Effect of Operating Conditions on Inclusion of Die Steel during Electroslag Remelting

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The current paper focuses on the effect of different operating conditions on the content of inclusions and cleanliness of remelting ingots. For these investigations, eight ingots were remelted with two slag amount and with two current intensity under otherwise comparable remelting conditions. A two-dimensional (2D) coupled mathematical model was employed to simulate the velocity field, solidification and inclusion motion for a system of electrode, slag and ingot in electroslag remelting (ESR) processes, to reveal the inclusion removal mechanism. The results showed that the content of large-sized inclusions in ESR ingot was decreased by approximately 66.18% when the slag amount was increased from 17.85 kg to 20.50 kg. Because of the increase of slag amount, the metal and slag flow faster and the maximal velocity increases by 10.3%, thus there is an increasing trend in trajectories of inclusions (i.e., inclusion motion) in slag pool resulted from the stronger natural convective flow, which is beneficial for the inclusion removal. When the average current was increased from 4 kA to 5 kA, the content of large-sized inclusions in ESR ingot was decreased by approximately 51.38%. Because of the increasing of current, the flow in the middle of the slag pool becomes stronger and the maximal downward velocity increases by 2.7%, thus there is an increasing trend in the renewal rate of the metal film surface due to the stronger washing by slag flow, which can promote the inclusion removal.

KEY WORDS: electroslag remelting; inclusion content; cleanliness; slag amount; current; numerical simulation; H13 die steel.

1. Introduction

The increasing demand for producing high-quality steel has led to the stringent control of steel cleanliness, including oxygen content and inclusions, which has been attracting great attention in recent years.1–11) Non-metallic inclusions in steel are often the initiation source of fatigue crack which seriously affect the service life of steel, particularly the existence of large-sized inclusions are extremely detrimental to fatigue life and service performance despite their limited numbers.4–6) Electroslag remelting (ESR) is one of the most important and effective ways to control the cleanliness and solidification structure of steel.7,8)

ESR process is caused by means of an alternating current (AC) or a direct current (DC) which flows through the high-resistivity slag pool from the electrode to the ingot, thus generating Joule heating in the electrode and slag. The Joule heating at the electrode periphery is significantly greater than that in the center, owing to the higher current density concentrated at the electrode periphery induced by the skin effect;9,10) in addition, the maximum of Joule heating is at the electrode/slag interface region around the electrode tip, which provides the necessary energy to melt consumable electrode. During the ESR process, the consumable electrode tip immersed in slag with a high temperature will melt, and then thin liquid metal films come into being when the thermal energy supplied by Joule heating accumulated to a certain degree. The newly formed thin liquid metal film on the surface of electrode tip is washed away driven by the Lorentz force and the gravity force and gradually accumulates on the nose of the electrode tip. When the drop on the nose of the electrode tip reaches the critical size (normally 1 to 10 mm in diameter11), necking effect will happen leading to the formation of small droplets. At last, the droplet will detach from the electrode tip when it reach a certain mass, and thereafter travel through the slag pool into the metal pool in the ways of individual droplets or the continuous stream of liquid steel. Thus, the molten steel is refined by both physical and chemical ways during its interaction with the molten slag at the molten steel-slag interface.11,12)

Variety of factors impact the cleanliness level of ESR ingot including steel quality and grade of the consumable electrode, remelting atmosphere, slag amount and its composition, remelting current, furnace voltage, cooling rate, remelting rate, input power, filling ratio, magnetic field, etc., in which the operating conditions is a critical aspect to improve steel cleanliness.13) Extensive research studies

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2. Experimental

2.1. Experimental Procedure

The H13 electrode materials were produced by using an MgO crucible of a induction melting furnace. After induction melting, the molten steel was cast into ingots, which were thereafter used as consumable electrodes in ESR experiments. The chemical composition of the produced H13 consumable electrode is shown in Table 1. The produced ingots were divided into two groups (i.e., 1E and 2E) and designated as 1E-1, 1E-2, 1E-3, 1E-4, 2E-1, 2E-2, 2E-3, and 2E-4, respectively.

The schematic diagram of experimental apparatus is shown in Fig. 1. The inner diameter and outer diameter of water-cooled copper mold is 300 mm and 335 mm respectively. The length of water-cooled copper mold is 1500 mm. The remelting processes were conducted in the water-cooled copper mold, fixed on a similarly cooled copper baseplate. The water flow pressure in the mold and baseplate was about 0.6 Mpa during remelting.

The individual experimental condition for each experiment is summarized in Table 2. In order to investigate the

Table 1. Chemical composition of the H13 consumable electrode (mass%).

<table>
<thead>
<tr>
<th>C</th>
<th>Si</th>
<th>Mn</th>
<th>Cr</th>
<th>Mo</th>
<th>V</th>
<th>Ni</th>
<th>P</th>
<th>S</th>
<th>Al</th>
<th>Cu</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.42</td>
<td>1.20</td>
<td>0.32</td>
<td>4.59</td>
<td>1.20</td>
<td>0.85</td>
<td>0.16</td>
<td>0.025</td>
<td>0.008</td>
<td>0.025</td>
<td>0.11</td>
</tr>
</tbody>
</table>

Table 2. Experimental conditions in each ESR experiment.

<table>
<thead>
<tr>
<th>Group</th>
<th>Exp. No.</th>
<th>Ingot No.</th>
<th>Slag amount/ kg</th>
<th>Current/ kA</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>1T</td>
<td>1T-1</td>
<td>1ESR-1</td>
<td>17.85</td>
<td>3.5–4.5</td>
<td>To study the influence of slag amount.</td>
</tr>
<tr>
<td>1T</td>
<td>1T-2</td>
<td>1ESR-2</td>
<td>20.50</td>
<td>3.5–4.5</td>
<td></td>
</tr>
<tr>
<td>1T</td>
<td>1T-3</td>
<td>1ESR-3</td>
<td>20.50</td>
<td>3.5–4.5</td>
<td>To study the influence of current.</td>
</tr>
<tr>
<td>1T</td>
<td>1T-4</td>
<td>1ESR-4</td>
<td>20.50</td>
<td>4.5–5.5</td>
<td></td>
</tr>
<tr>
<td>2T</td>
<td>2T-1</td>
<td>2ESR-1</td>
<td>17.85</td>
<td>3.5–4.5</td>
<td>To study the influence of slag amount.</td>
</tr>
<tr>
<td>2T</td>
<td>2T-2</td>
<td>2ESR-2</td>
<td>20.50</td>
<td>3.5–4.5</td>
<td></td>
</tr>
<tr>
<td>2T</td>
<td>2T-3</td>
<td>2ESR-3</td>
<td>20.50</td>
<td>3.5–4.5</td>
<td>To study the influence of current.</td>
</tr>
<tr>
<td>2T</td>
<td>2T-4</td>
<td>2ESR-4</td>
<td>20.50</td>
<td>4.5–5.5</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 1. Schematic diagram of the electroslag remelting.
effect of operating conditions on the steel cleanliness of ESR ingot, eight experiments in two groups (i.e., 1T and 2T) were conducted with two slag amount (17.85 kg and 20.50 kg) and two current intensity (3.5–4.5 kA and 4.5–5.5 kA) under otherwise comparable remelting conditions. The electrodes in group 2T can be regarded as almost identical to group 1T. The premelted slag (65 mass pct CaF$_2$, 30 mass pct CaO, 5 mass pct Al$_2$O$_3$) was roasted at 500$^\circ$C in a dry box for at least 5 hours to remove the moisture in slag before the ESR experiment. All the electrode were remelted without a protective atmosphere.

After electroslag remelting, the produced ESR ingots in experiments 1T-1, 1T-2, 1T-3, 1T-4, 2T-1, 2T-2, 2T-3, and 2T-4 were designated as 1ESR-1, 1ESR-2, 1ESR-3, 1ESR-4, 2ESR-1, 2ESR-2, 2ESR-3, and 2ESR-4, respectively.

2.2. Inclusion Analysis

The small steel samples of 15 mm $\times$ 15 mm $\times$ 15 mm that were taken from 1ESR-1, 1ESR-2, 1ESR-3, 2ESR-1, 2ESR-2 and 2ESR-3 were prepared for inclusion analysis, respectively. Statistical analysis was conducted on the number and size distribution of microscopic inclusions through metallographic examination (i.e., optical microscopy and image analysis). Cleanliness index ($I$) was introduced to comprehensively assess the content of non-metallic inclusions in steel, which is defined as follows:

$$I = \frac{\sum_{i=1}^{k} (d_i \times n_i)}{B \cdot S \cdot N}$$

where $B$ is the equivalent diameter of inclusions; $I$ is the cleanliness index, i.e., the number of inclusions with equivalent diameter of $B$ per unit area in sample; $d_i$ is the average diameter of inclusions in different size ranges (in radius) of 0–2.5 μm, 2.5–5 μm, 5–10 μm and 10–15 μm, $i = 1, 2, 3...$; $n_i$ is the number of inclusion in each size range, $i = 1, 2, 3...$; $N$ is the number of observation field, and fifty view fields in each steel sample were randomly selected to analyze the size and number of inclusions in different size ranges by an image analyzer; $S$ is the total area of a single observation field.

In particular, to find large-sized inclusions, the big steel samples (Diameter: 50 mm, Length: 150 mm) were taken from the consumable electrode and ESR ingot for investigation. Electrolytic extraction method was performed to reveal the individual large-sized inclusions with size larger than 50 μm in steel. The big steel sample was immersed in electrolyte (i.e., FeSO$_4$ and FeCl$_2$ solution), and then electrolyzed with the applied anode current less than 7 A. Stainless steel slice was taken as cathode. The temperature of electrolyte was maintained below 35$^\circ$C during electrolysis. The description of the electrolytic extraction method was also described in detailed elsewhere.$^{4,19}$ After steel matrix dissolved in solution, the remaining slimes was collected into a suspended bag, and then washed. Then, the nonmetallic inclusions which remain in the slimes were elutriated with ethanol, and then collected and analyzed for the weight by Electronic balance.

3. Numerical Simulation

Figure 2 shows the mesh model at the initial state and the simplified calculation flowchart of the coupled Meltflow mathematical model used in the current work. The main physical properties, geometrical and operating conditions used in the simulation are shown in Table 3. The data used for the simulation are mainly the thermophysical properties of slag and metal, the geometry of the furnace (i.e., slag, ingot, electrode, and mold) and the operating conditions (i.e., the electric current profiles and power input). Table 4 shows the particle conditions used in the simulation. The particle conditions used in the simulation are mainly the the size, relative density and initial arrangement of particle conditions. All thermophysical properties were compared and performed with information reported in the literature and adjusted by comparison of the experiment.

A two-dimensional commercial software (MeltFlow) was employed to simulate the velocity fields, solidification and inclusion motion for a system of electrode, slag and ingot in
electroslag remelting processes. The theories and details for the computer simulation model have been described in the work\textsuperscript{20,21} by Kanchan Kelker who developed the software “Meltflow”. This program has been applied for the analysis and simulation of the microstructural characteristics with different alloys and process conditions.\textsuperscript{20,22}

### 4. Results and Discussions

#### 4.1. Chemical Compositions of the Inclusions

In order to determine the chemical compositions of large-sized inclusions, the collected large-sized inclusions were detected by using scanning electron microscopy and energy dispersive spectroscopy (SEM-EDS). Figure 3 shows the SEM image and EDS analysis results of the large-sized inclusions observed in the ESR ingot. Table 5 shows the chemical compositions of inclusions. Four types of inclusions were detected in the ESR ingot. The large-sized inclusions observed in ESR ingot are mainly CaO–SiO$_2$–MgO–Al$_2$O$_3$–TiO$_2$ and CaO–SiO$_2$–Al$_2$O$_3$. Meanwhile, a few TiO$_2$–MgO–Al$_2$O$_3$ and CaO–SiO$_2$–MgO–Al$_2$O$_3$ inclusions were found.

#### 4.2. Effect of Slag Amount on the Content and Removal of Inclusion

The effect of slag amount on the content of large-sized inclusions in ESR ingots and their removal ratio are shown in Fig. 4. It is clear that: (i) in the case of the remelting performed with the adding of increased amount of slag (Exp. No. 1T-2), the content of large-sized inclusion can be reduced by 75.77% from the original inclusion content, \textit{i.e.}, 2.27 mg/10 kg, in the ESR ingot 1ESR-1 (Exp. No. 1T-1) to 0.55 mg/10 kg in the ESR ingot 1ESR-2 (Exp. No. 1T-2); (ii) in the case of the remelting performed with the adding of increased amount of slag (Exp. No. 2T-2), the content of large-sized inclusion can be reduced by 56.59% from the original inclusion content, \textit{i.e.}, 2.58 mg/10 kg, in the ESR ingot 2ESR-1 (Exp. No. 2T-1) to 1.12 mg/10 kg in the ESR ingot 2ESR-2 (Exp. No. 2T-2). It was confirmed on the basis of experimental results that the large-sized inclusions content of H13 ESR ingot can be reduced by about 66.18% by increasing the slag amount from 17.85 kg to 20.50 kg.

Figure 5 shows the effect of slag amount on cleanliness index of ESR ingots. It is clear that: (i) in the case of the remelting performed with the adding of increased amount of slag (Exp. No. 1T-2), the cleanliness index of steel can be reduced by 18.70% from the original cleanliness index, \textit{i.e.}, 12.03 ind·mm$^{-2}$, in the ESR ingot 1ESR-1 (Exp. No. 1T-1) to 9.78 ind·mm$^{-2}$ in the ESR ingot 1ESR-2 (Exp. No. 1T-2); (ii) in the case of the remelting performed with the adding of increased amount of slag (Exp. No. 2T-2), the cleanliness index of steel can be reduced by 15.60% from the original cleanliness index, \textit{i.e.}, 11.35 ind·mm$^{-2}$, in the ESR ingot 2ESR-1 (Exp. No. 2T-1) to 9.58 ind·mm$^{-2}$ in the ESR ingot 2ESR-2 (Exp. No. 2T-2). It was confirmed on the basis of experimental results that the cleanliness index of H13 ESR

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**Table 3.** Physical properties, geometrical and operating conditions of the current ESR system.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrode (diameter/height), m</td>
<td>0.25/0.6</td>
</tr>
<tr>
<td>Ingot (diameter/height), m</td>
<td>0.32/1.5</td>
</tr>
<tr>
<td>Slag (diameter/height), m</td>
<td>0.32/0.080, 0.091, 0.190</td>
</tr>
<tr>
<td>Slag (weight), kg</td>
<td>17.85/20.5/40</td>
</tr>
<tr>
<td>Current amplitude, KA</td>
<td>4, 5, 10</td>
</tr>
<tr>
<td>Voltage, V</td>
<td>50</td>
</tr>
<tr>
<td>Frequency, Hz</td>
<td>50</td>
</tr>
<tr>
<td>Electric conductivity of ingot, $\sigma_{m}$, $\Omega^{-1}$ m$^{-1}$</td>
<td>9.215 × 10$^{5}$</td>
</tr>
<tr>
<td>Electric conductivity of slag, $\sigma_s$, $\Omega^{-1}$ m$^{-1}$</td>
<td>220</td>
</tr>
<tr>
<td>Density of electrode, kg/m$^3$</td>
<td>7 800</td>
</tr>
<tr>
<td>Density of slag, kg/m$^3$</td>
<td>2 790</td>
</tr>
<tr>
<td>Density of ingot, kg/m$^3$</td>
<td>7 800</td>
</tr>
</tbody>
</table>

**Table 4.** Particle conditions used in the simulation.

<table>
<thead>
<tr>
<th>Size, mm</th>
<th>Relative density, $\rho_p/\rho_m$</th>
<th>Initial arrangement of particle conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.1, 0.3, 0.5, 0.7, 5.0</td>
<td>0.5, 0.95, 1.0, 1.16, 2.0</td>
<td>Particles originating at three uniformly spaced locations in the metal pool under the electrode shadow just below slag-metal interface</td>
</tr>
</tbody>
</table>

**Table 5.** Chemical compositions of large-sized inclusions (mass%).

<table>
<thead>
<tr>
<th>No.</th>
<th>Mg</th>
<th>Al</th>
<th>Si</th>
<th>Ca</th>
<th>Ti</th>
<th>O</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2.0</td>
<td>5.1</td>
<td>27.5</td>
<td>–</td>
<td>–</td>
<td>65.5</td>
</tr>
<tr>
<td>2</td>
<td>–</td>
<td>1.7</td>
<td>45.4</td>
<td>4.0</td>
<td>19.7</td>
<td>62.4</td>
</tr>
<tr>
<td>3</td>
<td>0.8</td>
<td>2.5</td>
<td>10.7</td>
<td>–</td>
<td>–</td>
<td>54.1</td>
</tr>
<tr>
<td>4</td>
<td>–</td>
<td>11.9</td>
<td>27.8</td>
<td>6.1</td>
<td>–</td>
<td>54.1</td>
</tr>
<tr>
<td>5</td>
<td>1.0</td>
<td>3.5</td>
<td>12.3</td>
<td>3.8</td>
<td>23.8</td>
<td>55.5</td>
</tr>
<tr>
<td>6</td>
<td>–</td>
<td>12.1</td>
<td>30.8</td>
<td>10.0</td>
<td>–</td>
<td>47.1</td>
</tr>
<tr>
<td>7</td>
<td>–</td>
<td>10.0</td>
<td>24.7</td>
<td>10.9</td>
<td>–</td>
<td>54.5</td>
</tr>
<tr>
<td>8</td>
<td>1.0</td>
<td>7.0</td>
<td>33.0</td>
<td>2.5</td>
<td>–</td>
<td>56.5</td>
</tr>
</tbody>
</table>

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![Fig. 3. SEM image and EDS analysis results of large-sized inclusions observed in ESR ingot.](image-url)
ingot, it can be seen from the above experimental results shown in Figs. 4, 5 and 6 that the decrease ratio of the large-sized inclusions is relatively higher than that of the microscopic inclusions in H13 ESR ingot with the increase of slag amount during remelting. It indicates that the bigger inclusions are more easily to be removed than smaller ones. Similar to the current findings, Bouris et al.\textsuperscript{23) }reported that the larger inclusion can be more easily removed than smaller one.

Therefore, it can be concluded from the above experimental results that removal of inclusion and the cleanliness of the ESR ingot can be promoted by increasing the slag amount during ESR process.

Four reasons may account for this result. The first possible reason may be that less electrode has been oxidized by air due to less surface area surrounding the electrode exposed to area and lower surface temperature of the slag pool under the condition of large slag amount. When the amount of slag is increased, the thickness of the molten slag pool and the electrode immersion depth in slag pool are increased leading to the decrease of the electrode oxidation at high temperature. In addition, it is well known that surface temperature of electrode for large slag amount is lower than that for small slag amount due to greater electrode immersion depth and low surface temperature of slag pool, which lead to different oxidation conditions. Oxide scale caused by oxidation of electrode surface at high temperature is the main reason for the increase of oxygen and inclusion content in steel during ESR. Therefore, electrode surface oxidation is considered to play a greater role according to the above experimental results.

Secondly, the depth of the molten slag pool increases with the increase of slag amount. This indicates that the residence time for the metal droplet travelling through the molten slag pool increases with the increase of slag amount. Thus, the interaction time between metal droplets and molten slag increases with the increase of slag amount, which is conducive to inclusion removal.

The third one is that the flow of the slag pool becomes stronger for large slag amount. Varying the slag amount dramatically affects both the simulated distribution of the Joule heating and thermohydrodynamic behavior of the slag phase; it then logically influences the flow of slag. A two-dimensional model (MeltFlow) was employed to simulate the velocity fields and inclusion motion for a system of electrode, slag and ingot in electroslag remelting processes. Figure 7 provides the velocity fields and inclusion motion in liquid fraction fields of the ingot with different slag amount. Information about the behavior of inclusions and the trajectories of all inclusions at the time instant are noticed in the slag pool and metal pool. Color (red, green, blue, yellow, cyan) is used for trajectories of inclusions of density number (1, 2, 3, 4, 5) respectively. Particles originating at three uniformly spaced locations in the metal pool under the electrode shadow just below slag-metal interface. For each density, trajectories for each size are written for all originating locations. The trajectories are written for the left and the right r-x planes around the ingot centerline for easy visualization using layout files. Thus, each type of inclusion has two trajectories written for it. The two zones for each type of particle correspond to the densities, sizes,
and locations of particles. The motions of inclusions are tracked before they enter the mushy zone or completely dissolve in the molten pool. Inclusions would be expected to experience very different buoyancy and drag forces due to the difference between the slag and molten pools in density and viscosity when they travel through the slag-metal interface. The time required for inclusions to attain their terminal velocities depends on these external forces and their inertia forces. Information about the behavior and trajectories of all inclusions at the time instant are observed in the slag pool and metal pool, as shown in Fig. 7. It can be seen from Fig. 7 that the large light inclusions are expected to vibrate and drift across the slag-metal interface between the slag and the metal pool, while light small inclusions are expected to float into the slag pool and swirl around in it with the flow of slag depending upon their relative particle densities. The relative density of main inclusions calculated from chemical composition in Table 5 is approximately 0.5. By comparing the density calculated from the chemical composition and the simulation results in Fig. 7, it also can be seen that there is an increasing trend in trajectories of inclusions (i.e., inclusion motion) in slag pool with the increase of slag amount. This is especially the case for inclusions with relative density of 0.5 (i.e., the relative density of the main large-sized inclusions in the ESR ingot).

Figure 8 shows the distribution of the two phases, velocity field and metal pool profile of the ingot with three added slag amount at 5 400 s. A liquid sump is expected to be formed with the freezing of metal driven by the heat extracted by the cooling water. The V-shaped metal pool is formed due to the weaker cooling intensity at the bottom of the ingot comparing to that at the edge of the ingot, as shown in Fig. 8. Comparing to the metal pool, the slag pool is more thermodynamically unstable and has stronger velocity field with magnitude larger than that in the molten pool. The predicted velocity vector in the metal pool and slag pool represents the magnitude and direction of the local velocity. It should be mentioned here that no-slip boundary condition on all solid boundaries is used for an implicit treatment of the slag-metal interface which is assumed to be flat. A pair of vortex is observed in each side of the slag pool. One is at the center of the slag pool, while the other is at the vicinity of the slag pool/mold interface. The former is considered to be resulted from the cooling by contact with the electrode tip and the divergence of the electric current lines caused by the local predominance of electromagnetic forces (EMF) in the center of the slag. While the latter is created by the natural convective flow caused by a local principal large radial temperature gradient due to the loss of heat extracted by the cooling water at the lateral wall. It can be seen from Fig. 8 that with more slag is added, the downward flows near the mold surface becomes stronger. This is mainly attributed to the loss of more heat extracted by the cooling water at the lateral wall due to the increase of slag pool/mold interface area caused by the increase of slag amount. Because of the increase of slag amount, the metal and slag flow faster, resulting in a more vigorous circulation cell, and the maximal velocity increases from 0.0482 m s$^{-1}$ to 0.0532 m s$^{-1}$. The momentum and heat transfer are reinforced by the increase of slag amount. Thus the thermohydrodynamic conditions of the slag pool is expected to be improved by the stronger natural convective flow, which is beneficial for inclusion removal. Therefore, it can be inferred from Figs. 7 and 8 that the mass transfer, reaction kinetics and chemical interactions between inclusions and slag is promoted by the stronger natural convective flow with the increase of slag amount, which is conducive to inclusion removal.

Moreover, it can be inferred from Fig. 8 that the liquid metal pool is shallowed for large slag amount. Varying the slag amount dramatically affects both the effect of the Joule heating and heat extracted by the cooling water; it then logically influences the depth of liquid metal pool. Concerning the metal pool profile, metal pool depth results from the combined action of Joule heating and cooling by contacting with the water-cooled mold.

The metal pool depth varies during remelting with changes in the operating conditions, while it is always determined by the combination and competition between these two actions, as indicated in Fig. 8, in which the distribution
of the two phases, velocity field and metal pool profile of the ingot with three added slag amount, at 5 400 s of remelting, are illustrated. In the close vicinity of the slag/pool interface, the metal flows along the radial direction toward lateral wall and, thereafter, from there downwards driven by the combined action of Lorentz force, molten slag pressure and temperature gradient. Most of the heat is thus taken away from the metal flow by the cooling water resulted from its interaction with the water-cooled mold. As a result, the metal gets “colder” and denser. Afterwards, the denser metal flows downwards the bottom along the slanting solidification front. Finally, a metal pool is expected to be formed as the melting proceeds due to the stronger cooling action of the lateral wall than that of the bottom. In the liquid pool, the fluid motion is low which is mainly driven by buoyancy, due to the weaker cooling along the slanting solidification front than that along the lateral wall.

Clearly, the larger the slag amount, the stronger the Joule heating effect. This is mainly linked to the fact that, for a constant electrical power consumption, the power applied in the slag pool increases with an increase in the slag amount. Figure 9 shows the circuit diagram of ESR process. Wherein, \( R_1 \) is constant; \( R_2 \) is determined by the attributes of electrode; \( R_3 \) is determined by slag composition and slag amount (i.e., the depth of liquid slag pool); \( R_4 \) is determined by the characteristics of the ESR ingot. As it can be seen from Fig. 9, the value of the slag resistance is determined by the slag amount. Actually, the slag resistance is a direct consequence of electrical current flows the slag pool, which can be roughly calculated by the following classical equation,

\[
R_{\text{slag}} = \frac{h_{\text{slag}}}{\sigma_{\text{slag}} S} \tag{2}
\]

where \( h_{\text{slag}} \) is the height of the slag layer, \( \sigma_{\text{slag}} \) is electrical conductivity of the slag, \( S \) is the cross-sectional area of slag layer between the electrode and the ingot. Because the surface through which the electrical current flows is the same in all cases, the only parameter modified by a variation in the slag amount is the slag height. Therefore, the slag resistance increases hyperbolically when the slag amount increases in ESR process. It also can be seen from Fig. 9, the value of the power applied in the slag pool is determined by the slag resistance under the condition of the same electrical power consumption in the circuit diagram. Therefore, it can be inferred that the Joule heating of slag pool profoundly increases with the increase of slag amount when the power system is maintained constant during ESR process. However, the interaction between the slag pool and water-cooled mold also becomes stronger due to the increase of the slag/mold interface area, and the amount of heat taken away by the cooling water outweighs that created by the newly generated Joule heating. Consequently, the liquid metal pool becomes “colder” with the increase of slag amount leading to the decrease of the metal pool depth. Thus the molten pool is expected to become shallower with the increase of slag amount, which is conducive to inclusion removal.

However, the amount of slag is not the bigger the better. The selection of slag amount is also related to the given material and ingot size. It has been widely accepted that the selection of slag amount used for the ESR process is mainly based on the following empirical formula,\(^{24}\)

\[
G_{\text{slag}} = (4\% - 5\%)G_{\text{ingot}} \tag{3}
\]

\[
G_{\text{slag}} = S_{\text{mould}} H_{\text{slag}} \rho_{\text{slag}} \tag{4}
\]

\[
H_{\text{slag}} = f_{\text{slag}} D_{\text{mould}} \tag{5}
\]

where \( G_{\text{slag}} \) is the amount of slag, \( G_{\text{ingot}} \) is the weight of ingot, \( S_{\text{mould}} \) is the cross-sectional area of slag layer, \( \rho_{\text{slag}} \) is the density of slag, \( H_{\text{slag}} \) is the height of slag layer, \( f_{\text{slag}} \) is the coefficient of the slag height, \( D_{\text{mould}} \) is the diameter of mould. The calculation value of the slag amount derived from the empirical formula can be used as a reference value, whereas the optimum slag amount must be determined according to the used equipment and actual conditions of the local electric power system.

From the above analysis, it can be found that in the conditions of a certain electric power system and process permitting under the current ESR process, the appropriate increase of slag amount from 17.85 kg to 20.5 kg considering the relationship between slag amount and mould size can reduce the electrode oxidation at high temperature and promote inclusion removal in ESR process, and thus help to improve the ESR ingot cleanliness.

4.3. Effect of Remelting Current on the Content and Removal of Inclusion

Figure 10 shows the effect of current on the large-sized inclusion content in ESR ingots. It is clear that: (i) in the case of the remelting performed with the increased current of 4.5–5.5 kA (Exp. No. 1T-4), the content of large-sized inclusion can be reduced by 54.44% from the original inclusion amount, i.e., 2.59 mg/10 kg, in the ESR ingot 1ESR-3 (Exp. No. 1T-3) to 1.18 mg/10 kg in the ESR ingot 1ESR-4 (Exp. No. 1T-4); (ii) in the case of the remelting performed with the current of 4.5–5.5 kA (Exp. No. 2T-4), the content of large-sized inclusion can be reduced by 48.32% from the original inclusion amount, i.e., 2.98 mg/10 kg, in the ESR ingot 2ESR-3 (Exp. No. 2T-3) to 1.54 mg/10 kg in the ESR ingot 2ESR-4 (Exp. No. 2T-4). It was confirmed on the basis of experimental results that the content of large-

![Fig. 9.](image-url)
sized inclusions of H13 ESR ingot can be reduced by about 51.38% through increasing the current from 3.5–4.5 kA to 4.5–5.5 kA. Therefore, it can be concluded that the inclusion removal was promoted with the increase of remelting current during ESR process.

The removal of non-metallic inclusion occurred mainly during liquid metal film formation, its gradual coalescence into droplet at the consumable electrode tip and subsequent falling through the slag, especially the earlier stage of droplets formation where the removal efficiency of non-metallic inclusion is more pronounced due to the following reasons: First, as current flows from the electrode, through the slag to the ingot, the electrode tip is heated to a high temperature above the melting point of the electrode, even up to 2 000°C. Thus original solid inclusions in consumable electrode can be generally turned into liquid, and thereafter absorbed by the slag at slag–metal pool interface; Second, an extreme thin layer of liquid metal film (normally 50–200 μm) is expected to be formed at the consumable electrode tip resulted in the relatively high ratio of surface/interface area to volume between the slag and liquid metal film and, hence, the relatively larger contact area between metal and slag, thus the inclusion particles stand greater chance of interacting with molten slag in liquid metal film; Third, the formation of liquid metal film and subsequently coalescence of droplets took much longer time than molten drops falling through the slag layer. Therefore, it can be concluded that high chemical reaction temperature, large contact area between metal and slag, long chemical reaction time in this stage is favorable for the chemical reaction dynamics of inclusion removal.

Based on the above considerations, three reasons may account for this result. The first possible reason may be that the interaction between molten steel and slag is more sufficient for large remelting current. Wang et al. built a transparent experimental model to visualize the ESR process, and investigated the effects of different strength of the remelting current on the droplet evolution. Their experimental results show that the tips become sharper and sharper with the increase of remelting current as shown in their experimental results. Predictably, the average diameter of droplets decreases with the increase of remelting current during ESR process according to their experimental results. This is mainly due to the fact that the interfacial tension between slag and steel decreases with the increase of slag temperature caused by the increase of remelting current. In addition, the droplet radius decreases with the decrease of the interfacial tension between slag and steel according to the equation given by Campbell. Consequently, this decrease in diameter will result in the increase of slag/droplet interface area. Moreover, it can be inferred that less droplets drip simultaneously due to the more pointed electrode tip under large remelting current condition. Thus the number of droplets for large remelting current is less than that for small remelting current condition, the size of droplets is smaller than that for small remelting current; then interaction between slag and droplets is more sufficient for the large remelting current condition. Furthermore, according to the classical work of Wang et al that the surface area of electrode tip immersed in slag pool increases with the increase of current in ESR process. This is due to the fact that the area of the electrode in contact with the slag increases with the increase of current. Consequently, this increase in surface area of electrode tip will result in the increase of liquid metal film-slag interface. Based on the above discussions, it can be concluded that the average radius of droplets decreases and the area of liquid metal film-slag interface increases with the increase of remelting current. This fact indicates that the contact area between molten steel and slag increases with the increase of remelting current during ESR process, which is beneficial to the chemical reaction dynamics of inclusion removal.

The second one is that the flow in the middle of the slag pool becomes stronger for large remelting current. Varying the remelting current dramatically affects both the simulated distribution of the velocity field and the electromagnetic force (EMF) and thermohydrodynamic behavior of the slag phase; it then logically influences the flow of slag pool. A two-dimensional model (MeltFlow) was employed to simulate the velocity field and solidification for a system of electrode, slag and ingot in electroslag remelting processes. Figure 11 shows the distribution of the two phases, velocity field and solidification of the ingot with three applied remelting current at 5 400 s. A pair of vortex is observed in each side of the slag pool. One is at the center of the slag pool, while the other is at the vicinity of the slag pool/mold interface. The former is considered to be resulted from the cooling by contact with the electrode tip and the divergence of the electric current lines induced by the local predominance of electromagnetic forces (EMF) in the center of the slag. While the latter is created by the natural convective flow caused by a local principal large radial temperature gradient due to the loss of heat extracted by the cooling water at the lateral wall. It can be seen from Fig. 11 that with more current is applied, the flow in the middle of the slag pool becomes stronger, and the maximal downward velocity increases from 0.0165 m s⁻¹ to 0.0169 m s⁻¹. This fact indicates that the renewal rate of the metal film surface due to the washing by slag flow would become stronger with the increase of remelting current. In addition, the migration speed of inclusions from the inside of liquid metal to the steel-slag interface is determined by the renewal rate of the
metal film surface during the droplet formation period at the electrode tip.32) Thus, it is expected that the migration speed of inclusions from liquid metal internal to the steel-slag interface would be quickened with more current is applied. Consequently, the interaction frequency between slag and inclusion in liquid metal film (i.e., the opportunity for inclusions to directly contact with molten slag) increases with the increase of current, which is beneficial to inclusion removal.

The third one is that the interfacial tension is lower for large remelting current. Obviously, the temperature of slag pool increases with the increase of remelting current. Thus, both the interfacial tension between slag and steel and that between slag and inclusion decrease with the increase of remelting current as discussed above, which is conducive to inclusion removal at the steel-slag interface.

However, the remelting current is not the bigger the better. With further increase of remelting current, it is expected that the dynamic conditions for inclusion flotation during ESR process would become poor if the the remelting current is too large. The two-dimensional model (MeltFlow) was employed to simulate the velocity fields and inclusion motion for a system of electrode, slag and ingot in electroslag remelting processes. Figure 12 provides the velocity fields and inclusion motion in liquid fraction fields with different applied remelting current. Information about the behavior of inclusions and the trajectories of all inclusions at the time instant are noticed in the slag pool and metal pool. Color (red, green, blue, yellow, cyan) is used for trajectories of inclusions from liquid metal internal to the steel-slag interface as shown in the case of 10 000 A. It can be inferred that those inclusions would be left in ESR ingot due to the rapid advance of the solidifying front during solidification.

This might be attributed to the fact that, with an excessive amount of remelting current application, the molten pool is expected to be changed into deep and wide concave shape. Wang et al. investigated the effect of the current on the steel cleanliness, content and removal ratio during ESR process.

5. Conclusions

The effect of operating conditions (i.e., slag amount and current) on the steel cleanliness, content and removal ratio of inclusions, especially those large-sized inclusions during electroslag remelting of H13 die steel was investigated. The mechanisms of inclusion removal during ESR process were each type of inclusion has two trajectories written for it. The two zones for each type of particle correspond to the densities, sizes, and locations of particles. The motions of inclusions are tracked before they enter the mushy zone or completely dissolve in the molten slag. Information about the behavior and trajectories of all inclusions at the time instant are observed in the slag pool and metal pool, as shown in Fig. 12. By comparing the density calculated from the chemical composition and the simulation results in Fig. 12, it can be seen that a large proportion of the inclusion with relative density of 0.5 (i.e., the relative density of the main large-sized inclusions in the ESR ingot) would sink into the metal pool and cannot float to the steel/slag interface as shown in the case of 10 000 A. It can be inferred that those inclusions would be left in ESR ingot due to the rapid advance of the solidifying front during solidification.
also discussed, based on experimental results along with thermohydrodynamic analysis. To sum, important conclusions can be drawn as follows:

(1) When the slag amount was increased from 17.85 kg to 20.50 kg, the content of large-sized inclusions in ESR ingot was decreased by approximately 66.18%, the number of inclusions decreases and the cleanliness index of H13 ESR ingot was decreased by about 17.15%. The influence of the increasing of slag amount on the decrease ratio of bigger inclusions was more pronounced than it was for the small inclusions (radius of 0–15 μm). Under the current ESR process, it is suggested that the appropriate increase of the remelting current from 4 kA to 5 kA can promote the inclusion removal, thus improve the cleanliness of H13 ESR ingots.

(2) Because of the increase of slag amount, the metal and slag flow faster and the maximal velocity increases from 0.0482 m s\(^{-1}\) to 0.0532 m s\(^{-1}\), thus the momentum and heat transfer are reinforced by the increase of slag amount. There is an increasing trend in trajectories of inclusions (i.e., inclusion motion) in slag pool resulted from the stronger natural convective flow with the increase of slag amount, which is beneficial for the inclusion removal.

(3) When the average current was raised from 4 kA to 5 kA, the content of large-sized inclusions in H13 ESR ingot was decreased by about 51.38%. Because of the increasing of current, the flow in the middle of the slag pool becomes stronger and the maximal downward velocity increases from 0.0165 m s\(^{-1}\) to 0.0169 m s\(^{-1}\), thus the washing of metal film surface by slag flow is reinforced by the increase of current. There is an increasing trend in the renewal rate of the metal film surface due to the stronger washing by slag flow with the increase of remelting current, which can promote the inclusion removal.

(4) A large proportion of the inclusion with relative density of 0.5 (i.e., the relative density of the main large-sized inclusions in the ESR ingot) would sink into the metal pool and can not float to the steel/slag interface in the case of 10 kA. Under the current ESR process, it is suggested that the appropriate increase of the remelting current from 4 kA to 5 kA can promote the inclusion removal, thus improve the cleanliness of H13 ESR ingots.

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