Specific Distribution of Thermal Effects of Graphite Forming Reactions in High-strength Cast Iron

The dependence of thermal effects of reactions in the formation of inclusions of vermicular and globular graphite is substantiated in the article. The calculation of thermal effects according to the heat of formation of reagents in cast iron and their dependence on the temperature factor is given. The distribution of temperatures and carbon content in the eutectic shell is shown, increasing the compact inclusion of graphite due to diffusing carbon from the melt through the austenitic shell.

high-strength cast iron, globular graphite, vermicular graphite, crystallization, thermal effect of reactions

Statement of the problem. High-strength cast irons continue to keep a leading position amongst structural materials in modern engineering. This group of cast irons is characterized by high strength and durability indicators at the level of carbon steels with high level of foundry properties (liquidity, segregation). All the facts show the increase of demand for the use of high-strength cast iron for the manufacture of machine parts and mechanisms. Modern technologies applied in the production of high-strength cast iron make it possible to obtain a set of enhanced properties, which are achieved by modifying the melts with spheroidizing elements and forming graphite inclusions of compact form, as well as alloying. At the same time, the issues of the mechanism of forming globular graphite are still debatable and do not allow reaching a generally accepted theoretical explanation of the phenomena.

It is known that cast iron has thermophysical characteristics, which strongly depend on the nature of the forming structure, the parameters of which depend on local crystallization conditions. Physicochemical processes occurring during crystallization of high-strength cast iron complicate the explanation of the mechanism of formation of inclusions of spherical and compact forms of graphite. The varied hypotheses of the formation of globular graphite and current possibilities of computer simulation do not lead to a generalized theoretical explanation of the phenomena that occur during graphitization. The discovery of the
mechanism of the processes of formation of globular graphite will open possibilities for controlling the structure and properties of high-strength cast iron. That allows developing effective technological processes for obtaining cast products for various purposes.

**Analysis of recent research and publications.** Modern research methods create new opportunities to confirm or disprove the validity of one or another hypothesis about the formation mechanism of globular and vermicular graphite in high-strength cast iron.

There is a number of works [2] in which the formation of globular graphite (after modifying the melt with magnesium-containing additives) is associated with "gas bubbles" of magnesium vapour. However, such hypotheses do not explain the formation mechanism of vermicular graphite inclusions in magnesium iron. Attempts are made to transform the bubble theory with new experimental data, for example, the discovery of new modification of carbon - fullerenes and their detection in iron-carbon alloys [3].

Studies on the formation of graphite nucleus [11-13] show that the centre of the graphite globule contains Ca, Mg, S, Si, and O, therefore, calcium and magnesium sulphides (CaS, MgS) are surrounded by an oxide layer and are considered as substrates for graphite nucleation. With the increase of magnesium in the melt, the number of magnesium-containing compounds increases and they affect the formation of the microstructure of the matrix, graphite and inter-granular boundary. Growth of graphite globules, their size and shape are determined by such factors as: the nature of the nucleus; the ability of the nucleus to adsorb impurities and gases and the rate of carbon diffusion.

The authors in the work [7] associate the formation of a globular graphite form with the purification of the melt from surfactants of sulphur, oxygen and other impurities. As a result of this cleaning, the ratio of surface tension on the interface surfaces between the melt and the base and prismatic planes of graphite inclusions changes. This theory allows explaining the difference in the formation of globular and vermicular graphite, but does not explain a number of issues. They are the correlation between the refining and spheroidizing action of chemical elements, the influence of the rate of crystallization of the melt on the formation of graphite.

A number of researchers [8, 9] associated the formation of compact forms of graphite with their growth in the process of eutectic crystallization due to the diffusion of carbon through the austenitic shell. The calculations confirmed the sufficiency of the eutectic crystallization time of the modified melt for the growth of compact inclusions due to carbon diffusion through the austenitic phase. It is noted that the conditions of heat transfer from the surfaces at the crystallization fronts on which exothermic reactions occur: crystallization of austenite and separation of graphite out of a liquid or solid carbon solution in iron, play a significant role in the processes of graphite formation during the crystallization of cast iron. The results of computer simulations indicate that these thermal effects can have a significant impact on the development of diffusion and liquidation processes.

**Statement of the objective.** Thus, the purpose of this work is to clarify the role of thermal effects of reactions in the process of crystallization of austenite and the formation of graphite inclusions in modified high-strength cast iron.

**Main material.** The studies were carried out on samples of high-strength cast iron, which was smelted in an induction furnace, the chemical composition of which corresponded to the ВЧ 500-2 brand (DSTU 3925-99). The melt was modified in the ladle by sandwich method to Mgresidual portion = 0.035% by VL 63 (O) ligature with Mg = 6.3%, Ca = 1.8%, Si = 46%, Fe – the rest.

Slides preparations were carried out according to conventional methods. Metallographic analysis was performed using a MIM-7 microscope.

In the first phase of the study, the modified cast iron was poured into graphite crucibles with the capacity of 1 kg and 5 kg. After a period of time which is sufficient for the formation of crust up to 10 mm thick on the surface of the crucible, the non-crystallized liquid
residue in the centre of the crucible was poured out and subsequent rapid cooling of the crystallized crust with aqueous brine was carried out. Microstructure studies have shown that the placed inclusions of globular graphite, even directly near the inner surface of the crust are surrounded by a ferrite shell. Inclusions of vermicular graphite come to the surface of the section with the ends surrounded by ferrite only on the sides. This indicates that the ends of the inclusions of compact and vermicular graphite in some periods of the crystallization process had contact with the melt. At the same time, it is noted that hardened specimens in the end region, the inclusions of vermicular graphite have a significantly smaller thickness (up to 3 microns), compared to the thickness of the inclusions in the zones that are farther from the crystallization front. This indicates a significant role in further formation of the diffusion growth of inclusions of compact graphite through the austenitic shell. Thus, after the formation of the nucleus of globular graphite, it is completely surrounded by its austenitic shell, and when forming vermicular graphite it is only partial.

According to the method of calculation of thermal effects of reactions on the heat of the formation of reagents in cast iron, on the basis of the initial data given in [10], a thermodynamic calculation of thermal effects of reactions was carried out:

\[ \text{Fe} \rightarrow \text{Fe}^{\text{tv}}; \quad [\text{C}]_{\text{Fe}} \rightarrow \text{C}^{\text{gr}}; \quad [\text{C}]_{\text{Fe}} \rightarrow \text{Fe}_{3}\text{C}. \]

The calculations of thermal effects of the given reactions in the cast iron were carried out by the formula:

\[ K = a + b \cdot 10^{-3} T + c \cdot 10^{5} T^{-2} \text{Дж/моль} \cdot \text{К} \quad (1) \]

The change of enthalpy was defined:

– for the reaction \( \text{Fe}^{\text{p}} \rightarrow \text{Fe}^{\text{tv}} \): \( \Delta H^{0}_{298} = 3 \Delta H^{0}_{\text{Fe}^{\text{tv}}} - 3 \Delta H^{0}_{\text{Fe}^{\text{p}}} = 0 \);

– for the reaction \( [\text{C}]_{\text{Fe}} \rightarrow \text{C}^{\text{gr}} \): \( \Delta H^{0}_{298} = \Delta H^{0}_{\text{C}^{\text{gr}}} - \Delta H^{0}_{[\text{C}]_{\text{Fe}}} = 0 \);

– for the reaction \( [\text{C}]_{\text{Fe}} \rightarrow \text{Fe}_{3}\text{C} \): \( \Delta H^{0}_{298} = \Delta H^{0}_{\text{Fe}_{3}\text{C}} - \Delta H^{0}_{[\text{C}]_{\text{Fe}}} = 22,61 \text{ кДж/моль.} \)

The coefficients of temperature dependency of the change of heating capacity \( \Delta C_{p} \) in the process of reaction \( \text{Fe}^{\text{p}} \rightarrow \text{Fe}^{\text{tv}} \) were:

\( \Delta a = 14,11 - 41,87 = -27,76; \quad \Delta b = 29,27 - 0 = 29,27; \quad \Delta c = 1,80 - 0 = 1,80. \)

And the change of heating capacity of \( \Delta C_{p} \) system, as a result of the reaction \( \text{Fe}^{\text{p}} \rightarrow \text{Fe}^{\text{tv}} \) was:

\[ \Delta C_{p} = -27,76 + 29,27 \cdot 10^{-3} T + 1,80 \cdot 10^{5} T^{-2} \text{ кДж/моль} \cdot \text{К.} \]

The coefficients of temperature dependency of the change of heating capacity in the process of reaction \( [\text{C}]_{\text{Fe}} \rightarrow \text{C}^{\text{gr}} \) were:

\( \Delta a = 16,75 - 0 = 16,75; \quad \Delta b = 4,27 - 0 = 4,27; \quad \Delta c = -8,37 - 0 = -8,37. \)

Heating capacity change of \( \Delta C_{p} \) system as a result of reaction \( [\text{C}]_{\text{Fe}} \rightarrow \text{C}^{\text{gr}} \) was defined by the ratio:

\[ \Delta C_{p} = 16,75 + 4,27 \cdot 10^{-3} T + (-8,37) \cdot 10^{5} T^{-2} \text{ кДж/моль} \cdot \text{К.} \]

For the reaction \( [\text{C}]_{\text{Fe}} \rightarrow \text{Fe}_{3}\text{C} \) the calculated coefficients of temperature dependency of the change of heating capacity were:

\( \Delta a = 82,23 - 0 = 82,23; \quad \Delta b = 83,73 - 0 = 83,73; \quad \Delta c = 0 - 0 = 0. \)

The change of heating capacity of \( \Delta C_{p} \) system of the reaction \( [\text{C}]_{\text{Fe}} \rightarrow \text{Fe}_{3}\text{C} \):

\[ \Delta C_{p} = 82,23 + 83,73 \cdot 10^{-3} T + 0 \cdot 10^{5} T^{-2} \text{ кДж/моль} \cdot \text{К.} \]

Calculated heating effect \( \Delta H^{0}_{1} \) of the reaction \( \text{Fe}^{\text{p}} \rightarrow \text{Fe}^{\text{tv}} \) for the temperature interval 1103…1401оС was \( \Delta H^{0}_{1103\rightarrow1401} = 43,368 \text{ кДж/моль} \).

Similarly, heating effects for the reaction \( [\text{C}]_{\text{Fe}} \rightarrow \text{C}^{\text{gr}} \) were defined in the temperature interval 1152…1348оС and for the reaction \( [\text{C}]_{\text{Fe}} \rightarrow \text{Fe}_{3}\text{C} \) in the temperature interval 1170…1374оС. The results of the assessment of heating effects of the reactions are shown in
fig. 1 in the form of histogram.

1 – reaction \( Fe_p \rightarrow Fe_{tv} \); 2 – reaction \([C]_{Fe} \rightarrow Crp\); 3 – reaction \([C]_{Fe} \rightarrow Fe_{3C}\)

Figure 1 – Dependency of thermal effects of reactions on temperature factor

Source: developed by the author

Mole particles (content) of Fe and C in cast iron equal 96% and 4% respectively:
\[
m_{Fe} = m_{(cast \ iron) \cdot m \ particle (Fe)} = 100 \cdot 0.96 = 96 \ g;
\]
\[
m_{C} = m_{(cast \ iron) \cdot m \ particle (C)} = 100 \cdot 0.04 = 4 \ g.
\]

Molar mass of Fe and C are: \( M_{Fe} = 56 \ g/mole \); \( M_{C} = 12 \ g/mole \).

The quantity of Fe and C in (100 g) cast iron, that is their molar particle are:

\[
n_{Fe} = \frac{m_{Fe}}{M_{Fe}} = \frac{96 \ g}{56 \ g/mole} = 1.714 \ \text{mole};
\]
\[
n_{C} = \frac{m_{C}}{M_{C}} = \frac{4 \ g}{12 \ g/mole} = 0.333 \ \text{mole}.
\]

The amount of heat released during the crystallization of the austenitic phase along the boundary of the melt-austenite separation is 76.69 kJ per 1 kg of melt. Accordingly, the amount of heat released on the austenite-compact graphite inclusion boundary is 15.39 kJ per 1 kg of melt. This nature of the distribution of heat release areas determines that the surrounding inclusion of compact graphite is the solid austenitic phase and it will have a substantially higher temperature than the surrounding melt (Fig. 2).

The existence of a continuous overheated austenitic shell is possible due to the reduced carbon content and, consequently, an increase in the solidus temperature. In its turn, it creates the conditions for the intense diffusion of carbon through austenite from the melt of the eutectic composition to the surface of graphite inclusions. Special thermophysical conditions of
formation of austenitic shells lead to high content of silicon and reduced manganese, which results in the formation of ferrite shells around the inclusions of compact graphite.

Conclusions.
1. The calculation of thermal effects affecting to the heat of the formation of reagents in cast iron was carried out and their dependence on the temperature factor was established.
2. The role of thermophysical processes in the formation of inclusions of vermicular and globular graphite with their growth due to the diffusion of carbon from the melt through the austenitic shell has been experimentally and theoretically substantiated.

References
р) вуглецю з вузлами вермикулярного графіту парциальної термічної обробки вуглецю з вузлами вермикулярного графіту парциальної термічної обробки вуглецю з вузлами вермикулярного графіту парциальної термічної обробки вуглецю з вузлами вермикулярного графіту парциальної термічної обробки вуглецю з вузлами вермикулярного графіту парциальної термічної обробки вуглецю з вузлами вермикулярного графіту парциальної термічної обробки вуглецю з вузлами вермикулярного графіту парциаль