

THE CASTING PROCESS AND HIGH TEMPERATURE OXIDATION RESISTANCE OF HIGH CHROMIUM CAST IRON GRATE BAR

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Abstract:

The solidification process of a high chromium cast iron grate bar used for sinter machine was simulated, and high temperature oxidation resistance was also investigated. The simulation result shows that sequence solidification can be achieved and no shrinkage cavity and porosity were observed. Based on the analysis of the microstructure, it could be known that the grate bar was well protected by the Fe_3O_4 , Fe_2O_3 and Cr_2O_3 oxide films at temperatures lower than $800^\circ C$.

1 Introduction

A grate bar is an important part of sintering machine. When used, it usually has to bear very complex working conditions like strong impact force, high temperature and severe wear, which make it easier to get damaged [1-3]. In the past, the grate bar was widely manufactured by common gray cast iron, which was found to have many quality problems such as poor high temperature oxidization resistance, short service life, and high consumption, etc. In order to improve the quality, a lot of research work has been done on the grate bar materials. Many kinds of cast iron have been developed, such as silicon nodular iron, high chromium cast iron, and multi-elementally alloyed high chromium cast iron [4]. Among the irons, high chromium cast iron is considered to be the most excellent wear resistant and high temperature oxidation resistance material [5]. For the high content of chromium, Cr_2O_3 membrane will be formed on the surface, which can help to prevent or delay the oxidation. Therefore, high chromium cast iron grate bars usually present

good thermal stability and oxidation resistance [6]. Ramírez-Ramírez et al. [7] did investigation on high temperature oxidation behavior of high-chromium white cast iron in atmospheres that include dry air and a mixture of dry air and water vapor. They pointed out that the presence of water vapor and the chromium content has an obvious effect on oxidation rate by a mechanism that involved chromium depletion of oxide as a result of hydroxide formation. Efremenko et al. [8] studied the oxidation and decarburization behavior of 14.55 wt pct Cr-cast iron at $1000^\circ C$ to $1150^\circ C$ in a dry atmosphere, and decarburization phenomenon appeared with the form of secondary and eutectic carbide dissolution accompanying oxidizing heating. Zhang et al. [9] deemed that the oxide type in high-chromium cast iron depends on chromium content, and best oxidation resistance can be obtained with chromium content of 15wt.%.

A kind of high chromium cast iron was designed and used for a grate bar. The grate bar was manufactured by lost foam casting, because it can promote surface smoothness and reduce subsequent

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machining. To prolong the working life of the bar, casting process was simulated to eliminate the defects using View-cast software, and effect factors of high temperature oxidation were also discussed.

2 Numerical simulation of casting and solidification

2.1 The chemical composition design

Chromium is the main alloy element, which forms Cr_2O_3 membrane on the surface at high temperature to improve the heat and oxidation resistance of the casting. It is also worthwhile to mention that the ratio of Cr/C has an effect on the type of carbide. When the content of Cr is higher than 11% and rate of Cr/C is higher than 3.5%, M_7C_3 carbide forms in the high chromium iron [10]. The hardness of M_7C_3 carbide is as high as 1300HV-1800HV, which is useful for improving strength and wear resistance of the iron. Therefore, appropriate content of Cr and ration of Cr/C are expected to get more M_7C_3 carbide.

The designed chemical composition of high chromium iron used for the grate bar is shown in Table 1.

Table 1. The designed chemical composition of high chromium iron

Element	C	Cr	Si	Mn	P	S
Content (wt.%)	1.6-2.4	15-18	1.5-2.2	≤ 1.0	≤ 0.1	≤ 0.05

2.2 Casting process design of the grate bar

Grate bar is a small and simple-structure casting. It weighs about 3.7 Kg with the average thickness of about 15 mm. Considering the requirement of batch production, high dimension accuracy, high quality and integrated performance, lost foam casting was used to take place of the traditional sand casting [11].

In order to improve the production efficiency, three-layer modeling of castings was adopted for this case. The castings were connected with the ingrates and six ingrates with the section diameter of 40 mm were settled. The gating system is shown in Fig. 1. 42 castings are arranged evenly on the six ingrates. The size of the sprue and ingrate are 40 mm×40 mm

and $\phi 40$ mm. The pouring temperature is controlled to be 1570°C.

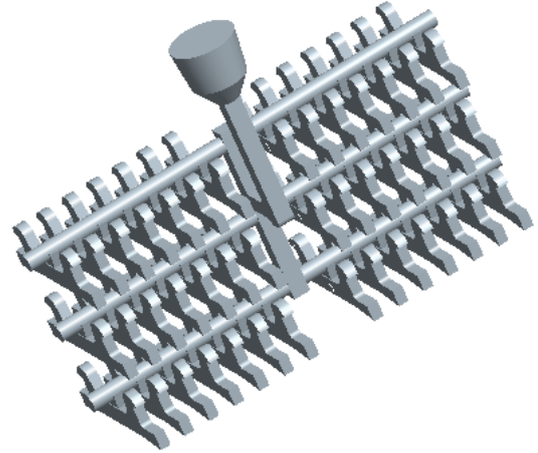


Figure 1. Three-dimensional figure of the Grating system.

2.3 Simulation of casting and solidification of the bar

View Cast is useful software that is effective to simulate the solidification temperature field, solidus fraction, mould-filling time and velocity field during the filling process. Usually, it is also used to predict casting defects and optimize gating system [12, 13]. Before simulation, three-dimensional model is generated by Pro/E and is exported in *.stl file format. The casting and gating system are divided into 2 million grids. The initial and gating temperatures are set at 25°C and 1570°C respectively, and negative pressure is -400 KPa. The simulation results of mould-filling and solidification are represented respectively in Fig. 2 and Fig. 3.

Figure 2 presents the six stages of the mould-filling. The diagram demonstrated that the gating process is smooth and there is not turbulence in the pouring process.

As shown in Fig. 3, when the solidification time was 135 seconds, both sides of the casting began to solidify. After 335 seconds, the solidification area expanded from lateral to internal gradually. As the size of the casting is small with very thin thickness, the solidification rate was very fast, and complete solidification in the castings was obtained after 415 seconds. However, there was still incomplete solidification in the middlesprue, ingrate and runner, which led to more easily feeding.

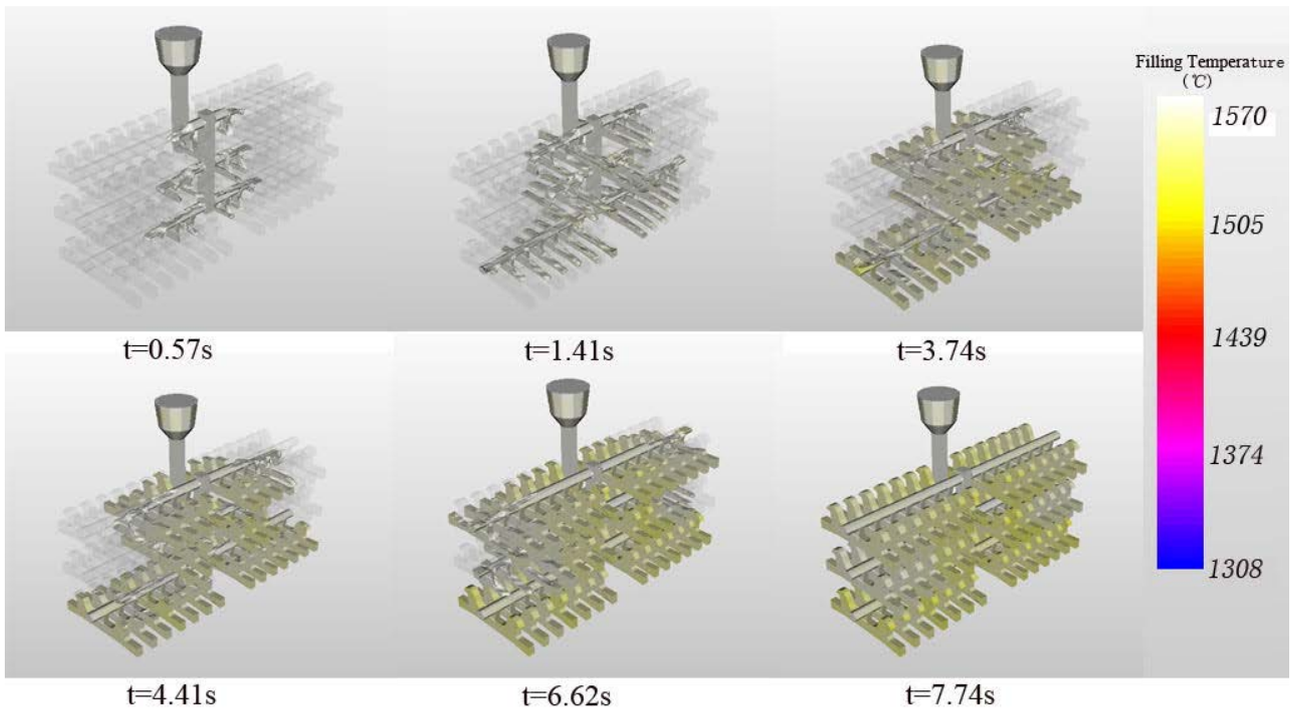


Figure 2. The simulation results of mould-filling after different times.

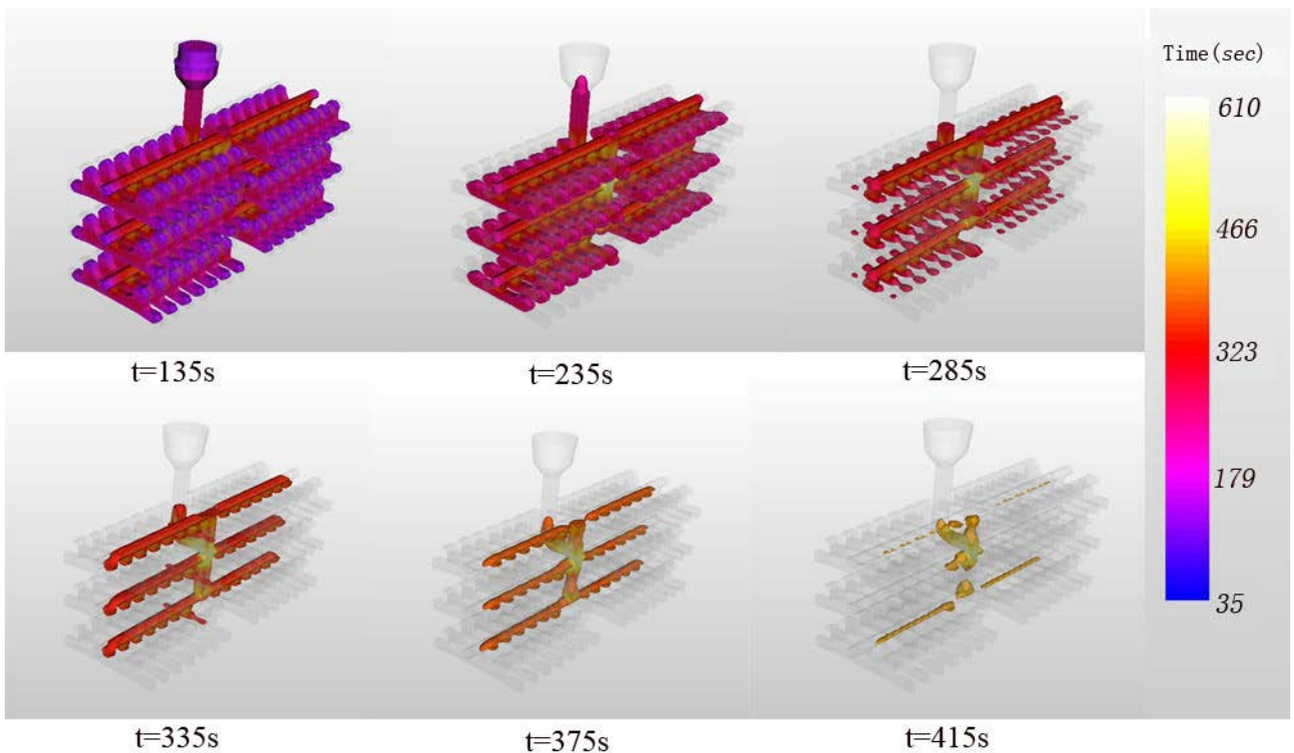


Figure 3. The simulation results of solidification after different times.

The simulation prediction of defects distribution in the castings solidified is completely illustrated in Fig. 4. The transparency in the figure indicates that

the metal content is above 95% and that these parts are deemed as non-defective. It can also be found that large scale of shrinkages exists in the ingrate

and sprue cup, while none in the casting. Therefore, the designed casting process is suitable for manufacture of the grate bar. Fig. 5 shows the grate bar produced by the designed casting process. It can be observed that surface smoothness of the casting is excellent.

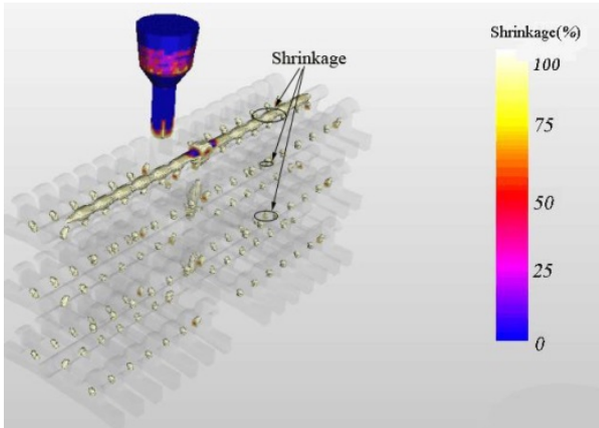


Figure 4. The simulation forecast map of defects in grate bar.



Figure 5. Morphology of grate bar casted by the designed process.

3 High temperature oxidation resistance of the bar

3.1 High temperature oxidation test method

While sintering machine is working, the grate bars are in the alternating hot and cold working environment. The bar may be heated to the temperature in range of 700°C to 900°C, which requires well high temperature oxidation resistance. Thus, the temperature of high temperature oxidation test was set between 700°C and 900°C.

Before the test, samples cut from the bar were firstly ground into the size of 4 mm × 10 mm × 20 mm. All samples were cleaned by ultrasonic with acetone solution and anhydrous ethanol orderly. Then these samples were divided into 8 groups, and each group had three samples. Second, the crucible roasted for 1 hour at 150°C for holding samples. The crucible and the sample were numbered together and weighted using an analytical balance with the accuracy of 0.0001 g. At last, the crucible containing a sample was put into a furnace to be tested. Four groups were respectively heated to 600°C, 700°C, 800°C and 900°C, inside a high temperature resistance furnace protected by argon gas, while the other four without the protected gas. After being kept for 5 hours, the crucibles were taken out and were air cooled to room temperature. Again the crucibles with the sample were weighted together.

According to the national standard (GB/T 13303-91), the oxidation rate would be calculated as the following formula.

$$K = \frac{m_l - m_o}{s_o \cdot t} \quad (1)$$

Where K is weight gain on per square meter at per hour, m_l is the weight of the crucible containing sample after tested, m_o is the weight of that before tested, S_o is the superficial area of sample and t is the keeping time at the tested temperature. Based on the test data, average weight gains of the samples on per square meter at per hour are calculated, shown in Table 2.

Table 2. Average weight gains of the samples tested at different temperatures with or without protected gas

Experimental conditions	Test Temperature	Average weight gain on per square meter at per hour
	°C	g/m ² •h
Non protected	600	0.67076
	700	0.56127
	800	0.90679
	900	3.45811
Protected by argon gas	600	0.08990
	700	0.07425
	800	0.13708
	900	0.69240

3.2 The analysis of high temperature oxidation resistance

It is obvious to see from Table 2 and Figure 6 that the weight gain on per square meter at per hour is 0.67076 g after being kept at 600°C for 6 h and cooled to room temperature without argon protection. When the temperature was raised to 700°C, the oxidation rate is slightly decreased to be 0.56127 g/m²•h. When the test temperature reached 800°C, oxidation rate is 0.90679 g/m²•h, which is about 35% higher than that at 600°C. The oxidation rate is sharply increased to be 3.45811 g/m²•h, about 5 times of that at 600°C, with raising the test temperature to 900°C.

At high temperature, iron is easily oxidized on the surface of the sample. And the formation of FeO, Fe₃O₄ or Fe₂O₃ oxide is related to the temperature. Below the temperature of 560°C, the oxide film mainly consists of Fe₃O₄ and Fe₂O₃, which have higher density and small lattice constant. The film can prevent the diffusion of oxygen atoms and then delay the oxidation of iron. When the temperature is over 560°C, FeO layer will be formed under the oxide films of Fe₃O₄ and Fe₂O₃. FeO layer has a simple lattice structure, which is the omission solid solution of iron atoms. It is easier for oxygen atoms to diffuse through FeO layer combining with iron atoms and aggravate iron oxidation [14].

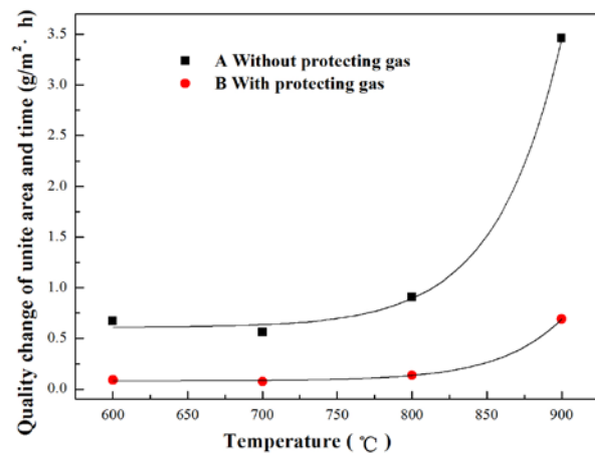


Figure 6. Oxidation rates of the tested samples with or without protection by argon gas under different temperatures.

Moreover, chromium content of the experimental grate bar was very high, Cr atoms would also combine with oxygen atoms, then Cr₂O₃ oxide film formed. Cr₂O₃ oxide film firmly combined with the casting and prevented the diffusion of oxygen atoms. Furthermore, Cr element raised the formation temperature of FeO. As a result, the oxidation resistance temperature was promoted. By the corporate protection of Fe₃O₄, Fe₂O₃, Cr₂O₃ films, the sample had well oxidation resistance at temperature in range of 600°C to 800°C, without protected argon gas. The oxidation rate of the sample was delayed to be only about 0.56127 g/m²•h to 0.90679 g/m²•h, as shown in Fig. 6.

SEM was used to investigate the oxide film formed at 800°C, shown in Fig. 7. It can be observed that the content of Cr sharply increases in the area where the scanning distance is 1.6 μm, while the content of Fe is less. When the scanning distance reaches about 7.2 μm, the content of Cr decreases rapidly, while that of Fe increases sharply. It is obvious that a 5.6μm-thick oxidation layer existed between resin and matrix of the sample. Variations of the concentrations of Cr, Fe and O indicate that the outer and the inner layer of the oxide film were consisted of equivalent amount of iron oxide and chromium oxide. However, in the middle layer, the film was mainly chromium oxide, considered as Cr₂O₃, while that of the iron oxide was few. It is thought that this kind of middle layer of the oxide film is conducive to delay the diffusion of oxygen and improve high temperature oxidation resistance of the sample.

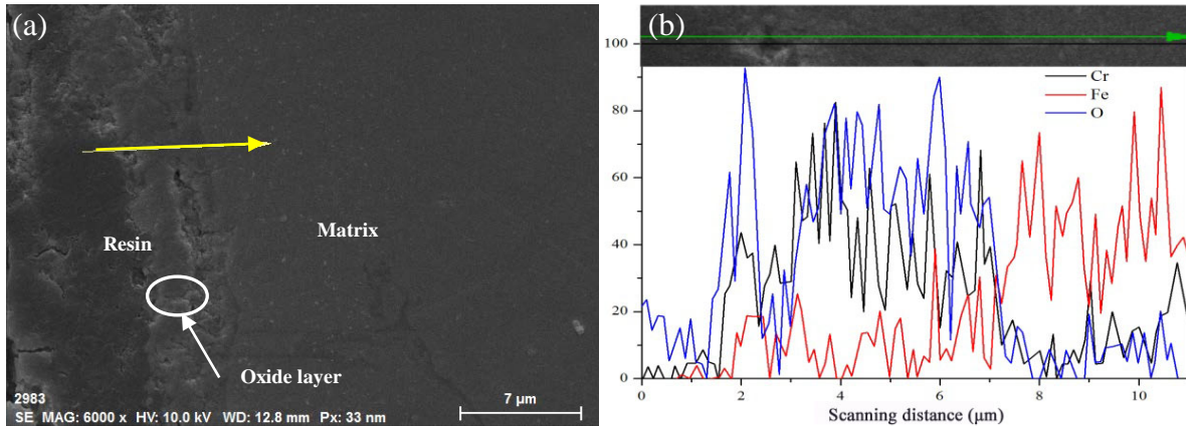


Figure 7. Linear SEM-EDS scanning of the oxide layer formed on the sample tested at 800 °C without protected gas.

The result of Linear SEM-EDS scanning of the oxide film formed at 900°C is shown in Fig. 8. It can be seen that the thickness of the oxide film is about 11.5 μm (scanning distance from 1.5 μm to 13 μm), much thicker than that formed at 800°C. It has also been found that the concentration of iron is about equal to that of chromium in the majority of the oxide film, which reduces the compactness. In the inner layer close to the matrix, the concentration of iron is higher than that of chromium, which is considered to be mainly consist of FeO. This kind of oxide film can not prevent the diffusion of oxygen, leading to severe oxidation of the bar. This phenomenon is analyzed to be caused by dissolution of chromium. Since chromium is a strong carbide

forming element, it will dissolve in cementite to form alloy cementite with the sample heated at 900°C. As a result, Cr₂O₃ membrane reduced and broke gradually, then oxygen atoms diffused inside, leading to rapid oxidation and consequent significant increase of the weight gain. When oxidation temperature was high, the frequent contact, collision, and friction between the grate bar and sintering material result in the fast growth, abscission, and sprout of oxide film on the surface of the grate bar. At last, the grate bar breaks down. In order to improve the service life of the grate bar, working temperature must be controlled strictly.

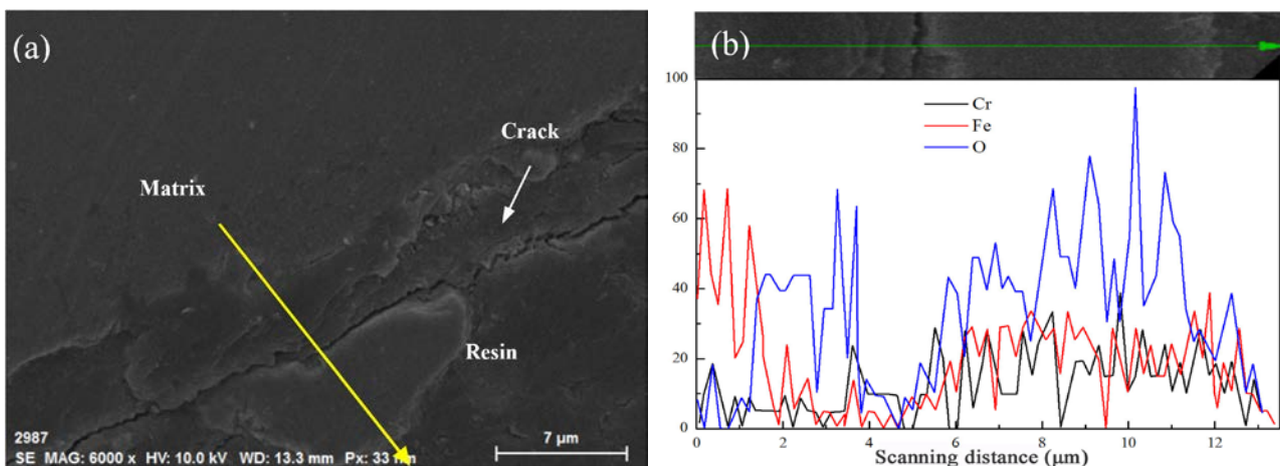


Figure 8. Linear SEM-EDS scanning of the oxide layer formed on the sample tested at 900 °C protected by argon gas.

In the test, it has also been found that the weight gain is lower by near an order of magnitude on the sample protected by argon gas, compared with the non-protected, shown in Fig. 6(b). Only when the test temperature was raised to 800°C, the oxidation rate increased gradually. The oxidation rate at 900°C increased substantially, similar to that at air atmosphere, and reached 0.69240 g/m²·h. Fig. 9 is the linear SEM-EDS scanning of the oxide layer cross section under 900°C with

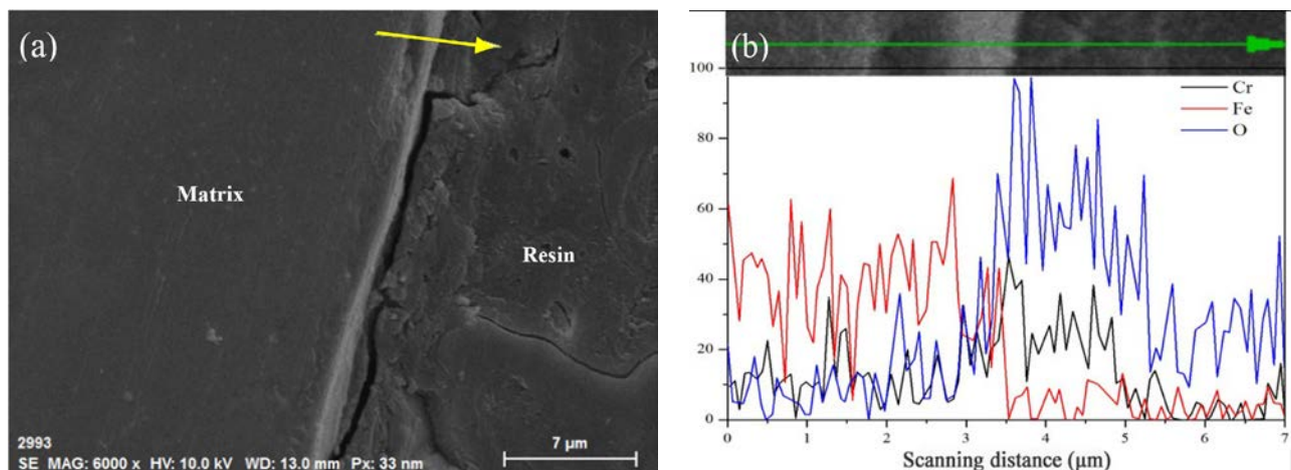


Figure 9. Linear SEM-EDS scanning of the oxide layer formed on the sample tested at 900 °C without protected gas.

Conclusion

A kind of high chromium cast iron was designed and used for grate bar. In order to reduce or eliminate the cast defects, casting process was designed and simulated. High temperature oxidation resistance of the manufactured bar was also investigated according to the working conditions. The main results are shown as follows:

- (1) Chemical composition of the grate bar was shown as follow: 1.6%~2.4% carbon, 15%~18% chromium, 1.8%~2.2% silicon, ≤1.0% manganese, ≤0.01 phosphorus, ≤0.05 sulphur.
- (2) The simulation results show that the casting process parameters and pouring system size can satisfy the requirement of a high chromium grate bar, with the ingrate diameter of 40 mm, the middle sprue, runner and sprue size of 40 mm × 40 mm. Defects of shrinkage cavity and porosity can be eliminated in the grate bar casting.
- (3) The high temperature oxidation test shows that weigh gain increases slowly with the temperature increase from 600°C to 800°C, while increases

sharply at temperature of 900°C. Oxidation mainly occurs in heating and insulation process and is notably affected by temperature and the consequent constitute of the oxide film.

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