Analysis of Micro-Segregation of Solute Elements on the Central Cracking of Continuously Cast Bloom

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Abstract: On the basis of the Brody–Flemings model and modified Voller–Beckermann model, an analytical model of micro-segregation is established by considering the actual solidification cooling conditions of bloom. According to the developed model, the interdendritic solute distribution at the origin of the cracking gap is obtained. It is found that both phosphorus and sulfur have quite severe segregation, but both carbon and manganese have slight segregation; these results agree well with the semiquantitative analysis results of the scanning electron microscope (SEM). At the same time, the interdendritic segregation leads to an enhanced increase in the temperature range of crack formation; correspondingly, the possibility of cracking significantly increases and, thus, element segregation is the internal cause of crack formation. On the other hand, taking into account heat transfer, phase transformation, and metallurgical pressure, the strain of the solid shell is revealed through finite element software. When the solid shell thickness is equal to the distance of 90 mm between the opening point of the crack and the inner arc side, the tensile strain of the solid front is much bigger than the critical strain, which meets the external cause of crack formation; therefore, reasons for the cracking of blooms are successfully found.

Keywords: solute element; micro-segregation; central crack; analytical model; the strain of the solid shell

1. Introduction

Mathematical models can be used to analyze the quality of metallic materials [1]. For continuous casting bloom, central cracks are one of the major defects which appear during the casting process [2]. It is very important to predict and control the central crack in the production of high-quality and defect-free products in the steel industry. According to a previous investigation [3,4], the conclusion obtained was that the interdendritic solute redistribution has a significant impact on the formation of central crack. Carton steel has two typical characteristic temperatures under the high-temperature condition: the liquid impenetrable temperature (LIT) and the zero ductility temperature (ZDT). Many studies [5–9] support the point that internal cracks usually form in the temperature range from the LIT to the ZDT [9], and the solid fractions of 0.9 and 1.0 are believed to correspond to the LIT and ZDT, respectively. In view of the above, analyzing the solute micro-segregation and its effect on the LIT and the ZDT has practical significance for the control of internal cracks. Under the cooling rate of 0.5 °C/s, Kobayashi [10] gave the redistribution of solute elements C, Si, Mn, P, S, and Fe–C phase diagrams under nonequilibrium solidification using a mathematical model with the δ/γ phase transformation taken into account. Ueshima [11] assumed that the transverse cross-section of dendrite was approximated by a regular hexagon, and the interdendritic solute distribution with δ/γ phase transformation

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was analyzed during the solidification. Depending on the model of Ueshima, under the cooling rate of 0.17 °C/s and the dendrite spacing of 1000 µm, Kim [3] discussed in detail the effect of solute micro-segregation in a continuously cast strand on surface longitudinal cracks.

At present, there are few papers focusing on the investigation of solute micro-segregation at the end of the columnar crystal zone of blooms. In view of the severe central cracks of blooms in domestic corporations, it is very necessary to find the formation mechanism of the cracks. In this paper, using the developed analytical model, the segregation of solute elements at the origin of the cracking gap located at the end of columnar crystal zone is obtained by taking into consideration the actual solidification cooling conditions, and the effect of solute segregation on the temperature range of crack formation from the LIT to the ZDT is discussed in detail. In addition, the strain of the solid shell during solidification is given through the finite element software (MSC Marc, MSC Software Corporation, Newport Beach, CA, USA), so as to find the reasons for the formation of central cracks.

2. Analytical Model

In this model, taking into account the Brody–Flemings model and the modified model suggested by Voller and Beckermann, the governing equations of the model were obtained as follows [12,13]:

\[ C_{L,i} = C_{0,i} \left[ 1 - \left( 1 - 2\alpha_i' k_i \right) \cdot f_S \right] (k_i - 1)/(1 - 2\alpha_i k_i), \]  
\[ \alpha_i' = \alpha_i + \frac{1}{2} \exp\left(-\frac{1}{\alpha_i} \right) - \frac{1}{2} \exp\left(-\frac{1}{2\alpha_i} \right), \]  
\[ \alpha_i' = \alpha_i + \alpha^\prime_i, \]  
\[ \alpha^\prime_i = 0.1, \]  
\[ t_f = \frac{T_L - T_S}{C_R}, \]

where \( C_{L,i} \) is the concentration of solute element \( i \) in liquid phase (wt. %), \( C_{0,i} \) is the initial concentration of solute element \( i \) in liquid phase (wt. %), \( k_i \) is the equilibrium partition coefficient of solute element \( i \), \( f_S \) is the solid fraction, \( \alpha_i \) is the back-diffusion parameter, \( \alpha_i' \) is the modified value of \( \alpha_i \), \( D_{S,i} \) is the diffusion coefficient of solute element \( i \) in solid phase (cm²/s), \( t_f \) is the local solidification time (s), \( T_L \) is the liquidus temperature (°C), \( T_S \) is the solidus temperature (°C), \( C_R \) is the cooling rate (°C/s), and \( L \) is the length scale of the micro-segregation domain (µm).

In the present study, the length scale was the final primary dendrite arm spacing \( \lambda \), which was calculated as a function of cooling rate \( C_R \) and nominal steel carbon content (wt. %) from the following equation [14]:

\[
\begin{cases}
\lambda = 278.748(C_R)^{-0.206}([\%C])^{-0.316} + 2.033([\%C]) & 0 < [\%C] \leq 0.15 \\
\lambda = 278.748(C_R)^{-0.206}([\%C])^{-0.0189} - 0.492([\%C]) & 0.15 < [\%C] \leq 1.0
\end{cases}
\]

The liquidus and solidus temperatures depend on steel composition as follows [15]:

\[ T_L = T_{pure} - \sum_i m_i \cdot C_{0,i}, \]
\[ T_S = T_{pure} - \sum_i n_i \cdot C_{0,i}, \]

where \( T_{pure} \) is the fusion temperature of pure iron (1536 °C), and \( m_i \) and \( n_i \) denote the slopes of the liquidus and solidus line of each solute element in the pseudo-binary Fe-phase diagram, respectively.
The solid/liquid interface temperature $T_{\text{int}}$ is obtained by summing the contributions of all alloying elements.

$$T_{\text{int}} = T_{\text{pure}} - \sum_{i} m_{i} C_{L,i},$$  \hspace{1cm} (10)

where $C_{L,i}$ is the interface element concentration in the liquid, corresponding to a given solid/liquid interface. When $f_{s} = 1.0$, $T_{\text{int}}$ is just the solidus temperature.

Two solid phases, the $\delta$-ferrite phase and the $\gamma$-austenite phase, occur naturally during the solidification of steel [16]. For carbon contents lower than 0.53%, when the liquid steel cools down slightly below the liquidus temperature, the solid nucleates and grows as the $\delta$ phase until solidification is complete at the low carbon content; however, for the intermediate carbon steel, once the carbon content exceeds 0.53% in the residual liquid, the solid $\gamma$ phase starts to form around the periphery of the $\delta$-phase dendrites until the completed solidification. For carbon contents over 0.53%, the solid nucleates from the melt as the $\gamma$ phase, which grows until the end of solidification. The enrichment of solute elements in the interdendritic region during solidification causes a switch from $\delta$-phase to $\gamma$-phase solidification once the carbon concentration exceeds 0.53%. This change suddenly lowers the diffusion rates, which increases micro-segregation and further lowers the solidus temperature. This behavior has an important effect on the metal properties that control micro-segregation, because of the very different solidification parameters in the two crystal phases (as shown in Table 1 [15]).

**Table 1.** Equilibrium partition coefficients, diffusion coefficients, and liquidus line slopes of the solute elements.

| Element | $k^{\gamma L}$ | $k^{\delta L}$ | $D^{\delta}$ (cm$^2$/s) | $D^{\gamma}$ (cm$^2$/s) | $m$ (C%) | $n$ (C%) |
|---------|----------------|----------------|-----------------|-----------------|---------|---------|
| C       | 0.19           | 0.34           | $0.0127 \times 10^{-52870.5/RT}$ | $0.076 \times 10^{-67419.9/RT}$ | 78.0    | 415.3   |
| Si      | 0.77           | 0.52           | $8.0 \times 10^{-16173.6/RT}$ | $0.3 \times 10^{-16360.6/RT}$ | 7.6     | 12.3    |
| Mn      | 0.76           | 0.78           | $0.76 \times 10^{-145808.5/RT}$ | $0.055 \times 10^{-16209.4/RT}$ | 4.9     | 6.8     |
| P       | 0.23           | 0.13           | $2.9 \times 10^{-149505.4/RT}$ | $0.01 \times 10^{-118788.8/RT}$ | 34.4    | 124.5   |
| S       | 0.05           | 0.035          | $4.56 \times 10^{-139447.7/RT}$ | $2.4 \times 10^{-145156.2/RT}$ | 38.0    | 183.9   |

Notes: R is the gas constant of 1.987 cal/mol K, and T is the temperature in Kelvin.

In order to incorporate the $\delta$/$\gamma$ transformation into the present model, it is assumed that only the carbon concentration controls this transition and other components do not influence the transition. The solid fraction at which the transition occurs can be calculated by using Equation (1) [17].

$$f_{S,\delta \rightarrow \gamma} = \frac{1}{2a_{i}k_{i}} \left[ 1 - \frac{0.53}{C_{0}} \left( \frac{1 - 2a_{i}k_{i}}{(1 - k_{i})} \right) \right],$$  \hspace{1cm} (11)

where the ferritic data should be employed for $a_{i}$ and $k_{i}$. For getting the segregation distribution corresponding to any solid fraction higher than $f_{S,\delta \rightarrow \gamma}$, a virtual zero concentration must be obtained. This could lead to the same concentration in residual liquid at the transition point by using Equation (1) and the solidification parameters of the austenitic phase. The corresponding calculation equation is expressed as follows:

$$C_{0,i} = C_{L,i} \left[ 1 - \frac{(1 - 2a_{i}k_{i}) f_{S,\delta \rightarrow \gamma}}{(1 - k_{i})} \right],$$  \hspace{1cm} (12)

where the austenitic data should be employed for $a_{i}$ and $k_{i}$.

In order to obtain the accurate liquidus and solidus temperatures, firstly, an estimate of the local solidification time is done by evaluating the equilibrium liquidus temperature and solidus temperature from Equations (8) and (10), respectively. From this initial guess, the value of $T_{s}$ is improved with a few iterative steps through Equations (1)–(3) and (10) with $f_{s} = 1.0$, until consistent values are obtained for $t_{f}$, $C_{L,i}$, and $T_{\text{int}} = T_{s}$.

If carbon content is less than 0.53% and $f_{s} = 1$, then $\delta$-phase properties are used in the previous procedure. If carbon content is greater than 0.53% and $f_{s} = 0$, then $\gamma$-phase prop-
erties are used throughout. If none of these conditions are met, then both phases may exist; thus, the procedure is repeated firstly for the δ phase. Once the carbon concentration of the residual liquid exceeds 0.53%, the virtual zero concentration is obtained, and the following γ-phase properties are adopted until completed solidification of the residual liquid.

After finding $T_s$, the previous procedure should be repeated to find $T_{int}$ corresponding to any other desired $f_s$ value without the need for further iteration. In practice, it is often desirable to find the relationship between the entire solid fraction and the temperature. Thus, it can be convenient to solve the above equations by incrementing the solid fraction in steps of 0.01.

3. Validation of Analytical Model

In order to assess the validity of the analytical model, under the steel compositions and cooling conditions (shown in Table 2) used in the unidirectional solidification experiment, the model predictions are compared with the experiment results suggested by Matsumiya [18].

Table 2. Steel compositions and cooling condition for the calculated model.

| C  | Si | Mn | P  | S  | $C_R$ (°C/s) |
|----|----|----|----|----|-------------|
| 0.13 | 0.35 | 1.52 | 0.016 | 0.002 | 0.045       |

As shown in Figure 1, it can be seen that the model calculations are in reasonable agreement with these measurements, which indicates that this model can accurately calculate the concentration of each element during solidification; thus, the segregation distribution of solute elements is predicted.

![Figure 1](image-url) Comparison of (a) manganese and (b) phosphorus concentration in the liquid phase calculated and experimental measurements.

4. Analysis of Solute Segregation on the Central Cracking

The etching print of the bloom transverse section with typical central cracks is shown in Figure 2a. According to the statistical data from the etched prints of massive blooms, it is found that the central cracks all appear at the inner arc side, the distance between the opening point of the crack and the inner radius of the surface mainly concentrates in the scope of 90–105 mm, and the width of the cracks mainly occurs in the range of 0.5–2 mm.
After the typical central crack samples are mechanically ground, polished, and etched using picric acid, they are visualized using a video camera to observe the relationship between the crack and the columnar or equiaxed crystal. As shown in Figure 2b,c, the columnar crystal of crack samples is developed such that the length of the primary dendrite arm spacing reaches approximately 1000 μm. In addition, the central cracks all occur in the gap between two adjacent primary dendrites at the end of the columnar crystal zone, thus exhibiting typical cracking along the gap between the columnar crystals. Chemical compositions of the crack sample (Table 3) were detected using XRF (sequential X-ray fluorescence spectrometer, ZSXPrimus IV).

Table 3. Chemical compositions of crack sample (wt. %).

| Steel Grade     | C   | Si  | Mn  | P   | S   | Cr  | Mo  | Ni  |
|-----------------|-----|-----|-----|-----|-----|-----|-----|-----|
| 49MnVS3         | 0.475 | 0.373 | 0.734 | 0.009 | 0.0483 | 0.161 | 0.007 | 0.18 |

As described in Figure 3, with the increasing solid fraction, the segregation of both carbon and manganese is quite slight, whereby they only have a little enhancement at the final stages of solidification. However, for phosphorus and sulfur, their segregation is very severe, especially at the final stages of solidification, whereby the maximum index values reach 16.6 and 33.3, respectively. The segregation distributions of elements near the crack are shown in Figure 4 using the line scanning of SEM (field-emission scanning electron microscopy, Apreo S HiVac). Here, it is clear that the fluctuation peaks of carbon, phosphorus, sulfur, and manganese all appear at the cracking gap, which means that all of these four elements have a certain amount of enrichment at the cracking gap, basically coinciding with the results calculated using the segregation model.

According to the investigation of previous researchers [19,20], central cracks generally form in the temperature range from the LIT to the ZDT under high temperature, and the solute segregation has a significant influence on this temperature range. However, the difference between the LIT and the ZDT is 69.1 °C when segregation of the elements is considered, but it is only 7.9 °C without segregation. Therefore, solute segregation facilitates an enhanced increase in cracking. From the previous line scanning analysis, it can be seen that carbon, phosphorus, sulfur, and manganese all have a certain amount of enrichment at the cracking gap, while the segregations of phosphorus and sulfur are especially serious at the final stages of solidification. Hence, the element segregation leads to an obvious increase in the temperature range of crack formation; thus, the conclusion can be obtained that the solute segregation is the internal reason for crack formation.
Figure 3. (a) Concentration and (b) segregation indices of elements C, P, and S in liquid phase at the mushy zone.

Figure 4. Line scanning of SEM in the crack band.

Subsequently, the crack samples were ground, polished, and etched using a nitric acid alcohol solution in which the alcohol occupied 4 wt.%. Then, they were placed in an optical microscope to observe the structure. It can be seen from Figure 5 that the cracking and extending directions of the crack were always along the grain boundaries, thus exhibiting a typical fracture mode along the grain boundaries.
Figure 5. Morphology of central crack at etching print ((a) typical fracture crack a; (b) typical fracture crack b).

The fracture morphologies of the cracks are of great help to find the formation temperature of cracks. As shown in Figure 6, it is clear that the fracture surfaces of the cracks are so smooth that there are no dimples to be found, which is very similar to the fracture surface which forms in the mushy zone. This indicates that the temperature range of crack formation should be at the mushy zone, denoting steel cracking with a low melting point.

Figure 6. Photograph of surface on central crack by SEM ((a) enlarged fracture surfaces of the cracks; (b) fracture surface morphology of cracks).

In order to analyze the mechanism of crack formation, the following assumption is considered: the position with the shortest distance from the inner radius is just the opening point. Depending on the statistical data from the etched prints of massive blooms, when the bloom begins to crack, the solid shell thickness is within the scope of 90–105 mm. At this time, the thickness of 95 mm is selected when the distance from the meniscus is about 13.5 m. Then, the strain of the solid shell is simulated by using finite element software (MSC Marc), where the factors included heat transfer, phase transformation, and metallurgical pressure. The thermo-elastic–plastic constitutive equation is as follows:

\[
\varepsilon_{ij} = \varepsilon_{ij}^e + \varepsilon_{ij}^i + \varepsilon_{ij}^T,
\]

(13)

where \(\varepsilon_{ij}^e\) is the elastic strain, \(\varepsilon_{ij}^i\) is the inelastic strain, and \(\varepsilon_{ij}^T\) is the thermal strain. In the analysis, the von Mises yield criterion is used to describe the thermo-elastic–plastic property. As can be seen in Figure 7, a positive value denotes tensile strain, and a negative value indicates compressive strain. Clearly, tensile strain near the solid front reaches the maximum value of 2.77–3.22%. Otherwise, Hiebler [21] gave the relationship between the critical strain and the carbon equivalent by summing up the results of previous studies, and the experimental data used in Figure 8 were obtained under the typical strain rate of a continuous casting process. According to Equation (14) and the chemical composition of steel 49MnVS3, the carbon equivalent content of 0.452% and the Mn/S index of 15.2 are obtained; therefore, the critical strain of this steel grade is approximately 0.50%, which is
much smaller than the tensile strain of the solid front. Therefore, it is very easy for tearing to occur, thus forming a central crack. The carbon equivalent can be calculated as follows:

\[ C_p = C + 0.02Mn + 0.04Ni - 0.1Si - 0.04Cr - 0.1Mo. \]  

(14)

Figure 7. Simulation analysis of the strain of the solid shell for the opening point of the crack.

During the solidification process of bloom, columnar crystals are developed such that the interdendritic segregation is quite severe. On the one hand, enriched solute elements exist in the grain boundaries and form inclusions and others, thereby greatly reducing the high-temperature strength and the high-temperature plasticity of the dendrite grain boundaries [22]. Correspondingly, the critical strain value of steel significantly decreases, thus causing increased brittleness of grain boundaries and extending the path of the crack. On the other hand, interdendritic segregation reduces the melting point of the dendritic liquid film, which leads to the freezing temperature of steel significantly decreasing and
the difference between the LIT and the ZDT obviously increasing; thus, the possibility of racking is increased. During the solidification process of bloom, when the solid shell thickness grows to approximately 95 mm, the tensile strain of the solid front becomes much larger than the critical strain of blooms. This easily results in cracking of the solid front, which extends along the grain boundaries. In addition, once cracking of the solid front happens, the solute-rich liquid ahead of the solidification interface is immediately sucked into the cracking gap, accompanied by the solidification of liquid steel. Moreover, due to the effect of negative pressure, the liquid steel containing the enriched impurities is continuously sucked into the cracks, which leads to the cracking gap extending to the center of the bloom, thus forming a severe central crack.

5. Conclusions

(1) Taking into account the Brody–Flemings model and the modified Voller–Beckermann model, an analytical model of micro-segregation was established by considering the actual solidification cooling conditions of bloom, and the accuracy of the model was verified using the results of directional solidification experiments.

(2) Through the developed analytical model, the solute segregation at the opening point of the crack was analyzed. The segregation of both phosphorus and sulfur was quite severe, while that of carbon and manganese was relatively light, which agrees well with the semi-quantitative analysis results from the SEM. Moreover, interdendritic segregation led to an increase in the temperature range of crack formation; therefore, it was concluded that element segregation is the internal cause of crack formation.

(3) Taking into consideration heat transfer, phase transformation, and metallurgical pressure, the strain of the solid shell was obtained through finite element software (MSC Marc). It was found that, when the solid shell thickness grows to approximately 95 mm, the tensile strain of the solid shell is much bigger than the critical strain, representing the external cause of crack formation.

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