilibrium under the conditions that are equivalent to those found in situ, within the brine pores and channels of sea ice. A more complete description of the mineralogical evolution of sea ice that these experiments could provide would allow for a quantitative assessment of the contribution of sea ice minerals to the processes driving global warming and cooling cycles, both historically and during the present day.

5.3.2. Ocean worlds in the Solar System Among the ice and ocean worlds of the Solar System Europa is thought to be unique because observations and models suggest its ocean is in direct contact with its rocky interior. Therefore the potential chemical composition of the Europan ocean can be constrained through possible weathering/leaching reactions and the assumed composition of its carbonaceous chondrite core. From observational data, the salinity of Europa’s subsurface ocean is believed to
be 3-15 g/kg (Hand & Chyba, 2007) and a model Na-Mg-Ca-Cl-SO$_4$ solution was prepared gravimetrically with the following ionic composition in mol kg$^{-1}$H$_2$O: Na = 1.630, Mg = 2.929, Ca = 0.0064, SO$_4$ = 3.5964 and Cl = 0.308. The solution is cooling at the same rate as the seawater and brine samples and the expected phases at various temperatures include MS7, CaSO$_4$·2H$_2$O (gypsum), Na$_2$SO$_4$·10H$_2$O (mirabilite), MS11, water ice and NaCl·2H$_2$O (hydrohalite). However, as we have already shown with the MS11 results obtained thus far, precipitate behaviour may not be fully described by the current thermodynamic model.

Other possible future targets include Titan which, aside from Earth, is the only Solar System body with a thick N$_2$ atmosphere and the only moon to retain a substantial atmosphere. This makes Titan home to a complex atmospheric chemistry and the only other solar system body with stable liquid currently on its surface. The combination of oxygen-bearing molecules and favourable conditions for organic haze formation means that atmospheric processes could yield molecules of prebiotic interest. Similarly, a combination of organics and liquid water in a subsurface ocean and methane and ethane in surface lakes and oceans makes Titan a prime target for furthering our understanding of prebiotic chemistry, habitability and the potential for life to exist beyond Earth (Horst, 2017). The presence of a saline subsurface ocean was confirmed by the Huygens probe that descended to the surface (Béghin et al., 2012) and by radio Doppler tracking (Iess et al., 2012), while analysis of Titan’s gravity and topography constrain the subsurface ocean density to be slightly higher than pure water, suggesting the presence of salts (Mitri et al., 2014), with ammonium sulphate ((NH$_4$)$_2$SO$_4$) being considered a likely constituent (Fortes et al., 2007; Grindrod et al., 2008). Such an ocean would have a freezing temperature of approximately $-23^\circ$C (Grindrod et al., 2008), which is well within the cold cell’s specification.
6. Conclusions

We have presented the design of a cold cell for long-duration studies of mineral precipitation from cold aqueous environments and demonstrated its performance. Along with this we reported on a commissioning experiment using the MgSO$_4$·H$_2$O system showing its capability for uncovering new scientific phenomena. Specifically we have observed under slow 0.3 °C/day cooling that, as well as the predicted hexagonal water ice and MgSO$_4$·11H$_2$O phases, metastable MgSO$_4$·7H$_2$O and a disordered ice-rich phase are also co-formed. Following this system for one year, using the beamline I11 Long-Duration Experiments facility at the Diamond Light Source (UK), we observed a complex low-temperature dynamic behaviour, relating to the differential growth rates of the two sulphate phases, the time dependent growth and variation of the disordered phase caused by changes in ionic concentration brought about by the precipitation of the sulphates. We have also discussed how data quality for future experiments can be improved and outlined some of the current and planned experiments for this facility.

Cold cell experiments will produce the first in-situ observations of mineral formation during the evolution of brine chemistry in aquatic environments that are characteristic of oceans on Earth and other planetary objects, and are undergoing freezing over environmentally relevant timescales. Such novel results will contribute to our understanding of the global processes that occur on, and shape, such bodies and the results will determine how closely solution chemistry is linked to thermodynamic equilibrium and to mineral solubilities currently predicted by the extrapolation of relevant data down to sub-zero temperatures.

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Table 1. Ocean worlds of the Solar System grouped according to current evidence (drawn from the literature, though see reviews by Husseman et al., 2006; Nimmo & Pappalardo, 2016; Lunnie, 2017 and Mann, 2017). The first group are where extensive multi-channel observations, remote sensing by lander’s and flyby missions unequivocally point to liquid oceans. The second is where calculations and models using current observational data support the likelihood of there being an ocean, while the third category is where objects, based on known observational properties could support a (mostly subsurface) ocean if certain conditions were met, e.g. ocean of sufficient salinity to act as anti-freeze, or internal heating occurs. This latter category also includes the larger trans-Neptunian objects (TNOs), where surface ices are observed, a few examples of which are listed.

**Supported by observation**

| World   | Description                                      |
|---------|--------------------------------------------------|
| Earth   | Circulating surface oceans, polar ice caps.       |
| Mars    | Polar ice, evidence of global ancient surface ocean. |
| Ceres   | Evidence of ancient, possibly surviving, sub-surface ocean, |
| Europa  | Ice crust, evidence for sub-surface ocean.         |
| Ganymede| Ice crust, may have stacked oceans separated by ice layers. |
| Callisto| Evidence shows global sub-surface fluid, possibly water. |
| Enceladus| Ice crust, evidence (plumes) of saline sub-surface ocean. |
| Dione   | Ice crust, evidence for global sub-surface ocean.  |
| Titan   | Ice crust, evidence for saline global ocean.       |
| Pluto   | Plausible interpretation of limited observational data. |

**Predicted by theory and modelling**

| World   | Description                                      |
|---------|--------------------------------------------------|
| Triton  | Water is lesser constituent of surface ices, ocean predicted. |
| Mimas   | Sub-surface ocean proposed to explain orbital wobble. |
| Charon  | Possible evidence supporting frozen sub-surface ocean. |
| Oberon  | Ice on surface, sub-surface ocean considered possible. |
| Orcus   | Evidence of ancient sub-surface ocean, may have survived. |
| Titania | Possible global sub-surface ocean.                |

**Plausible under certain conditions**

| World   | Description                                      |
|---------|--------------------------------------------------|
| Tethys  | Pure water ice surface, internal structure unknown. |
| Rhea    | Sub-surface ocean plausible from known properties. |
| TNOs    |                                                  |
| 2007 OR10 | Bulk density, surface colour and spectroscopic evidence consistent with liquid surface water |
| Haumea  | activity, possibly due to tidal heating caused |
| Quaoar  | orbital interactions with moons. Models suggest |
| Makemake| heating by radioactive decay could support sub-surface |
| Eris    | oceans in many larger TNOs.                      |
| Sedna   |                                                  |
Table 2. Lattice parameters of the various MgSO₄·nH₂O phases used as starting values for fits to cold cell data. Values taken from International Center for Diffraction Data PDF 4 database.

| phase | space group | a (Å)    | b (Å)    | c (Å)    | α (°)     | β (°)     | γ (°)     |
|-------|-------------|----------|----------|----------|-----------|-----------|-----------|
| MS1   | A2/a        | 7.511    | 7.611    | 6.921    | 90.00     | 116.17    | 90.00     |
| MS2   | P212121     | 8.8934   | 12.440   | 8.488    | 90.00     | 90.00     | 90.00     |
| MS3   | Pbca        | 10.930   | 12.420   | 8.200    | 90.00     | 90.00     | 90.00     |
| MS4   | P21/n       | 7.902    | 13.594   | 5.920    | 90.00     | 90.89     | 90.00     |
| MS5   | P-1         | 6.335    | 10.550   | 6.075    | 99.17     | 109.88    | 75.00     |
| MS6   | A2/a        | 24.442   | 7.216    | 10.119   | 90.00     | 90.00     | 90.00     |
| MS7   | P212121     | 11.910   | 12.020   | 6.870    | 90.00     | 90.00     | 90.88     |
| MS11  | P-1         | 6.732    | 6.792    | 17.293   | 88.22     | 89.49     | 62.66     |

Table 3. Parametric MgSO₄·11H₂O lattice parameters at absolute zero, X₀, Einstein temperature, θₑ, and polynomial coefficients, eₙ for the constant E given by equation (2) taken from Fortes et al. (2008); lattice parameters refined from data collected at -23.3 and -28.2 °C compared to calculated values at the same temperature using the parametric values for the lattice parameters at absolute zero.

| parametric | a (Å)    | b (Å)    | c (Å)    | α (°)     | β (°)     | γ (°)     |
|------------|----------|----------|----------|-----------|-----------|-----------|
| X₀         | 6.72740(5) | 6.78187(5) | 17.31589(14) | 88.2126(9) | 89.4499(9) | 62.6109(8) |
| θₑ         | 340(3) | 179(11) | 80(6) | 79(170) | 195(109) | 185(23) |
| e₀         | 6.9(1)E-2 | 9.6(19)E-3 | -3.2(5)E-2 | 1.0(7)E-2 | 1.1(1)E-1 | 2.1(6)E-1 |
| e₁         | 0      | 9.8(4)E-5 | 2.2(5)E-4 | -5(9)E-4 | -1.7(18)E-3 | -9.1(39)E-4 |
| e₂         | 0      | 0       | -9.4(24)E-7 | 3(6)E-6 | 1.5(13)E-5 | 1.8(8)E-6 |
| e₃         | 0      | 0       | 1.6(4)E-9 | -6(15)E-9 | -3.8(31)E-8 | 0 |

| cell at -23.3 °C | a (Å) | b (Å) | c (Å) | α (°) | β (°) | γ (°) |
|------------------|-------|-------|-------|-------|-------|-------|
| calculated       | 6.75  | 6.81  | 17.28 | 88.1  | 89.4  | 62.6  |
| refined          | 6.72(2)| 6.77(2)| 17.29(3)| 87.8(1)| 89.1(2)| 62.2(2)|
| % difference     | 0.44  | 0.59  | 0.06  | 0.34  | 0.34  | 0.64  |

| cell at -28.2 °C | a (Å) | b (Å) | c (Å) | α (°) | β (°) | γ (°) |
|------------------|-------|-------|-------|-------|-------|-------|
| calculated       | 6.75  | 6.81  | 17.29 | 88.2  | 89.5  | 62.7  |
| refined          | 6.78(2)| 6.88(2)| 17.36(4)| 86.9(2)| 89.8(2)| 62.8(3)|
| % difference     | 0.44  | 1.03  | 0.40  | 1.47  | 0.34  | 0.15  |
Fig. 1. CAD drawing of the I11 Long-Duration Experiments sample table showing main components. Each sample stage hosts an individual experiment; the area detector moves along the table to a pre-defined distance as each experiment is brought successively in and out of the X-ray beam (see Murray et al. 2017 for full details).
Fig. 2. Top and middle: CAD schematics showing front and rear (beam exit and entrance) sides of cold cell design (top); internal construction (middle); Bottom photograph of cell installed on the I11 beamline LDE facility.
Fig. 3. Screenshot showing cell temperature (red, top trace) and chiller temperature (blue, bottom trace) during script-controlled cooling ramp covering a one month period. The difference between measured sample temperature and chiller temperature is initially \( \sim 5 \) C, widening slowly as the temperature decreases.
Fig. 4. Cell temperatures at data collection dates. Gaps are due to synchrotron shut-
downs. Arrows A, B and C indicate 30, 70 and 184 days (T = -12.7, -23.3 and -27.7
°C respectively) for comparison with data shown in Figs. 6 and 7.
Fig. 5. Area detector image collected at point $B$ (70 days, $T = -23.3^\circ C$, $\lambda = 0.49392$ Å, detector distance = 335.097 mm) in Fig. 4 showing presence of diffuse scattering.
Fig. 6. Diffraction patterns of MgSO₄·H₂O system obtained after (bottom to top) 30, 70 and 184 days (T = -12.7, -23.3 and -27.7 °C; λ = 0.49383, 0.49392 and 0.49412 Å; detector distance = 335.153, 335.097 and 335.072 mm respectively). Patterns correspond respectively to points A, B and C in Fig. 4 and show increase in diffuse background scatter.

Fig. 7. Diffraction background signal at 12.5 °2θ normalised to the integrated intensity of the CeO₂ standard as a function of time. Arrowed points as per Fig. 4.
Fig. 8. Area detector image collected at point C (184 days, T = -27.7 °C; \( \lambda = 0.49412 \) Å, detector distance = 335.072 mm) in Fig. 4 shown in negative to emphasise the symmetry of the diffuse scatter, which exhibits very similar symmetry to diffuse scatter observed in hexagonal ice with static disorder (Wehinger et al., 2014).
Fig. 9. Ratio of the area of the diffraction fit attributed to (a) MS11, (b) MS7 to the area of the first diamond peak from the cell window. Dotted lines are 2-point moving averages to highlight trend.
Fig. 10. Phase diagram for H$_2$O-MgSO$_4$ system (after Hogenboom, 1995; Peterson & Wang, 2006 and Fortes et al., 2008). Red vertical dashed line represents the 8 wt% starting composition used in this work, black square superimposed on this represents the achieved base temperature of the cell.
Fig. 11. Predicted detector resolution as a function of distance from the sample for two beamsizes defined by a square slits of dimension 400 and 300 μm towards the centre of two area detectors: Thales Pixium 4343 with pixel size 150 μm and Perkin Elmer 4343 with pixel size of 100 μm.
Fig. 12. CeO$_2$ 1 1 1 peak measured ($\lambda = 0.49319$ Å) for different nominal detector distances and beam size combinations. Strongest trace (solid line): detector distance 350 mm, beam size 400 $\times$ 400 $\mu$m; middle two traces: (dash and dash-dot-dot) detector distance 350 mm, beam size 300$\times$300 $\mu$m and detector distance 800 mm, beam size 300 $\times$ 300 $\mu$m respectively; weakest trace (dash-dot): detector distance 800 mm, beamsiz 300 $\times$ 300 $\mu$m.

**Synopsis**

A sample environment cell for long-duration X-ray diffraction studies of mineral precipitation in very slow-cooling aqueous environments on Earth and other planetary bodies is described. The Results of a year-long commissioning experiment monitoring the dynamics of a freezing MgSO$_4$-H$_2$O solution at -28 °C in which meridianite and epsomite are observed to form are reported.