GRAIN REFINING ALUMINIUM ALLOYS BY THE SAME-ALLOY ROD

D.G. Eskin and F. Wang

Brunel University London, BCAST
Uxbridge, United Kingdom
(*Corresponding author: dmitry.eskin@brunel.ac.uk)

ABSTRACT

It is well known that it takes some time for the solid phase to completely dissolve upon melting, especially inside the defects of insoluble particles, e.g. oxides. Until then the oxides remain active solidification substrates in the case of subsequent solidification. It is also known that ultrasonic melt treatment causes grain refinement through activation and dispersion of solidification substrates (one of the mechanisms) and also accelerates the dissolution of solid metal in the melt. In this study we combine these effects and demonstrate that the introduction of an alloy rod into the matrix melt of the same composition results in significant grain refinement, this effect being increased by the ultrasonic vibration of the rod. The achieved grain size is comparable to that obtained by a standard Al–Ti–B grain refiner. All samples were cast using a standard TP-1 mould to enable correct comparison. The effect of the temperature range of the rod introduction, as well as the effect of ultrasonic vibrations are discussed.

KEYWORDS

Grain refinement, Ultrasonic processing, Solidification, Nucleation, Dissolution
INTRODUCTION

The fact that by introducing solid metal into the melt one can achieve structure refinement is rather well known. As early as in the 1930s–1950s a series of papers and patents have been published, showing the benefits for grain refinement when dissolving solid metal in the melt before solidification (Danilov and Neimark, 1938; Scheil, 1956). Later on this way of structure refinement was developed further and dubbed “suspension casting” or introduction of “internal chills” (Madyanov, 1969; Zatulovsky, 1981). The solid metal was added in a form of cut wire, cut sheet, or powder, with the resulting grain refining, elimination of columnar grain structure in ingots and castings of steel, copper and aluminium alloys. The underlying mechanisms have been suggested as (1) rapid cooling of the melt due to the latent heat consumption upon melting of the solid metal with the resultant melt undercooling and (2) introduction of many solidification substrates in the form of crystal fragments and active non-metallic inclusions (Danilov & Neimark, 1938; Balandin, 1973). A number of patents have been filed where either the same solid alloy or a master alloy with additions (e.g. Ti for Al alloy) is introduced in the amounts up to 50% (typically less than 10%) into the melt close to the liquidus temperature (Schmidt, 1966; Talbot & Soller, 1973, Krupp, 1974; Bondarev, 1979). The main reason that this technique is not widely used in industry is the possible incomplete dissolution of the solid parts introduced into the melt with ensuing inhomogeneous as-cast structure and potential defects. Also the selection of the temperature range where the technology works the best is not clear.

Another well documented means of controlling the grain structure of as-cast metals is ultrasonic melt processing (Eskin, 1965; Abramov, 1972; Abramov, 1998; Eskin & Eskin, 2015). Although very powerful in grain refinement, this technology suffers from instrumental issues, i.e. the ultrasonic tool (sonotrode) that is used for direct introduction of high-frequency vibration into the melt is subject to cavitation erosion and (if made of metal) gradual dissolution. At the same time, some schemes that used a “consumable” sonotrode have been suggested in ultrasonic welding (wire feeding) and in electro-slag remelting (vibrating electrode) as reviewed elsewhere (Abramov, 1972; Abramov, 1998). Such a scheme opens some new avenues in ultrasonic melt processing as it eliminates the issue of sonotrode material selection. It is also well known (Eskin, 1965) that ultrasonic vibrations significantly accelerate the dissolution of solid metals in the melt as well as promote rapid mixing of solutes (Brodova, 2002).

In this study we attempted to combine these two technologies to achieve a synergetic effect.

EXPERIMENTAL

An AA6082 alloy was used as a base material (1.16% Si, 0.7% Mg, 0.98% Mn, 0.26% Fe, 0.17% Cr, 0.04% Ti) and a 9-mm rod of the same alloy was used for additions. In some experiments a standard Al5Ti1B rod (LSM) was used (0.2% addition). A total of 10% of AA6082 rod was introduced in each experiment.

When ultrasound was applied, the rod was inserted tightly into a hole in a steel sonotrode in the half-wavelength position and then fed into the melt while the ultrasonic transducer was on. The sonotrode was not in contact with the melt. As a result the longitudinal oscillations of the sonotrode were transformed into flexural oscillations of the rod with the resultant shorter wavelength and larger amplitudes. Experiments in water showed that cavitation conditions were achieved at the tip of the rod as well as in nodes along the rod length. A 5kW ultrasonic generator and a 5 kW water-cooled transducer (Reltec) were used at a power of 3 kW.

The melt was prepared in clay-graphite crucibles (0.5 kg charge) in an electric resistance furnace with melt temperature up to 770°C. Rod introduction was made either without or with ultrasonic vibrations applied into the melt in a temperature range with the final pouring temperature 670°C. Reference samples were prepared by adding the same amount of rod into the melt that was then superheated to 750°C, left in the furnace for 30 min and then cooled in air and cast at 670°C. Casting was done in a standard TP-1 mould (Aluminum Association, 1990).
The castings were then sectioned along the vertical centre plane and etched for structure examination. Macroetching was done in a 10% NaOH water solution after grinding. To reveal grain microstructure smaller samples were cut, ground and polished using standard procedures and then anodised in a 5% HFB₄ water solution at 20 VDC. The grain structure was subsequently examined in an optical microscope in a polarised light. Statistical analysis of the measured grain size was performed and the average values are reported.

RESULTS AND DISCUSSION

Reference samples (a number of those had been produced) showed a coarse columnar grain structure as shown in Figure 1 with the average grain size 630 µm.

![Image](image1.png)

Figure 1. Macro- (a) and micro- (b, c) grain structure of a reference sample cast at 670°C in a TP1 mould

First experiments when the rod was introduced without or with ultrasound in the temperature ranges 740 to 715°C and 740 to 710°C did not produce any significant results with very coarse mixed columnar and equiaxed structure formed. This confirmed the importance of the temperature range needed for the desired effect. Although the introduced rod was quick to dissolve, the relatively high temperature apparently prevented the survival of solid fragments as well as facilitated deactivation of solid inclusions (only 10–15°C above liquidus is needed for that (Bondarev, 1979).

Following experiments were performed in the temperature ranges 720 to 670°C and 710 to 670°C. The decrease in the introduction temperature yielded a remarkable change in the grain structure even when the rod was introduced without ultrasonic vibrations, Figure 2, the average grain size 250 µm. It can be noted that the slight decrease in the starting melt temperature gives some additional grain refining effect (228 µm), most probably due to the better survival chances of crystal fragments and active inclusions.

Ultrasonic oscillations applied to the rod further refined the grain structure as illustrated in Figure 3. This might be due to the better distribution of crystal fragments in the melt as well as to the additional activation of insoluble inclusions by ultrasonic cavitation (Eskin and Eskin, 2015; Wang, 2018). At the same time the grain size obtained upon rod introduction with ultrasonic oscillations in a wider temperature range was finer than in a narrower temperature range, i.e. 123 µm and 165 µm, respectively. As ultrasonic oscillations may facilitate additional mechanisms of grain refinement such as activation of inclusions and fragmentation of solid crystals, their effect is time-dependent, i.e. the longer the ultrasonic cavitation and vibration works, the better the result. Therefore, slight increase in the rod introduction time in a wider temperature range is beneficial for the grain refinement.
Figure 2. Effect of rod introduction into the melt in the temperature range 720 to 670°C (a, c) and 710 to 670°C (b, d, e) on the grain structure of the alloy cast in a TP1 mould.

Figure 3. Effect of rod introduction with ultrasonic vibrations into the melt in the temperature range 720 to 670°C (a, c) and 710 to 670°C (b, d, e) on the grain structure of the alloy cast in a TP1 mould.
Figure 4. Effect of Al5Ti1B grain refiner addition and same-alloy rod introduction into the melt in the temperature range 710 to 670°C on the grain structure of the alloy cast in a TPI mould: a, macrostructure; b, c, microstructure

Figure 5. Grain sizes achieved in experiments: (a) the effects of the same-rod introduction, ultrasonic oscillations and the temperature range of introduction and (b) comparison of the same-alloy rod introduction with the standard grain refinement by master alloy addition

The standard procedure of grain refinement of aluminium alloys is the addition of an AlTiB master alloy. In order to compare the grain refining effects that we have achieved with the same-alloy rod introduction, we did additional experiments with the standard Al5Ti1B grain refining master alloy addition. Two experiments were performed, i.e. addition of the grain refiner to the base alloy and same-alloy rod introduction into the grain refiner master alloy. The obtained grain sizes were rather small (112–123 µm) and comparable with those obtained upon same-alloy rod introduction with ultrasonic oscillations, Figure 4. Although the addition of rod to the grain refined alloy gave a slight additional decrease in the grain size (112 µm vs 123 µm), it was not statistically significant. These results demonstrate that under given cooling conditions (TPI mould) the addition of 2% grain refiner as well as the addition of the same-alloy rod give the maximum number of solidification sites that can be realised and produce grains. Therefore the achieved grain size 112–123 µm is the smallest possible and does not depend on the way how the active solidification sites are introduced. Figure 5 summarises the results of grain size measurements for all studied cases.

CONCLUSIONS

1. The grain refining effect upon introduction of a same-alloy rod into the melt prior to casting has been confirmed.
2. The temperature range of the same-alloy rod introduction should be close the casting temperature to assure that the generated solidification sites remain active.
3. The application of ultrasonic vibrations to the same-alloy rod gives additional grain refining effect that is most probably related to the known effects of cavitation on the activation and fragmentation of solidification sites.

4. The observed grain refining by the same-alloy rod introduction with the ultrasonic oscillations applied is similar to that achieved by the standard grain refining procedure.

5. The results show a promising technology of using the same-alloy rod instead of the grain refining rod in continuous casting processes.

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REFERENCES


