Early pore formation in aluminium foams studied by synchrotron-based micro-tomography and 3D image analysis

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Abstract

The mechanisms behind pore formation in aluminium alloys that are in commercial use for making foam components are still not understood. It is accepted that they play a key role for the quality of the evolving pore structure. In order to shed light on early pore formation, aluminium foams in low expansion stages made by the powder-compact melting process were investigated by synchrotron-based hard X-ray micro-tomography and quantitative image analysis. The calculated spatial correlation functions between the gas source (TiH₂ particles) and the nucleating pores show that the location of pore formation and the spatial gas source distribution for AlSi7 and Al-Si-Cu alloys are essentially independent. Therefore, in the alloys investigated, the position of the melt formation is found to determine the pore nucleation rather than the spatial distribution of TiH₂. This behaviour explains the known positive effect on the pore structure achieved by adding copper.

Keywords: micro-tomography, image analysis, CT, aluminium foam, aluminium alloy
I. INTRODUCTION

Replacement of engineering structures, e.g. in automobiles, by lightweight aluminium foam has been predicted several times in the past, see Ref. [1] or more recently [2], but many years after these predictions the market for aluminium foam is still small [3]. One reason for this is that the available aluminium alloy foams still do not have the desired properties, partially owing to the fact that during foaming the evolving cellular structure contains more heterogeneities than would be expected from purely statistical scatter and more than are acceptable in practice. Therefore, research has been carried out aiming at improving the uniformity of cell structure. To understand these irregularities, it is necessary to study the nucleation of pores and their evolution in early stages of foam evolution.

This paper solely focuses on foaming of aluminium alloys based on metal powders: metal and blowing agent powders are mixed and compacted, after which the compacted mixture is heated to a temperature high enough to ensure the melting and foaming of the alloy [4]. The blowing agent particles – usually TiH$_2$ – start to release gas (hydrogen) that leads to pore nucleation and subsequent pore growth. Mostly, highly expanded foams are studied whilst not much work has been done on early stages of foam expansion when the solid precursor is just starting to increase its volume.

It was noticed early in metal foam development that the different alloys in use showed different foaming behaviour, pore shapes and size distributions. Two of the alloys used for the production of metal foam components at that time – the wrought alloy 6061 and the casting alloy AlSi7 – were studied by Duarte et al. [5], Baumgärtner et al. [6] and Mosler et al. [7]. Especially for AlSi7, extremely anisotropic foaming behaviour was found. Bellmann et al. [8] defined the problems, suggested suitable methods for investigating early stages of foaming and distinguished between two types for pore formation in metal foams:

**type-I:** local pore formation occurring at the location of individual blowing agent particles,

**type-II:** non-local formation at other locations in the precursor material.

In a further step, AlSi7 was studied in more detail using synchrotron X-ray microtomography to determine non-destructively the size distribution of the pores in early stages as well as to analyse their morphology and connectivity [9]. A further advance was the description of the different pore formation mechanisms in both AlSi7 and 6061 alloy by employing micro-tomography with higher spatial resolution, and the observation not only
of different pore morphologies but also of different correlations between the positions of the TiH$_2$ particles and the pore volume \cite{10, 11}. While in 6061 the TiH$_2$-particles were preferentially found near the inner pore surfaces, they were distributed almost randomly in AlSi7. It was concluded that in 6061 the pores are locally inflated around blowing agent particles (type-I), whereas in AlSi7 the evolving hydrogen migrates to weak points in the matrix and creates pores there non-locally (type-II). The suspicion that these weak points are defined by the Si particles in the pressed powder mixture (Al+Si) \cite{7} was confirmed later by holographic tomography \cite{12}. A further continuation of the work was undertaken subsequently and came in two parts. First, as the method of determining spatial correlations was based on standard morphological transformations, leading to a coarse grid of analyzable distance values around the pore volume, the mathematical routines were reviewed. A new approach to access the distance values around the pore space was implemented and applied to 6061 and AlSi7 alloy foam in early stages, thus demonstrating the new method \cite{13}. Second, and this is the topic of the present paper, new high resolution data sets were analysed using the refined algorithm. Moreover, a number of new samples were manufactured for imaging with lower resolution but greater field of view in order to increase the statistical significance of the conclusions derived. As in the meantime alloys from the alloy system Al-Si-Cu had emerged as preferred materials for industrially produced foam sandwich cores \cite{14, 15}, characterised by low melting points and more uniform pore size distributions, two alloys from this system were included in this study.

II. METHODS

This paragraph describes the preparation of the different specimens that were analysed as well as the experimental and mathematical approaches used to investigate early pore formation in aluminium foams.

A. Sample preparation

The specimens investigated were made by foaming powder compacts, i.e. blends of elemental metal powders and TiH$_2$ powder. Table I summarises the processing parameters used for the investigations in this work, namely composition, blowing agent content, pressing temperatures and pressures and foaming temperature.
Some of the samples were prepared at the Fraunhofer Institute (IFAM), Bremen (Germany) and were cut to dimensions that the foams could be imaged with synchrotron microtomography (\(S\mu\)CT) employing a high spatial resolution around 1 \(\mu\)m. The parameters that were varied included the alloy composition (AlSi7 vs. AlSi6Cu4), TiH\(_2\) content (0.5 or 1 wt.\%) and compaction temperature (200°C and 450°C). The lower compaction temperature was known to lead to insufficient compaction and expansion, while the higher temperature produces a virtually dense precursor material with a residual porosity \(\leq 0.75\%\), see Ref. [5]. The results presented in this work are restricted to those samples series, where at least four different expansion stages (see below) could be successfully investigated by \(S\mu\)CT. The samples in this series will be called IFAM-AlSi (1 and 2) and IFAM-AlSiCu. In all cases, the foaming process was triggered by heating the obtained precursor material in a pre-heated furnace (600°C). Quenching the samples after pre-defined varying dwell times in the furnace conserves pore structures in various expansion states [10, 16]. The highest temperatures reached during foaming can be estimated by comparison with measurements on samples of the same composition foamed in a previous work [5]. Depending on the expansion stage this temperature ranges from 576°C to 585°C.

A second series of samples (TUB-AlSi and TUB-AlSiCu) was later prepared at Berlin University of Technology (TUB, Germany) in order to be investigated with \(S\mu\)CT employing spatial resolutions around 10 \(\mu\)m and correspondingly larger sample volumes and statistics. The production procedure was slightly improved compared to the manufacture of the IFAM samples, as higher compaction pressures could be applied. The foaming device used allowed more precise temperature and time control. Samples were quenched after expansion to different expansion levels in ice water [17]. The highest temperature reached in these experiments was measured by a thermocouple inserted into the sample for the Al-Si-Cu alloy (unlike in the previous IFAM experiments) and was about 525°C for AlSi6Cu10, or slightly higher. For each parameter set of the TUB series, samples were produced by using non-treated TiH\(_2\) and a heat-treated TiH\(_2\) (3 h at 480°C in air) that helps to avoid premature hydrogen evolution [18].

Samples for \(S\mu\)CT were prepared by electrodischarge machining (IFAM) as well as by sawing (TUB). The samples consisted of rods with a square cross section of 1×1 mm\(^2\) for the high resolution experiments and around 6×6 mm\(^2\) for the lower resolution.
B. Synchrotron micro-tomography

Experiments were carried out using the micro-tomography facilities at the European Synchrotron Radiation Facility’s ID19 (ESRF) and the BAMline located at the light source BESSY at the Helmholtz Centre Berlin [19, 20]. Owing to the availability of higher photon flux density we scanned selected samples with highest resolution at ESRF and performed scans of a broader range of larger specimens with moderate resolutions at BESSY. Both facilities employ conventional high precision sample manipulators for the alignment and rotation of the specimen before as well as during the tomography scan. For high resolution pixel detectors, the indirect approach based on the concepts of W. Hartmann et al. as well as F. Busch and U. Bonse is used [21, 22]. Here, a scintillating screen (powder or single crystal) converts X-rays into visible light. This luminescence image is captured by coupling a CCD chip optically with the scintillator. The indirect detection concept using thin single crystal film scintillators makes it possible to reach resolutions in the sub-micrometre range [23].

The high resolution scans at ESRF were done utilizing a pixel size of 0.7 μm, leading to a true spatial resolution around 1.5 μm and a field of view of 1.4×1.4 mm² (due to the 2048×2048 pixel ’FReLoN type 2000’ CCD camera used [24]). The maximal sample volume to be analysed with this setting would be (1.4 mm)³. The synchrotron beam was monochromatised to 17 keV which allowed for easy subsequent image analysis (see next section). 1400 projection images were recorded during 180° scans in order to ensure high quality reconstruction of the tomographic images [19].

Tomographic scans with a moderate resolution of around 12 μm (10% of the modulation transfer function, determined via a knife-edge scan) were performed at BESSY. As CCD camera, a Princeton Instruments, type VersArray:2048B was used, with an effective pixel size of 3.5 μm (2048×2048 pixels, effective 7×7 mm² field of view, (7 mm)³ maximal sample volume to be analysed). Depending on the sample dimensions, the synchrotron beam was monochromatised to photon energies between 20 keV and 25 keV. Typically, 720 to 900 projection images were recorded during 180° scans [25].

Tomographic reconstructions were carried out using the conventional filtered-backprojection algorithm. As software we chose the software package PyHST developed by the SciSoft group at ESRF [26] as it allows for the use of self-written filters as plugins. In order to improve the reconstruction quality, two filters – a median filter to reduce noise and
a ring artefact filter based on a bandpass applied to the sinograms prior to backprojection – were coded as plugins in cooperation with the SciSoft group, see e.g. Ref. 27.

C. 3D image analysis

Synchrotron micro-tomography delivers volume data sets as 3D matrix of numbers that can be interpreted as grey-scales and thus appear as images. When working in absorption contrast mode, each matrix value represents the local attenuation of a voxel (acronym for ‘volume picture element’) which is related to the density and the attenuation coefficient of the material [28]. The grey-scale information can be used to identify and subsequently separate different material phases inside the volume images, provided that a significant difference in the attenuation behaviour of the individual phases is given. For example, one can distinguish between aluminium and TiH$_2$ or Cu, while there is almost no absorption contrast between aluminium and silicon as, e.g., at an X-ray energy of 17 keV the attenuation coefficients are: 13.5 cm$^{-1}$ (Al), 15.3 cm$^{-1}$ (Si), 110.8 cm$^{-1}$ (Ti), 471.3 cm$^{-1}$ (Cu) [29]. In order to separate the different phases into independent Boolean images, where voxels belonging to the selected material are given the foreground value ’1’ and the remains are set to the background value ’0’, we use a grey-scale threshold hysteresis in combination with a region growing algorithm [30]. Here, two thresholds are used to define a range in the image’s histogram where voxels definitively belong to one phase, e.g. the pores. A second pair of thresholds defines a larger range in the histogram, selecting voxels that might belong to this phase. In successive stages the algorithm extends the voxels already identified as belonging to one phase, e.g. pores, by those that, a) contain a grey value in the second range defined and, b) are direct neighbours of the already identified phase. The required parameters are optimised manually in order to acquire Boolean images with minimal amounts of artefacts and noise. For some data sets, the morphological transformations open and closure are applied to the Boolean images to further reduce noise and artefacts [30, 31].

The aim of our image analysis is to identify spatial correlations between the positions of the blowing agent particles and the positions of the pores inside aluminium foams at different expansion stages. The analysis is based on the Boolean images introduced above, here specifically the Boolean images of the pore space and the blowing agent particles. Our algorithm dilates the Boolean pore structure, creates the difference between the dilated
and the original Boolean pore image and then calculates the density of the blowing agent particles in this differential volume, see Fig. 1. This procedure is repeated in successive steps until the dilated pore volume encompasses 99% of the total image volume. If the blowing agent density is high during the first dilation steps (close to the pore surface) and then decreases with ongoing dilation, there is a correlation between the pore surfaces and the blowing agent particles, identifying type-I behaviour, as the majority of the blowing agent particles is located close the the pores. If the density remains constant for all dilation steps then there is no spatial correlation between pores and blowing agent, indicating type-II behaviour, cf. e.g. Fig. 7.

Sometimes the density varies in an irregular way or even slightly grows as the distance to the pore surface increases. There is no reason to believe that TiH$_2$ particles should be preferentially found at a non-zero distance from the pores, at least none we are aware of. One might speculate that such apparent 'negative correlations' between pores and TiH$_2$ densities stem from coarse spherical TiH$_2$ particles that are situated at the interfaces and have their maximum cross section at a distance of half their diameter of, say, 20 μm. We prefer the viewpoint that such variations are unphysical and class them as 'no correlation'. The presence of various differently absorbing phases around the pores could give rise to spurious contributions to the distribution function. This viewpoint is supported by the observation that these 'negative correlations' occur in intermediate stages of evolution, i.e. between 'unfoamed' and 'fully molten'.

Care has to be taken to treat the interfaces between the pores and the metal. To avoid possible artefacts, a layer around each pore space of the thickness of the FWHM of the detector’s point spread function is discarded from the analysed density data. A more detailed description of the algorithm including a sound mathematical discourse has been published elsewhere [13].

III. RESULTS

In this section, the 3D image analysis of volume data sets acquired at ESRF and BESSY will be presented in two parts, each dedicated to one of the two alloy families investigated. The high resolution scans of selected samples are introduced first, followed by the analysis of the intermediate resolution scans that are greater in number and underline the statistical
relevance of our observations.

A. AlSi7 foams

The AlSi7 samples labelled 'IFAM' were scanned with high resolution of around 1.5 μm. The four specimens of the series IFAM-AlSi-1 compacted at 450°C (60 s, 75 s, 90 s and 120 s dwell-time in furnace) show porosities of 0.31%, 0.43%, 0.46% and 7.43% as determined by 3D image analysis, see Fig. 2. The onset of foaming from the still dense material to the pore-containing state is therefore fairly immediate, taking place between 90 s and 120 s after heating begins. Tomographic slices of the five IFAM-AlSi7-2 samples compacted at 200°C (0 s, 60 s, 75 s, 90 s and 120 s dwell-time in furnace) are displayed in Fig. 3. The respective (initial) porosities of 15.8%, 11.4%, 15.2%, 18.3% and 14.5% are significantly higher and more constant compared to the first four due to the lower compaction efficiency at 200°C. Already the unfoamed precursor contains porosity when this compaction temperature – generally regarded as insufficient for making foams – is applied, whereas the 'normal' compaction temperature leads to a virtually dense precursor.

After the creation of Boolean images the correlation analysis algorithm was applied, the results of which are displayed in Fig. 4 for the specimens compacted at the higher temperature. The quantity calculated is the blowing agent particle density (normalised to the mean density in the foam matrix) as a function of distance to the closest pore surface. For the unfoamed precursor, no value can be given since this sample is nearly dense and no meaningful pore volume exists to which the TiH$_2$ density can be related. The range for which the correlation function can be given varies in Fig. 4 and in the subsequent figures in this paper. For geometrical reasons, the available pore space can be dilated more before meeting adjacent pores in dense samples. For the first 3 stages of foaming – all still associated with porosities well below 1% – the density function does not show any sign of preferential positioning of TiH$_2$ particles with respect to the surface of the pores. This means that the tiny first pores nucleate at any distance from the TiH$_2$ particles. Only in the last sample of the series (120 s dwell time, 7.4% porosity) is a high density visible close to the pore space (spatial correlation) that then decays as the distance from the nearest pore surface increases, i.e. in this sample most of the pore volume is close to a blowing agent particle.

For 90 s dwell time and also for some more data sets in the rest of the article, there is
even a slight increase of density as the distance to the pore space increases. Such increases
do not correlate with any direct observation of the foams by microscopy and, together with
minor variations as in the first two curves, will be considered irrelevant in the following,
although no clear reason for these variations could be identified.

The correlation has also been observed for the other series, see Fig. 5, where the com-
paction temperature was deliberately set too low. The difference here is that the unfoamed
state already contains 15% porosity and therefore can be used for the correlation analysis.
The density curve is almost horizontal for this sample as it should be for a random powder
mixture, and the fluctuations still observed provide an estimate for the error of the analy-
sis. Heating for 60 s and 75 s does not change the picture very much. Except for a slight
indication of the artefact already mentioned there is no sign of big changes in the distribu-
tion function. No sign is found that TiH₂ particles preferentially segregate near the pore
surfaces. Only after longer heat treatments (90 s and 120 s in the furnace), the density of
the blowing agent particles close to the pore space increases at the cost of the locations far
away from the surface, i.e TiH₂ is segregated to the surface.

In order to validate these findings based on a small number of low-volume samples we
investigated further AlSi7 foam samples, the TUB-AlSi series. These larger samples were
imaged with a lower resolution (corresponding to a larger pixel size and, hence, a larger
field of view of the detector), while the resolution was still sufficient to distinguish between
correlated and uncorrelated TiH₂ distributions. Of the 23 samples in this series, 12 can
be considered as early stages with porosities ranging from 1% to 6%, while the remaining
11 samples are in more expanded foaming stages with porosities between 15% and 50%.
Typical tomographic slices can be seen in Fig. 6. Our analysis revealed that in 11 out
of these 12 AlSi7 foam specimens in early stages no segregation of TiH₂ particles to the
pore surfaces can be observed, while, on the contrary, all of the more expanded AlSi7 foam
samples showed a clear segregation. In other words, in the early stages of foam expansion of
this alloy the pores are generated and inflated anywhere in the sample without necessarily
being close to the gas-generating TiH₂ particles (type-II), whereas in later stages most of
the TiH₂ particles are found very close to the pores’ surfaces (type-I).

Two typical correlation plots from the TUB samples showing the two limiting cases can
be found in Fig. 7. In the early foaming stage corresponding to 1.5% porosity, the density
of the blowing agent particles is only weakly dependent on the distance to the pore space,
indicating the absence of segregation. In the more extended stage – 28% porosity – the density distribution of the blowing agent particles shows a clear spatial correlation between TiH$_2$ and the pore space, see also Fig. 6 and 8. The increase in the curve corresponding to the lower density (from 0.7 to 1.2) is again seen as an artefact and is indeed markedly lower than the decrease in density for the more expanded sample from 1.85 to 0.35. For a complete listing of tomographic slices and correlation plots see 'Appendix D' of Ref. 25.

The nature of type-II pores in AlSi7 foam can be better understood by taking a look at cut, polished, and etched specimens by light microscopy. Etching makes it possible to distinguish between former powder particles more easily. In Fig. 8(a) a metallographic image of an AlSi7 foam from the TUB batch in the early stage – 1.5% porosity – is shown. In the vicinity of the TiH$_2$ particles pores can only be found if silicon particles are also in the neighbourhood, while the majority of pores is located directly adjacent to silicon particles or silicon particle clusters, i.e. the silicon particles define the positions of the pores. In contrast, in an extended stage with 28% porosity, see Fig. 8(b), most of the then much larger pores are located around or close to blowing agent particles.

B. AlSi6Cu4 and AlSiCu10 foams

Four IFAM-AlSiCu specimens with the nominal composition AlSi6Cu4 + 0.5 wt.% TiH$_2$ were imaged with the high-resolution (around 1.5 μm) setup at ESRF. The respective dwell-times in the pre-heated furnace for these samples were 0 s, 60 s, 90 s and 120 s, i.e. the precursor and three foamed states were investigated. Tomographic slices of the 0 s, the 90 s and the 120 s sample are displayed in Fig. 9. Because of the high stopping power of pure copper the volume images of the 0 s and the 60 s samples contain so many artefacts in the form of white streaks that these images cannot be interpreted more than qualitatively. After 90 s, see Fig. 9(b), obviously most of the copper has been dissolved in the melting aluminium. At the positions of former copper particles a moderately absorbing mixed-phase and first pores can now be seen. In addition, several TiH$_2$ particles in the slice can be identified between the positions of former copper particles. Pores only seem to be inflated at these positions appearing in light grey, i.e. the copper particles define the positions of the pores. The volume image of the specimen after 120 s of heat-treatment was the only one that was artefact-free, see Fig. 9(c), and allowed for a correlation analysis.
We separated all highly absorbing material (TiH₂ and Al-Si-Cu mixed phase, about 0.32% mass fraction / 0.20% volume fraction) into one Boolean image, the pore space (6.1% porosity in this sample) into another. The result of the correlation analysis is plotted in Fig. 10. Two peaks can be distinguished: one close to the pore surface and a second one at a distance of approximately 29 μm from the pore surface. From the tomographic slices in Fig. 9 we identify the first maximum with the mixed-phase (Al-Si-Cu), while the second maximum is mainly caused by the TiH₂ particles in the foam. While being more difficult to interpret, the tomographic scans of the IFAM AlSi6Cu4 + 0.5 wt.% TiH₂ samples point towards type-II behaviour, i.e. pores are preferentially surrounded by Cu-rich aluminium melt, while the blowing agent particles can be at some distance from the pores and in fact even preferentially around 30 μm away from the gas/metal interfaces. Unlike AlSi7, the interpretation of these correlation curves requires some previous knowledge taken from metallographic sections – see below – because the value for the X-ray absorption coefficient for the mixed phase (Al-Si-Cu) can be very similar to that of TiH₂.

In analogy to the previous section, we scanned a larger batch of samples at moderate resolutions and higher sample volumes at BESSY, the TUB-AlSiCu series. The best and most consistent results were obtained for an alloy with an increased copper content that will be presented here. Altogether, volume images of 16 AlSi6Cu10 + 1.0 wt.% TiH₂ could be acquired and analysed. In 5 out of 16 samples the porosity is between 4% and 11% (early stage) and the TiH₂ contrast high enough so that a correlation analysis is possible. None of these 5 samples shows a spatial correlation in the sense of segregation of the blowing agent to the pore surface, thus indicating type-II pores. In Fig. 11 an exemplary plot of the correlation analysis from this sample series can be found. Here, the Al-Si-Cu peak is not visible as in Fig. 10, possibly due to the lower resolution of the imaging system used and the higher Cu content in this alloy that increases the signal from the Cu-rich phase. The density of highly absorbing material is nearly constant throughout the volume and no spatial correlation between this material and the pore space can be found. Again, for a complete listing of tomographic slices and correlation plots, see ’Appendix D’ of Ref. 25.

As for AlSi7, metallographic images are used to get additional information. In the image showing the raw precursor material, see Fig. 12(a), the round copper particles, silicon particles as well as aluminium are visible. After 90 s of heat-treatment, see Fig. 12(b), the copper particles have dissolved in the evolving melt and in their positions the first pores
have now appeared. We performed EDX analysis in the vicinity of such an early pore in the IFAM sample, heat treated for 90 s, see Fig. 13. The high content of copper found close to and inside the pore confirms the observation that the pore surfaces are decorated with copper-rich eutectic melt. The metallographic image of the IFAM sample heat-treated for 120 s given in Fig. 12(c) illustrates the type-II nature of the pores in AlSi6Cu4 foam – early pores inflate at the positions of former copper particles, sometimes far away from the TiH2 particles.

Although the number of tomographic data sets that could be used for correlation analysis was much lower for this alloy than for AlSi7 – mainly because we could not identify the blowing agent with certainty in many of the samples – we find firm evidence that pores are of type-II in this foam too, but no transition to type-I is observed in later stages of foaming.

IV. DISCUSSION

The experimental evidence provided by the work presented here confirmed earlier observations that pore formation in metallic foams varies with the type of aluminium alloy chosen. A new alloy type – ternary Al-Si-Cu alloys – was investigated and yet another pore formation type found. The simple direct pore inflation mechanism identified in alloy 6061 alloys by Helfen et al. [10] and explained schematically in Fig. 14(a) was not found for any of the alloys studied here.

For AlSi7, the earlier observation that pores are not formed directly around the blowing agent particles as would be expected, is confirmed by the correlation analysis (Figs. 4 and 5) and the two-dimensional metallographic sections. The idea that most pores form in the vicinity of silicon particles is supported by metallography (Fig. 8 and Ref. [7]) – but only on a non-representative basis – and by the holotomographic correlation analysis presented by Helfen et al. [12]; but only two samples were studied there. The combined evidence, however, now makes it very likely that the reason for this pore formation mechanism is that pores open at the weakest point in the metallic matrix which in this case is the Al/Si interface. As gas is being generated by the TiH2 particles, this gas can be transported to other parts of the foamable precursor, e.g. along the boundaries between the former powder particles, or through microchannels in the oxide films around these particles etc. There it can create a gas pressure even at locations some tens of μm away from the blowing agent.
particle. As soon as the pressure exceeds the delamination threshold of the Al/Si interface, which is significantly reduced due to local melting of Al/Si eutectic, a pore opens there.

A new finding of the current work is that there is a transition from type-II to type-I behaviour during the evolution from an early stage to a later one which, however, is still far away from full expansion. The exact reason for this transition is not clear. Possibly, in the course of the formation of pores that in AlSi7 are very jagged and often crack-like, a system of dense interconnected cracks evolves that then propagates through the entire sample. Helfen et al. [9] have found that this crack system can be very extensive and interconnects a large volume of the sample. If, in addition, the cracks propagate preferentially along the TiH₂ particles and not through the bulk of aluminium, most of the TiH₂ particles would then be very close to the next open volume and create the type-I behaviour observed. This can only be explained by an expansion of the open void space that eventually reaches all TiH₂ particles (visualised in Fig. 14(b)).

This idea can also provide an explanation for the poor foamability of insufficiently pressed powders such as the ones used in Figs. 3 and 5. In these samples compacted at 200°C, the change from type-I to type-II behaviour takes place earlier than in the samples compacted at 450°C, see Fig. 2, and the correlation subsequent is much more pronounced. As the inter-particle bonding is weaker in the insufficiently pressed sample, the opening and propagation of cracks is more easily initiated by the internal pressure built up by the evolving blowing gas. The blowing agent particles are then connected to the large network of cracks and pores from an earlier stage which in turn leads to a more serious loss of blowing gas.

The behaviour of the Al-Si-Cu alloy is different from that of the AlSi7 alloy. Both alloys share the feature that pore formation is not at the blowing agent site – i.e. type-II – but the weakest point in the matrix, which is not the Al/Si interface this time but the ternary eutectic melt that is formed at much lower temperatures than the binary eutectic in AlSi7, namely at 525°C compared to 577°C. As the sample expands further, the evolving liquid continuously wets the aluminium grains and significantly reduces the excessive formation of cracks [32]. Therefore, the pores can continuously grow in a fairly spherical way within the increasingly large pool of liquid metal.

In a wider context, these findings explain why Al-Si-Cu foams actually exhibit better foaming behaviour than AlSi7 and have replaced AlSi7 as core materials for aluminium foam sandwich panels [15]. First of all, the blowing agent releases gas at temperatures at
which the matrix starts melting and then delivers gas into a pool of liquid where it can form round pores – governed by surface tension – instead of the gas forcing the still solid material apart and propagating cracks through the material as happens in AlSi7. The situation in AlSi7 is also more unfavourable because melting takes place at a higher temperature (577°C) than in Al-Si-Cu (525°C) and therefore the hydrogen pressure generated by the decomposing blowing agent is higher [18], [33]. Although such cracks round off after the metal has melted, as X-ray radiographic images show [34], they may lead to partial gas losses in the heating stage and give rise to the more heterogeneous pore morphology and size distribution generally observed in AlSi7 foams.

Looking into the future, the foaming behaviour of aluminium alloys could be further improved by tailoring the alloy composition so that either the ternary eutectic temperature is lowered – by further alloying elements – or the amount of melt immediately available after melting is increased by tuning the composition of the ternary alloy [35]. This would allow the blowing agent to feed gas into the evolving liquid continuously, thus keeping the pores spherical.

V. CONCLUSIONS

We confirmed a previous observation that AlSi7 foam develops type-II pores in the first stage, i.e. the pores form at the weakest links in the pressed powder, the Al/Si interfaces. An improved statistical analysis based on 27 samples and a quantitative evaluation based on correlation analysis has been given. A new finding is the change from type-II to type-I pores in the following stage of foaming in AlSi7.

In addition to Al-Si, two alloys of the common and industrially applied Al-Si-Cu alloy family haven been investigated. In these alloys, the pores first form along the liquid ternary eutectic around individual copper particles which is then the weakest part of the material. These pores are therefore of type-II and remain so even in later stages with porosities up to ≈ 10%.

The sequence of pore formation with rising temperature in three different alloys (including 6061 from the literature) is summarised in Fig. 14 and can be described as follows:

6061: hydrogen gas from blowing agent evolves $\rightarrow$ local pressure build-up around particles $\rightarrow$ melting of matrix $\rightarrow$ pore expansion around TiH$_2$ particles [10].
**AlSi7:** hydrogen gas from blowing agent evolves → debonding at Al/Si interfaces and local melting of Al/Si eutectic → hydrogen pressure expands pore space at Si particles → further opening of pore space that eventually reaches most of the TiH₂ particles → complete melting of matrix.

**Al-Si-Cu:** ternary eutectic melts → hydrogen gas from blowing agent evolves → pores nucleate along ternary melt pools and expand.

An interpretation of the favourable foaming characteristics of Al-Si-Cu alloys is now accessible and the criteria for selecting further alloys are clearer.

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[25] Rack A. Charakterisierung komplexer Materialsysteme mittels Synchrotron-Tomographie


APPENDIX A: TABLES

TABLE I: Overview of sample series investigated, including their foaming and imaging parameters.

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<tr>
<th>sample series</th>
<th>composition</th>
<th>blowing agent</th>
<th>compaction</th>
<th>foaming</th>
<th>image resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>IFAM-AISi-1</td>
<td>AlSi7</td>
<td>0.5 wt.% TiH2</td>
<td>450°C, 120 MPa</td>
<td>600°C</td>
<td>~ 1 μm</td>
</tr>
<tr>
<td>IFAM-AISi-2</td>
<td>AlSi7</td>
<td>1.0 wt.% TiH2</td>
<td>200°C, 120 MPa</td>
<td>600°C</td>
<td>~ 1 μm</td>
</tr>
<tr>
<td>IFAM-AISiCu</td>
<td>AlSi6Cu4</td>
<td>0.5 wt.% TiH2</td>
<td>450°C, 120 MPa</td>
<td>600°C</td>
<td>~ 1 μm</td>
</tr>
<tr>
<td>TUB-AISi</td>
<td>AlSi7</td>
<td>1.0 wt.% TiH2</td>
<td>450°C, 295 MPa</td>
<td>650°C</td>
<td>~ 10 μm</td>
</tr>
<tr>
<td>TUB-AISiCu</td>
<td>AlSi6Cu10</td>
<td>1.0 wt.% TiH2</td>
<td>400°C, 295 MPa</td>
<td>600°C</td>
<td>~ 10 μm</td>
</tr>
</tbody>
</table>
FIG. 1: Schematic representation of the correlation analysis applied, based on the segmentation of the grey scale image (left column) into Boolean images representing, from top to bottom of middle column, the pores, the metallic matrix, and the blowing agent plus – for Al-Si-Cu alloys only – highly absorbing parts of the metallic matrix. Only the latter and former are used for the final analysis (right column), whereas the Boolean image of the metallic matrix is not needed. The ranges given for the absorption coefficient $\mu$ are approximate and correspond to a photon energy of 17 keV.
FIG. 2: Tomographic slices of the IFAM-AISi-1 foam series (AlSi7+0.5wt.% TiH₂ hot compacted at 450°C) with pores in dark grey, metallic matrix in grey and blowing agent particles in white. Porosities with respective dwell-times in the furnace are, a) 0.31% (60 s), b) 0.43% (75 s), c) 0.46% (90 s), d) 7.4% (120 s). Orientation of compaction and foaming are perpendicular to the image plane. See also Fig. 4.
FIG. 3: Analogous to Fig. 2 for 200°C compaction temperature (IFAM-AISi-2 series). Porosities with respective dwell-times in the furnace are, a) 15.8% (0 s), b) 11.4% (60 s), c) 15.2% (75 s), d) 18.3% (90 s), e) 14.5% (120 s). Orientation of compaction and foaming are perpendicular to the image plane. See also Fig. 5.
FIG. 4: Spatial correlation between pore volume and TiH$_2$ particles of the A1Si7+0.5 wt.% TiH$_2$ samples (IFAM-A1Si-1 - precursor material hot compacted at 450°C, $\mu$CT volume images acquired with high spatial resolution). The TiH$_2$ density is normalised to the mean density found in the foam matrix. Data for distances < 1.5 µm omitted due to the limitation given by the resolution. Graphs are arranged in the same way as in Fig. 2.
FIG. 5: Same as Fig. 4 for samples hot compacted at 200°C (IFAM-AISI-2, SµCT volume images acquired with high spatial resolution). Data for distances < 1.5 µm omitted due to the limitation given by the resolution. Graphs are arranged in the same way as in Fig. 3.
FIG. 6: Sample slices from the AlSi7+0.5 wt.% TiH2 (TUB-AlSi) foam series with heat-treated TiH2 used as blowing agent (SµCT volume images acquired with moderate spatial resolution). The pores are dark, metallic matrix grey and the blowing agent appears in white. a) 1.5% porosity, b) 28% porosity, cf. Figs. 7, 8. Orientation of compaction and foaming are parallel to the image plane [25].
FIG. 7: Correlation analysis of the two samples from the TUB-AISi foam series shown in Fig. 6. Data for distances < 10 \( \mu m \) omitted due to the limitation given by the resolution.
FIG. 8: Metallographic image of AlSi7 + 0.5%TiH$_2$ (TUB-AlSi) samples in two foaming stages, a) early stage with 1.5% porosity (discussed also in Ref. 13), b) later stage with 28% porosity. Both samples were etched with 0.5% HF. Aluminium particles appear in light grey, silicon particles in medium density grey and TiH$_2$ particles are marked with circles, cf. Figs. 6, 7.
FIG. 9: Tomographic slices of IFAM-ALSiCu specimens (ALSi6Cu4 + 0.5 wt.% TiH2, hot compacted at 450°C, SμCT volume images acquired with high spatial resolution). White represents Cu, light grey TiH2, medium grey metal, dark grey pores. a) raw precursor material with strong artefacts originating from the copper particles, b) state after 90 s of heat treatment showing that copper has been largely dissolved and first pores have appeared, c) state after 120 s showing no more Cu, the remaining highly absorbing particles are TiH2 (6.1% porosity). See also Fig. 10.
FIG. 10: Correlation analysis of an IFAM-AlSiCu specimen (AlSi6Cu4 + 0.5 wt.% TiH₂ hot compacted at 450°C) after 120 s of heat-treatment: the density of highly absorbing material (aluminium-copper alloy and TiH₂) is plotted vs. its distance to the pore surface. See also Fig. 9(c). Data for distances < 1.5 μm omitted due to the limitation given by the resolution.
FIG. 11: Correlation analysis of a TUB-AISiCu specimen (AISi6Cu10 + 1.0 wt.% TiH$_2$). The density of the blowing agent particles is almost constant for all distances to the pore surface, indicating a lack of clear spatial correlation between the positions of the pores and the TiH$_2$ particles. Data for distances < 10 µm omitted due to the limitation given by the resolution.
FIG. 12: Metallographic images of some IFAM-AISiCu samples (AISi6Cu4 + 0.5 wt.% TiH$_2$, hot compacted at 450°C, sample surfaces etched with 0.5% HF), a) raw precursor material (0 s heat-treatment), b) 90 s of heat treatment, c) 120 s of heat treatment. Copper appears in orange, silicon in blueish-grey, aluminium in white-grey aluminium and pores are dark. TiH$_2$ particles are marked by red circles in (c). See also Fig. 9.
FIG. 13: SEM close-up of the sample shown in Fig. 12(b), EDX analysis was performed at positions marked, red cross: 57 wt.% aluminium, 12 wt.% silicon and 31 wt.% copper, green cross: 94 wt.% aluminium, 4 wt.% copper and 2 wt.% silicon. The large white particle at the left border is copper, the one to the right of the pore (in the centre of the image) was identified as silicon particle.
FIG. 14: Schematic summary of pores formation in different alloys including alloy 6061 studied elsewhere [10, 11, 12, 13]. The predominant pore formation type is given for each of the states and the images and figures supporting a given representation.