Grain refinement in a AlZnMgCuTi alloy by intensive melt shearing:
A multi-step nucleation mechanism

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Abstract
Direct chill (DC) cast ingots of wrought Al alloys conventionally require the deliberate addition of a grain refiner to provide a uniform as-cast microstructure for the optimisation of both mechanical properties and processability. Grain refiner additions have been in widespread industrial use for more than half a century. Intensive melt shearing can provide grain refinement without the need for a specific grain refiner addition for both magnesium and aluminium based alloys. In this paper we present experimental evidence of the grain refinement in an experimental wrought aluminium alloy achieved by intensive melt shearing in the liquid state prior to solidification. The mechanisms for high shear induced grain refinement are correlated with the evolution of oxides in alloys. The oxides present in liquid aluminium alloys, normally as oxide films and clusters, can be effectively dispersed by intensive shearing and then provide effective sites for the heterogeneous nucleation of Al3Ti phase. As a result, Al3Ti particles with a narrower size distribution and hence improved efficiency as active nucleation sites of α-aluminium grains are responsible for the achieved significant grain refinement. This is termed a multi-step nucleation mechanism.

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1. Introduction

In order to underline the importance of grain refinement, in direct chill (DC) casting of aluminium alloys, grain refinement can [1–3] interact with or influence the end product mechanical properties, its formability during thermo-mechanical processing, the occurrence of surface defects during downstream processing, the homogenization treatment kinetics and the anodizing properties. During casting itself, grain refinement directly influences the formation of porosity and hot tearing. The microsegregation scales as well as the macrosegregation features, especially when related to grain transport phenomena, are strongly influenced by the grain refining strategy.

Al–Ti–B master alloys as grain refiner additions are commonly added to melts for the DC ingot production of wrought aluminium alloys. However, only about 1% of the added particles successfully nucleate aluminium grains and this low efficiency is undesirable not only for its immediate cost implications, but also because refiner particles may themselves be detrimental in the final microstructures [2] particularly for products intended for extrusion, deep drawing or high performance structural applications [3].

Grain refinement theories proposed so far can be classified as either related to an enhanced heterogeneous nucleation mechanism or to a growth restriction mechanism. Nucleation based theories include phase diagram theory, peritectic hulk theory, hypernucleation theory [4,5] and duplex nucleation theory [6,7]. However, most of the nucleation based theories have been discredited and their greatest weakness is that they do not apply to both foundry and wrought alloys [8]. Growth restriction is generally related to the high constitutional undercooling of solute elements and is usually characterised by the growth restriction factor (GRF) [8–12]. The theoretical basis of the growth restriction theory is that restriction of the growth of already nucleated grains permits further nucleation in the undercooled melt until the total latent heat release is sufficient to cause recalescence and the loss of undercooling. Greer and co-workers developed the free growth model to understand the physical mechanism behind the low efficiency of commercial grain refiners based on the Al–Ti system [13–15]. This showed that the size distribution of grain refiner particles plays a key role on the resultant grain refinement effect.

Oxides are inevitably present in liquid metals and alloys in the form of oxide films and/or oxide clusters that can cause severe difficulties during casting, and become inclusions and other related defects resulting in the degradation of the mechanical properties of cast components. Extensive effort has been made to remove oxides from liquid metal prior to casting using both chemical and physical
Recently, the melt conditioning by advanced shear technology (MCAST) process has been developed for conditioning liquid metal under intensive forced convection before solidification and it has been demonstrated that naturally occurring oxides can enhance heterogeneous nucleation for microstructural refinement in commercial Mg–Al alloys. Oxide clusters and films are composites, whereas nano-scale oxide particles are embedded in a liquid metal matrix. Such oxide particles can be effectively dispersed by intensive forced convection and can be potent sites for heterogeneous nucleation since there is a good crystallographic match with the nucleated solid crystalline phases.

In this paper, we have used a specially designed experimental procedure with a lower cooling rate during solidification to demonstrate the grain refinement effect of intensive melt shearing. A standard TP-1 mould and a wedge-shaped copper mould were used to assess the grain refinement induced by intensive melt shearing under varying cooling rates. A pressure filtration technique combined with SEM observation and quantitative metallographic analysis was used to investigate the evolution of oxides and correlate it with the achieved grain refinement. Based on these experimental observations, a multi-step nucleation mechanism is proposed.

![Fig. 1. Calculated solid fraction and volume fraction of Al3Ti in a AlZnMgCuTi alloy using Factsage software.](image1)

![Fig. 2. Flow chart of specially designed solidification experiment using mild steel mould.](image2)
2. Experimental

The experimental AlZnMgCuTi alloy had the following nominal composition: Al 10.5 wt%; Zn 10.5 wt%; Mg 2 wt%; Cu 1.6 wt%; Fe 0.35 wt%; Si 0.15 wt%; and Ti 0.2 wt%. The reason for choosing this particular composition with the titanium level at 0.2 wt% is to ensure the formation of the Al3Ti phase on solidification. Assessment of the liquidus temperature of the alloy could be done by adding the binary peritectic contribution of Ti to the liquidus of the hypoperitectic multicomponent alloy. However, Fortier et al. [24] mentions that for a 7046A alloy the formation temperature of Al3Zr is 20 °C higher than that in the Al–Zr binary alloy. The same might be true for a 7046A alloy the formation temperature of Al3Zr is 20 °C higher than that in the Al–Zr binary alloy. The same might be true for titanium aluminium in a binary vs. a multicomponent alloy.

 Indeed, Factsage calculations suggest that the equilibrium temperature for Al3Ti formation in this alloy is 729 °C (Fig. 1), in accordance with the trend published for Al3Zr [24]. Depending on the nucleation conditions Al3Ti precipitates may form at quite variable undercooling or even supercooling [25].

For all the cases with melt conditioning, the shearing temperature of the MCAST unit was set at 650 °C and the temperature of the melting furnace was set at 750 °C. Melt conditioning was carried out for 60 s at a shearing speed of 500 rpm.

To eliminate the influence of pouring operation on the resulting solidification microstructure, a specially designed experimental procedure is shown in the experimental flow chart in Fig. 2. A mild steel mould, of which the dimensions and the corresponding cross section for grain size assessment are shown in Fig. 3a, was preheated to 650 °C and then cast into a preheated mild steel mould after pouring for 60 s. For the alloy without melt conditioning, the melt was cooled to 650 °C (from 750 °C) and isothermally held at 650 °C for 30 min, and was then transferred into a preheated mild steel mould. After the melts were transferred into two mild steel moulds, respectively, the two mild steel moulds were isothermally held at 650 °C for additional 3 min in a well preheated electrical resistance furnace. During isothermal holding, the temperature of the melts was monitored by a K-type thermocouple to make sure the temperature of the melt was within 650 ± 2 °C. The thermocouple used was calibrated against the melting temperature for pure Al (99.9 wt.%). The two mild steel moulds were then taken out from the holding furnace and air cooled to room temperature.

A standard TP-1 mould [22] and a wedge-shaped copper mould were used to evaluate the grain refinement effect of melt conditioning under intermediate and high cooling rates, respectively. The geometry of the wedge-shaped copper mould sample is shown in Fig. 3b. Three different processing schemes were used for TP-1 samples with the same casting temperature all at 650 °C, as presented in Fig. 4. For the sample without melt conditioning, the melt was air cooled to 650 °C from 750 °C and then cast into
a TP-1 mould (scheme A in Fig. 4); for the samples with melt conditioning, two different melt conditioning processes were adopted. One was sheared at 650°C isothermally for 60 s after the melt temperature had been air cooled to 650°C and then poured into MCAST unit (scheme B in Fig. 4). The other one was sheared immediately after the melt was poured into MCAST unit from 750°C, including two consecutive stages for 60 s in total: continuous cooling and shearing from 750 to 650°C and subsequent isothermal shearing at 650°C (scheme C in Fig. 4). In Fig. 4, the calculated volume fraction of Al₃Ti against temperature by Factsage software is also included (same as Fig. 1) to facilitate the understanding of the development of Al₃Ti during different processes. For the wedge-shaped copper mould samples, schemes A and C were used for the cases with and without melt conditioning, respectively.

The metallographic samples for optical microscopy and scanning electron microscopy (SEM) were prepared using standard metallographic procedures. The samples were sectioned as shown in Fig. 3a and b. The TP-1 sample was sectioned perpendicular to its axis 38 mm from its base. The samples for grain size measurement

Fig. 5. Polarized light micrographs of (a) edge area and (b) middle of non-sheared mild steel mould sample; (c) edge zone and (d) middle of sheared mild steel mould sample.

Fig. 6. Grain structures in TP-1 samples: (a) air cooled to 650°C from 750°C and cast without shearing (scheme A in Fig. 4), grain size: 123 ± 24 μm; (b) air cooled to 650°C and then sheared (scheme B in Fig. 4), grain size: 120 ± 21 μm; and (c) poured into MCAST unit at 750°C and then sheared (scheme C in Fig. 4), grain size: 48 ± 3.5 μm.
were anodized with Barker’s reagent (4% HBF4 in distilled water) and then viewed under polarized light using a Zeiss optical microscope with an Axio Vision 4.3 image analysis system. The mean linear intercept technique was used to measure the grain size.

A model alloy Al–20Zn–0.05Ti was subjected to pressure filtration to analyse oxides in the melt with/without melt conditioning. The reason for choosing this alloy chemistry is to decrease the liquidus of the Al–Zn alloy to the same as the experimental wrought alloy and with a similar oxide formation and a simplified composition. The filtration crucible was preheated to \(300–350\,^\circ\text{C}\), to reduce heat loss during transfer of the liquid alloy. Filtered samples containing the concentrated oxides within the metal (\(\sim 5\,\text{mm in thickness}\)) in contact with the filter were sectioned, mounted and polished for metallographic examination. Scanning electron microscopy (SEM) was carried out with a field emission gun Zeiss Supera 35 machine, equipped with an energy dispersive spectroscopy (EDS) facility and operated at an accelerating voltage of 5–20 kV.

3. Results

3.1. Specially designed experiment demonstrating grain refinement effect by intensive melt shearing

Fig. 5 shows the microstructures of sheared and non-sheared mild steel mould samples showing that the grain size was refined after intensive melt shearing. The grain size was decreased to about 260 \(\mu\text{m}\) after intensive melt shearing compared to 620 \(\mu\text{m}\) for non-sheared sample. The chilled grains adjacent to the mould wall and coarse equiaxed grain structure were observed in the non-sheared sample (Fig. 5a and b). For the sheared sample, chilled grains adjacent to the mould wall were not observed and the grain structures both near the edge and in the interior were finer and more uniform as shown in Fig. 5c and d.

3.2. Grain size as a function of cooling rate with and without shearing

Fig. 6 shows typical optical micrographs viewed under polarized light of the TP-1 samples cast at 650 \(^\circ\text{C}\), with and without melt shearing. For the sample without melt conditioning (processing scheme A in Fig. 4), the average grain size was 123 ± 24 \(\mu\text{m}\) (Fig. 6a); for the sample in Fig. 6b, after the melt temperature was air cooled to 650 \(^\circ\text{C}\), then poured into MCAST unit and melt conditioned (scheme B in Fig. 4), the grain size was 120 ± 21 \(\mu\text{m}\), almost the same as in Fig. 6a; however, in the case of the sample melt conditioned with melt directly transferred into the MCAST unit from 750 \(^\circ\text{C}\) (scheme C in Fig. 4), the grain size was reduced to 48 ± 3.5 \(\mu\text{m}\) (Fig. 6c).

Fig. 7 shows the effect of shearing time on grain size of TP-1 samples with the alloy melts directly transferred into the MCAST unit (scheme C in Fig. 4). The plot of grain size against shearing time fits an exponential decay function, showing that the grain size reaches a plateau after a period of rapid decrease beyond a critical value, that is about 10–20 s in the current study.

![Fig. 7. Grain size as a function of shearing time in TP-1 samples with the alloy melts directly transferred into the MCAST unit (scheme C in Fig. 4).](image)

![Fig. 8. Grain structures in the tip part of wedge-shaped samples of experimental alloy: (a) without shearing (scheme A in Fig. 4), showing columnar dendrites in the location close to the edge of sample; and (b) with shearing (scheme C in Fig. 4), showing equiaxed grains throughout the cross section of sample.](image)

![Fig. 9. Grain size as a function of cooling rate, with and without shearing.](image)
The grain refinement effect of intensive shearing in the tip part of wedge-shaped samples, where a very high cooling rate of 1000 °C/s [26] was achieved, can be seen in Fig. 8. There was an almost total absence of columnar dendritic grains at the edge of the melt conditioned sample (Fig. 8b) in contrast to the readily observed columnar grains at the edge zone of the sample without melt conditioning (Fig. 8a).

Fig. 9 compares the sensitivity of grain size to the variation of cooling rate with and without melt conditioning. It can be seen clearly that the grain refinement effect by intensive melt shearing occurs across the cooling rate range examined and is more pronounced under the conditions of lower cooling rate during the solidification process.

3.3. Characterisation of Al₃Ti intermetallics and oxide particles

Detailed SEM observations were carried out using TP-1 samples with and without melt conditioning (schemes C and A in Fig. 4). Fig. 10a shows that Al₃Ti intermetallic particles can be readily observed within the interior of refined grains. Comprehensive SEM observations indicated that there is an association between the intermetallic Al₃Ti particles and oxide particles, as shown in Fig. 10b, where a typical Al₃Ti intermetallic particle is associated with an oxide particle located in the centre of an α-aluminium grain. The size distribution of Al₃Ti intermetallic particles in TP-1 samples with (scheme C in Fig. 4) and without (scheme A in Fig. 4) melt conditioning is plotted in Fig. 11, showing that the Al₃Ti particles with melt conditioning have a narrower size distribution compared to the sample without melt conditioning. For instance, the size frequency of the Al₃Ti particles frequently observed within α-aluminium grains in the sheared TP-1 sample was 78.3% in the range of 6–20 μm, compared to the value for the sample without shearing of only 42.7%.

Figs. 12 and 13 show oxides in the alloy melt of Al–20Zn–0.05Ti, with and without melt conditioning. Without melt conditioning, oxides in alloy melt are in the form of oxide films or oxide clusters, as shown in Fig. 12a-c. Within films or clusters, crystalline oxide particles with sub-micron size are held together by capillary force [20] and each individual particle is surrounded by other particles rather than the aluminium matrix (Fig. 12c). The SEM-EDXS spectrum shown in Fig. 12d suggests that the oxides formed in the model alloy could be Zn-spinel. After melt conditioning, it can be seen from Fig. 13 that oxides are in the form of dispersed individual particles, which are embedded in the aluminium alloy matrix.

4. Discussion

For non-sheared samples, the grain size was coarser and chill zones of finer grain size were found compared to the sheared samples where the grain size was finer and uniform across the entire sample. This difference in grain structure can be understood as without melt conditioning, nucleation behaviour was triggered by thermal undercooling through the mould wall, whilst after melt conditioning, nucleation occurred throughout the whole volume of the sample by enhanced heterogeneous nucleation and resulted in homogeneous solidification.

Fluid flow during melt conditioning has been analysed previously [27–29]. The MCAST unit has a pair of screws co-rotating inside a barrel. The screw profiles are fully intermeshing and self-wiping and fluid flow is characterised by a high shear rate and a high intensity of turbulence, with a cyclic variation of shear rate. During the melt conditioning, there is an enormous amount of changing interfacial area, providing enhanced heat transfer and strong dispersive mixing. Thus, the melt with intensive shearing is extremely uniform in terms of composition fields, thermal conditions and dispersed individual oxide particles.
For an oxide particle to act as a nucleus for Al₃Ti there should be good crystallographic matching. Al₃Ti has a tetragonal structure based on an ordered cubic close packed (ccp) structure [14]. Each of the oxides possibly formed in the experimental alloy melt, γ-Al₂O₃, MgO, ZnAl₂O₄ and MgAl₂O₄, has a similar cubic structure [30–33]. Based on the reported lattice parameters and the expression of the interatomic spacings misfit \( f = \left[\frac{(d_2 - d_1)}{d_2}\right] \times 100 \), where \( d_2 \) and \( d_1 \) are interatomic spacings of Al₃Ti and each individual oxide phase, respectively, the estimated interatomic misfit \( f \) along the closely packed directions between Zn-spinel/Mg-spinel and Al₃Ti are only 0.84/0.92%, respectively. This means that both Zn-spinel and Mg-spinel could be good substrates for the nucleation of the Al₃Ti phase. In a much more recent investigation on the formation of oxides in Al–6Zn–X Mg (X = 0 and 2 wt%) alloys, Zn-spinel and Mg-spinel were detected on the Al–6Zn and Al–6Zn–2Mg alloy samples, respectively [34]. In our experimental wrought alloy, the oxide formed is more likely to be Mg-spinel due to the relatively high magnesium content. With melt conditioning, dispersed Mg-spinel particles in the alloy melt may act as active nucleation sites for Al₃Ti particles. When the alloy melt was poured into the MCAST unit from 750 °C (scheme C in Fig. 4) in the current study, the existing oxides in the alloy melt were dispersed within a few seconds. Based on the thermodynamic calculation using Factsage software, the equilibrium formation temperature of Al₃Ti intermetallic phase in the experimental alloy can be estimated as 729 °C (Fig. 1). Hence Al₃Ti would be anticipated to precipitate below this temperature. Due to the good crystallographic match between oxide and Al₃Ti phases as discussed above, Al₃Ti particles would preferably nucleate on oxide particles at relatively low undercooling. This applies for all the melt conditioned samples in the present study except for the case, where the melt temperature was air cooled to 650 °C and then poured into the MCAST unit (Fig. 6b, scheme B in Fig. 4). Because of the homogeneous thermal and compositional fields resulting from melt conditioning, Al₃Ti particles tend to be equiaxed and with a narrower size distribution, as evidenced by the quantitative metallographic analysis results shown in Fig. 11. Without melt conditioning, the nucleation of Al₃Ti was mainly triggered by thermal undercooling and as a result, Al₃Ti particles exhibited a faceted morphology and with a wider size distribution, as indicated in Fig. 11.

The respective roles of temperature gradient and nuclei density on the columnar to equiaxed transition (CET) have been well established since Hunt’s pioneering work published in 1984 [35]. The role of a grain refining additive is, in essence, to promote the CET [3]. In the modelling work on grain refinement by addition of

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Fig. 12. SEM images showing typical morphology of oxides in non-sheared and filtered sample of Al–20Zn–0.05Ti with: (a) low and (b) high magnification images of oxide film, and (c) oxide clusters, both of which consist of individual oxide particles with sub-micron size; (d) a typical EDS spectrum of individual oxide particles marked by “A” in (c), suggesting it possibly being the Zn-spinel phase.

Fig. 13. Typical SEM image of oxides in sheared and pressure filtered sample of Al–20Zn–0.05Ti showing individual dispersed oxide particle uniformly distributed in Al matrix.
inoculants [13,15], the isothermal melt model was adopted in which macroscopic temperature gradients were absent, which is a reasonable assumption for air-cooled steel moulds adopted in the specially designed procedure (Fig. 2). At higher gradients such as in the case of wedge-shaped samples a higher nuclei density is necessary in order to obtain the CET (Fig. 8).

With melt conditioning, dispersed oxide particles and thereby enhanced nucleation of Al3Ti intermetallic particles with a narrower size distribution will enhance the heterogeneous nucleation of α-aluminium phase. Greer and co-workers developed the free growth model, based on the Al–Ti–B grain refiner system in which TiB2 particles have high potency of inoculant for α-aluminium [13,15]. It was concluded that the grain refinement achieved is determined by a free-growth condition in which growth undercooling, $\Delta T_{fg}$, is inversely proportional to the particle diameter of inoculants, $d$:

$$\Delta T_{fg} = \frac{4\pi}{\sigma \Delta S_s} d$$

where $\sigma$ is the solid–liquid interfacial energy and $\Delta S_s$ is the entropy of fusion per unit volume. More importantly, in their modelling, based on the understanding of recalcensce limiting the number of active nucleation particles and therefore the number of grains, the size distribution of grain refiner particles rather than the size in single fixed value was treated as input. To achieve this, spatially dispersed grain refiner particles throughout the bulk melt are necessary and either settling or agglomeration behaviour of grain refiner particles should be avoided. In other words, the existence of potent nucleating agents may not necessarily lead to grain refinement in the solidified microstructure. For effective grain refinement of a given alloy composition, the potent nucleating particles need to have sufficient numbers, favourable particle size and size distribution. In the present work, intensive melt shearing can disperse the oxide films and clusters into individual oxide particles that promote the nucleation of Al3Ti particles on these oxide particles. As a consequence, Al3Ti particles with a compact and equiaxed morphology and a narrower size distribution were obtained, which in turn enhanced heterogeneous nucleation of α-aluminium grains throughout the entire volume of alloy melt. As a result, significant grain refinement effects can be achieved by intensive melt shearing. This grain refinement mechanism is termed a multi-step nucleation mechanism.

The argument of the enhanced heterogeneous nucleation by Al3Ti through a multi-step nucleation mechanism is also supported by the result shown in Fig. 6b, where Al3Ti had formed before the melt was poured into the MCAST unit at 650 °C (scheme B in Fig. 4), so that intensive melt shearing (dispersed oxide) had no influence on the formation of Al3Ti and hence no effect on the grain refinement. In terms of the influence of melt conditioning time when the melt conditioning time exceeds a critical value, 10–20 s, a sufficient number of individual dispersed oxide particles have been achieved for the heterogeneous nucleation of Al3Ti particles. Further, increasing the melt conditioning time does not contribute to an enhanced heterogeneous nucleation of Al3Ti particles on oxide particles and its size distribution. Therefore the grain size cannot be refined further (Fig. 7).

5. Concluding remarks

(1) Grain refinement has been successfully achieved in an experimental wrought Al2ZnMgCuTi alloy by intensive melt shearing. The grain refining effect is more pronounced under the conditions of lower cooling rate during solidification processing.

(2) With intensive melt shearing, the oxide films are dispersed and uniformly distributed in the alloy melt. These oxide particles can then act as active heterogeneous nucleation sites for Al3Ti intermetallic particles providing shearing occurs during the formation of the Al3Ti phase.

(3) Following the dispersion of oxide particles by intensive melt shearing, a narrower size distribution of Al3Ti particles is produced providing many active sites for α-aluminium nucleation resulting in a finer grain size. This multi-step nucleation process, oxide to Al3Ti to α-aluminium, is shown to be significantly enhanced by intensive melt shearing during the formation of the Al3Ti phase.

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