CARBIDIC BAINITIC AND AUSFERRITIC DUCTILE CAST IRON

ŽELIWO SFEROIDALNE BAINITYCZNE I AUSFERRYTYCZNE Z WĘGLIKAMI

This article presents new kinds of carbidic ductile cast iron with different microstructures of the metal matrix. This cast iron was obtained using the Inmold method nodularisation which guarantees strong refining of graphite and the metal matrix components. A different microstructure of the metal matrix of the cast iron was obtained without any thermal treatment (unwrought) by a suitable composition of alloy additives. It was shown that by adding molybdenum, chromium, nickel and copper it is possible to obtain in the cast iron metal matrix consisting of upper bainite, its mixture with lower bainite or ausferrite in the casts with the wall thickness of 3÷25 mm. The process of cast iron crystallization is presented and described with the help of the thermal and derivative analysis (TDA) curves. It also shows the thermal effects from transformation of austenite in solid state.

Keywords: Carbidic Ductile Cast Iron, Bainite, Ausferrite, Thermal and Derivative Analysis (TDA)

W artykule przedstawiono nowe rodzaje żeliwa sferoidalnego z węglikami o różnej mikrostrukturze osnowy metalowej. Żeliwo to otrzymywano stosując sferoidyzację metodą Inmold, zapewniającą dużą liczbę kulek grafitu i rozdrobnienie składników osnowy metalowej. Różną mikrostrukturę osnowy metalowej żeliwa otrzymywano bez stosowania obróbki cieplnej (w stanie surowym) poprzez odpowiednią kombinację ilościową dodatków stopowych. Wykazano, że dodatek molibdenu, chromu, niklu i miedzi w żeliwie sferoidalnym pozwala uzyskać osnowę metalową złożoną z bainitu górnego, jego mieszaninę z dolnym lub ausferrytu w odlewach o grubości ściany 3÷25 mm. Proces krystalizacji żeliwa przedstawiono i opisano za pomocą krzywych analizy termicznej i derywacyjnej (ATD). Pokazano efekty cieplne od przemiany austenitu w stanie stałym.

1. Introduction

The dynamic technical progress sets clear requirements to the materials used for production of machine and device elements. It creates a necessity to work out and use new materials with high mechanical and application properties. It also concerns the ductile cast iron whose world production in 2010 was 22.6 million tons and constituted 25% of the world cast production [1]. In comparison to the previous year it increased by 3.2 million tons. It speaks to the fact that this material is still one of the most eagerly used by material engineers for machine and device elements.

At present the most wear resistible ductile iron is ADI. The microstructure of the metal matrix of this cast iron consists of ausferrite which is the mixture of bainitic ferrite and austenite in the quantity up to 40%. It is obtained using thermal treatment of the casts based on austenitizing and further austempering in the range of temperatures of about 250÷400°C [2].

The bainitic and ausferritic microstructure can also be obtained without using any thermal treatment (unwrought) using a suitable composition of the alloy additive [3÷5]. This technology is however less investigated. Such elements as Mo, Cr, Ni and Cu influence, with different intensity, on the austenite stability in the pearlitic and bainitic range, that is why their suitable composition enables obtaining the metal matrix in ductile cast iron consisting of upper bainite, its mixture with lower bainite or ausferrite during continuous cooling. In bainite, unlike ausferrite, there are carbides separated from the carbon-saturated austenite and ferrite. It is also possible to obtain in cast iron carbides precipitated from a liquid which increases its wear and adhesive wear resistance [6].

In connection with the abovementioned the aim of the current work was to present the possibilities of obtaining of the bainitic and ausferritic microstructure in ductile cast iron by introducing molybdenum, chromium, nickel and copper. The process of cooling and crystallization was shown using the TDA curves. It was stated that it was possible to register the thermal effect when austenite was transforming into upper bainite or ausferrite.

2. The methodology of the research

The original cast iron was melted in the inductive furnace type PI 30 by Elkon with the frequency 3.8 kHz and capacity of 30 kg. The furnace charge was a pig iron with
the chemical composition presented in Table 1 and scrap iron from S235JR steel with the chemical composition according to PN-EN 10025:2002U norm.

### TABLE 1

<table>
<thead>
<tr>
<th>Chemical composition, wt. %</th>
<th>C</th>
<th>Si</th>
<th>Mn</th>
<th>P</th>
<th>S</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>4.44</td>
<td>0.69</td>
<td>0.025</td>
<td>0.046</td>
<td>0.009</td>
</tr>
</tbody>
</table>

The content of silicon and manganese was regulated by additives of FeSi75 and FeMn75 correspondingly. Technically pure: Cr, Mo, Ni and Cu were used as alloy additives.

The chemical composition of the cast iron was tested in the emissive spectrometer with the spark excitation SPECTROMAXx by SPECTRO Analytical Instrument GmbH. The chemical composition presented in Table 1 and scrap iron and their eutectic carbon equivalent C$_{\text{e}}$ and the eutectic saturation coefficient S$_{\text{e}}$.

### TABLE 2

<table>
<thead>
<tr>
<th>Group No</th>
<th>Chemical composition, wt. %</th>
<th>C$_{\text{e}}$ wt. %</th>
<th>S$_{\text{e}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Cr 0.34·Si−0.027·Mn−0.063·Cr+0.074·Cu+0.053·Ni−0.015·Mo</td>
<td>1.49</td>
<td>4.98</td>
</tr>
<tr>
<td>II</td>
<td>Cr 0.30·Si−0.06·Cr+0.18·Cu+0.09·Mn−0.21·Ni</td>
<td>1.20</td>
<td>4.70</td>
</tr>
<tr>
<td>III</td>
<td>Cr 0.32·Si−0.07·Cr+0.18·Cu+0.09·Mn</td>
<td>1.10</td>
<td>4.70</td>
</tr>
<tr>
<td>IV</td>
<td>Cr 0.30·Si−0.07·Cr+0.18·Cu+0.09·Mn</td>
<td>1.10</td>
<td>4.70</td>
</tr>
</tbody>
</table>

The average concentration of P and S in the cast iron was 0.04% and 0.01% correspondingly. The content of Mg in the cast was within the range of 0.04±0.06%.

The eutectic saturation coefficient was calculated according to the formula:

$$S_{\text{e}} = \frac{C_{\text{e}}}{C_{\text{e, cast}}}$$

where:

- $C_{\text{e}}$ – complete content of carbon in the cast iron, wt. %,
- $C_{\text{e, cast}}$ – carbon content in the eutectic according to the stable system, wt. %.

The nodularisation of the cast iron was carried out by Inmold method. The scheme of the bench for cast iron nodularisation and TDA curves registration is shown in Figure 1.

The master alloy (5) is placed in the reaction chamber (4) situating in the ingate system behind the pouring gate (3). The chemical composition of the master alloy is given in Table 3. The master alloy was divided into four groups differing by its chemical composition presented in Table 1 and scrap iron and their eutectic carbon equivalent $C_{\text{e}}$ and the eutectic saturation coefficient $S_{\text{e}}$.

### TABLE 3

<table>
<thead>
<tr>
<th>Chemical composition, wt. %</th>
<th>Si</th>
<th>Mg</th>
<th>Ca</th>
<th>RE$^*$</th>
<th>Al</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>40.0±49.0</td>
<td>3.0±6.5</td>
<td>0.3±1.0</td>
<td>0.3±1.4</td>
<td>0.5±1.2</td>
</tr>
</tbody>
</table>

$^*$ – rare earth elements.

The microstructure of the cast iron was tested on the polished sections etched by nital magnified $\times$ 200 and $\times$ 1000 on the metallographic microscope Eclipse MA200 by Nikon.
3. The research results

3.1. Transformations which take place during crystallization and cooling of ductile cast iron

3.1.1. Non-alloy ductile cast iron crystallization

In order to estimate the influence of the alloy additives on the temperature of the phase transformations and microstructure of the cast iron on the basis of the TDA curves non-alloy cast iron was analyzed (group I, Tab. 2).

In connection with this, Figure 3 (a÷d) presents the representative TDA curves (a, b) and microstructure (c, d) of ductile cast iron with the composition: 3.64% C; 2.56% Si; 0.33% Mn.

![Image of TDA curves and microstructure](image.png)

In follows from the chemical composition analysis that it is hypereutectic cast iron ($C_e = 4.57\%$), however, on the derivative curve there is no thermal effect from the initial graphite crystallization. This phenomenon is described in more details in the work [8]. The DEFH thermal effect comes from the crystallization of the austenite + nodular graphite eutectic. The temperature recalescence is $2\,^{\circ}C$. At the temperature of $1113\,^{\circ}C$ (point H, Fig. 3a) the cast iron finishes crystallization and its microstructure consists of nodular graphite and austenite.

It follows from Fig. 3b that at the temperature of $785\,^{\circ}C$ (point M) the transformation starts in the solid state. The high temperature shows that it starts from the $\gamma \rightarrow \alpha + C_gr$ transformation. Maximum liberation of heat due to austenite transformation takes place at the temperature of $729\,^{\circ}C$ (point N). The transformation ends at the temperature of $695\,^{\circ}C$ (point O, Fig. 3b). In follows from Fig. 3c that after cooling to the ambient temperature the microstructure of the cast iron consists of nodular graphite, ferrite and pearlite. Regarding the small cooling rate of the cast iron in the TDA probe, a double thermal effect from the $\gamma \rightarrow \alpha + P$ (pearlite) transformation is not noticed. The surface percent of the ferrite in cast iron matrix after the end of the crystallization process is about 55%.

3.1.2. Carbidic ductile cast iron crystallization

As it was mentioned, in order to obtain bainite or ausferrite in cast iron without using thermal treatment it is necessary to introduce the alloy additives which, when dissolved in austenite, have an influence on its hardness. To obtain upper bainite in the metal matrix a proper combination of molybdenum and nickel should be used. In connection with this, Figure 4 (a÷d) presents the representative TDA curves and microstructure of the ductile cast iron with upper bainite and carbidies from group II (Tab. 2). In Table 4 the values describing the characteristic points in the TDA curves of studied types of ductile cast iron with carbides and obtained type of metal matrix microstructure are presented.

The chemical composition shows that this is slightly hypo-eutectic ($C_e = 4.20\%$) and that is why its crystallization starts from forming of austenite seeds and their further growth, it is seen in the derived curve $dt/d\tau = \Gamma(\tau)$ in the form of a thermal effect described by points AB (Fig. 4a).
Fig. 4. (a÷d). The TDA curves (a, b) and microstructure (c, d) of carbidic ductile cast iron with the chemical composition: 3.46% C; 2.39% Si; 0.07% Mn; 2.13% Mo; 0.66% Ni.

Then, within the temperature range of 1145÷1078°C the eutectic mixture consisting of austenite and nodular graphite crystallizes (thermal effect BDEFH, Fig. 4a). The synergic additive Mo and Ni caused a decrease in the temperature of eutectic crystallization in relation to the non-alloy cast iron by about 15°C. Carbides M3C form after the end of eutectic γ + graphite crystallization. It is caused by simple microsegregation of molybdenum whose atoms are “moved” by the dendrites of austenite and eutectic cells front. For this reason the last portions of the liquid cast iron are enriched with molybdenum, so the tendency of the cast iron to crystallization increases according to the metastable system. On the crystallization curve the thermal effect is caused by the carbides crystallization described by the HKL points. The cast iron finishes its crystallization at the temperature of 996°C (point L, Fig. 4a). The microstructure of the cast iron at this temperature consists of nodular graphite, austenite and carbides placed on the borders on the eutectic cells.

A certain quantity of Mo dissolved in the austenite during cooling precipitates in the form of carbide Mo2C as a result of variable dissolving of the Mo in the austenite. The remaining quantity of the molybdenum causes a significant increase in the austenite stability within the range of pearlite transformation and insignificant in the bainitic zone. In connection with this, at the temperature of 554°C the transformation γ → BU (upper bainite) starts (point M, Tab. 4, group III). In order to underline the thermal effect from this transformation the time axis is presented in the logarithmic system. The thermal effect from the γ → BU transformation ends at the temperature of 380°C. At the room temperature the microstructure of the cast iron situated in the TDA probe and containing about 2.0% Mo and 0.7% Ni consists of nodular graphite, upper bainite and carbides M3C. Regarding a different cooling rate of certain walls of the sample cast the surface percent of the carbides changed from about 12 to 4% correspondingly in the 3 and 25 mm walls.

Figure 5 (a, b) presents the microstructure of carbidic ductile cast iron containing about 2.0% Mo and 1.5% Ni (group III, Tab. 2).

Regarding the fact that this cast iron is hypereutectic (Ce = 4.62%), its crystallization starts from formation in the liquid iron of the graphite seeds whose size is bigger than the critical and their further growth in the form of the balls. Similar to the non-alloy cast iron the thermal effect from this process does not happen. The DEFH points described the thermal effect from the crystallization of the γ + Cpe eutectic. The increase in the carbon and nickel content with the similar concentration of molybdenum caused an increase in the temperature of the end of eutectic crystallization by 22°C (point H, Tab. 4, group III). It is caused by their graphitization process. Carbides M3C crystalized within the temperature range of 1100÷1001°C (points HKL, Tab. 4, group III). Within the temperature range of 529÷377°C there is a slight thermal effect coming from the austenite transformation (points M and N, Tab. 4, group III). A bigger concentration of nickel in comparison to the previously described cast iron caused a significant decrease in the temperature of the beginning of the γ → BL (lower bainite) transformation (by about 40°C). It follows from the presented data that the increase of the nickel and carbon concentration in the cast iron decreases the temperature of the beginning of the austenite → upper bainite transformation; however, it does not influence the temperature of this transformation end. A relatively high temperature in point N means that the thermal effect from the γ → BL transformation
The chemical composition: 3.75% C; 2.40% Si; 0.33% Mn; 1.41% Cr; 0.51% Mo; 1.53% Cu.

The microstructure of carbidic ductile cast iron with the chemical composition: 3.75% C; 2.40% Si; 0.33% Mn; 1.41% Cr about 1.5% Cu and 0.5% Cr enables obtaining the ausferrite. In the cast iron comprising about 2.0% Mo and 1.5% Ni consists of nodular graphite, ausferrite and carbides placed on the borders of the eutectic cells. The surface percent of lower bainite is from 10 to 3% correspondingly in the wall 3 and 25 mm. The surface percent of carbides is from about 10 to 3% correspondingly in the wall 3 and 25 mm.

In the cast iron containing about 1.5% Mo adding of about 1.5% Cu and 0.5% Cr enables obtaining the ausferritic microstructure of the metal matrix without using thermal treatment. The microstructure of that kind of the cast iron (group IV, Tab. 2) are presented in Figure 6 (a, b).

It comes from the transformation happens implosively, that is why the thermoelement, due to its inertia, is not able to register it. It follows from Fig. 5a, b that the microstructure of the metal matrix of the cast iron comprising about 2.0% Mo and 1.5% Ni consists of nodular graphite, ausferrite and carbides placed on the borders of the eutectic cells. The surface percent of lower bainite is about 55%. The surface percent of carbides is from about 10 to 3% correspondingly in the wall 3 and 25 mm.

The research results presented in the current work enable coming to the following conclusions:

- the synergic additive of about 2.0% Mo and 0.7% Ni enable obtaining upper bainite and carbides in the unwrought metal matrix of ductile iron,
- an increase in the quantity of nickel to about 1.5% causes the austenite transformation into the mixture of upper and lower bainite in the casts with the wall thickness of 3±25 mm,
- introduction of about 0.5% Cr, 1.5% Mo and 1.5% Cu to the cast iron enable obtaining ausferrite without using thermal treatment,
- on the derivative curve there is a thermal effect from the austenite transformation into upper bainite and ausferrite.

Acknowledgements

Scientific project financed from means of budget on science in years 2009÷2012 as research project N N508 411437

REFERENCES


Received: 10 April 2012.