

# Distribution of brine in grain boundaries during static recrystallization in wet, synthetic halite: insight from broad ion beam sectioning and SEM observation at cryogenic temperature

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Received: 7 January 2011 / Accepted: 24 May 2011  
Springer-Verlag 2011

**Abstract** We report observations from room temperature three major processes. First, dissolution on one side of the static recrystallization experiments (annealing times from minutes to year) of cold-pressed, synthetic, coarse-grained wet sodium chloride, prepared by broad ion beam polishing and SEM observations at cryogenic temperature to observe the grains bounding the grain boundary brine. When both sides of a grain boundary are able to develop low-index facets in a thick brine film, the resulting impingement connected in 2D sections along grain boundaries. Another boundary is interpreted to be immobile and may prevent part of the brine is in isolated brine inclusion arrays along the new grain from migrating into a deformed neighbor. When one side of a faceted boundary consists of low-index migrating brine-filled grain boundaries. Most of these crystallographic planes and the other side passively follows boundaries are mobile because the aggregate is coarsening into its faceted shape along irrational surfaces, the boundary is immobile. We interpret that the boundaries without observable brine films, films (15 nm) and brine inclusion arrays are healed and producing solid-solid grain boundaries without resolvable immobile. Evolution of grain boundary structure involves brine films.

Communicated by J. Hoefs.

Electronic supplementary material The online version of this article (doi:10.1007/s00410-011-0656-x) contains supplementary material, which is available to authorized users.

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**Keywords** Static recrystallization · Halite · BIB-cryo-SEM · Brine-filled GB · In situ GB microstructures

## Introduction

The presence of brine in halite strongly affects its mechanical and transport properties (Urai et al. 2008). Solution-transfer creep and dynamic recrystallization are dramatically enhanced by the presence of brine at grain boundary, with major changes in rheology. In addition, rock salt forms seals for hydrocarbon accumulations, aquitards, and chemical barriers. Therefore, a detailed knowledge of grain boundary microstructures in wet halite is fundamental for modelling and prediction of deformation and transport in a wide range of geological environments (Hudec and Jackson 2007; Schoenherr et al. 2007; Desbois et al. 2010) as well as for the storage of gas, anthropogenic carbon dioxide, and radioactive waste (Pinckney 1985).

Staudtmeister and Rokahl 1997, Langer 1993, Brest et al. 2010) in deep geologic formations.

In evaporite minerals, the structure of mobile brine-filled grain boundaries and their influence on recrystallization processes were intensively investigated in theoretical and experimental studies (Urai et al. 1986, Brantley et al. 1990, Drury and Urai 1990, Hickman and Evans 1991, 1995, Gratier 1993, Heidug and Leroy 1994, Lehner 1995, Holness and Lewis 1997, Spiers and Schutjens 1999, Visser 1999, Ghoussoub and Leroi 2001, De Meer et al. 2002, 2005, Den Brok et al. 2002, Watanabe and Peach 2002, Lohkaemper et al. 2003, Renard et al. 2004, Schenk and Urai 2004, 2005, Van Noort et al. 2006, 2008).

However, interpretation of results from these studies is still a matter of debate because of the difficulty to observe the morphology of in situ wet grain boundaries at a resolution of nanometers (De Meer et al. 2005, Desbois et al. 2008). One recently explored method is to rapidly freeze the samples to very low temperatures, which effectively quenches the brine-filled grain boundaries without crystallization of the brine, followed by high-resolution elec-

tron microscopy at cryostatic temperatures (Schenk et al. 2006, Holzer et al. 2007, 2010, Desbois et al. 2008, 2009). Cryo-SEM can be combined with a Focussed Ion Beam (FIB) milling to prepare smooth, damage-free surfaces (Holzer et al. 2007, 2010, Desbois et al. 2008, 2009, Holzer and Cantoni 2011). Commercial FIB tools in SEM are based on Gallium ion sources (1 pA to 50 nA) to produce typical cross-sections of a few  $\mu\text{m}^2$ , with serial sectioning for the investigation of microstructures in 3D (Holzer et al. 2007, 2010, Desbois et al. 2008, De Winter et al. 2009).

Broad ion beam (BIB; up to few mA, argon source) cross-sectioners are commercially available as stand-alone machines, to produce polished cross-sections at room temperature of much larger area than the FIB. Because the

BIB source is not focused, a shielding plate is placed on the top surface of the sample in order to create a flat cross-section. BIB has two main advantages: (1) it is potentially less damaging since it is based on noble gas source and it is able to produce polished surfaces up to few  $\text{mm}^2$  about hundred times bigger than those produced by FIB. This last feature is of particular interest for geomaterials since it is better to the typical length scale range of microstructures and representative elementary area.

Schenk and Urai (2004), studied wet, coarse-grained (200–355  $\mu\text{m}$ ), compacted samples of synthetic sodium chloride which undergo recrystallization and grain growth at room temperature to produce large (up to 300  $\mu\text{m}$ ) strain-free grains growing into deformed old grains. The new grains are frequently euhedral with mobile grain boundaries moving at time-averaged velocities up to 6  $\mu\text{m}/\text{s}$ . Arguments in favor of the existence of brine-enriched zones on grain boundaries were the following: (1) the

efflorescence from such boundaries upon drying, (2) euhedral shape of the growing grains, and (3) the high migration rates. However, details of the brine distribution and its influence on grain boundary migration were only partly resolved, because brine was removed during sample preparation. In a later study, Schenk et al. (2006) studied brine-filled grain boundaries in similar coarse-grained samples by cryo-SEM on freeze-fractured surfaces. They found evidence for brine at the boundaries of the primary recrystallizing grains indicating the existence of brine films with a thickness between a few nm and 30 nm. In this last study, the observation of brine-filled grain boundaries was difficult because of the surface roughness induced by

fracturing the sample during preparation. The process of brine-assisted grain boundary migration was interpreted to occur simultaneously with surface energy-driven grain boundary healing, which dramatically reduces grain boundary mobility (Hickman and Evans 1991, Lehner 1995, Visser 1999, Ghoussoub and Leroi 2001, Schenk and Urai 2004, 2005, Schleder and Urai 2005, Van Noort et al. 2008).

This contribution reports of experiments that were identical to Schenk and Urai (2004) and Schenk et al. (2006) to study the distribution of brine in moving grain boundaries during static recrystallization in synthetic, coarse-grained wet sodium chloride, annealed at room temperature over periods between a few minutes and 1 year. Here, we used a novel and unique BIB-cryo-SEM instrument combining in situ broad ion beam (BIB) cross-sectioning and state-of-the-art SEM cryo-SEM to study brine into grain boundary.

The BIB-cryo-SEM instrument

The BIB-cryo-SEM instrument used in this contribution combines a state-of-the-art field emission SEM for high-resolution imaging (SE and BSE detectors, EDX system), with an argon broad ion beam excavation system and a cryogenic workstation (shock-freezing stage, cryo-preparation chamber with cold knife and coating system). A detailed description of all facilities included in this instrument as installed at RWTH Aachen University (Germany) is given in Online Resource 1.

After freezing the samples and inserting into the cryo-SEM, (Fig. 1), the shielding plate is positioned onto the sample surface in the region of interest by nanomotor positioning. The BIB is placed at 70  $\mu\text{m}$  from the electron beam. In condition of milling, the BIB, with beam diameter around 800  $\mu\text{m}$  in size, irradiates both the region of interest and the edge of the mask designed with a slope of 20. One half of the BIB is masked by the shielding plate, and the part of the beam, which is not

Table 1 Overview of samples investigated in this contribution

Sample	Number of samples	Starting grain size ( $\mu\text{m}$ )	Annealing time <sup>a</sup>
20 min	2	200 $\pm$ 355	18 min 15 min
2 h	2	200 $\pm$ 355	1 h, 54 min 1 h, 56 min
2 days	2	200 $\pm$ 355	2 days, 2 h, 7 min 1 day, 23 h, 46 min
1 week	2	200 $\pm$ 355	5 days, 22 h, 30 min 6 days, 23 h, 6 min
2 weeks	2	200 $\pm$ 355	14 days, 19 h, 10 min 14 days, 15 min
1 month	2	200 $\pm$ 355	24 days, 19 h, 20 min 30 days, 18 h, 8 min
1 year	3	200 $\pm$ 355	1 year <sup>b</sup> 1 year, 2 months

Fig. 1 Principle of in situ BIB milling into the SEM. The original wet sample is first vitrified in slushy nitrogen using the cryo-unit and then transferred inside the SEM by the cryo-preparation chamber. Once the sample is placed on the SEM sample stage, a titanium blade is positioned over the region of interest using a 3 axis Cartesian nanomotor. The argon ion beam irradiates both the region of interest in the sample and the edge of the mask designed with a slope of 20°. Beneath the edge of blade, the density of ions is maximal and the sputter beam is parallel to the blade edge. This configuration allows creating a vertical 2D cross-section directly beneath the titanium blade suitable for imaging microstructures at SEM resolution (beam)

masked, excavates the sample producing a cross-section. After milling, the cross-section is suitable for SEM investigations.

### Samples and sample preparation

Samples are compacted synthetic polycrystals from commercial halite powder (NaCl content 99.9%) prepared as in Schenk and Urai (2004). After treating for 30 s in an ultrasonic bath, the initial salt/brine mixture (grain size in the range of 200 $\pm$ 355  $\mu\text{m}$ ) was poured into the cylindrical die of a uniaxial compaction apparatus. Subsequently, the pressure on the upper piston was raised to 150 MPa in a few minutes and maintained for 5 min. Resulting samples are dense cylindrical disks with a diameter of 1 cm and a height of 1 $\pm$ 1.5 mm. Produced samples were then annealed (from about 20 min. to more than 1 year, see Table 1) in salt/brine-saturated atmosphere in order to allow static recrystallization of samples (Schenk and Urai 2004).

BIB-cryo-SEM experiments were performed on 15 different samples (Table 1) following the same procedure of sample preparation and imaging: (a) the in situ wet sample is mounted on the appropriate sample holder at room temperature; (b) plunged frozen in slushy nitrogen, stirring to minimize the Leidenfrost effect (Desbois et al. 2008); (c) via the cryo shuttle operating at cryogenic temperatures and vacuum conditions, the sample is transferred from the

The annealing was carried out at room temperature slushy nitrogen container to the cryo-preparation chamber; (d) the surface of sample is coated with platinum (15 A, 30 s.) at cryo-temperature; (e) transfer from the cryo-preparation chamber to the SEM sample stage maintained at cryo-temperature via the same cryo shuttle previously mentioned; (f) the cross-section is then cut at cryo-temperature, on frozen/stabilized samples using the BIB gun for 3 h at 8 kV acceleration voltage; (g) when cross-section is complete, SEM imaging is typically performed at 3 kV for a working distance of around 16 mm and aperture of 30  $\mu\text{m}$ , on non-coated cross-section.

To reveal the crystal orientation patterns, the EBSD analysis (Electron Back Scattered Diffraction) on JEOL JSM 7000F SEM (Gemeinschaftslabor fuer Elektronenmikroskopie, RWTH Aachen) with EDAX-TSL DigiView III detector equipped with OIM Data Collection/OIM Analysis (version 5.2) software was employed. The patterns were acquired at acceleration voltage of 25 kV, probe current  $\sim$  15 nA, and working distance 29 mm. Several regions of interest from BIB section of a sample annealed for 4 days were scanned in an area of about 300  $\mu\text{m}$  at a magnification of 750 with a step size of 0.25  $\mu\text{m}$ . Because EBSD tool is not available in BIB-cryo-SEM machine, EBSD analysis was performed on sample cut at 80C for 3 days.

### Results

#### Cross-section overviews and surface quality

Figure 2 presents typical cross-sections obtained at 8 and 30 kV (BIB acceleration voltage) under cryogenic conditions. All samples were cut in a plane, which contains the

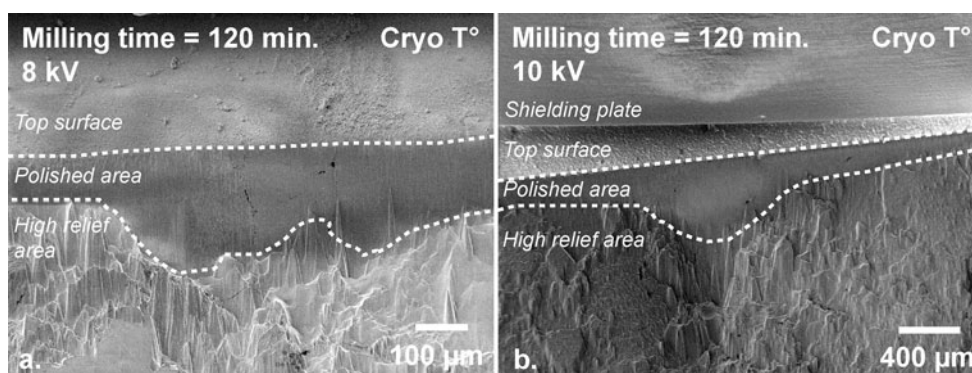


Fig. 2 Overview of typical cross-sections produced by in situ BIB pseudo-Gaussian shape reflecting the ion density in the beam. High milling (120 min) into the SEM at cryo-temperature at acceleration relief areas correspond to material regions being removed voltage of 8 kV (a) and 10 kV (b). Cross-sections have a typical

direction of compression. Resulting cross-sections have pseudo-Gaussian shape reflecting the ion density distribution in the beam split by the shielding plate and the typical high relief in the areas being removed. For NaCl, milling rates of about 350 μm/hr at 10 kV, 150 μm/hr at 8 kV, and 60 μm/hr at 6 kV are typical values.

The quality of cross-sections is reproducible. At the scale of overview (Fig. 2), cross-sections appear flat; at higher magnifications (Fig. 4, 5, 7 and Online Resource 2), surfaces show curtaining parallel to the beam direction with a regular wavelength in the range of 1 μm and a topography in the range of few nanometers. In flat section of the curtains, microstructures down to 15 nm

In previous contributions dealing with cryo-SEM on wet rock salt (Schenk et al 2006, Desbois et al 2008), in situ brines were sublimated to help with the detection of brine distribution. This technique is also called freeze-drying. Freeze-dried brine has a characteristic foam-like morphology (Schenk et al 2006, Desbois et al 2008), which can be used to decorate and thus detect brine. In this contribution, brine also develops a foam texture (Fig. 5, 7 and Online Resource 2).

#### Interface nomenclature

To facilitate the description of grain boundaries, we propose to introduce the following nomenclature (Fig. 3). The HH boundary is a boundary between two halite grains, with no visible structure at the scale of observation. This means that at low magnification, boundaries, which can be shown to contain a brine film, are called HH. The HBH boundary, where there is resolvable brine between two halite grains at the scale of observation. The HDB boundary is the contact between halite and brine. Therefore, a brine-filled boundary consists of two HB boundaries (HB BH=HBH) and B.1 in Online Resource 2).

Fig. 3 Interface nomenclature. The HBH boundary is a interface with resolvable brine between two halite grains, and HH boundary is a boundary between two halite grains, with no visible structure, both at the scale of observation. In case of HBH boundary, the shape of the boundaries can be faceted (straight, indicated by subscript f) or non-faceted (curved, subscript n)

The shape of the boundaries can be faceted (straight, indicated by subscript f) or non-faceted (curved, subscript n). Again, this is defined at the relevant scale of observation.

Thus, a boundary with resolvable brine, which is faceted on one side and curved on the other side of the brine film, is called a HBH<sub>n</sub> boundary. The same structure at much lower magnification is called a HH boundary (Fig. 4).

#### Grain microstructures

At the scale of a few micron, in the whole sample, (Fig. 5a, b, and B.1 in Online Resource 2), halite grains up to 0.1 mm in size with serrated HBH<sub>n</sub> boundaries containing up to 1 μm thick brine films are common. The shape of these grains varies from elongated to equiaxed (Fig. 5a, b

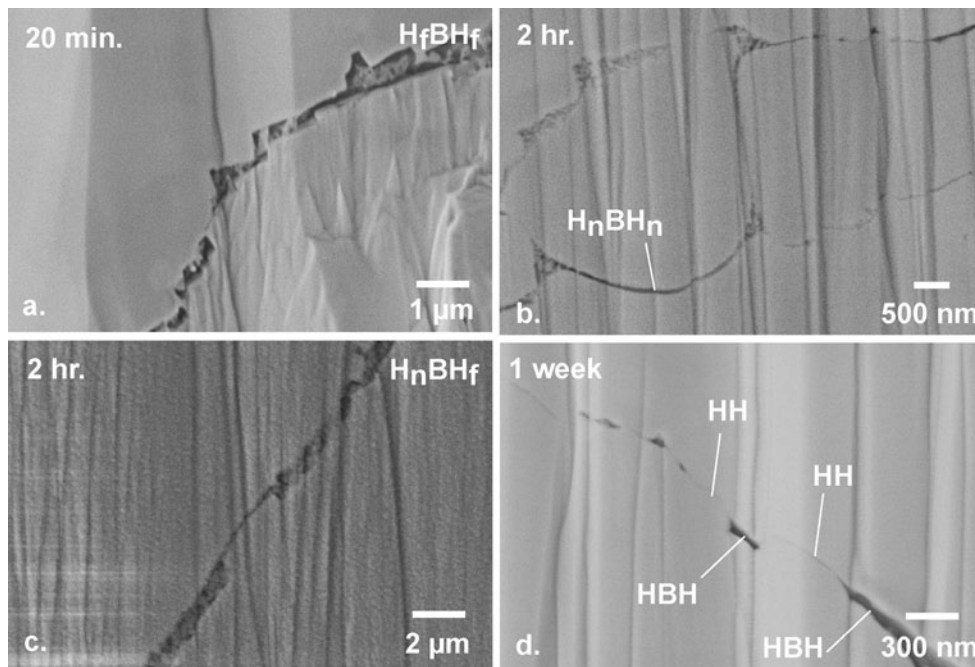


Fig. 4 SE micrographs of different halite-brine interfaces detected by magnification. Interfaces are described by the following abbreviations:  $H_fBH_f$ ,  $H_nBH_n$  boundary with low wavelength curved interfaces;  $H_nBH_f$ ,  $HBH$  interfaces.  $c$   $H_nBH_f$ ,  $d$   $HH$  and  $HBH$  boundaries at high magnification. Interfaces are described by the following abbreviations:  $H$  halite,  $B$  brine,  $n$  non-faceted,  $f$  faceted (see also Fig. 3)

Estimates of the total amount of brine in the samples using our images are in agreement with the porosities measured (2.5%) by Schenk and Ura (2004). Independently of annealing time, brine filled all space between halite grains (there was no evidence for gas inclusions in the brine).

For all annealing times, continuous brine-filled grain boundaries are frequent, with variable thickness from about 15 nm (limit of resolution) to more than 1 μm (Figs. 4, 5, 6 and B.2 in Online Resource 2) over long sections (up to few hundreds micrometers). The thickest brine-filled continuous grain boundaries (1 μm) bound the largest grains (Fig. 5c, d, and Online Resource 2). These small euhedral grains with  $H_nB$  contacts (Figs 5a, b, 7g and B.1 in Online Resource 2), while brine films with thicknesses below 1 μm are usually at interfaces involving at least one  $HBH$  contact (Fig. 5e, f, and B.3 in Online Resource 2). Exceptions are curved  $H_nBH_n$  boundaries with high wavelength (Fig. 4b and B.3.f in Online Resource 2) with typical brine film thickness 50 nm.

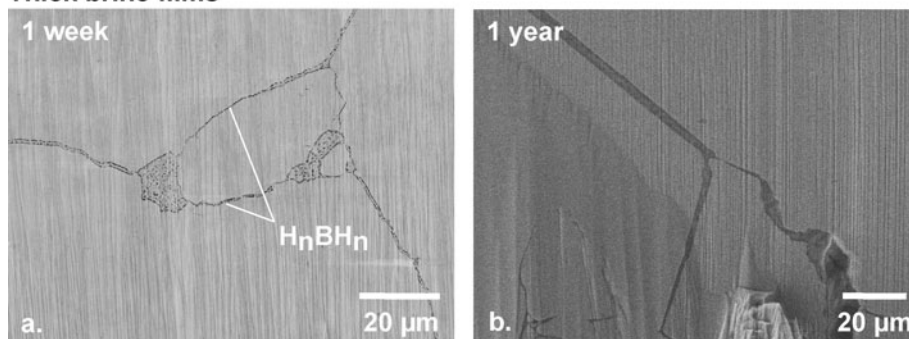
A second class of grains bounded by facets is defined by a larger grain size (2 μm) and more complex facet structures (Fig. 5e, f, and B.3 in Online Resource 2). These euhedral grains have multiple facets defining morphologies from flat (Fig. 5e and B.3.a, b, c, e, f in Online Resource 2) to round (Fig. 5f and B.3.a, c, f in Online Resource 2) and step-shaped (Figure B.3.d in Online Resource 2). Irregular brine pockets between grains are bound by  $HBH$  boundaries. The geometry of  $HBH$  boundary impingement controls the morphologies and area (up to  $1 \mu m^2$ ) of these irregular brine pockets (Fig. 5e, f, and B.3 in Online Resource 2). Adjacent irregular brine pockets can be connected by very small brine-filled throats down to 15 nm thick controlled by facets developed by the host grains (Fig. 5e, f and B.3 in Online Resource 2).

### Brine distribution

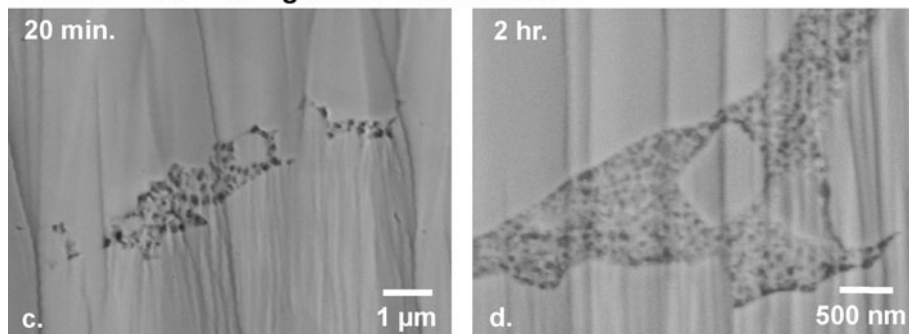
At higher magnification (below one micron, Fig. 7), brine-filled grain boundaries show strong variations in the morphology of the halite-brine contacts. We observed all possible types  $H_nBH_n$ ,  $H_fBH_f$ ,  $H_nBH_n$ , and  $HH$  (Fig. 4). In contrast to irregular brine pockets, there are intergranular brines inclusions (IBI, size smaller than 500 nm) and  $HHH$  grain boundaries, which are not defined by  $HBH$

Fig. 5 SE micrographs of grain boundaries interfaces taken at different annealing times and depict networks of thick brine films (1 μm). c and d show small euhedral halite grains which appear floating in brine-filled grain boundaries identified in Fig. 4. These small new grains are frequent for short annealing times and f show halite grains with multiple euhedral edges. These grains are slightly bigger than those presented in c and d and exhibit more complex face structures according to the Hartman-Perdok theory for cubic crystals. Interfaces are described by the following abbreviations: *H* halite, *B* brine, *n* non-faceted, *f* faceted (see also Fig. 4). For additional pictures, please check also the Online Resource 2

**Thick brine films**



**Small euhedral halite grains in brine-filled GB**



**Halite grains with multiple euhedral facets in brine-filled GB**

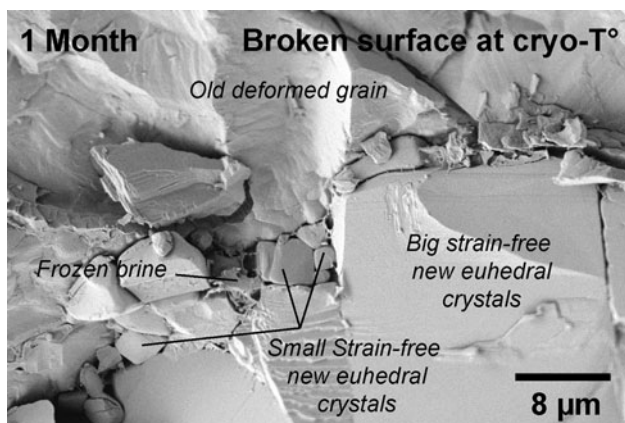
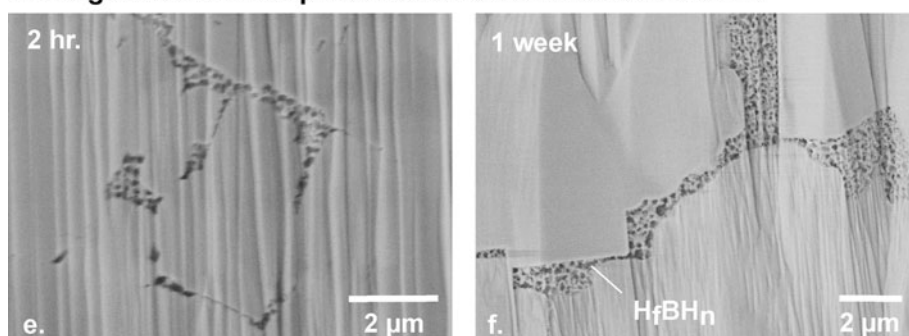


Fig. 6 SE micrograph of a freeze-fractured broken sample annealed for 1 month. This micrograph depicts a contact between a big strain-free new euhedral crystal growing toward a old deformed grain. Small strain-free euhedral crystals are also present into the grain boundary

boundaries (Fig. 7). These have three main different characteristic morphologies as follows: (i) "elongated-shaped" IBI have a very high aspect ratio with rounded (Fig. 7a) or sharp tips (Fig. 7b); (ii) "triangle-shaped" IBI have also a high aspect ratio (Fig. 7c, d); and (iii) sub-micron arrays of "quadrangle-shaped" IBI (Fig. 7e). Sometimes, a very thin line (20 nm thick) is resolvable connecting the arrays of IBI (Fig. 7a-f). Unfortunately, since the thickness of this very thin line is in the range of the SEM resolution, it is not clear whether this is brine-filled.

In addition to the first three types, there is an additional fourth type: the intragranular "quadrangle-shaped" brine inclusion (Fig. 7g). This last type differs from others because they are significantly bigger (1 μm in size), are not aligned with other similar IBI, and are never linked to visible "very thin line" as described previously.

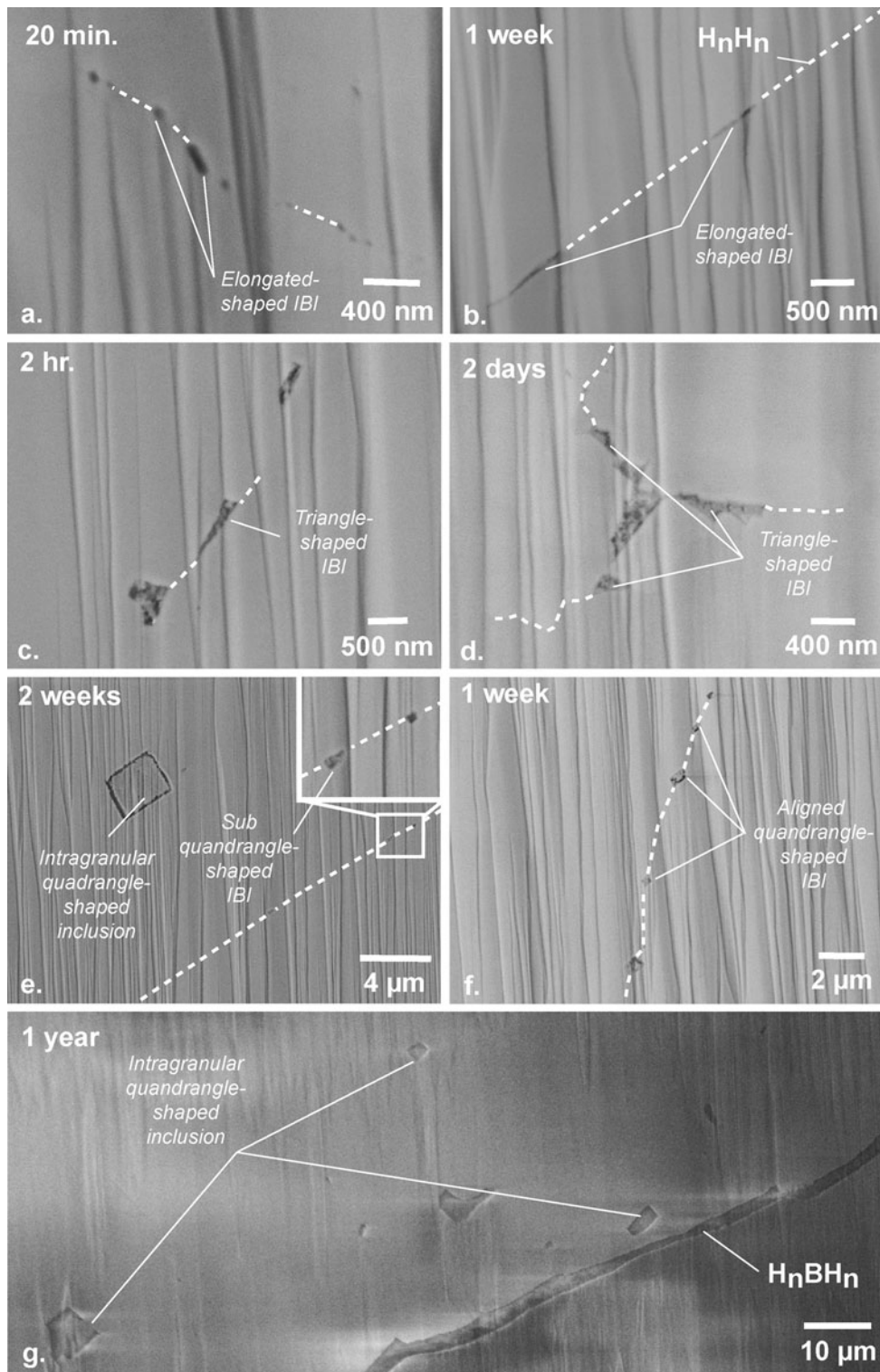


Fig. 7 SE micrographs of brine-filled isolated brines inclusions (IBI; grain boundary whose different morphologies are controlled by the  $\sim 500$  nm in size) with different characteristic morphologies and relative crystallographic orientation of grains at contact. The intragranular big quadrangle-shaped IBI (in size) are also identified (e and g) and are interpreted as IBI left over migrating to grain boundary. White-dashed lines indicate HDH interfaces. Interdevelop during grain boundary healing. All of these IBI are faces are described by the following abbreviations: a) non-faceted, b) faceted (see also Fig. 3)

Each of the four different IBI described above is presented since produced 2D cross-sections optimize the microstructural interpretations, which are difficult to reach when the time but intragranular quadrangle-shaped brine inclusions (topographies of surfaces are rough of few ten micrometers in size for the case of broken samples). In addition, and

## Discussion

### Quality of milled surfaces

In his current state, our BIB-cryo-SEM does not produce comparable quality surfaces to those of commercial stand-alone BIB cross-sectioners (Desbois et al. 2009, Loucks et al. 2009), since cross-sectioned surfaces presented hereon, it is obvious that the BIB locally heats the sample exhibit curtaining which can mask and distort the very small microstructures (15 nm). The curtaining effect is known to be due to heterogeneous milling rate, driven by heterogeneities in the material. In commercial stand-alone BIB cross-sectioners, this is minimized by rocking the sample during milling along the axis perpendicular to the plane to be polished and milling at different angles. Sample rocking during milling at room temperature was recently implemented in our system. Preliminary runs produced perfect cross-sectioned surfaces without curtaining (Fig. 8). In the near future, sample rocking will be also implemented under cryogenic conditions. However, considering the limitations just discussed, many of the relevant grain boundary structures could be resolved in the development stage of the machine when sample rocking was not available, either at a scale much larger than the characteristic curtain diameter or at a scale smaller than the characteristic curtain diameter. Moreover, even in its actual state, the BIB-cryo-SEM instrument provides a better insight into pores, interfaces and fluid-related microstructures than previous experiments performed on similar broken samples (Schenk et al. 2006).

Because samples were not freeze-dried during preparation during milling since frozen brine exhibits the characteristic foam pattern typical of freeze-drying (Schenk et al. 2006, Desbois et al. 2008). Freeze-dried brine during FIB milling was also observed in Desbois et al. (2008). In the future, heating of the frozen sample during BIB milling will be minimized, by lowering the current density, optimizing the distance between the shielding plate, and by cooling the mask. However, in case of wet rock salt, freeze-drying of brine is also a useful indicator for in situ brine (Schenk et al. 2006, Desbois et al. 2008).

Development of euhedral facets  
Along the original grain margins (up to few hundreds of micrometers in size), grains with multiple facets are present (Fig. 4a, c, 5e, f, 8, and B.3 in Online Resource 2). As discussed in Schenk and Urabe (2004), the origin of such faceting is due to the deformation of original halite powder in uniaxial compaction apparatus (see section 3.1.1 and sample preparation 1.00), which produced gradients of defect density and microcracks favorable to nucleation of

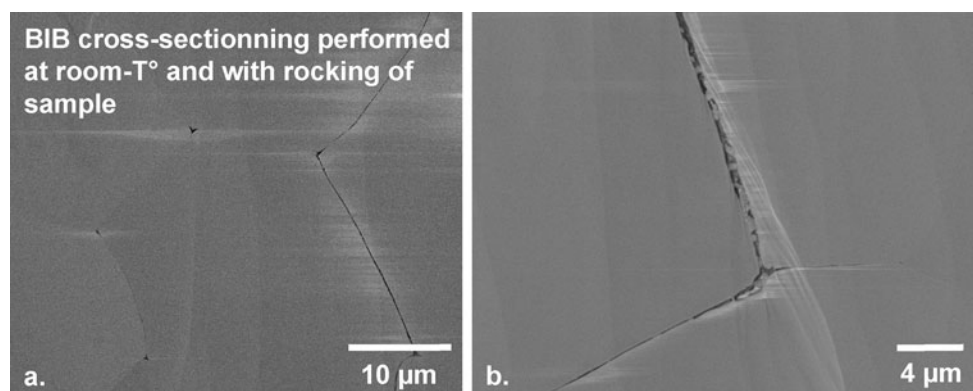


Fig. 8 SE micrographs of surfaces produced by in situ-SEM BIB boundary at euhedral grain edge is filled with halite recrystallized by cross-sectioning at room temperature with sample rotating (along axis perpendicular to the ion beam) during milling. This hardware vacuum. Surfaces are smooth and flat surface without critical improvement allows creating perfect surfaces without curtaining. curtaining  
a Grain boundary network exhibiting small triple junctions  
A grain



Fig. 9 a SEM (BSE) and b EBSD micrographs of the same local grain fabric in dried sample annealed for 4 days. Micrographs show faceted new crystals with low-index facets (euhedral growth) and free of lattice distortion. In contrast, in a.1 and a.2, the grain at right side is lattice distortion-rich pointing to old deformed grain. At the scale of observations, the numbers 1, 2, and 3 in circles indicate  $H_f H_{fr}$ ,  $H_n H_n$ , and  $H_f B H_f$  interfaces, respectively.  $H_f H_{fr}$  interfaces reflect the growing of new grains at the expense of old deformed grain forming an irrational grain boundary,  $H_n H_n$  interfaces are interpreted as healed contact between two new crystals, and  $H_f B H_f$  interfaces as euhedral facets growing into fluid-filled pore space. *White cube* exhibits three-dimensional representation of the crystallography of the respective grain showing the dominance of grain boundaries compatible with equivalent  $\{001\}$  facets. Interfaces are described by the following abbreviations: *H* halite, *B* brine, *n* non-faceted, *f* faceted, *i* irrational, *r* rational (see also Fig. 3 and section 4.1 on mobility of grain boundaries).

strain-free grains with  $H_f B H_f$  boundaries (Humphreys and Hatherly 1996). Such newly nucleated crystals initially develop crystal faces because they grow in a brine environment (as described by Hippert and Egydio-Silva 1996 in quartz mylonite). Due to differences in dislocation density, some old grains are interpreted to have a slightly higher solubility, providing the material for growth of the face can be identified by F, S, and K indicating a flat face ( $H_f B$  facets elsewhere. This may explain why some original grains do not exhibit faceting (Fig. 4c, 5a, b, 7g, and B.1 in Online Resource 2).

Because they are fully surrounded by brine, the small euhedral grains in the core and mantle structures (Fig. d, 6 and B.2 in Online Resource 2) are interpreted to have grown epitaxially from very fine grains stuck on starting material (and thus not removed by sieving) or from small grain fragments formed during cataclasis of the starting material. Euhedral new grains exhibit multiple facets (Fig. f and B.3 in Online Resource 2) whose structure was shown to correspond to the Hartman-Perdok theory for flat (F) faces (Hartman and Perdok 1955; Urai et al. 1986; Van Noort et al. 2006; Schenk et al. 2006). Considering three periodic bond chains (PBC), three different types of face can be identified by F, S, and K indicating a flat face (parallel to at least two PBCs), a stepped face (parallel to one PBC), and a kinked face (not parallel to any PBCs), respectively. Our conclusions are in agreement with EBSD-based analysis of euhedral crystals in these samples (Fig. 9 and e.g. Schenk et al. 2006). Evolution of brine-filled grain boundaries At all stages of annealing,  $H_f B H_f$  boundaries are systematically brine filled with no evidence of gas inclusions in

brine, confirming that samples were brine saturated throughout the experiment.

The majority of brine is connected in 2D sections along the grain boundaries from 15 nm up to few micrometers thick (Figs. 4, 5a, b and B.1 in Online Resource 2). This is consistent with earlier observations in experiments (Schenk et al. 2005; Van Noort et al. 2006) based on studies of post-mortem roughness of pressure dissolved surfaces (grain boundaries in natural domal salt continuous brine films as grain indenter deforming under pressure solution) in much less frequent (Spiers et al. 1986; Urai et al. 1986). Further work is needed to resolve this discrepancy, which is of special importance for problems of nuclear waste storage in domal salt, since continuous brine-filled grain boundary networks can decrease dramatically the flow stress and sealing performance of rock salt (Littke et al. 2008). Another part of the brine is present in irregular brine pockets bounded by facets (Fig. 5e, f, and B.3 in Online Resource 2), intergranular arrays of IBI, and intragranular quadrangle-shaped brine inclusions (Fig. 7). In our interpretation, the formation of HH boundaries directly confirms that a given boundary was moving at the moment that it was quenched for BIB-cryo sectioning. The growing of new euhedral crystals (Figs. 4c, 5e, f, and B.3 in Online Resource 2). The development of facets according to the Hartman-Perdok theory closes progressively continuous HBH boundaries and irregular brine-filled pockets by the healing of the boundaries as described by Urai et al. (1986), Spiers et al. (1990), Hickman and Evans (1991). The three different types of IBI (elongated shaped, triangle shaped, and quadrangle shaped, Fig. 7) are interpreted as very late stage of grain boundary healing. In majority, these IBI are not connected to each other at the ultimate resolution of the SEM (i.e., above 15 nm). However, sometimes thin lines are present in the queue of these inclusions (Fig. 7) as HH boundaries. At this resolution, it is not possible to distinguish true solid-solid boundaries from a boundary with a 1 nm thick brine film.

Intragranular quadrangle-shaped brine inclusions (Fig. 7e, g) are much larger (typical few micrometers in size) than the three IBI discussed earlier, are poorly aligned, and no HH boundaries are visible at the immediate vicinity, suggesting that these are leftover brine inclusions behind migrating grain boundary. The frequency of these intragranular quadrangle-shaped brine inclusions is relatively high at long-term annealing since samples have completed their static recrystallization (Schenk and Urai 2004; Schmatz et al. 2010).

As in Fujiwara et al. (2009) for growing Si crystals, the variation of IBI morphologies address to the control of crystallographic orientations in grain impingement. Following on the arguments of Van Noort et al. (2006), the systematics of Hartman-Perdok theory is used to interpret that: (i) elongated-shaped IBI and intragranular quadrangle-shaped brine inclusions originate from the convergence of two F faces; (ii) quadrangle-shaped IBI reflect the contact of a S or K face with an other S or K face; and (3) triangle-shaped IBI correspond to the convergence of one F face with one S or K face. Previous contributions (Hickman and Evans 1991, 1995; De Meer et al. 2006) but in current models of equilibrated grain boundary roughness of pressure dissolved surfaces (grain boundaries in natural domal salt continuous brine films as grain indenter deforming under pressure solution) in much less frequent (Spiers et al. 1986; Urai et al. 1986). Further work is needed to resolve this discrepancy, which is of special importance for problems of nuclear waste storage in domal salt, since continuous brine-filled grain boundary networks can decrease dramatically the flow stress and sealing performance of rock salt (Littke et al. 2008). Another part of the brine is present in irregular brine pockets bounded by facets (Fig. 5e, f, and B.3 in Online Resource 2), intergranular arrays of IBI, and intragranular quadrangle-shaped brine inclusions (Fig. 7). In our interpretation, the formation of HH boundaries directly confirms that a given boundary was moving at the moment that it was quenched for BIB-cryo sectioning. The growing of new euhedral crystals (Figs. 4c, 5e, f, and B.3 in Online Resource 2). The development of facets according to the Hartman-Perdok theory closes progressively continuous HBH boundaries and irregular brine-filled pockets by the healing of the boundaries as described by Urai et al. (1986), Spiers et al. (1990), Hickman and Evans (1991). The three different types of IBI (elongated shaped, triangle shaped, and quadrangle shaped, Fig. 7) are interpreted as very late stage of grain boundary healing. In majority, these IBI are not connected to each other at the ultimate resolution of the SEM (i.e., above 15 nm). However, sometimes thin lines are present in the queue of these inclusions (Fig. 7) as HH boundaries. At this resolution, it is not possible to distinguish true solid-solid boundaries from a boundary with a 1 nm thick brine film.

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It is, perhaps, surprising to find thick boundaries in samples annealed for one year (Fig. 7g). On the other hand, there is no evidence that the few% brine present in the samples after compaction was expelled by a sintering-like process during annealing, so that the brine must have been redistributed in the samples. Considering that some grain boundaries did heal during annealing, the presence of the thick boundaries is interpreted to be a necessary part of the annealing process. Even if there was a potential difference across these boundaries, these thick boundaries are interpreted to have a very low mobility due to the long diffusion path across the thick grain boundary. (Urai et al. 1986b)

Model for evolution of brine distribution during static recrystallization in coarse-grained halite

In what follows, we attempt to compile our results and interpretations (Fig. 10) in a model for the evolution of brine distribution in wet sodium chloride aggregates during static recrystallization.

From the initial compaction to the complete recovery of samples, steps are as following:

1. The initial compaction of polycrystalline salt plugs produced abundant intracrystalline defects and microcracks favorable to nucleation of strain-free grains at these sites, together with much smaller grains, which were not removed by sieving. (A. in Fig. 10). Primary recrystallization then proceeds in an isotropic stress field of zero effective stress in fully saturated samples under static conditions without intracrystalline recovery processes inside the deformed grains since annealing was performed at room temperature (Schenk and Urai 2004). There are no external forces to increase the normal stress on brine-filled grain boundaries. Euhedral grains grow into deformed grains by brine-assisted grain boundary migration (Fig. 9).
2. Driven by very small, local differences in brine concentration, some deformed grains are interpreted to grow low-index facets (B. in Fig. 10). This rim of undeformed, faceted halite can locally prevent the migration of a neighbor into this grain but it still needs to be checked using EBSD measurement.
3. Therefore, the evolution of microstructure over time involves three main grain boundary contacts: (1) contact between two old deformed grains (Inset B.1

Fig. 10 Model for evolution of brine distribution during coarsening of chloride sodium aggregate during static recrystallization. The spheres point to impingement sites in B.4 (in C). See details in text

in C. Fig. 10), (2) contact between an old deformed grain and a grain developing low-index faces (Inset B.2 in Fig. 10), and (3) contact involving two grains developing low-index faces (Insets B.4 and B.5 in Fig. 10).

If the grain boundary involves two old deformed grains with thick brine films ( $H_1BH_n$ ), the grain boundary has a very low mobility.

If the grain boundary consists of a deformed grain and a new grain with low-index faces ( $H_1BH_p$ ), and the grain boundary is brine-filled (50 nm), the new grain grows into the deformed grain, driven by reduction of dislocation density, and facilitated by the fact that the inside of the faceted boundary is irrational (Insets B.2 and B.3 in Fig. 10; Passchier and Trouw 2005; Schenk and Urai 2004; Humphreys and Hatherly 1996).

If the grain boundary involves two grains developing low-index faces ( $H_1BH_{fr}$ ), both crystals continue to grow until they impinge. Our interpretation is that part of the boundary becomes solid-solid (HH) (Insets B.4 and B.5 in Fig. 10).

5. Driven by the incipient of grain boundary healing process, brines at impingement evolve as isolated inclusions (C. in Fig.10): (a) elongated-shaped IBI appear at interface of two F faces, (b) aligned quadrangle-shaped IBI are at contact of a S or K face with an other S or K face, and (c) triangle-shaped IBI involve one face with one S or K face. Intragranular big quadrangle-shaped IBI are leftover brine inclusions behind migrating grain boundary.

## Conclusions

- € BIB-cryo-SEM allows direct in situ investigations of brine-filled grain boundaries in wet halite polycrystals, resolving brine films in grain boundaries down to 15 nm thicknesses.
- € Our observations are consistent with earlier work on brine-assisted grain boundary migration in wet halite and provide new information on the evolution of grain boundary structure.
- € Brine-filled grain boundary migration is strongly affected by the development of facets. These facets can be rational or irrational and lead to migration, impingement, and healing of the grain boundary.
- € Grain boundary healing locally produces boundaries in which our instrument cannot resolve brine film. We interpret these to be healed, solid-solid boundaries.
- € Some brine-filled grain boundaries remain several micrometers thick over annealing periods of over a year.

**Acknowledgments** We thank the Deutsche Forschungsgemeinschaft for supporting our project and funding the BIB-cryo-SEM instrument (Project UR 64/9-2). We are also grateful to Dr. Georg Koschek, Dr. Christian Jaeger and Mr. Hans-Rolf Schreiber from Carl Zeiss NTS GmbH, and Dr. Sebastian van Offern from Klocke Nanotechnik GmbH for their work to help us to develop the BIB-cryo-SEM instrument. We thank Prof. Dr. F. Michael Meyer from Institut für Mineralogie und Lagerstättenlehre at RWTH Aachen University for hosting the instrument; and Franz Gmüder from the Geological Institute at RWTH Aachen University for his help with installing the instrument.

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