Growth behavior of hydrogen micropores in aluminum alloys during high-temperature exposure

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https://hdl.handle.net/2324/1813034
Growth behavior of hydrogen micropores in aluminum alloys during high-temperature exposure

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Abstract

X-ray microtomography was used to observe hydrogen micropores and their growth behavior at high temperatures in several aluminum alloys. High-density micropores were observed in high-purity Al-Mg alloys, but their density and volume fraction were much lower in pure aluminum. Our results have revealed that the growth behavior of micropores is dominated by Ostwald ripening. About 53 % of hydrogen is trapped in micropores in Al-Mg alloy with low hydrogen content, making micropores the predominant hydrogen trap site. Although total hydrogen content is similar to that in the alloy, the ratio of hydrogen trapped in micropores is below 7 % in pure aluminum. This difference is attributable to the lack of hydrogen precipitation sites in pure aluminum. Although the overall amounts of hydrogen at dislocations and grain boundaries are small in all the materials, the occupancies for these trap sites were concluded to be very high.
1. Introduction and background

The solubility of hydrogen in aluminum is extremely small, at only $10^{-6}$ atomic fraction near the melting point, and decreases rapidly with temperature [1]. Hydrogen content extrapolated from the high temperature side is $3.9 \times 10^{-4}$ cm$^3$ at room temperature, equivalent to the volume of hydrogen gas at 0 °C and 1 atm per 100 g of aluminum [1]. Measured hydrogen content ranges, however, from about 0.11 ~ 0.60 cm$^3$ / 100 g Al in wrought aluminum alloys [2] and 0.21 ~ several cm$^3$ / 100 g Al for cast aluminum alloys [2, 3]. These are three to four orders of magnitude larger than the above hydrogen solubility. [2]. These values may be attributable to the well-known hydrogen solubility gap at the melting temperature of aluminum [4], as a result of which solid aluminum contains supersaturated hydrogen just after solidification.

Supersaturated hydrogen must be partitioned between interstitial solution and trap sites. Hydrogen trap sites in high-purity aluminum include lattice interstices, dislocations and vacancies [5]. In the case of aluminum alloys, solute atoms, Guinier-Preston zones and subsequent metastable and stable precipitates, and high-angle grain boundaries also behave as trap sites [5, 6]. It has been claimed that there is no evidence to suggest that subgrain boundaries trap hydrogen, due to its low-energy dislocation structure [6]. It is interesting to note that trapped hydrogen is not readily lost during storage at room temperature, or even after a combination of high-temperature exposure and hot working in the case of wrought aluminum alloys. It has been reported that the thin but robust oxide film at the surface plays an important role in impeding the release of hydrogen from solid aluminum [1]. Russell has reported that a thermally-formed oxide film drastically reduces the permeability of aluminum membranes to hydrogen [7]. The above-mentioned extraordinary hydrogen content in solid aluminum can therefore be attributed to such hydrogen occlusion due to the presence of a surface oxide layer, plus the contributions of various trap sites.
Porosity is commonly observed in both wrought and cast aluminum alloys [1,2,10,8]. The porosity formation mechanisms are the precipitation of hydrogen gas or volumetric shrinkage upon solidification. Porosity can therefore be characterized as another hydrogen trap site in aluminum alloys. It has been reported, using a small-angle X-ray scattering technique, that hydrogen-vacancy defects form platelets 15 nm in radius and 7 nm in thickness lying on specific crystallographic planes [9]. This may be the direct formation mechanism of micropores in aluminum. On the other hand, Anyalebechi et al. have categorized the nucleation of micropores into several mechanisms [10], claiming that dislocation loops, formed from clustered vacancies, provide energetically favorable sites for the nucleation of hydrogen micropores as well as oxide inclusions and coarse constituent phase particles.

Ichimura et al. have investigated the effects of micropores on hydrogen diffusivity and solubility, concluding that the decrease in hydrogen diffusivity and the increase in solubility are attributable to the presence of molecular hydrogen in pores [11]. Other researchers have examined hydrogen concentration in pure aluminum, drawing the conclusion that a large amount of hydrogen is trapped in vacancy clusters or pores on a microscopic scale [12]. Thermal desorption spectroscopy (TDS) was employed by Outlaw et al. [8]. Although their technique does not provide a direct measurement of the hydrogen trapped within each pore, they have claimed that in pure aluminum, more than 99% of hydrogen is trapped in micropores. Percentage porosity has, in fact, been closely studied, and proves to depend on the hydrogen content in aluminum alloy [1,13,14], strongly suggesting that micropores in aluminum alloys are filled and pressurized with molecular hydrogen. Another point to note is that high-purity aluminum usually shows much lower porosity for a given hydrogen content [14]. It can be inferred that this difference is associated with the kinds and density of available trap sites in high-purity aluminum. It will be examined fully later in the present experiment.

The mechanical properties of aluminum alloys are sometimes sensitive to the presence of micropores [14]. In response to this, the relationship between metal working and micropore healing
has been well documented [1,14,15]. Wang et al. have reported the necessary hydrostatic pressure
and holding time to realize the healing of micropores. On the other hand, it has been claimed that
interdendritic porosity in cast aluminum alloys does not close completely in the course of metal
working. Instead, it is flattened, thereby reducing its short-transverse mechanical properties [1].
This so-called secondary porosity has been reported to remain unchanged through extensive
defformation by hot and cold rolling [1]. Direct observation of healing/reopening during plastic
working and subsequent heat treatments will be undertaken in another paper [16].

Numerous experimental reports are available in the literature that report micropore growth at
high temperatures [3,10,17,18]. However, in the conventional cross-sectional or surface
observations, micropores are easily filled with abrasive powders during sample preparation
processes for the observations, such as cutting and polishing [19,20,21,22,23], resulting in
uncertainty and inaccuracy in micropore observation. In the present study, synchrotron X-ray
microtomography with recently enhanced resolution levels [24] was applied to clarify the above
issues. Also, the high resolution 3-D/4-D (i.e., 3-D plus time axis) imaging readily enables the
visualization of individual microstructural features, providing a unique opportunity to quantitatively
analyze the growth, shrinkage and annihilation behavior of each pore. This is clearly an
improvement on the procedures applied in the available literature, which allow only macroscopic
evaluations.

2. Experimental methods

2.1 Sample preparation

Aluminum of 99.999% purity (Sumitomo Chemical Co., Ltd.) and pure magnesium were
used to prepare Al–5.5 mol% Mg alloys. The purity of the magnesium was 99.97% (Ube Co., Ltd.)
and 99.98% (Timminco Co., Ltd.) for alloys MH/HH (as hereinafter described) and alloy LH (as
hereinafter described), respectively. Crucibles for the former and latter alloys were made of alumina at 99% purity (TEP Co., Ltd.) and for electrode-grade graphite (Shin-Nihon Tanso Co., Ltd.), respectively. The Al–5.5 mol% Mg binary alloy was melted in an atmosphere of argon and cast into iron molds (for simplicity, mol% is expressed as % below). Here, a melting atmosphere of argon was applied to avoid hydrogen contamination. This alloy is hereinafter called alloy LH to indicate its low hydrogen content. Another Al–5.5% Mg alloy, melted in ambient laboratory air, is hereinafter called alloy MH. The details of the chemical compositions of alloy LH and MH are available elsewhere [25] (Alloys LH and MH correspond to Alloy-3 and Alloy-4, respectively, in the reference paper.). These ingots were homogenized at 703 K for 18 h in an argon atmosphere. A part of the alloy MH ingot was remelted in ambient laboratory air. Before remelting, TiH₂ was added to a crucible with a cover to raise the hydrogen partial pressure during melting. TiH₂ decomposes below the melting point of the aluminum alloy. The alloy melt was protected from contamination with any remaining Ti by placing an appropriate divider plate in between. The material is hereinafter called alloy HH. All the Al–5.5 mol% Mg alloys contained impurity silicon of 0.003 ~ 0.004 mass %. No other impurity such as Fe, Na and Ca was detected. Aluminum at 99.999% purity, hereinafter called 5N-Al, was also used for comparison purposes. To avoid contamination, no homogenization treatment was applied to 5N-Al.

The three Al-Mg alloys were then exposed at 823 K for 0 to 270 h to observe any changes in micropore morphology. Changes during the homogenization treatment at 703 K for 18 h were also observed only for alloy HH. A specimen 15 mm in length and 0.6 x 0.6 mm in cross section and one 7 mm in length and 0.05 x 0.05 mm in cross section were machined from the ingots and used for projection-type and imaging-type microtomography observations, respectively.

The vacuum fusion method was utilized to measure the total hydrogen content of each material. Dislocation density was measured using Keh’s method [26] on {100} faces observed with a transmission electron microscope operated at 200 kV. Grain size was measured using an optical microscope after etching polished samples by means of Barker’s method.
2.2 Tomographic imaging

X-ray tomography was performed at the undulator beamline BL47XU of the SPring-8 synchrotron radiation facility. A monochromatic X-ray beam with a photon energy of 20 keV, produced by a liquid nitrogen-cooled Si (111) double crystal monochromator, was used for the projection-type microtomography. A beam diffuser rotating at 600 rpm, packed with graphite powder, was employed to modify the high coherence of the X-rays. An image detector was positioned 20 mm behind the sample, making the imaging system sensitive to phase modulation in addition to the classical absorption contrast [27]. The image detector consisted of a cooled 4000 (H) x 2624 (V) element CCD camera (effective pixel size of the camera: 5.9 µm), which was used in 2 x 2 binning mode, a scintillator (Lu2SiO5:Ce) and a relay lens (×20). In total, 1500 radiographs, scanning 180 degrees, were obtained in 0.12-degree increments. The entire cross-section of the specimen and a region about 622 µm high were captured on the CCD camera.

Imaging-type microtomography, which has state-of-the-art high resolution, was performed at 8 keV. A schematic illustration of the setup is available elsewhere [28]. The system consists of the beam diffuser, high precision stages, a center beam stop, a condenser zone plate (CZP), a pinhole as an order-sorting aperture for the CZP, a Fresnel zone plate (FZP) and the X-ray image detector. The accuracy of the sample rotary stage, which is one of the key issues in determining the whole resolution of the imaging CT, showed less than ±70 nm wobble during 360° rotation [28]. The CZP has a concentric grating pattern with an even zone width, and was utilized as an illumination device for Köhler’s illumination under a quasi-parallel beam from the synchrotron radiation. First-order diffraction was used for illumination purposes. The FZP was used as an objective. The focused beam size of the 1st-order diffraction of the FZP was measured as 120 nm in full width at half maximum, almost equal to the diffraction-limited spot size defined by Rayleigh’s criterion. Both CZP and FZP were fabricated using an electron-beam lithography technique. The detector consisted of a thin phosphor (P43, Gd2O2S:Tb+) screen, tandem camera lens and the cooled CCD camera (as
above). The CCD camera was also used in the 2 x 2 binning mode. The exposure time for each projection was 500 msec (at 8 keV). Total scan time for 1,500 projections was about 30 minutes.

2.3 3-D image reconstruction and subsequent image analysis

Image slices were reconstructed from a series of projections based on the conventional filtered backprojection algorithm. The grey value in each dataset was calibrated such that the linear absorption coefficient of 0 ~ 20 cm⁻¹ fell within an 8-bit grayscale range between 0 and 255. Isotropic voxels with 0.474 and 0.088-µm edges were achieved in the reconstructed slices for the projection- and imaging-type setups, respectively.

To estimate the volume of each micropore at sub-voxel accuracy, pentagonal faceted iso-intensity surfaces were computed from the volumetric data set using the conventional Marching Cubes algorithm [29]. A threshold value to obtain binary images was chosen as a median between the linear absorption coefficient peaks of air and the aluminum matrix. To suppress inaccuracies originating from image noise, only micropores over 6.668 voxels in volume were counted as micropores in the quantitative analysis.

Figure 1 (a) illustrates a two-dimensional (2-D) grey value distribution on a virtual slice where three micropores and some particles, which are seen in black and white respectively, are embedded in a grey aluminum matrix. In general, light and dark fringes appear at each micropore/matrix and particle/matrix interfaces due to Fresnel diffraction [30]. A pair of light and dark fringes must also appear at a particle/micropore interface if there is such an interface in the materials used. The existence of such an interface implies that pre-existing particles act as heterogeneous nucleation sites. Due to the existence of such fringes, however, it is hard to confirm if micropores are attached directly to particles or there is intervening matrix aluminum in between, as seen in Fig. 1 (b), which is the result of segmenting the virtual cross-sectional image shown in Fig. 1 (a). For closer examination, segmented micropores were expanded in 3-D, as shown in Fig. 1 (c). The ratio of micropores nucleated heterogeneously on particles was estimated by changing the degree of
expansion.

3. Micropore growth behavior at high temperatures

3.1 Observation with projection-type setup

Figure 2 shows the 3-D reconstructed images of micropores in as-cast, homogenized and heat-treated conditions in alloy HH. Note that only micropores are extracted, while underlying aluminum and dispersion particles are not displayed in Figure 2. In the as-cast condition, interdendritic shrinkage cavities with complex shapes are seen together with small, high-density micropores. During the homogenization treatment at 703 K for 18 h and subsequent high temperature exposure at 823 K for 90 h, the shrinkage cavities gradually shrunk and spheroidized as is indicated in circle A. Instead, small and high-density micropores were grown, making overall spatial distribution relatively homogeneous, as seen in Fig. 2 (c). Such growth might be attributable to the creep deformation of aluminum around the micropores due to the presence of high hydrogen pressure in the micropores.

Similar observations for alloys MH and LH and 5N-Al are shown in Figs. 3 – 5. It appears that although, as is shown later, the hydrogen content of the two alloys is similar according to the macroscopic hydrogen analysis, alloy MH is thought to contain more significant shrinkage cavities than alloy HH. On the other hand, alloy LH exhibits much fewer shrinkage cavities than alloys MH and HH. It is interesting to note that micropores in alloys LH and MH show angular shapes after being exposed to 823 K for 90 h. Micropores in alloy HH also exhibited slightly angular shapes after exposure for 270 h. A closer inspection revealed that the angular micropores exhibit octahedral shape with some type of crystallographic orientation relationship with the underlying aluminum matrix. It can be inferred that if hydrogen trapped in micropores is released from the alloys, micropores with lower internal gas pressure reach an equilibrium between internal gas pressure and surface energy, and change their shape to reduce the surface energy. Since the \{111\} planes have the
lowest surface energy of all the low-index planes in aluminum [31], it is reasonable to assume that micropores are prone to form octahedra surrounded by eight \{111\} planes, making them energetically stable in aluminum. Similar faceted pores have been observed in a Nb-Zr alloy implanted with He, which exhibited 18-sided polyhedral pores bounded by (100) and (110) faces [32]. The alignment of the angular micropores over a relatively large volume is attributable to the very large grain size of the Al-Mg alloy used (typically 100 ~ 400 \(\mu\)m). There is a marked difference in the appearance of the micropore distribution in 5N-Al compared to the Al-Mg alloys. Only two ellipsoidal and coarse micropores were observed in 5N-Al, and their appearance did not significantly change, even after exposure to 823 K for 90 h, as shown in Figure 5.

Figure 6 shows magnified views of micropore distributions for alloy MH and 5N-Al. A variety of behaviors, such as the growth, coalescence, shrinkage and annihilation of the micropores, are demonstrated in this tiny volume. For example, pore A was annihilated after 10 h of heat treatment. Pore B grew up to 30 h and then coalesced with a neighboring pore. Pore C also appeared to grow up to 30 h and then self-annihilated. Meanwhile, pore D showed monotonic growth up to 90 h. Fundamentally, micropores can grow by two different mechanisms: one is Ostwald ripening and the other is migration and coalescence. The former mechanism needs gas re-solution and the latter is likely to occur as a result of gas surface diffusion and/or vacancy volume diffusion. Detailed inspection of the tomographic volumes in this study did not reveal any migration of micropores. A more quantitative discussion on this issue will appear later. Although it is not seen in Figure 6, newly-initiated small spherical micropores were also seen in other regions. One coarse micropore and another very small micropore beside it are seen in Figure 6 (b). Both of these micropores showed very gradual shrinkage behavior during the period of 90 h, but it might be conceivable only after 90 h has elapsed. The change in the shape of micropores from spheroid to octahedral is also clearly confirmed in Figure 6.

3.2 Observations with the imaging-type setup
Figure 7 shows a high-resolution 3-D image of material HH that has been captured after homogenization treatment at 703 K for 18 h. In this figure, a translucent image of the specimen shown in grey has been superposed on the micropore image shown in red. It can be confirmed that there are no micropores smaller than 1 µm. Since the resolution of the projection-type microtomography such as used for Figs. 2 – 6 is approximately 1 µm, it can be concluded that all the micropores embedded in the materials used can be observed using projection-type microtomography. Subsequent quantitative analysis will therefore be performed solely on the basis of the microtomography images taken using the projection-type setup.

3.3 Results of 3-D quantitative analysis

Figures 8 ~ 10 show the results of the 3-D quantitative analysis. The number density of the micropores was 16.1, 50.7 and 54.7 x 10^{12} m^{-3} for alloys LH, MH and HH, respectively. The number densities markedly decrease with exposure at 823 K in the cases of alloys LH and HH, while alloy MH showed a rapid increase up to 30 h and then began to decrease. The number density of micropores for 5N-Al remained at less than one-tenth of those of the Al-Mg alloys throughout the high-temperature exposure. It seems to be consistent with the past literature that high-purity aluminum shows much lower porosity for a given hydrogen level [14]. Indeed, the reported difference in volume fraction between 99.99 % aluminum and aluminum alloys, such as 99.2 % aluminum and Al-4.6% Cu alloy, for hydrogen content of 0.3 cm^3 / 100 g Al is 8 ~ 10 times, which is even larger than the difference between alloy LH and 5N-Al (0.29 ~ 0.3 cm^3 / 100 g Al) shown in Fig.10 [14]. Average micropore diameter increased rapidly in the cases of alloys LH and HH, whereas it was almost flat in alloy MH and 5N-Al, as shown in Figure 8. In alloys LH and HH, the size distributions changed from a narrow distribution, centered at about 3 µm with similar diameter to the as-homogenized condition, to a broad distribution peaking at 8 ~ 13 µm after exposure at 823 K for 90 h, as shown in Figure 9.

These patterns of number density and size distribution suggest that the micropore growth
behaviors in alloys LH and HH are dominated by Ostwald ripening. In alloy HH, the average diameter began to decrease again at the longest time period, as seen in Figure 8. This suggests the possibility of a decrease in hydrogen content during long exposure. Alloy MH 9 (Figure 9) showed a relatively irregular size distribution. The singular behavior in size and number density observed in alloy MH might be attributable to its characteristic pore distribution: an enormous shrinkage cavity, several hundred micrometers across, as seen in Fig. 3, which almost reaches the length of a diagonal line for the tomographic region of interest. The volume fraction of micropores shown in Fig. 10 increased rapidly after 5 h had elapsed at 823 K in all the Al-Mg alloys. This time period is close to the standard solution treatment time for aluminum alloys. Meanwhile, volume fraction decreased monotonically in 5N-Al, in marked contrast to the Al-Mg alloys. These results indicate that the prolongation or repetition of heat treatment might inevitably cause a degradation in mechanical properties due to the growth of micropores. It seems most likely that micropores can be healed during cold and/or hot plastic working before being exposed at the solution treatment temperature. Complementary studies on this will be available in a companion paper [16].

4. Discussion

4.1 Nucleation sites for micropores

Figure 11 shows variation in the ratio of micropores directly attached to particles, \( P_h \), after each micropore has been dilated at one-voxel intervals. If micropores and particles both independently nucleate and no preferential sites are available for them, the distance between a micropore and the nearest particle would tend to be randomly distributed. In this case, \( P_h \) would be expected to increase moderately with the dilation width, \( w_d \). However, if there are micropores heterogeneously nucleated on particles, the presence of light and dark fringes in between, caused by the Fresnel diffraction, might cause an apparent separation between particles and micropores. In this case, \( P_h \) is expected to increase rapidly until \( w_d \) reaches the width of the light and dark fringes and then
decelerate rapidly to a more moderate rate of increase, dominated by statistical encounters with neighboring particles. Figure 11 illustrates this tendency, suggesting the presence of heterogeneously nucleated micropores. The intersection point of two auxiliary lines in Fig. 11 gives $P_h$ of 59.6% at a $w_d$ of about 2.0 µm (i.e., four voxels).

The first Fresnel zone size is given as $(\lambda d)^{0.5}$, where $\lambda$ is the wavelength of the X-rays and $d$ is the sample/detector distance [33]. $\lambda = 0.062$ nm and $d = 20$ mm in this study, resulting in a $(\lambda d)^{0.5}$ value of 1.1 µm. Since a pair of light and dark fringes is seen in the tomographic images, it would be reasonable to take the second Fresnel zone size, which is almost equivalent to the obtained $w_d$ value. It can therefore be concluded that the majority of the micropores have been nucleated on pre-existing particles. Since the resolution level for current projection-type tomography is about 1 µm, there would appear to be a large number of sub-micron particles in the materials used that are not visible using the current projection-type setup. The obtained $P_h$ value of 59.6% would therefore be more or less an underestimated value, and the actual ratio would be much higher.

As was described in the Introduction, it has been reported that nanometer-sized platelets form from hydrogen-vacancy defects [9] and that dislocation loops form from clusters of vacancies [10], providing energetically favorable sites for the nucleation of hydrogen micropores. This means that the formation mechanism of micropores is homogeneous nucleation. According to the present analysis, however, at least 60% of micropores have been formed heterogeneously on constituent particles. There is little evidence within the present study that vacancies are involved in the formation process of micropores in any form. As is shown in Table 3, total hydrogen content is almost identical between alloy LH and 5N-Al. Nevertheless, the present study reveals a marked difference between them in the number density and size of micropores. It might also support the above hypothesis that micropores are formed heterogeneously on particles. In other words, heterogeneous nucleation sites appear to be necessary for hydrogen to form micropores, and if not, micropore formation is effectively suppressed. This will be analyzed later where hydrogen, which has failed to form micropores, is trapped in pure aluminum.
4.2 Micropore growth mechanism

It would be reasonable to assume that the observed micropores have been in thermal equilibrium where surface tension, \( \gamma \), is balanced by an opposing gas pressure inside a micropore, \( P \), as follows:

\[
4\gamma / d = P
\]

(1)

where \( d \) is the micropore radius [34]. Two possible growth mechanisms under such thermal equilibrium would be micropore migration/coalescence and Ostwald ripening [35]. For the former mechanism to operate, micropores should migrate via atom transport through surface diffusion, vacancy volume diffusion or vapor transport through gas [35]. According to Nichols [36], mean micropore diameter, \( \bar{d} \), and number density, \( \rho_p \), are expressed as follows if ideal gas behavior and spherical micropores are assumed:

\[
\bar{d} \propto (D M_h t)^{1/n}
\]

(2)

\[
\rho_p \propto C_h (D M_h t)^{-2/n}
\]

(3)

where \( D \) is the diffusion coefficient of aluminium, \( M_h \) is total hydrogen content and \( t \) is time. On the other hand, according to the model for Ostwald ripening proposed by Greenwood et al. [37], \( \bar{d} \) and \( \rho_p \) are expressed based on the same assumptions as Eqs. (2) and (3), as follows:

\[
\bar{d} \propto (D_h C_h t)^{1/2}
\]

(4)

\[
\rho_p \propto M_h (D_h C_h t)^{-1}
\]

(5)
where $D_H$ is the diffusion coefficient of hydrogen in aluminium and $C_{H}$ is the hydrogen concentration in the aluminum matrix. Eqs. (2) – (5) indicate that in the case of migration- and coalescence-dominated growth, both $\bar{d}$ and $\rho_p$ are functions of $M_H$. Meanwhile, if Ostwald ripening dominates, only $\rho_p$ is a function of $M_H$, and $\bar{d}$ is independent of $M_H$.

Table 1 shows $\bar{d}$, $\rho_p$ and $M_H$ for the three Al-Mg alloys in the as-homogenized condition. The reason that only the as-homogenized condition is provided for the analysis is that a decrease in $M_H$ might be expected during long exposure at 823 K, as was suggested in Figure 8. It can be seen in Table 1 that $\bar{d}$ is nearly constant, while $\rho_p$ shows some dependency on $M_H$, suggesting that Ostwald ripening is predominant in the growth behavior of micropores in the present materials. As indicated in Eq. (5), $\rho_p$ is not linearly proportional to $M_H$. This is likely due to the limited size range of micropores provided for in the quantitative analysis.

Markworth et al. have also analyzed the asymptotic coarsening kinetics in the case of Ostwald ripening and shown asymptotic size distribution functions [38]. According to their analysis, the maximum micropore diameter, $d_{\text{max}}$ is expressed as follows.

$$d_{\text{max}} \approx 2.36\bar{d}$$

(6)

Table 2 shows the $d_{\text{max}}/\bar{d}$ ratio calculated using the tomographic data. The values just after the homogenization treatment range from 2.2 to 2.3, averaging 2.23. A similar tendency has been confirmed by the present authors for a commercial wrought aluminium alloy [39]. Agreement with Markworth’s analysis also supports the predominance of Ostwald ripening, which is also in qualitative agreement with the observations shown in Figs. 2 – 6.

4.3 Hydrogen trapped within micropores

Hydrogen gas trapped within micropores might be amenable to measurement using
conventional analytical methods such as electron-energy-loss spectroscopy (EELS) and ultraviolet absorption spectroscopy (UVAS) as energy shifts of hydrogen excitations [40]. Huang diffuse scattering (HDS) has also been applied to visualize strain fields around pores [40]. However, there are certain size ranges for which these techniques are suited. For examples, EELS and UVAS cover 1 ~ 100 nm, while HDS can be applied to micropores smaller than 10 nm [40]. Therefore, micropores observed in the present study cannot be measured with these techniques. In addition, these techniques provide only macroscopically averaged information, and hydrogen trapped in individual micropores is out of reach of measurement.

It has been reported for very small pores ($d < 100$ nm) that equilibrium pressure and gas density can no longer be described by the ideal gas law [35]. However, this is not the case: all the micropores observed in the present study can be analyzed with reasonable accuracy using the ideal gas law. Above $T_m/2$ ($T_m$: melting temperature), thermal vacancy absorption is rapid enough to maintain the gas pressure within micropores at the equilibrium value characterized in Eq. (1) [40]. This is because the equilibrium concentration of vacancies is approximately three orders of magnitude higher than the solubility of hydrogen at the melting point of aluminium [41]. By combining Eq. (1) with the ideal gas law, hydrogen trapped in all the micropores, $C_{\text{pore}}$ (mol), is given by summing the hydrogen content in every micropore as follows:

$$C_{\text{pore}} = \sum \left( \frac{2\pi\gamma d_i^2}{3RT} \right)$$  \hspace{1cm} (7)

where $d_i$ is the diameter of the $i$th micropore, $R$ is the gas constant and $T$ is temperature. $\gamma$ was assumed to be 1.16 N/m, which is the measured value for the (111) face [42]. Since the (111) face has the minimum energy in aluminum, the calculation yields conservative lower limit values. The diameter of each micropore is known from the tomographic data, so $C_{\text{pore}}$ can therefore be readily obtained, as shown in Table 3. Here, the total hydrogen trapped is expressed as the equivalent
volume of hydrogen gas at 0 °C and 1 atm per 100 g of aluminum, $M_{H}^{\text{pore}}$. It's important to note that micropores should be in thermal equilibrium as eq. (1) to obtain $C_{\text{pore}}$. Since micropores are expanded due to creep deformation of the aluminium matrix during the exposure at 823 K as shown in Figs. 8 and 9, the analysis has been applied only to the materials without the exposure. It is noteworthy that about 53 % of hydrogen is trapped within the micropores in alloy LH, while although the total hydrogen content is similar, it is less than 7 % in 5N-Al. This difference may be attributable to the lack of hydrogen precipitation sites in the pure aluminium, as discussed above. $M_{H}^{\text{pore}}$ is also a low 3 ~ 9% for alloys MH and HH. Since the total hydrogen in these alloys was very high, it can be inferred that the way that hydrogen is partitioned into possible trap sites would be different if the total hydrogen content is high.

As stated earlier, Outlaw et al. performed a thermal desorption experiment [8], concluding that, in pure aluminum, more than 99% of hydrogen is trapped in micropores. This is starkly inconsistent with the results obtained in this study. In their technique, a gas pulse observed before the melting of aluminum was assumed to correspond to the desorption of hydrogen from all the pores. It is not clear at this moment whether the inconsistency can be traced to their indirect measurement method or to marked difference in the state of the materials used.

4.4 Estimation of hydrogen trapped at each trap site

Available trap sites for hydrogen in aluminum are lattice interstices, dislocations, vacancies, solute atoms, precipitates and high-angle grain boundaries. Of these, the vacancy and precipitate can be neglected for the materials used because they are analyzed after being cooled slowly from the homogenization temperature. In 5N-Al, grain size is a few tens of millimeters, suggesting that no grain boundaries are present in the present tiny tomographic sample. Microstructural parameters associated with the trap sites are listed for the materials used in Table 4.

Smith et al. have performed TDS experiments for hydrogen pre-charged aluminum alloys with a varying thermal ramp rate, estimating binding energies for various trap sites based on Arrhenius
plots [43]. According to them, trap binding energy, $E_b$, for the interstitial lattice, dislocations and high angle boundaries is 0, 31.7 and 35 kJ/mol, respectively. No hydrogen trapping at sub-grain boundaries was identified in their study. The $E_b$ value for dislocations in 99.999% aluminum has been also reported by Young et al. to be 27.3 kJ/mol [4], which is close to the above value. The binding energy between hydrogen and solute magnesium atoms has been measured by Leger et al. to be $E_b < 5.8$ kJ/mol in high purity Al-Mg alloys [44]. Smith et al. have analyzed trapped hydrogen in each trap site together with trap coverage with hydrogen [43]. In the following part, a similar procedure is applied to measuring hydrogen partitioning in the materials used. Hydrogen stored in normal interstitial lattices and other trap sites should, according to Oriani’s theory, always be in equilibrium as follows [45]:

$$\frac{\theta_T}{1-\theta_T} = \theta_L \exp\left(\frac{E_b}{RT}\right)$$

(8)

where $\theta_L$ and $\theta_T$ denote the occupancy of the interstitial sites and trap sites, respectively. Total hydrogen concentration, measured in hydrogen atoms per unit volume, $C^T_H$, is the sum of the hydrogen stored in a normal interstitial lattice and all the trap sites, corresponding to the total hydrogen content listed in Table 3:

$$C^T_H = \theta_L N_L + \sum \theta_{T_i} N_{T_i} + C_{\text{pore}}'$$

(9)

where $N_L$ and $N_{T_i}$ (atoms per unit volume) respectively denote trap densities in the normal interstitial lattice and $i$th trap sites other than micropores. $C_{\text{pore}}'$ represents total hydrogen atoms per unit volume trapped in micropores, which can be converted from $C_{\text{pore}}$. Table 5 shows the values of the trap density estimated from the measured dislocation density, grain size and the concentration of solute magnesium. Trap site occupancy and the concentration of hydrogen for each trap site are
listed in Table 5. In this analysis, $E_b$ for solute magnesium was assumed to be its upper limit, i.e., 5.8 kJ/mol, and $E_b$ values for other trap sites were taken from Smith’s paper [43].

Due to the relatively low dislocation density and coarse grain size in the Al-Mg alloys used, the hydrogen trapped at these trap sites is limited. It is also worth noting that the trap site occupancy values are high for dislocations and grain boundaries, ranging from 0.357 to 0.970 and 0.677 to 0.992, respectively. The total amounts of hydrogen trapped in micropores are similar in the three Al-Mg alloys. Since the total hydrogen content is significantly lower in alloy LH than in alloys MH and HH, micropores are the major hydrogen trap site in alloy LH, higher than the interstitial lattice by a factor of 2.46. In general, it is well known that apparent hydrogen diffusivity, $D_a$, is reduced with $D_a = D_L / (1 + \phi)$ [46], where $D_L$ is the ideal diffusivity of hydrogen in aluminum and $\phi$ is the ratio of hydrogen between the trapped sites and interstitial lattices given by $\phi = (\Sigma \theta_i N_i + C_{pore}) / \theta_L N_L$ from Eq. (9). The result in Table 5 indicates that $D_a$ is reduced by about 80% in alloy LH due to the high ratio of hydrogen trapped in the microstructural defects.

Although hydrogen trapped in micropores is different by a factor of 3.4 in alloys MH and HH, similar ratios (about 62%) of the total hydrogen are located in the interstitial lattices in both alloys. This is in contrast to the common knowledge that the true interstitial lattice solubility in aluminum is relatively small compared to other solid FCC metals. It can be inferred that excess hydrogen in solid aluminum is predominantly partitioned to micropores, and the occupancies for interstitial lattices, dislocations and grain boundary sites are elevated with further increases in hydrogen concentration. The reason that more hydrogen is not held in micropores might be mutual interaction between neighboring micropores.

Comparison of alloy LH and 5N-Al, which have a similar total hydrogen content, indicates that if micropores cannot precipitate on some preferential sites, dislocations store a large amount of hydrogen, bringing levels close to those of the high-hydrogen alloys MH and HH. It has been reported that the formation of blisters is suppressed by applying cold deformation of 20 ~ 60% [47]. This may be consistent with the high trap site occupancies at dislocations, which are close to unity.
for alloys MH and HH. Cold deformation inevitably leads to an increase in dislocation density, as a result of which the capacity for hydrogen storage at dislocations would be significantly expanded. Grain boundary-trapped hydrogen concentration has been measured and analyzed, and in the literature it is associated with hydrogen embrittlement behavior [48,49]. There has been a certain amount of controversy concerning grain boundary trapping, since it has sometimes been reported that hydrogen segregation is not observed experimentally [4,50]. The estimated value for grain boundary trapping has yielded the lowest value in each material, which is one to two orders of magnitude smaller than the other trap sites. It might be reasonable to suppose that even a small amount of hydrogen trapped at grain boundaries can have a strong influence on fractures, but since a grain boundary is a 2-D defect, the absolute amount of hydrogen stored at grain boundaries was confirmed to be very low in the present analysis.

4.4 Insights into practical material manufacturing processes

In general, high-temperature heat treatments are performed on aluminum alloys with the aim of the eliminating segregation and coarse intermetallic particles and/or solutionization. However, in the light of the findings of the present study, heat treatment conditions should be controlled such that both homogenization/solutionizing (positive effects) and the suppression of micropore growth (negative effects) are satisfied in an optimal balance from a practical point of view. For examples, a shorter heat treatment might sometimes bring a better result. The fundamental mechanism of micropore growth and the hydrogen partitioning behavior into micropores, which have been clarified in the present study, would provide valuable insights for advanced property controls of this type. Since wrought aluminum alloys experience some plastic working before usage, it is indispensable to assess the shrinkage and annihilation behaviors of micropores in subsequent metal working processes. The results of the direct observation of healing/reopening during plastic working and subsequent heat treatments performed in another paper [16] should also be taken into account to make the heat treatment control more efficient.
It is also interesting to note that micropore formation is suppressed in the absence of alloying elements. In practical aluminum alloys with some alloying elements, the existence of numerous micropores would exert some effects on mechanical properties. For example, it has been reported that tensile strength decreases rapidly if porosity increases up to 0.5 ~ 1% [1]. The high resolution 3-D/4-D imaging also enables the visualization of individual microstructural features, providing a unique opportunity to reveal the contribution of each pore on fracture. It will be reported by the present authors in another paper [51], providing more detailed information on the microstructural control in the heat and thermo-mechanical treatments.

5. Summary

Synchrotron X-ray microtomography was used to observe hydrogen micropores and their shrinkage, annihilation, initiation and growth behaviors during high-temperature exposure in high-purity Al-Mg alloys with three different hydrogen contents and in 99.999% pure aluminum. Ultra-high-resolution imaging tomography was also used to identify the minimum pore size in the materials.

High-density micropores were observed in the Al-Mg alloys, while the density and overall volume fraction of micropores were much lower in pure aluminum. At least 60% of micropores had been formed heterogeneously on constituent particles. There is little evidence within the present study that vacancies are involved in any way in the formation process of micropores. It was clarified that the growth behavior of micropores is dominated by Ostwald ripening. Hydrogen trapped in all the micropores was estimated, assuming a) that a thermal equilibrium exists between surface tension and b) that the internal gas pressure in the micropores obeys the ideal gas law. Unlike macroscopic measurements such as the thermal desorption spectroscopy, this study would be the first direct and local approach to estimate hydrogen trapped at micropores. Combining with
some theoretical considerations on hydrogen trapping, the hydrogen trapped at the other trap site was also estimated. It is noteworthy that about 53 % of hydrogen is trapped within micropores in the Al-Mg alloy with low hydrogen content, while although the total hydrogen content was similar, it was less than 7 % in pure aluminum. This difference is attributable to the lack of hydrogen precipitation sites in pure aluminium. Hydrogen partitioning to the normal interstitial lattice and other trap sites was also analyzed, revealing that micropores are the major hydrogen trap site in the Al-Mg alloy with low hydrogen content. It can be inferred that excess hydrogen in solid aluminum is predominantly partitioned to the micropores, and that occupancies of other trap sites are elevated with further increases in hydrogen concentration. It has been also clarified that the rate of trap site occupancies in dislocations and grain boundaries is very high, although the absolute amount of hydrogen trapped at grain boundaries is one to two orders of magnitude smaller than at the other trap sites.

Acknowledgements

The synchrotron radiation experiments were performed with the approval of JASRI through proposal numbers 2006A1056, 2007A1618 and 2007B1078. This work was undertaken as part of a Grant-in-Aid for Scientific Research (A) from the JSPS, as subject No. 20246102. The authors thank Dr. Katsumi Koyama for analyzing the hydrogen content of the materials used. One of the authors (HT) is also grateful for the support of the Light Metal Educational Foundation.

References


[34] Talbot DEJ, Granger DA. J Inst Metals 1963-64;92:290-297


[38] Markworth AJ. Metall Trans 1973;4:2651

Met 2008; in press.

[40] Trinkaus H. Radiation Effects 1983;78:189-211


[42] Baba Y. Proc ICSMA, Supplement to Trans JIM 9 1968;: 356


Caption list

Fig. 1 Explanation for detecting micropores heterogeneously nucleated on intermetallic compound particles. Each pore is expanded one to thirteen voxels in 3D. (a) 2-D tomographic slice of a reconstructed volume, (b) segmented pores and particles and (c) pore image after the expansion. Here, pore C is identified to be isolated, and pores A and B are identified to nucleate on particles P1~P3.

Fig. 2 3-D perspective views of micropores in alloy HH in various heat-treatment conditions. Note that only micropores were extracted and shown here.
Fig. 3 3-D perspective views of micropores in alloy MH in two heat-treatment conditions. Note that only micropores were extracted and shown here.

Fig. 4 3-D perspective views of micropores in alloy LH in two heat-treatment conditions. Note that only micropores were extracted and shown here.

Fig. 5 3-D perspective views of micro-pores in 5N-Al in two heat-treatment conditions. Note that only micropores were extracted and shown here.

Fig. 6 Enlarged 3-D perspective views of micropores in alloy MH and 5N-Al in various heat-treatment conditions. Note that only micropores were extracted and shown here.

Fig. 7 3-D perspective view of micropores in alloy HH observed after being exposed at 703 K for 18 h. The imaging-type microtomography set-up was used to obtain this ultra-high resolution image. Note that submicrometer micro-pore is not seen in the region.

Fig. 8 Variations in average diameter of micropores in alloys LH, MH, HH and 5N-Al during exposure at 823 K.

Fig. 9 Size distributions of micropores in alloys (a) LH, (b) MH, (c) HH and (d) 5N-Al in various heat-treatment conditions.

Fig. 10 Variations in volume fraction of micropores in alloys LH, MH, HH and 5N-Al during exposure at 823 K.

Fig. 11 Ratio of micropores heterogeneously nucleated on particles. The extent of dilation is varied
to see the effects of the Fresnel diffraction.

Table 1 Average size and number density of micropores together with total hydrogen content.

Table 2 Size ratio $d_{\text{max}} / d$ in alloys LH, MH and HH in various heat-treatment conditions.

Table 3 Measured hydrogen content together with the estimated content of hydrogen trapped within micropores.

Table 4 Microstructural parameters for hydrogen trapping sites; micropores, grain boundary and dislocation. Table 5 Estimated trap site coverage and the amount of hydrogen at each trap site.
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Fig. 2 3-D perspective views of micropores in alloy HH in various heat-treatment conditions. Note that only micropores were extracted and shown here.
(a) After homogenization at 703 K for 18 h

(b) 823 K - 90 h after (a)

Fig. 3 3-D perspective views of micropores in alloy MH in two heat-treatment conditions. Note that only micropores were extracted and shown here.
Fig. 4 3-D perspective views of micropores in alloy LH in two heat-treatment conditions. Note that only micropores were extracted and shown here.
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Table 1 Average size and number density of micropores together with total hydrogen content.

<table>
<thead>
<tr>
<th></th>
<th>Average diameter of micropores, $d / \mu m$</th>
<th>Number density of micropores, $\rho_p / \times 10^{12} m^{-3}$</th>
<th>Total H content, $M_H / ml/100g Al$</th>
</tr>
</thead>
<tbody>
<tr>
<td>HH (As-homogeneized)</td>
<td>3.6</td>
<td>54.7</td>
<td>5.12</td>
</tr>
<tr>
<td>MH (As-homogeneized)</td>
<td>3.5</td>
<td>50.7</td>
<td>4.94</td>
</tr>
<tr>
<td>LH (As-homogeneized)</td>
<td>3.6</td>
<td>16.1</td>
<td>0.30</td>
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Table 2  Size ratio $d_{\text{max}} / \bar{d}$ in alloys LH, MH and HH in various heat-treatment conditions.

<table>
<thead>
<tr>
<th></th>
<th>HH</th>
<th>MH</th>
<th>LH</th>
</tr>
</thead>
<tbody>
<tr>
<td>As-cast</td>
<td>1.9</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>As-homogeneized (AH)</td>
<td>2.2</td>
<td>2.2</td>
<td>2.3</td>
</tr>
<tr>
<td>5 h at 823 K after AH</td>
<td>2.0</td>
<td>2.9</td>
<td>2.6</td>
</tr>
<tr>
<td>10 h at 823 K after AH</td>
<td>2.5</td>
<td>3.2</td>
<td>3.3</td>
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Table 3: Measured hydrogen content together with the estimated content of hydrogen trapped within micropores.

<table>
<thead>
<tr>
<th>Material</th>
<th>H$<em>2$ in micropores, $M</em>{H}^{\text{pore}}$</th>
<th>Total H content, $M_{H}$ (ml/100gAl)</th>
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</thead>
<tbody>
<tr>
<td>HH As-cast</td>
<td>0.13</td>
<td>5.12</td>
</tr>
<tr>
<td>Al-5%Mg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HH As-cast</td>
<td>0.16</td>
<td>0.30</td>
</tr>
<tr>
<td>MH As-homogenized</td>
<td>0.45</td>
<td>4.94</td>
</tr>
<tr>
<td>LH As-homogenized</td>
<td>0.02</td>
<td>0.29</td>
</tr>
<tr>
<td>5N-Al</td>
<td>As-cast</td>
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</tr>
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Table 4 Microstructural parameters for hydrogen trapping sites; micropores, grain boundary and dislocation.

<table>
<thead>
<tr>
<th>Material</th>
<th>Volume fraction of micropores, $V_f$ (%)</th>
<th>Grain size, $\mu$m</th>
<th>Dislocation density, $\times 10^{13} \text{m}^{-2}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al-5Mg</td>
<td>HH As-cast</td>
<td>0.071</td>
<td>205.8</td>
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<tr>
<td></td>
<td>MH As-homogenized</td>
<td>0.229</td>
<td>360.4</td>
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<tr>
<td></td>
<td>LH As-homogenized</td>
<td>0.150</td>
<td>469.6</td>
</tr>
<tr>
<td>5N-Al</td>
<td>As-cast</td>
<td>0.024</td>
<td>-</td>
</tr>
</tbody>
</table>
Table 5 Estimated trap site coverage and the amount of hydrogen at each trap site.

<table>
<thead>
<tr>
<th>Materials</th>
<th>Trap sites</th>
<th>Trap density, $N_T$ / $\times 10^6$ sites/m$^3$</th>
<th>Trapp site occupancy, $\theta_T$</th>
<th>Trapped hydrogen, $C_T$/atoms H/m$^3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>HH As-cast</td>
<td>Interstitial</td>
<td>$5.85 \times 10^{22} = N_L$</td>
<td>$8.88 \times 10^{-5} = \theta_L$</td>
<td>$5.20 \times 10^{18}$</td>
</tr>
<tr>
<td></td>
<td>Solute Mg atom</td>
<td>$1.23 \times 10^{21}$</td>
<td>$9.21 \times 10^{-4}$</td>
<td>$1.33 \times 10^{18}$</td>
</tr>
<tr>
<td></td>
<td>Dislocation</td>
<td>$6.89 \times 10^{17}$</td>
<td>0.970</td>
<td>$6.68 \times 10^{17}$</td>
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<tr>
<td></td>
<td>Grain boundary</td>
<td>$8.47 \times 10^{16}$</td>
<td>0.992</td>
<td>$8.40 \times 10^{16}$</td>
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<td></td>
<td>Micro pore</td>
<td>-</td>
<td>-</td>
<td>$1.88 \times 10^{17}$</td>
</tr>
<tr>
<td>MH As-homogenized</td>
<td>Interstitial</td>
<td>$5.85 \times 10^{22} = N_L$</td>
<td>$8.46 \times 10^{-5} = \theta_L$</td>
<td>$4.95 \times 10^{18}$</td>
</tr>
<tr>
<td></td>
<td>Solute Mg atom</td>
<td>$1.23 \times 10^{21}$</td>
<td>$8.78 \times 10^{-4}$</td>
<td>$1.08 \times 10^{18}$</td>
</tr>
<tr>
<td></td>
<td>Dislocation</td>
<td>$2.54 \times 10^{17}$</td>
<td>0.968</td>
<td>$2.46 \times 10^{17}$</td>
</tr>
<tr>
<td></td>
<td>Grain boundary</td>
<td>$4.84 \times 10^{16}$</td>
<td>0.991</td>
<td>$4.80 \times 10^{16}$</td>
</tr>
<tr>
<td></td>
<td>Micro pore</td>
<td>-</td>
<td>-</td>
<td>$6.34 \times 10^{17}$</td>
</tr>
<tr>
<td>LH As-homogenized</td>
<td>Interstitial</td>
<td>$5.85 \times 10^{22} = N_L$</td>
<td>$1.55 \times 10^{-6} = \theta_L$</td>
<td>$9.07 \times 10^{16}$</td>
</tr>
<tr>
<td></td>
<td>Solute Mg atom</td>
<td>$1.23 \times 10^{21}$</td>
<td>$1.61 \times 10^{-5}$</td>
<td>$1.98 \times 10^{16}$</td>
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<td></td>
<td>Dislocation</td>
<td>$1.80 \times 10^{17}$</td>
<td>0.357</td>
<td>$6.43 \times 10^{16}$</td>
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<tr>
<td></td>
<td>Grain boundary</td>
<td>$3.71 \times 10^{16}$</td>
<td>0.677</td>
<td>$2.51 \times 10^{16}$</td>
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<tr>
<td></td>
<td>Micro pore</td>
<td>-</td>
<td>-</td>
<td>$2.23 \times 10^{17}$</td>
</tr>
<tr>
<td>5N-Al As-cast</td>
<td>Interstitial</td>
<td>$6.03 \times 10^{22} = N_L$</td>
<td>$4.09 \times 10^{-6} = \theta_L$</td>
<td>$2.46 \times 10^{17}$</td>
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<tr>
<td></td>
<td>Dislocation</td>
<td>$2.26 \times 10^{17}$</td>
<td>0.594</td>
<td>$1.34 \times 10^{17}$</td>
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<tr>
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<td>Micro pore</td>
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