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DETERMINATION OF THE THIXOCASTING TEMPERATURES OF AZ91D AND OTHER Mg ALLOYS USING A QUENCHING METHOD

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Abstract — The thixocasting injection temperature is a crucial processing parameter, frequently unavailable in the literature, especially for newly developed alloys. In this work, a simple and rapid method derived from the quenching method was developed to obtain an acceptable thixocasting injection temperature range for various magnesium alloys: AZ91D, AJ62x, AXJ530, MRI153 and ZAEX10440. Small specimens of each alloy were heated to appropriate temperatures within the semisolid range and quenched. Their microstructure was observed optically and the proportion of phase was measured using quantitative image analysis. The microsegregation was characterized and two different heating cycles prior to quenching temperature between liquidus and solidus. These results were found useful to obtain a practical injection temperature range for each alloy and to obtain good quality thixocast parts.

Résumé — La température d'injection pour la thixocoulée est un paramètre crucial de procédé qui n'est pas toujours disponible dans la littérature et ce, surtout dans le cas de nouveaux alliages développés en laboratoire. Dans cet article, une méthode simple et rapide, dérivée de la méthode de trempe, est proposée pour estimer un intervalle de température d'injection d'alliages de magnésium tels que AZ91D, AJ62x, AXJ530, MRI153 et ZAEX10440. Des spécimens de chaque alliage ont été chauffés à différentes températures dans l'intervalle semisolide et trempés. Leur microstructure a été observée en microscopie optique et analysée quantitativement par analyse d'image afin de mesurer la proportion de la phase solide primaire. La microségrégation a également été mesurée dans les phases après trempe et deux cycles thermiques différents ont été comparés. La fraction volumique apparente de la phase solide primaire a été déterminée en fonction de la température de trempe entre le liquidus et le solidus. À partir de ces résultats, un intervalle de température a été estimé pour chaque alliage afin d'obtenir par thixocoulée des pièces d'excellente qualité.

INTRODUCTION

Semisolid injection moulding methods are regarded as emerging fabrication processes for magnesium based components requiring excellent surface finish, dimensional accuracy and low porosity [1,2]. Among them, the thixocasting process that uses semisolid billets is particularly attractive since reduced quantities of material are necessary for experimentation of new alloys, testing their rheological and mechanical properties as well as their corrosion performance.

Various methods have been developed for the preparation of billets: extrusion [3], mechanical

stirring [1,4], electromagnetic stirring (EMS) [5], rapid slug cooling technology (RSCT) [6] and direct thermal method [7]. The specific rheological behaviour of semisolid billets depends on their microstructure [8] and also on the thermochemical properties of the alloy [8,9]: latent heat of melting, heat capacity, solidification range and liquid fraction sensitivity (df_I/dT).

The reheating cycle of billets, die temperature, ram speed and applied pressure are thixocast parameters that contribute to the overall quality of thixocast components [5]. The preheating temperature of billets is probably the most important parameter since it strongly influences the fraction of primary solid phase at injection temperature, the optimum solid ranging usually between 40-60 vol% [8-11]. Consequently, the knowledge of solidification curves of alloys, i.e., the relation between fraction of primary phase and temperature is necessary.

Theoretical solidification curves can be obtained by using two limiting models: 1) the lever rule which assumes complete thermodynamic equilibrium and 2) the Scheil relation which presupposes local equilibrium at the interface, complete diffusion in the liquid phase and no diffusion within the solidified phase. These two models have been used by many authors [2,12-17,] for comparison purposes but they are often difficult to use with the newly developed alloys for which thermophysical properties are unknown.

Ghosh et al. [2] have used differential scanning calorimetry (DSC) to determine experimentally the solidification curve of AZ91D magnesium alloy. However, a significant volatilization of zinc and magnesium at high temperature and contamination as well as oxidation problems associated with the low scanning rates prevented reliable measurements. Recently, Mirkovic et al. [13] have used DSC with sealed Ta-crucible to obtain the solid fraction of AZ91D alloy as a function of temperature assuming a simple linear dependence of heat evolution with solid fraction. They also proposed an in situ heat transfer model to further improve differential thermal analysis (DTA) and reduce smearing problems associated with thermal methods [18]. Results from DSC and DTA methods cannot discriminate between the liquid phase remaining entrapped within primary grains and the liquid phase formed between grains which directly influence rheological behaviour.

A more direct method which involves quenching of specimens from the semisolid condition and evaluation of the fraction of phases by quantitative image analysis has been used by many authors [12,13,18-25]. This so-called quenching method is relatively rapid and does not require expensive equipment. Moreover, no assumption about any thermophysical property of the alloys is required. However overestimation of the solid fraction caused by a substantial solid deposition on to the primary phase during quenching was reported frequently [12,13,18-25]. It was observed that high cooling rates (160 K/s) during quenching of small aluminum specimens can prevent this phenomenon efficiently [21]. Lower cooling rates (80 K/s) were also found acceptable for specimens containing a lower fraction of solid [21]. Mirkovic *et al.* [12,13,18] have used this quenching method with specimens of AZ91 alloys [12]. They also reported overestimation problems and indicated that the use of liquid nitrogen as quenching medium could not provide a very effective cooling medium, likely due to the formation of a gas blanket at the surface of the specimens during quenching [26].

In the present work, small semisolid specimens of magnesium alloys preheated at different temperatures were rapidly quenched to preserve as much as possible the primary phase formed initially. The microstructure of quenched specimens was characterized as a function of quenching temperature and microsegregation in the primary phase was measured. The volume fraction of the primary phase was determined as a function of quenching temperature producing calibration curves. Using these curves, the injection temperature of different magnesium alloys was estimated. These predicted temperatures were comparable to best injection temperatures used during the thixocasting experiments.

EXPERIMENTAL PROCEDURES

Composition and Properties of Alloys

The experimental method has been developed with the AZ91D magnesium alloy. Then it has been applied to four newly developed magnesium alloys (AJ62x, AXJ530, MRI153 and ZAEX10440) in order to determine the influence of temperature on the apparent fraction of the primary phase and estimate a practical injection temperature. The chemical composition of these alloys is given in Table I whereas their solidus and liquidus are given in Table II. The solidus-liquidus range was varying between 83 to 125 °C for Mg-Al based alloys, increasing regularly with the concen-

Composition (wt%)								
Alloy	Al	Zn	Sr	Mn	RE	Са	Ce	Mg
AZ91D	9.0	0.71	-	0.25	-	_	-	Bal.
AJ62x	6.0	0.01	2.6	0.43	-	-	-	Bal.
AXJ530	4.9	-	0.16	0.3	-	3.0	-	Bal.
MRI153	8.3	1.0	0.01-0.2	0.16	0.05-1.0	0.84	0.11	Bal.
ZAEX10440	3.6	10.3	-	0.1	3.8	0.21	1.81	Bal.

 Table I – Chemical composition of magnesium alloys

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Alloy	Solidus (°C)	Liquidus (°C)
AZ91D	470	595
AJ62x	517	612
AXJ530	524	607
MRI153	505	602
ZAEX10440	324	586

Table II – Liquidus and solidus of alloys [27,28]

tration of aluminium and more than 262 °C for the Mg-Zn based alloy.

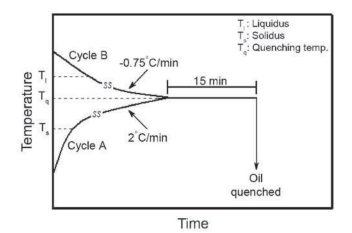
Specimens Preparation

Billets of 7.5 cm in diameter and 15.0 cm in length were prepared from ingots of magnesium alloys melted in a stainless steel crucible and protected using a CO_2 -0.5% SF₆ gas mixture with a positive pressure. The melt was held in the temperature range between 720 and 735 °C. Billets were gravity cast in a permanent mould preheated at about 350 °C and mounted on a massive copper heat sink [27]. Specimens of about 7.0 g were cut out from these billets and used to determine the influence of preheating temperature on the apparent volume fraction of the primary phase after quenching. These specimens were inserted into cylindrical steel crucibles having 2.0 cm in diameter, 2.5 cm in length and 2.0 mm of wall thickness.

Heating Cycles and Quenching Experiments

The cylindrical crucibles (filled with the 7.0 g specimens) were placed inside a larger stainless steel chamber electrically heated under a CO_2 -0.5% SF₆ protective gas (less than 0.1 litre per minute) assuring a positive pressure during melting. They were heated to the liquidus-solidus temperature range using one of two different heating cycles shown in Figure 1 and then quenched. A fine thermocouple was used to measure the temperature of each specimen within the crucibles. Using Cycle A, specimens were progressively heated to the quenching temperature (T_a) and then oil quenched. Using Cycle B, specimens were heated to 25 °C above their respective liquidus, cooled down to the quenching temperature (T_q) and then oil quenched. The quenching temperature (T_a) was maintained for 15 minutes for both cycles, a time sufficient to obtain a homogeneous temperature within the semisolid specimens.

Quenching was performed using canola oil which was chosen over conventional mineral oil and water for its excellent heat extraction capacity [29] and safety with respect with molten magnesium preventing



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Fig. 1. Schematic curves showing the heating Cycles A and B used prior to quenching magnesium alloy specimens (The given cooling and heating rates are average values within the liquidus-solidus range).

explosion of semisolid specimens [20]. For AZ91 specimens, the time between quenching temperature and solidus was less than 1.5 seconds as measured using a fine thermocouple inserted inside the specimens giving a cooling rate of about 80-90 K.s⁻¹. This cooling rate was confirmed by measuring the secondary dendrites arm spacings (SDAS) of the fine dendrites formed upon quenching between the primary α -Mg grains where the average aluminum concentration is about 15 wt%. The following relation proposed by [30] was used.

SDAS (
$$\mu$$
m) = 3.4 t_f^{0.46} (1)

where t_f is measured in seconds. Average SDAS of 4.5 μ m were measured within the rapidly quenched liquid regions. A local solidification time t_f of 1.8 seconds is then calculated yielding a cooling rate of about 75 K.s⁻¹ and a liquidus-solidus range of 130 °C being considered for the residual liquid. The uncertainty on quenching temperature was typically \pm 1 °C.

Determination of Volume Fraction of Primary Phase and Microsegregation

In quenched specimens, the apparent volume fraction of the primary phase was determined from the microstructure of quenched specimens using optical microscopy. The specimens were mechanically polished down to 0.1 μ m and then etched with the reagent developed by Maltais *et al.* [31]. The microstructure of each specimen was examined at 25X magnification and the volume fraction of primary grains was measured over six different fields using quantitative image analysis. In micrographs, the bright regions having a relatively large radius of curvature correspond essentially to the primary α -Mg phase. The dark regions containing α -Mg fine dendrites and secondary phases were formed upon quenching and originate from the solidification of the residual liquid phase.

After Cycle A, a liquid phase was entrapped within the primary grains. This liquid was neglected since it was not connected to the residual liquid surrounding the grains and consequently it was not available to improve flowing properties. In the present work, the volume of this entrapped liquid phase was estimated to about 2-3 vol%.

Microsegregation in quenched specimens was compared using elemental line scans across the primary phase and the rapidly solidified liquid phase. Semiquantitative concentration profiles of aluminum and zinc were obtained with an electron probe microanalyser (EPMA). Detection limits under analytical conditions were typically 0.1 wt% for zinc and 0.03 wt% for aluminum.

Thixocasting Tests with Billets

Billets of the five different magnesium alloys (AZ91D, AJ62x, AXJ530, MRI153 and ZAEX10440) were thixocast using a Buhler die-casting machine. They were first preheated for 8 minutes with a high frequency induction heating system [27]. They were then introduced into the injection sleeve and injected in an experimental box-like mould. Different injection temperatures were tested until the quality of the injected box was optimized [27].

RESULTS AND DISCUSSION

Microstructure of AZ91D Billet

The microstructure of the gravity cast AZ91D billets is shown in Figure 2. Massive $Mg_{17}Al_{12}$ secondary phase particles indicated by 1) were distributed in the α -Mg matrix and some Al-Mn particles indicated by 2) were observed. The grain size of specimens used for the heating and quenching experiments was about 215 µm.

Microstructure of Quenched Specimens

Optical micrographs of AZ91D specimens quenched from different semisolid temperatures are presented in Figure 3. The volume fraction of primary grains determined from image analysis is also given. Dark areas which are composed of fine eutectic constituents correspond to regions that were in the molten state at the moment of quenching. Bright and coarse areas show grains of the primary α -Mg grains. The microstructure of specimens quenched upon Cycle A

1 1 1 100 μm

Fig. 2. Microstructure of unetched AZ91D billets showing 1) massive $Mg_{17}AI_{12}$ particles and 2) fine Al-Mn particles dispersed within α -Mg matrix.

(Figures 3a and 3c) shows islands of β -phase entrapped within the primary α -Mg grains. These entrapped islands were formed above eutectic temperature. Local remelting occurred within the primary α -Mg grains in areas containing the highest concentration of aluminum and zinc. During quenching, these locally remelted areas have crystallized producing α -Mg phase and residual β -particles. The microstructure of specimens quenched upon Cycle B (Figures 3b and 3d) do not show such entrapped liquid.

Rapid epitaxial solidification onto the primary α -Mg grains was expected due to the limited cooling rate during quenching [12,13,18-26]. This solid deposition is visible in Figure 4 for specimens preheated using Cycle A where a thin but continuous layer of α -phase encloses particles of β -phase formed within α -Mg grain upon quenching. In specimens preheated using Cycle B, the dense α -Mg grains prevent any visual contrast that could reveal a rapid growth onto the primary solid phase.

Microsegregation in Quenched AZ91D Specimens

Evaluation of the microsegregation in quenched AZ91D specimens was performed using elemental line scans across the grains of primary α -Mg phase. The position of the line scans and the corresponding semiquantitative aluminum and zinc concentration profiles are shown in Figures 5 to 8.

The comparison of curves in Figures 6 and 8 shows that concentration profiles of Al and Zn are lower and more uniform across primary α -Mg grains formed during Cycle B. However, the concentration of Al and Zn is higher between primary grains of specimens treated using Cycle B. These concentration

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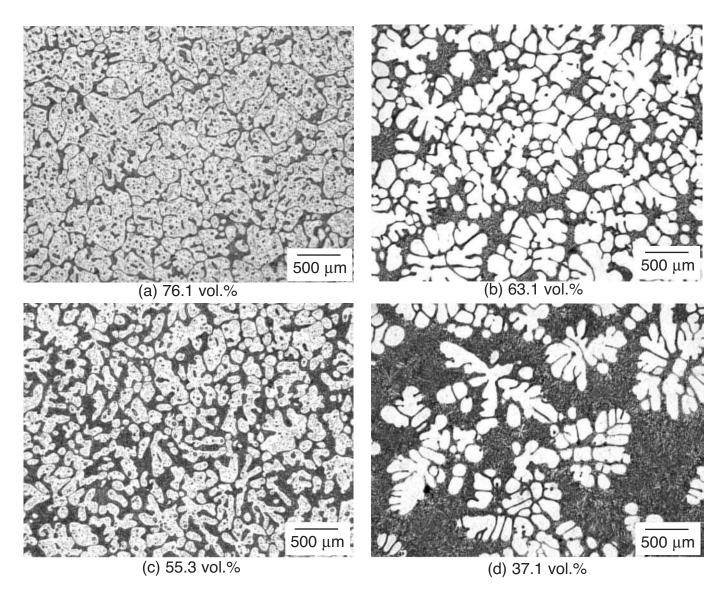


Fig. 3. Microstructures of AZ91D specimens quenched from the semisolid state: at 570 °C for specimens a) and b) following Cycles A and B, respectively and at 585 °C for specimens c) and d) following Cycles A and B respectively.

profiles are explained by the systematic rejection of solutes which occurred during the crystallization of α -Mg grains leading to solute rejection at the interface and diffusion into the surrounding liquid phase. The composition profile shows a homogeneous elemental distribution in primary grains where the average concentration of aluminum is about 3.5 wt% and zinc 0.2 wt%.

Grains formed during Cycle A experienced ripening after partial remelting of specimens. Many solute-rich liquid islands remained trapped within the primary α -grains. Solute diffusion occurred at the interface between the solid and liquid phases during heating Cycle A but somewhat less solute diffusion occurred since these specimens remained for a long period at a relatively lower temperature. Moreover, solute dispersion within the liquid phase was difficult due to the limited connectivity between liquid regions. The solute distribution within primary grains is consequently more heterogeneous due to the presence of many isolated particles of β -phase. In specimens quenched after Cycle A, the average concentration of aluminum in primary α -Mg grains is about 3.9 wt% and that of zinc is about 0.2 wt%.

Volume Fraction of Primary Phase in Quenched AZ91D Specimens

The volume of primary α -Mg grains measured in the polished cross-section of AZ91D specimens as a function of quenching temperature following Cycles A and B is presented in Figure 9. For comparison

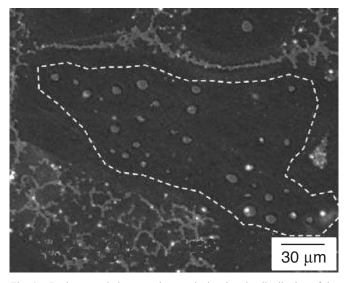


Fig. 4. Backscattered electron micrograph showing the distribution of the secondary phase formed within a grain of primary α -Mg phase during Cycle A in the AZ91D specimen.

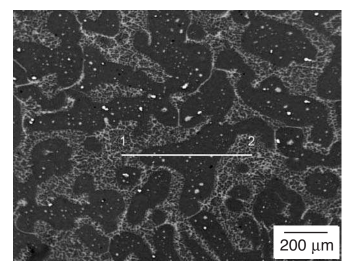


Fig. 5. Backscattered electron micrograph showing the position of the line scan across a primary α -Mg grain formed during Cycle A in the AZ91D specimen.

purposes, the theoretical temperature dependence of solid fraction in AZ91D as a function temperature is reproduced from Reference [2].

For a given quenching temperature, the volume fraction of primary α -Mg in specimens quenched after Cycle A is systematically superior to that formed after Cycle B. The volume of entrapped liquid, estimated at 2-3 vol% can not explain the observed difference. Moreover, since the temperature was monitored with a fine thermocouple embedded in semisolid specimens during heating cycles and quenching, a temperature lag effect is ruled out. So, the discrepancy between the two

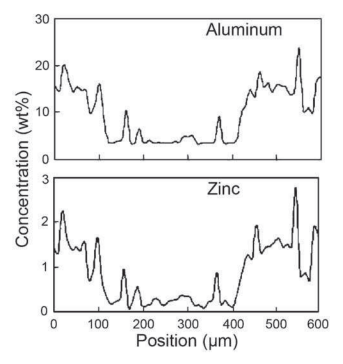


Fig. 6. Concentration profiles of aluminum and zinc across the primary α -Mg grain of the specimen shown in Figure 5 (see line scan $1 \rightarrow 2$).

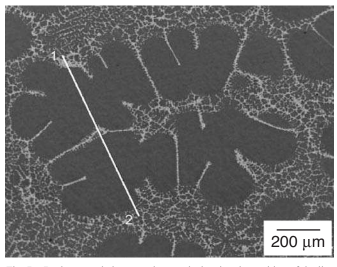


Fig. 7. Backscattered electron micrograph showing the position of the line scan across a primary α -Mg grain formed during Cycle B in the AZ91D specimen.

curves is explained as follows. Semisolid specimens obtained upon Cycle A were maintained for a long period at a relatively lower temperature prior to quenching compared to semisolid specimens obtained after Cycle B, which have formed upon cooling and at a relatively high temperature. Therefore, a less

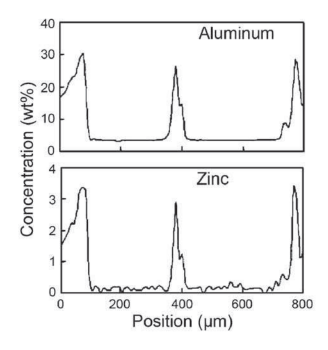


Fig. 8. Concentration profiles of aluminum and zinc across the primary α -Mg grain of the specimen shown in Figure 7 (see line scan $1 \rightarrow 2$).

extensive diffusion and a reduced redistribution of solute occurred in the solid phase during the heating portion of Cycle A and most of the solutes (Al and Zn) within primary α-Mg grains could not reach the rest of the liquid phase producing less solute in the liquid phase compared to specimens treated using Cycle B. In addition, during heating Cycle A, the gradual dissolution of the solid phase has produced a solvent build-up at the melting interface. This solvent accumulation (or solute depletion) increased locally the liquidus temperature slowing down the dissolution process until a more homogeneous dispersion was established in the interdendritic liquid phase. The greater volume of liquid observed in the specimens quenched after Cycle B is also in accordance with the lower aluminum concentration found within primary α-Mg grains of specimens quenched after Cycle B. The results obtained with Cycle B are in agreement with those of Carnahan et al. [32], although the latter show a wider dispersion.

For specimens treated using Cycles A and B, the volume proportions of primary α -Mg grains measured with quantitative image analysis are larger than those obtained theoretically from the solidification curve (Figure 9). This greater volume of apparent solid is ascribed to a rapid growth of the solid on to the primary α -phase during quenching [12,13,18-26]. Despite the rapid cooling from semisolid state, some crystallization on to primary α -Mg phase occurred.

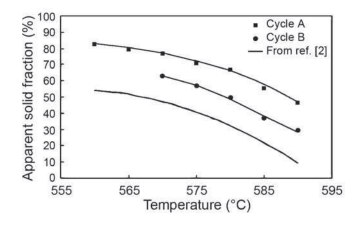


Fig. 9. Apparent volume fraction of the primary α -Mg in AZ91D specimens treated using Cycles A and B.

Practical Thixocasting Temperature for Billets

Shehata *et al.* [5] found experimentally that a temperature range of 575-580 °C is the most suitable for thixocasting AZ91D alloy. This temperature range corresponds to a fraction of the primary phase of approximately of 45-53 vol% according to calculations based on the Mg-Al phase diagram but to less than 30-40 vol% according to theoretical curves from Reference [2] (Figure 9). However, since the short preheating period prior to thixocasting provides far from equilibrium conditions for the liquid and solid phases, the actual proportion of the primary solid at thixocasting temperature is probably different from these estimates.

A more practical and empirical approach was considered based on the assumption that semisolid specimens containing comparable proportion of primary phase at quenching temperature also show a similar quantity of apparent primary phase after quenching. During quenching, a comparable growth of solid on to the primary solid is assumed to occur for the different magnesium alloys, assuming that similar heating and quenching conditions are maintained [23,24].

In the case of AZ91D specimens, a practical thixocasting temperature of 580 °C was found appropriate for AZ91D billets [2]. In the AZ91D specimens oil quenched from this temperature following Cycle B, there is about 50 vol% of apparent primary phase. So, it is assumed that the quenching temperature producing this fraction of the primary phase in other magnesium alloys would also indicate their corresponding appropriate thixocasting temperature.

It was found by experience that the close relation between the two calibration curves (based on Cycle A and Cycle B) allows using one or the other of the calibration curves and providing similar indications

Alloy	40 vol% apparent solid fraction temp. (°C)	60 vol% apparent solid fraction temp. (°C)	Practical injection temp. (°C)
AZ91D	585	573	580
AJ62x	603	588	595
AXJ530	597	587	592
MRI153	588	580	585
ZAEX10440	578	563	573

Table III – Quenching temperature of specimens containing 40 and 60 vol% apparent solid primary phase
following Cycle B and practical injection temperature for different magnesium alloys

about the best injection temperature. However, although Cycle A is apparently better at simulating the practical reheating cycle of billets prior to thixocasting, curves showing the apparent fraction of solid after Cycle A were not used to determine the thixocasting temperature of the other alloys since it was found more difficult to evaluate the volume fraction of the primary solid due to the relatively intricate distribution of entrapped liquid phases in specimens quenched after Cycle A.

Practical Injection Temperature of Other Magnesium Alloys

The above-described procedure for AZ91D alloy has been used to determine the volume fraction of the primary phase after quenching in newly developed AJ62x, AXJ530, MRI 153 and ZAEX10440 magnesium alloys for which no reliable thixocasting injection temperature was found in literature. Small specimens of these alloys were heated in the semisolid state using Cycle B and quenched. All specimens were polished and the apparent fraction of the primary solid phase was determined by quantitative image analysis as a function of quenching temperature.

Results presented in Table III indicate the quenching temperature for which 40 and 60 vol% apparent fractions of the primary phase were obtained following Cycle B. The practical injection temperature of successfully thixocast billets is also given for each alloy. It was found that the best injection temperature was close to about 50 vol% of apparent primary solid in specimens quenched after Cycle B [27].

CONCLUSIONS

A simple and rapid method has been developed to obtain the thixocasting injection temperature of different magnesium alloys. The apparent fraction of the primary α -Mg phase in quenched specimens was obtained as a function of the quenching temperature between the solidus and liquidus for each alloy. Two

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different heating cycles before quenching have been studied and calibration curves of the apparent solid fraction were obtained for the AZ91D magnesium alloy as a function of the quenching temperature and compared to literature.

Considering magnesium alloys have a comparable semisolid temperature range, it was found that a quenching temperature producing about 50 vol% of apparent primary phase in specimens quenched from the mushy state, indicate the practical injection temperature for thixocasting.

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