Effect of Pressure on Solidification Process and Mechanical Properties During Semi-Solid Casting by Computational Fluid Dynamics (CFD)

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Abstract: This paper focused on the development of Computational Fluid Dynamics (CFD) by a mathematical model based on the control volume method, which would enable efficient simulation of the semi-solid processing of complex industrial casting parts. Theoretical basis of the numerical simulation was briefly introduced. The latent heat was incorporated using the effective specific heat. The mass, momentum and enthalpy transport equation for each phase were solved. The application of the model allowed determining the temperature fields in the metal and the mold at 1 sec and the liquid fraction at time step 0.15 s with three different pressure regimes. The calculated pressure distribution and the evolution of liquid fraction through the material were examined and used to clarify their influence further investigated with tensile and hardness testing.

Keywords: Aluminum Alloy, Modeling, CFD, SEMI-Solid Casting, Solidification, Liquid Fraction, Mechanical Properties

1. Introduction

Semi-solid casting process, as its name indicates, is the technological procedure for metals and alloys at semi-solid state. This process was innovated at Massachusetts Institute of Technology (MIT) in the 70’ years of previous century [1-7]. So far, it has been the only casting process which gives the nodular structure instead of the dendritic one. One of its advantages is very high economic effectiveness, expressed in

i) accurate shape of castings, that required less subsequent machining;
ii) higher mold service and so on.

In general, the solid fraction in semi-solid technology varies from 5 to 60% depending on the technology involved. The semi-solid casting method can be devided on two principal kinds: rheo-casting and thixo-casting. Semi-solid casting method can be combined with others to develop new technologies such as rheo-die casting, compo-casting etc. Some authors named their technology “nano-casting” too.

In the case of rheo-die casting technology (i.e. combination of rheocasting with die casting) the turbulent flow of metal under high pressure that involves a lot of gas, resulting in the presence of porority in castings, is substituted by a laminar flow allowing more “peaceful” mold filling due to lower pouring temperature (between liquidus and solidus). However, the viscosity of alloy is certainly higher and, as the result, requires higher pressure. So, the pressure must play a very important role in this process, especially in the temperature distribution inside castings and in phase fraction during solidification process. Observation in real castings is not always a good way; instead, the numerical simulation with computational fluid dynamics is applied in our study.

In the early days of SSM and CRP developments [8-12], it was thought that one had to cool the liquid down into the two-phase region. In recent years, the projects sponsored by the National Research Program KC02-23/06-10 and by the Department of Science and Technology of Hanoi City, were carried out by the research team of the Key Laboratory of Metal Materials Technology to produce a semi-solid structure of the globular primary alpha phase via continuous rheo-casting and rheo-die casting [13-18].
2. Methodology

Computational fluid dynamics (CFD) [19] is a simulation method for fluid flow, heat and mass transfer and the phenomena related to solving the mathematical equations with numerical method. Figure 1 describes the elements in the melt bulk to be simulated via the properties such as flow rate $u$, pressure $p$, specific weight $\rho$ and energy $E$ in the fluid flow equation:

$$\frac{\partial}{\partial t} \int_V \rho f dV + \oint_{\partial V} \rho \vec{V} \cdot d\mathbf{A} = \int_V \rho \vec{F} \cdot d\mathbf{A} + \int_{\partial V} \rho \vec{S} \cdot d\mathbf{A}$$

Equation (1)

In the case of compressible as well as incompressible liquids the continuous equation (2) and mass conservation equation (3) can be applied:

$$\frac{\partial \rho}{\partial t} + \frac{\partial (\rho u)}{\partial x} + \frac{\partial (\rho v)}{\partial y} + \frac{\partial (\rho w)}{\partial z} = 0$$

Equation (2)

$$\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \vec{V}) = S_m$$

Equation (3)

where $\rho$ is specific weight, $\vec{V}$ is partial derivative, $t$ is time, $v$ is velocity and $S_m$ is the mass added to the continuous flow during phase transformation.

This model is applied to simulate the processes involving heat transfer and solid-liquid-vapor transformations, used in the FLUENT software. In this software the target function is enthalpy [20-22]. Enthalpy of a system can be calculated as

$$H = h + \Delta H$$

Equation (4)

$$h = h_{ref} + \int_{T_{ref}}^T c_p dT$$

Equation (5)

Liquid fraction can be determined as:

$$\beta = 0 \text{ if } T < T_{solidus} \text{ and } \beta = 1 \text{ if } T > T_{liquidus}$$

Equation (6)

$$\beta = \frac{T - T_{solidus}}{T_{liquidus} - T_{solidus}} \text{ if } T_{solidus} < T < T_{liquidus}$$

Equation (7)

Latent heat $\Delta H$ can be found as

$$\Delta H = \beta L$$

Equation (8)

where $L$ is latent heat during crystallization process of metals and alloys. The value of latent heat can vary from 0 (for solid) to $L$ (for liquid). In the case of the solidification of multicomponent alloy, the theoretical liquidus temperature can be calculated as follows:

$$T_{liquidus} = T_{melt} + \sum_{solute} m_i Y_i$$

Equation (9)

where $Y_i$ is weight fraction of $i$-solute, $m_i$ is the curvature for $Y_i$. For melting/solidification problems, the governing energetic equation can be written in the form:

$$\frac{\partial}{\partial t} (\rho H) + \nabla \cdot (\rho \vec{V}) = \nabla \cdot (k \nabla T) + S$$

Equation (10)

with $H$ is enthalpy, $\rho$ is specific weight, $\vec{V}$ is liquid velocity, $K$ is the distribution coefficient at solid/liquid interface, and $S$ is internal source of heat. The solution for temperature field is the irrelative solution of flow equations (1), (2) and (3), governing energetic equation (10) and equation for liquid fraction (7) [25].

By the solidification modeling with FLUENT one can take on account the presence of gas gap between the mold and the solidifying alloy, that creates the heat resistance at boundary, which reduce the heat transfer across boundary. [24]

Therefore, the heat flow across boundary, as shown in the Figure 1b, is:

$$q = \frac{T - T_w}{r^2 R_c (1 - \beta)}$$

Equation (11)

where $T_w$ is mold temperature, $k$ is heat conduction coefficient of liquid, $\beta$ is liquid fraction and $R_c$ is heat resistance, which is the inverse quantity of heat conduction coefficient.

3. Modeling of Process

The simulated product was the diesel engine cover RV125 (Figure 1), produced by rheo-die casting. Based on the product configuration and technology design, the appropriate cross-section was chosen for calculation and analyze. This cross-section passed the important parts of the product, such as the canal for oil lubrication, as well as the thick and thin wall of casting.

The numerical simulation to determine the stabilizing time of the alloy crystallization was carried out by used ANSYS FLUENT (ANSYS Workbench - Fluid Flow (FLUENT)) code [25]. The element mesh at curvature or at thin wall was finer for more accurate calculation. The finite element was triangle. The number of element in casting, moving mold and static mold is 7022, 19510, 13902, respectively (Figure 1b).

The chemical composition, thermal-physical properties of A356 are shown in Tables 1, 2 [26] and the technological parameters - in Table 3.

### Table 1. Chemical composition of A356 alloy.

<table>
<thead>
<tr>
<th></th>
<th>Cu</th>
<th>Mg</th>
<th>Mn</th>
<th>Si</th>
<th>Fe</th>
<th>Zn</th>
<th>Ti</th>
</tr>
</thead>
<tbody>
<tr>
<td>[%]</td>
<td>≤ 0.2</td>
<td>0.25-0.45</td>
<td>≤ 0.1</td>
<td>6.5-7.5</td>
<td>≤ 0.2</td>
<td>≤ 0.1</td>
<td>≤ 0.2</td>
</tr>
</tbody>
</table>
4. Results and Discussion

The results of simulation were shown in Figure 2, in which the perfect contact condition between the mold and the casting part was considered. Figures 2a, 2b and 2c show the temperature fields in the metal and the mold, for an initial mold temperature of 473 K at pressures of 1, 50 and 100 atm after one second. It can be seen that the position of the isotherms of the initial mold temperature and initial temperature of semi-solid melt lies on the values of 473 and 863 K. As seen, in the case of the pressure of 1 atm, temperature field of casting part was ranged from 650 to 840 K, the same results are obtained with 50 atm case and from 660 to 860 K for the case of 100 atm.

It can be seen also in the Figure 2, in the case of atmospheric pressure (1 atm), the temperature of the metal in the region 1 (thin sheet) of casting part was lower, varying from 590 to 600 K and descending from the center to the periphery. Solidification orientation seems to be similar to that by conventional casting methods. Temperature field in this case was depending on the geometrical structure of the casting parts.
In the cases of high pressures (50 and 100 atm), the change in heat flow distribution leads to the change of the crystallization direction, so the picture differs from the conventional casting methods. Crystallization process was relatively uniform in the whole of the casting part, the temperature distribution was therefore relatively uniform with higher cooling rate, which must have some effect on the grain size and crystal morphology.

Figure 3 shows the liquid fraction at 0, 15 s after pouring upon the pressures of 1, 50 and 100 atm. As seen, in the case of atmospheric pressure (1 atm), the solidification proceeds as usually from periphery toward center; the grain morphology is columnar-dendritic. The last region to solidify is center with a lot of defects such as porosity, shrinkage that cannot be eliminated afterward. In the case of higher pressure (50 and 100 atm) the directional solidification, according to the direction of pressure, can be observed, that allows control the solidification process by creating an appropriate solidification direction.

To clarify the phenomena, the simulation at regions II and
III under different pressures are taken place.

At region II, under atmospheric pressure (1 atm), the solidification process proceeds from periphery toward the center, and depends on the geometrical configuration of casting. Solidification rate at thin wall is higher than that of thick wall. Under 50 atm pressure, the solidification initiates near the gate (with the highest pressure), and does not depend on the geometrical configuration of casting. The same picture is in the case of 100 atm pressure, the only difference is that the solidification rate is slower. This creates more favorable conditions for transferring of pressure into region III; so, the solidification process there (the farthest region of casting) will happen under pressure, or, in other words, all regions of casting will solidify under high enough pressure, resulting in the finer microstructure.

The density (ρ) on casting with different pressures (1, 50 and 100 atm) showed that the density increased from $\rho = 2.21 \text{ g/cm}^3$ (with 1 atm) to $\rho = 2.59 \text{ g/cm}^3$ (with 100 atm) (the ASTM standard of 2.67 g/cm$^3$) [26]. In Figure 4, the results of the tensile test and the hardness on the casting specimen with pressures; A survey of three cases samples showed that the tensile strengths reached $(240 \div 255) \text{ MPa}$ and the hardness of the samples were greater than 90 HB.

![Figure 4. Mechanical properties on casting with pressures: 1 atm (1), 50 atm (2) and 100 (3).](image)

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5. Conclusions

The simulation of solidification of A356 alloy in rheo-die casting process with Computational Fluid Dynamics (CFD) was successful. The phase fractions were calculated under different pressures. The effect of pressure (50 and 100 atm) was observed as follows:

- The higher pressure applied, the higher solidification rate at thick part of the casting and the finer alloy microstructure obtained.
- The density (ρ) on casting showed that the density increased from 2.21 to 2.59 g/cm$^3$. The tensile strengths reached $(240 \div 255) \text{ MPa}$ and the hardness of the samples were greater than 90 HB.
- The solidification direction was changed, not from periphery to center, but from the part with higher pressure to that with lower pressure. This could result in reduction of such defects like porosity or shrinkage.

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References


