Raman tomography of natural air hydrates

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ABSTRACT. Cryo-Raman tomography allows us, for the first time, to determine accurate morphologies and volumes of natural air hydrates in Antarctic ice cores. The measurements show complex growth structures that are not accounted for by the available models of hydrate formation.

KEYWORDS: clathrate hydrates, crystal growth, ice core, ice rheology, paleoclimate

INTRODUCTION

Polar ice is a composite of ice, air and various impurities of atmospheric origin. This makes ice cores unique climate archives, because the air inclusions in the polar ice sheets are the only direct source for measuring and reconstructing the composition of the atmosphere of the past (e.g. Raynaud and others, 2000). To correctly interpret the findings, a firm understanding of the processes that may be able to change the gas content of the air inclusions is mandatory.

The precursors of polar ice, snow and firn, form a porous permeable material up to 120 m thick on top of the polar ice sheets. Atmospheric gases circulate through the connected pores and are entrapped at the transition from firn to ice, where the network of open pores closes off to form individual gas inclusions (air bubbles (ABs)) (Schwander and Stauffer, 1984). With time, the ABs move deeper into the ice sheet and start shrinking as the hydrostatic pressure of the overlying ice increases. When the gas pressure inside the AB exceeds the dissociation pressure of hydrate formation the ABs begin to convert to air hydrates (AHs). AHs are crystalline structures composed of water molecules and containing cavities suitable for housing one or two gas molecules at a time. In contrast to the known phase diagrams (Miller, 1973), AB-AH conversion in polar ice takes place over a large depth range (transition zone) of several hundred meters (Uchida and others, 2014). Microstructure mapping images of the EPICA Dronning Maud Land (EDML) (Antarctica) ice core show first AHs at ~720 m depth; ABs were observed to coexist with AHs down to \sim 1225 m depth (Bendel and others, 2013). The widely accepted reason for the thickness of the transition zone is the slow nucleation of hydrates, after which the remaining transition proceeds comparatively quickly (e.g. Shoji and Langway, 1982). This is supported by the fact that no intermediate stages of the AB-AH transition have been found in ice cores (Narita and others, 1999; Lipenkov, 2000; Kipfstuhl and others, 2001).

The influence of the AB–AH conversion processes on the measured gas contents has been indicated previously: Ikeda and others (1999) and Ikeda-Fukazawa and others (2001) presented results of Raman measurements on individual air inclusions that clearly show gas fractionation in the transition zone, where initial AHs have lower N_2/O_2 ratios (i.e. they are oxygen-enriched) and the late ABs much higher N_2/O_2 ratios (i.e. nitrogen-enriched) than anticipated. Furthermore, anomalies in mixing ratios and isotopic composition of O_2 , N_2 and argon were reported by Bender and others (1995). Lüthi and others (2010) showed significant deviations of CO_2 and O_2/N_2 ratios in and just below

the transition zone, that are caused by diffusion and fractionation processes during the AB–AH conversion. Little is known about these processes, and while a reduction of resolution in the transition zone helps with measuring CO_2 , the processes involved have to be understood to ascertain the correct storage of CO_2 in the climate archive.

The AHs produced by a direct conversion from ABs are called primary air hydrates (pAHs) (Kipfstuhl and others, 2001). Two types of this transformation have been proposed by Salamatin and others (1999):

- *Type A:* The hydrate nucleates and rapidly forms a closed hydrate shell, covering the entire bubble wall. In a second, much slower, step the remaining bubble volume is spherically filled with hydrate, where water molecules diffusing through the closed shell towards the growing AH are the rate-limiting factor.
- *Type B:* The AH does not cover the bubble perimeter but proceeds as a flat crystallization front through the AB. Water diffusion through AH is not needed in this case, which indicates this process is much faster.

Additionally, Kuhs and others (2000) discussed the production of monocrystalline or polycrystalline AHs through a single nucleus or polynucleation respectively. At the beginning of the transition zone, the packing density inside an AH is much larger than inside a bubble. Therefore, if the AH growth uses the gas content of the transforming bubble, the resulting AH will have a significantly smaller volume than the originating bubble. Differences in bubble pressure and diffusion of gases from neighboring bubbles or the ice matrix to the newly forming AH can influence the actual volume mismatch between AH and the original bubble (Salamatin and others, 1999), complicating the deduction of bubble radii from AH sizes.

While most AHs appear smooth-surfaced and very clear (smooth air hydrates, sAHs), AHs with visible structure have been found in all ice cores and have been described as 'cloudy', 'irregular' or 'graupel-like' (Shoji and Langway, 1982; Nakahara and others, 1988; Uchida and others, 1994a; Narita and others, 1999; Pauer and others, 1999, 2000; Lipenkov, 2000; Kipfstuhl and others, 2001). An example from the NorthGRIP ice core is shown in Figure 1a. Carefully chosen illumination conditions reveal threedimensional (3-D) structures inside the pAH (Fig. 1b). The genesis of these structures, which have up to now only been accessible to visual inspection, is still unclear. Salamatin and others (1999) suggested that clusters of structured AHs could represent an AH shell fragmented by hydrostatic



Fig. 1. AH from the NorthGRIP core: (a) transmitted light; (b) asymmetric illumination reveals 3-D structure.

pressure during the type A AH growth. The amount of structural data retrievable by microscopic methods is very limited, as no information about the actual 3-D structure can be obtained. While timescales of several thousand years are not adequately reproducible in the laboratory, numerous researchers have performed laboratory experiments on bubble–hydrate conversions (e.g. Uchida and others, 1992, 1994a; Ikeda and others, 1993; Shimada and Hondoh, 2004) and also Raman spectroscopy on synthetic air hydrates (Chazallon and others, 1998). We present for the first time measured data of the intricate 3-D structure of natural air hydrates.

METHODS

Samples

We used samples from the EDML ice core from 1005 m, 1048 m, 1083 m and 1084 m (all depths are inside the transition zone). The EDML core was drilled between 2001 and 2006 at Kohnen Station, Dronning Maud Land, East Antarctica (75°00' S, 0°04' E; 2882 m a.s.l.) within the framework of the European Project for Ice Coring in Antarctica (Oerter and others, 2004). The samples studied were drilled in the 2002/03 season and have been stored at -30° C ever since.

The samples were cut, microtomed and sublimated to produce clean surfaces (see Weikusat and others, 2012, for details). Overview images taken with a $10 \times$ lens and the automatic image stitching function of the WITec control software were used to locate suitable AHs for measurements.

Raman tomography

The novel Alfred Wegener Institute cryo-Raman system is equipped with fibre optics that allow for a split set-up where the microscope unit is in the cold laboratory and the controller and spectrometer are at room temperature. This set-up eliminates the need for sample transfer between the cold laboratory and a cooling cell mounted to the microscope. As such transfers inevitably expose the samples to large temperature gradients and moisture from the air they have been the most critical point in cryo-analysis. The cryo-Raman set-up allows for sample preparation in the same laboratory where the microscope unit is situated, and transfer to the system is virtually riskless. The set-up also provides much more stable surfaces and reduced sublimation conditions, allowing for long-time measurements of up to several days. The system consists of a WITec alpha 300 M+, UHTS 300 spectrometer using a grating with 600 grooves mm⁻¹, Peltier-cooled CCD detector, frequency-doubled Nd:YAG laser (532 nm), and Zeiss Epiplan 50× LWD objective (NA 0.35). The laser spot size for this set-up is ~1 μ m, which defines the spatial resolution limit of the measurements. The confocal hole is provided by the diameter of the light-transmitting fibre from microscope to spectrometer. We used a 100 μ m fibre which gives a good compromise between confocality and signal intensity. The optical fibre transmitting the laser to the microscope is polarization-maintaining, while the fibre to the spectrometer depolarizes the light for a better detection efficiency. The temperature in the cold laboratory, and thus the temperature of the sample, were held at -15°C for all measurements.

Hyperspectral large-area stack scans map a rectangular area of the sample at a given step size, where a full spectrum is acquired for each pixel. The microscope turret is then moved a predefined step downwards and another scan with the same parameters is acquired for the new depth. One image is generated from each scan in the stack. These images can then be used to create a 3-D reconstruction of the measured AH.

Data analysis

The Raman spectra of ice and air hydrate are very similar, both being crystalline structures made from H₂O molecules (Schicks and others, 2005). Additionally, due to the choice of the 100 µm optical fibre and the implied reduced confocality, there is always a small contribution from the ice matrix in every recorded spectrum. The differences used here to distinguish between ice- and air-hydrate phase are shown in Figure 2. The low-energy mode of ice at \sim 216 cm⁻¹ is shifted \sim 3 cm⁻¹ to lower values (\sim 213 cm⁻¹) in the hydrate while the band broadens considerably (FWHM (full width at half-maximum) in ice: $\sim 27 \text{ cm}^{-1}$; FWHM in AH: \sim 37 cm⁻¹). The shoulder at \sim 290 cm⁻¹ in the ice spectrum is absent from the AH spectrum. A broadband weighted center-of-mass filter from 180 to 340 cm⁻¹ was used to extract the phase boundaries between ice and air hydrate (WITec Project software). To obtain binary images (Fig. 3), pixels with a value $>250 \text{ cm}^{-1}$ were defined as hydrate, and pixels with lower values as ice. It is important to note that other parameters (e.g. the height of the O_2 and N_2 bands $({\sim}1552\,cm^{-1}$ and ${\sim}2326\,cm^{-1})$ or shifts in the OH region (\sim 3000–3600 cm⁻¹)) yield the same



Fig. 2. Differences in the Raman spectra of ice and air hydrate that were used for imaging (averages of 100 spectra with accumulation time of 0.3 s per spectrum). a.u.: arbitrary units.

results qualitatively, albeit with worse signal/noise ratio. The resulting maps were exported as binary images, with black representing AH and white, ice. The obtained slices were combined to produce a 3-D reconstruction of the measured AH. Automatic segmentation of the slices and initial mesh generation were performed with ImageJ (Abràmoff and others, 2004). Smoothing of the initial mesh and calculation of geometric parameters were done with Meshlab (Visual Computing Lab ISTI – CNR, 2013).

Background subtraction and fitting of Voigt functions to the Raman bands were done with the open-source software package FITYK 1.2.1 (Wojdyr, 2010). The intensities of the main vibrational bands of N₂ (~2326 cm⁻¹) and O₂ (~1552 cm⁻¹) were used to calculate N₂/O₂ ratios. To calibrate the instrument response, several measurements of room air were averaged, yielding a raw N₂/O₂ ratio of 2.97 \pm 0.04 (mean value and standard deviation of all air measurements). Assuming the laboratory air composition to be N₂/O₂ = 3.73, every calculated ratio was multiplied by a correction factor of 1.26.

N_2/O_2 ratios and crystal orientation

It has been shown that the intensity of the Raman bands is a function of the orientation of the AH crystal to the polarization direction of the incident laser (Ikeda and others, 1997). Ikeda and co-workers found values between 8.06 and 10.33 for the N₂/O₂ ratios measured at different laser polarization angles for measurements on a well-defined crystal plane of a monocrystalline AH from the Vostok (East Antarctica) ice core. However, the peaks for N₂ and O₂ scale homogeneously for most polarization angles (which is reasonable, as they inhabit the same cavities). Additionally, the laser used for this study was only ~90% polarized and we measured on an arbitrary plane of hydrates, so we assume that the effect is much less pronounced. To be on the safe side we should assume an additional margin of uncertainty of ~±1.1.



Fig. 3. Binary slices generated from the Raman stack scan of pAH-1 (black: air hydrate; white: ice). The relative depth is given for each image; the scale is given on the photograph (lower right). See Figure 4 for the corresponding 3-D reconstruction.



Fig. 4. pAH-1 (depth 1005 m). (a) Microscopic image (scale bar for all images). (b) 3-D reconstruction viewed in the same orientation (steps are an artifact of large z-step size). (c) Side view; large holes in the shell are evident. (d) Semi-transparent side view (the holes are to the left) with visible ice intrusion.

RESULTS

Structure and volumes

Figure 4a presents a microscopic image of a structured pAH (pAH-1) from 1005 m depth. It is almost perfectly round, with a diameter of \sim 130 µm. The typical 'cloudy' appearance and rough surface are directly visible. With a step size of $3 \mu m$ in the x-y plane and a z-step size of $10 \mu m$, the whole stack scan is comprised of 48355 full spectra. An acquisition time of 0.2 s per spectrum was used, amounting to ~3 hours total measurement time. The 3-D reconstruction is shown in Figure 4b–d. pAH-1 is not solid but incorporates a large ice intrusion that is connected to the surrounding matrix ice by two holes in the AH shell (Fig. 4c). The ice volume is asymmetrically distributed inside the AH (Fig. 4d). The volume of pAH-1 is $9.89 \times 10^5 \,\mu\text{m}^3$, and the volume of the ice intrusions is $2.12 \times 10^5 \,\mu\text{m}^3$ (Fig. 4d), giving an ice: AH ratio of 0.2. The combined area of the holes in the AH shell is $\sim 1687 \,\mu m^2$.

Figure 5a shows a microscopic image of a structured pAH (pAH-2) from 1048 m depth, exhibiting similar features to pAH-1 (Fig. 4a). The mapped section of the AH is indicated. The diameter of pAH-2 is ~134 µm. A volume of 55 µm × 165 µm × 150 µm was scanned with an *x-y* step size of 2 µm and a *z*-step size of 10 µm, totaling 34 031 full spectra. With an acquisition time of 0.2 s per spectrum the whole stack took ~2 hours to complete. Compared to pAH-1, the ice intrusion inside pAH-2 has a more complex structure. Holes in the AH shell that connect the ice intrusion to the matrix can be seen in Figure 5b and c. The AH volume in this section is $3.45 \times 10^5 \,\mu\text{m}^3$, and the ice intrusion volume is $1.07 \times 10^5 \,\mu\text{m}^3$, giving an ice:AH ratio of 0.31.

Figure 6a shows two connected AHs from 1084 m depth: a very clear oval sAH with smooth boundaries (sAH-1) above an irregular and structured pAH (pAH-3). Several solid inclusions (black dots) can be seen inside sAH-1. The longest diameter of sAH-1 is ~95 µm, and the diameter of pAH-3 is \sim 120 µm. The scan was done with a step size of 3 µm for all three directions, amounting to 212 333 full spectra. With an acquisition time of 0.3 s per spectrum, the total measurement time for the stack was ~ 18 hours. The 3-D reconstruction in Figure 6b exhibits a continouus ice intrusion through pAH-3 (the top part of sAH-1 was outside the scan range). Figure 6c and d clearly show that both AHs are connected. Figure 6d visualizes the dimensions of the ice intrusion through pAH-3. The volume of the ice intrusion is $9.17 \times 10^4 \,\mu\text{m}^3$. The volume of pAH-3 could not be calculated reliably due to the connection of both AHs. The volume of the measured connected structure is $1.59\times 10^6\,\mu\text{m}^3.$ As the top part of sAH-1 was outside the scan range, this calculation only yields a lower limit for the whole structure.

Figure 7a shows a smooth AH from 1084 m depth (sAH-2). The scan has a step width of 4 μ m for all directions and consists of 24000 full spectra. With an acquisition time of 0.3 s per spectrum the measurement time was ~2 hours. The 3-D reconstructions (Fig. 7b–d) show a slightly flattened AH with smooth outlines and no internal structuration. The volume of sAH-2 is 4.11 × 10⁶ μ m³.

Figure 8a shows a faceted sAH from 1084 m depth (sAH-3). The scan has a step size of 3 μ m for all directions and the stack consists of 23 466 full spectra. With an acquisition time of 0.3 s per spectrum the measurement time was ~2 hours. Multiple twinned octahedra are visible in the 3-D



Fig. 5. pAH-2 (depth 1048 m). (a) Microscopic image with highlighted scan area (scale bar for all images). (b) 3-D reconstruction viewed in the same orientation (steps are an artifact of large *z*-step size). (c) Side view of the section with evident ice inclusion. (d) Side view from the opposite side.



Fig. 6. Connected AHs pAH-3 and sAH-1 (depth 1084 m). (a) Microscopic images with focus on sAH-1 (left) and pAH-3 (right). (b) 3-D reconstruction viewed in the same orientation (top part of sAH-1 out of scan range; scale bar for (b–d)). Note the hole through pAH-3. (c) Side view reveals a connection of both AHs. (d) Semi-transparent side view with visible ice intrusion.



Fig. 7. sAH-2 (depth 1084 m). (a) Microscopic image. (b) 3-D reconstruction viewed in the same orientation (scale bar for (b–d)). (c) View perpendicular to (b). (d) View perpendicular to (c).

reconstruction (Fig. 8b–d). The volume of sAH-3 is $1.12 \times 10^5 \,\mu\text{m}^3$.

Figure 9a shows a plate-like sAH from 1083 m depth (sAH-4). The scan step size was 3 μ m for all directions. With an acquisition time of 0.3 s per spectrum and a total of 341 185 spectra the measurement time was ~28 hours. The dimensions of sAH-4 are 245 μ m × 125 μ m × 20 μ m; the volume is 6.5 × 10⁵ μ m³. The lower left part of sAH-4 that appears to be slightly tilted in Figure 9a is only partially connected to the rest of the AH (Fig. 9c).

N_2/O_2 ratios

To determine N₂/O₂ ratios, several spectra were acquired on different points for each AH and the resulting ratios were averaged. Table 1 shows the results for all presented AHs. The ratio for pAH-1 is 3.60 ± 0.07 , which is close to the atmospheric value of 3.7. pAH-2, pAH-3 and sAH-1 are distinctly enriched in nitrogen with ratios of 5.51 ± 0.15 , 4.84 ± 0.10 and 4.91 ± 0.15 , respectively. The connected AHs, pAH-3 and sAH-1, exhibit very similar ratios. The solid

and smooth sAH-2 has a distinctly higher ratio of 8.30 ± 0.20 ; the plate-like sAH-4 exhibits a similarly high nitrogen content, with an N₂/O₂ ratio of 8.09 ± 0.20 . The highest relative nitrogen content was found in the faceted sAH-3 with an N₂/O₂ ratio of 17.49 ± 0.70 .

The homogeneity of the mixing ratios inside an AH was checked for pAH-1 by averaging all spectra for each slice and determining the N_2/O_2 ratios from the averaged spectra. The values in Table 2 are rather homogeneous and show no systematic alterations with relative depth in the AH.

DISCUSSION

Structure and formation processes

The studied pAHs with complex internal structures and incorporated ice bodies are very round and seem to have clearly retained the shape of the originating bubble. A destructive collapse of the AH shell during growth is certainly more likely to form either irregular agglomerations of splinter-like pieces (as observed by, e.g., Salamatin and



Fig. 8. Faceted AH (sAH-3, depth 1084 m). (a) Microscopic image (scale bar for all images). (b) 3-D reconstruction viewed in the same orientation. (c) View perpendicular to (b). Several twinned octahedra are discernible. (d) View perpendicular to (c).



Fig. 9. Plate-like sAH-4 (depth 1083 m). (a) Microscopic image (scale bar for all images). (b) 3-D reconstruction viewed in the same orientation. (c) View perpendicular to (b).

others, 1999), or solid, volume-optimized AH bodies (and eventually sAHs), if the process is combined with subsequent recrystallization. We therefore suggest that the intrusion of ice into the former AB volume is a syngenetic and non-destructive process. Salamatin and others (1998) found that the rate-limiting step of post-nucleation AH conversion is the plastic deformation of the ice-AH volume around the bubble. They also deduced that AHs are harder than the surrounding ice. This was later confirmed by Durham and co-workers in a series of ice-hydrate rheology experiments, where both type I and type II hydrates with various guest molecules proved to be up to 20 times harder than ice under identical strain conditions (Durham and others, 2000, 2003, 2005, 2010). We propose the following path for the AB-AH transition yielding structured pAHs: The AH nuclei start growing into the bubble, without forming a closed hydrate shell around the whole bubble. With proceeding AH growth the pressure inside the remnant bubble decreases, causing the bubble volume to shrink and the surrounding ice to flow simultaneously into the former bubble volume. In this scenario the diffusion of water molecules is not inhibited by a closed hydrate shell and the whole growth process can happen very fast, similar to the Btype formation described by Salamatin and others (1999), yet the B-type transformation maintains a flattened interface between growing AH and gas phase throughout the process and should therefore result in solid AH bodies without internal structures. As AH growth can only continue as long as the bubble pressure is above the dissociation pressure of the AH, the viscosity of the surrounding ice is an important factor for the growth velocity. (If we assume a spherical geometry for the ice intrusion in pAH-1, we can gain a rough estimate for the timescales needed to fill this volume with ice: as a first approximation, we set the driving stress to the hydrostatic pressure, P = 9 MPa. The strain rate for single

Table 1. Averaged N_2/O_2 ratios and standard deviations

Name	Depth m	Fig.	N_2/O_2 ratio
pAH-1	1005	4	3.60 ± 0.07
pAH-2	1048	5	5.51 ± 0.15
pAH-3	1084	6	4.84 ± 0.10
sAH-1	1084	6	4.91 ± 0.15
sAH-2	1084	7	8.30 ± 0.20
sAH-3	1084	8	17.49 ± 0.70
sAH-4	1083	9	8.09 ± 0.20

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crystal ice can then be extracted from figure 2 of Duval and others (1983). As the orientation of the crystallite is unknown, we chose a medium rate of $\dot{\varepsilon} = 10^{-5} \text{ s}^{-1}$. To obtain the velocity, we integrated the strain rate, considering only the *x* direction for further simplification. With $x = 110 \,\mu\text{m}$, ice intrusion volume $V = 2.12 \times 10^5 \,\mu\text{m}^3$ and opening area of the hydrate shell $A = 1687 \,\mu\text{m}^2$, a time of t = 3.2 hours can be readily calculated. While this may give a lower boundary for the formation of pAH-1, it is by no means intended to represent a mathematical treatment of the microstructural flow of ice, which is well outside the scope of this publication.)

After all gas has been used up for the AH, an ice intrusion remains inside the AH. This process has been witnessed in laboratory experiments, even though the intrusion of ice into the forming hydrate was not recognized at that time (Uchida and others, 1994b; Shimada and Hondoh, 2004). Driven by a thermodynamic urge to reduce the surface energy, the AH can then recrystallize to form a smooth AH without internal structuration. The connected AHs seen in Figure 6 are an example of this process, where the pAH volume is diffusively redistributed in favour of a smooth and compact sAH body. This is further supported by the fact that the N₂/O₂ ratios of pAH-3 and sAH-1 are identical in the given error range. The impurities visible in sAH-1 could have acted as nucleation points for the sAH (Ohno and others, 2010). Microscopic images of more AHs undergoing such a metamorphism are provided by Pauer and others (2000). It is

Table 2. N_2/O_2 ratios from averaged spectra for each slice of pAH-1 (cf. Fig. 3)

Rel. depth	N_2/O_2 ratio	
μm		
0	3.54	
10	3.63	
20	3.57	
30	3.64	
40	3.69	
50	3.65	
60	3.64	
70	3.54	
80	3.48	
90	3.53	
100	3.69	
Average	$\textbf{3.60}\pm\textbf{0.07}$	

important to keep in mind that this is not the only process capable of forming smooth AHs; a direct conversion from AB to sAH has been demonstrated in laboratory experiments (Ikeda and others, 1993).

However, the shapes of sAH-3 and sAH-4 are evidently not a product of the above-mentioned processes. sAH-3 exhibits several twinned octahedra that clearly follow the cubic crystal structure of the AH, while sAH-4 closely resembles the so-called plate-like inclusions (PLIs), relaxation features that are known to form on the basal planes of ice crystals, following the hexagonal structure of ice Ih (Nedelcu and others, 2009).

The structured pAHs with ice intrusions are more susceptible to mechanical destruction than solid AHs. The frequently seen clusters of small AHs, described by Salamatin and others (1999) as a product of a collapsing AH shell during transition, could also have been formed by post-genetic separation due to ice deformation.

N_2/O_2 ratios

With the exception of pAH-1, which has a ratio close to the atmospheric value, all AHs measured for this study are enriched in nitrogen. This is contrary to other works that found the AHs in the transition zone of Dome Fuji, Antarctica, to be oxygen-enriched (Ikeda and others, 1999; Ikeda-Fukazawa and others, 2001). However, it has also been shown that ABs in the transition zone are enriched in nitrogen: Weikusat and others (2012) reported N₂/O₂ values up to 6 for EDML while Ikeda-Fukazawa and others (2001) found extreme values up to 14 in the Vostok core. As all ABs will have transformed by the end of the transition zone, it seems likely that the N_2/O_2 ratios measured in the AHs are due to the conversion from already nitrogen-enriched ABs. Exceptionally high N₂/O₂ ratios as measured in sAH-2 (8.30), sAH-4 (8.09) and sAH-3 (17.49) could also indicate a different path of hydrate formation, not necessarily implying equally nitrogen-enriched ABs. For sAH-3 and sAH-4, this is supported by the unusual morphologies, which are clearly not derived from a common air bubble. It is also worth noting that the solid AHs without ice intrusions investigated in this study show higher N2/O2 ratios than the cloudy pAHs, yet more data are needed for a statistical evaluation. Selective gas losses during the long storage time could also have altered the measured mixing ratios. A detailed interpretation of these unusually high values is outside the scope of this work and will be addressed in a dedicated publication.

CONCLUSION

The structured pAHs are a result of a fast conversion process, where the AH shell is not closed at any time and thus a fast conversion is possible during the whole process. Ice from the surrounding matrix flows to intrude the former AB volume, keeping the pressure in the remnant bubble above the AH dissociation pressure. The flow velocity is then an important factor for the conversion. On the microscale, the flow can be very fast (hours) and would therefore be on the same timescales as the formation processes seen in laboratory experiments (Uchida and others, 1992, 1994a; Ikeda and others, 1993; Shimada and Hondoh, 2004). The constant contact to the ice matrix also eliminates the need for (slow) diffusion of water molecules through the AH layer. Detailed investigations and modelling of the processes with a suitable tool (e.g. ELLE; Bons and others, 2008) could yield information about deformation and strain distribution on the microscale.

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