D Dephosphorization Kinetics and Reaction Region in Hot Metal during Lime Injection with Oxygen*

By Hitoshi ONO,** Tamenori MASUI*** and Hisashi MORI****

I. Introduction

Lately, the dephosphorization process of hot metal by injection of mixed flux of lime and iron oxide has been developed and adapted in commercial operation, and lime injection with oxygen is also tested at some plants.

The slagging rate of CaO has a great effect on the slag-metal reaction in the steel making process. For this reason, many fundamental and practical studies have so far been made on the mechanism of dephosphorization by CaO. Fe1O in {CaO-Fe(Mn)O}l and phosphorus, and the latter is fixed as (CaO-SiO2-Mn1O). Then, the dephosphorization reaction initiates between {CaO-Fe(Mn)O}l and phosphorus, and the latter is fixed as (CaO-SiO2-Mn1O). In hot metal, CaO starts to dissolve due to penetration of Fe1O and Mn1O. Then, the dephosphorization reaction initiates between (CaO-Fe(Mn)O)l and phosphorus, and the latter is fixed as (CaO-SiO2-P2O5)l in the reaction layers. Dephosphorization takes place mainly in the vicinity of the oxygen blowing nozzle.

The slagging rate of solid CaO decreases with increasing distance between the nozzle and the suspending position of CaO since carbon in hot metal consumes oxygen. The dephosphorization rate also decreases with increasing distance since Fe1O in (CaO-Fe(Mn)O)l is reduced by carbon.

It is concluded that the increases both in the slagging rate of solid lime and oxygen potential in the bath is necessary to enhance the dephosphorization rate and in this sense, the injection of iron oxide with oxygen is effective.

II. Experimental Method

I. Lime Powder Injection Test

1. The Methods Used for the Collection of Reaction Products by means of Al2O3 Tubes and Preparation of Samples

Slag sampling method is shown in Fig. 1. A MgO crucible was set in a high frequency induction furnace and pig iron having the composition shown in Table 1 weighing 10 kilograms was melted. The experiment was done at 1 350 °C. Lime powders were injected at the rate of 7.7 g/min for 13 min with a mixed gas (O2 4 l/min+Ar 3 l/min) through a copper nozzle provided in the bottom of the crucible. During injection, the floating reaction product (slag) adhered to Al2O3 tubes which were immersed in the bath at the positions 45 mm and 90 mm above the blowing nozzle.

A slag of 30 mm maximum diameter and about 50 mm long adhered around Al2O3 tubes as a result of bottom injection of lime. Lumps of slag sample were embedded in polyester resin which was solidified later. After that, doughnut-shaped specimens of 20 mm diameter and 10 mm thick were cut from the position near the end of slag adhered. After grinding with diamond paste, gold vapour deposition was done.

In view of the above-mentioned statement, an alumina tube and a sampler (see Fig. 2) were firstly immersed into hot metal, through which lime powder was injected with oxygen, and the floating reaction product was allowed to adhere to the tube or enter the sampler in order to examine the reaction process of CaO. Then, a single-crystal lime lump was immersed into hot metal melted in a crucible and oxygen was injected to examine the slagging process of CaO and the mechanism of dephosphorization. The reason why a lime crystal is used is that a single crystal is suitable to observe the reaction layer without the effect of voids and grainboundaries. Further, alumina tubes, steel samplers and single-crystal lime lumps were immersed into hot metal at various positions in order to know the region where the injected lime reacts.

The dephosphorization mechanism of hot metal and the region where dephosphorization takes place during lime injection with oxygen were discussed on the basis of the results of these experiments.

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Synopsis

To improve dephosphorization of hot metal by the lime injection process, it is necessary to clarify the dissolution mechanism of blown-in CaO in hot metal, the mechanism of dephosphorization reaction and the region where the reaction takes place. For this purpose, a single-crystal lime lump immersion test and a lime powder injection test were made. Lime lumps after immersion tests, reaction products adhered to Al2O3 tubes immersed into hot metal and collected slag samples by steel samples were analyzed by EPMA.

In hot metal, CaO starts to dissolve due to penetration of Fe1O and Mn1O. Then, the dephosphorization reaction initiates between (CaO-Fe(Mn)O)l and phosphorus, and the latter is fixed as (CaO-SiO2-P2O5)l in the reaction layers. Dephosphorization takes place mainly in the vicinity of the oxygen blowing nozzle.

The slagging rate of solid CaO decreases with increasing distance between the nozzle and the suspending position of CaO since carbon in hot metal consumes oxygen. The dephosphorization rate also decreases with increasing distance since Fe1O in (CaO-Fe(Mn)O)l is reduced by carbon.

It is concluded that the increases both in the slagging rate of solid lime and oxygen potential in the bath is necessary to enhance the dephosphorization rate and in this sense, the injection of iron oxide with oxygen is effective.
and electron scanning images by EPMA were obtained.

Slags adhered were subjected to X-ray fluorescence analysis to determine the contents of total Fe, CaO, SiO₂, MnO, P₂O₅, S, MgO and Al₂O₃, and the contents of metallic Fe, FeO and Fe₂O₃ were determined by ordinary chemical analysis.

2. Collecting Method of Reaction Products by means of Steel Samplers and Preparation of Samples

A MgO crucible was set in a high frequency induction furnace and low silicon pig iron (C: 4.33%, Si: 0.02%, Mn: 0.07%, P: 0.108%, S: 0.043%) weighing 50 kg was melted. The experiment was done at 1350 °C. A triple layered lance (stainless steel pipe of 4 mm diameter in the inner part, magnesia pipe in the middle part and graphite pipe in the outer part) was used for injection. Lime powder weighing 200 g was injected for 11 min with a mixed gas (O₂ 8 l/min + Ar 4 l/min). During injection, the floating reaction product (slag) was collected together with hot metal by the steel sampler (see Fig. 2) immersed in the bath. Two kinds of sampler (a) with and (b) without foot were used in order to collect the reaction product from two different positions. The sampler was quickly immersed into the bath up to the position where the bottom of sampler just reached the bottom of the crucible. Then, it was pulled up just after confirming that metal entered the sampler. Samples were subjected to the microscopic observation and the quantitative analysis by EPMA.

2. Single-crystal Lime Lump Immersion Test

A MgO crucible (inside diameter: 120 mm, depth: 250 mm) was set in a high frequency induction furnace and low-silicon pig iron (C: 3.14~3.79%, Si: 0.01~0.03%, Mn: 0.05~0.13%, P: 0.075~0.200%, S: 0.012~0.013%) weighing 10 kg was melted. The experiment was done at 1350 °C. A single-crystal lime lump of 5 mm × 5 mm × 30 mm was set at the position 20 mm above the tip of the oxygen top-blowing nozzle or three lumps were set as shown in Fig. 3 and those were kept in the immersed state for 60 s. Oxygen was top-injected at a rate of 37.5 l/min with a graphite lance 30 mm outer diameter, 8 mm

Table 1. Changes in composition of hot metal (%).

<table>
<thead>
<tr>
<th></th>
<th>C</th>
<th>Si</th>
<th>Mn</th>
<th>P</th>
<th>S</th>
</tr>
</thead>
<tbody>
<tr>
<td>Before flux injection</td>
<td>4.46</td>
<td>0.01</td>
<td>0.30</td>
<td>0.100</td>
<td>0.062</td>
</tr>
<tr>
<td>After flux injection</td>
<td>3.34</td>
<td>0.01</td>
<td>0.21</td>
<td>0.039</td>
<td>0.037</td>
</tr>
</tbody>
</table>
inner diameter and 500 mm long and bottom-blown at a rate of 8 l/min with copper tube 2 mm inner diameter. Further, iron oxide which is equivalent to 3.5 l/min of oxygen was added.

A single-crystal lime-lump after immersion was embedded in acrylic resin and cut off at the position 5 mm from its end, and the cut surface was ground with diamond paste. The ground surface was observed under an optical microscope to check the reaction layer. Gold vapour deposition was then done and the samples were subjected to line scanning by EPMA and the electron scanning images were obtained. Standard samples used for the quantitative analysis by EPMA are Fe, Mn, Al, Si, Mg, CaF₂, InP and FeS₂. Only absorption correction was made, because the greatest error is induced by absorption.

III. Experimental Results

1. Lime Powder Injection Test

1. Investigation of Slag Adhered to Al₂O₃ Tube

Table 1 shows the composition of hot metal before and after lime injection. Table 2 gives the average composition of slag adhered. It can be seen from Tables 1 and 2 that the percentage of dephosphorization is approximately 60 % and that the P₂O₅ content of slag adhered is as high as 6.8 %. Thus, it is evident that dephosphorization proceeds to a great extent by lime injection. The slag adhered to Al₂O₃ tube is characterized by the fact that it contains a large amount of metallic iron and small amounts of FeO and Fe₃O₄.

Photograph 1 shows the optical micrographs of the outer part (hot metal side) of slag adhered, where (a) and (b) are the photographs of the samples taken at the positions 90 mm and 45 mm above the oxygen blowing nozzle, respectively. In Photo. 1 (a) four phases were observed, namely metallic iron, CaS, Ca(Mn)O solid solution, and CaO–SiO₂–P₂O₅ solid solution as identified by EPMA.

On the contrary, CaO–SiO₂–P₂O₅ solid solution and CaO–Fe(Mn)O solid solution were observed in Photo. 1(b) but metallic iron was not observed in it.

Since the slag sampled at the position 90 mm above the oxygen blowing nozzle contains metallic iron particles but the slag sampled at the position 45 mm above the nozzle does not contain metallic iron, an investigation on the mechanism of the formation of iron particles was expected to provide a clue to the clarification of dephosphorization reaction. Therefore, a study was made on the distribution and composition of iron particles.

As shown in Photo. 1(a), it is the first feature that iron particles are very fine and are distributed only in the Ca(Mn)O solid solution. As given in Table 3, it is the second feature that the composition of iron particles (C=1.0~1.5 %, Mn=0.6~0.8 %, P=0.67 % [large particles]~0.01 % [small particles]) is greatly different from that of hot metal (C=4.46 %, Mn=0.30 %, P=0.10 %). Accordingly, it might be thought that {CaO–Fe(Mn)O}₃ is reduced by carbon in hot metal to iron particles containing C, Mn, P,

![Fig. 3. Experimental apparatus for single-crystal lime lumps immersion test.](image)

Table 2. Average composition of slag adhered. (%)  

<table>
<thead>
<tr>
<th></th>
<th>T₂O</th>
<th>MFe</th>
<th>FeO</th>
<th>Fe₃O₅</th>
<th>CaO</th>
<th>SiO₂</th>
<th>MnO</th>
<th>P₂O₅</th>
<th>S</th>
<th>MgO</th>
<th>Al₂O₃</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>21.2</td>
<td>18.8</td>
<td>0.82</td>
<td>2.51</td>
<td>45.3</td>
<td>1.31</td>
<td>5.27</td>
<td>6.84</td>
<td>0.91</td>
<td>5.82</td>
<td>2.96</td>
</tr>
</tbody>
</table>

Photo. 1.  
Micrograph of adhered slag.
etc. The reason why the phosphorus concentration of large iron particles is high is likely to be that these iron particles are formed by the reduction of high-phosphorus containing slag at the position near the oxygen blowing nozzle where both of $\text{CaO-Fe(Mn)}_O$ and $\text{CaO-SiO}_2-P_2O_5$ are the liquid phase and that they grow thereafter. Precipitation of the solid $\text{CaO-SiO}_2-P_2O_5$ phase containing phosphorus occurs when the FeO concentration decreases due to the reduction of molten slag, and it seems that small iron particles precipitate through the reduction by the remaining $\text{CaO-Fe(Mn)}_O$ melt of low phosphorus concentration. Therefore, these iron particles contain Mn and their phosphorus concentration is very low.

Thus, if many fine iron particles observed in slag are formed by the reduction of slag from the $\text{Ca(Mn)}_O$ phase, $\text{CaO-Fe(Mn)}_O$ molten slag of high FeO concentration is formed near the bottom blowing nozzle before reduction and the dephosphorization reaction is considered to proceed owing to the presence of this slag. On the other hand, it can be explained that at a position apart from the bottom blowing nozzle, the formation of CaO–Fe(Mn)$_O$O slows down due to the decrease in available oxygen and that at the same time, the Fe(Mn)$_O$O concentration decreases because of the reduction of formed CaO–Fe(Mn)$_O$O so that the dephosphorization reaction becomes difficult to occur.

2. Investigation of Slag Collected by Steel Sampler

Table 4 gives the analytical results of slag sampled from the top of the bath after the termination of injection. This slag is characterized by the existence of a large amount of metallic iron and by high concentrations of Al$_2$O$_3$ and MgO. High concentration of Al$_2$O$_3$ is originated from the use of alumina protection tube for thermocouple, and that of MgO is derived from the use of magnesia crucible. Then, the concentration of Al$_2$O$_3$ is important as a tracer to identify the sampling position.

Table 5 gives the results of a quantitative analysis of these slags by EPMA. Figure 4 shows the electron scanning images of the slag taken at the position 10 mm above the blowing nozzle. Since Al$_2$O$_3$ concentration of No. 5 slag in Table 5 is high, it seems to be a top slag which is recirculated into bath. Other slags are judged to have been sampled at a fixed position.

The reason why the total concentration is less than 100 % seems to be that the samples of slag are porous or rough. Figure 4 shows the relationship between the sampling positions of slag and the concentration of FeO.

The FeO concentrations in Fig. 4 are shown as a percentage after the total percentage is corrected so as to be 100 %.

It is evident in Photo. 2 that the dephosphorization reaction has already taken place near the blowing nozzle. It is also shown that the FeO concentration of the slag sampled near the blowing nozzle is high, whereas that of the slag sampled at a position apart from the nozzle is low. As previously mentioned, these facts support the postulation that (CaO–Fe(Mn)$_O$)$_O$ is formed near the blowing nozzle, and the Fe(Mn)$_O$O concentration decreases through the reduction by carbon on its floating.

2. Single-crystal Lime Lump Immersion Test

Photograph 3 shows a micrograph of the section of

Table 3. Composition of iron particles in adhered slag analysed by EPMA (%).

<table>
<thead>
<tr>
<th>Size of iron particle (μm)</th>
<th>C</th>
<th>Si</th>
<th>Mn</th>
<th>P</th>
<th>S</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>1.55</td>
<td>0.03</td>
<td>0.18</td>
<td>0.271</td>
<td>0.019</td>
</tr>
<tr>
<td>35</td>
<td>1.97</td>
<td>0.02</td>
<td>0.87</td>
<td>0.667</td>
<td>0.021</td>
</tr>
<tr>
<td>15</td>
<td>1.11</td>
<td>0.03</td>
<td>0.62</td>
<td>0.065</td>
<td>0.028</td>
</tr>
<tr>
<td></td>
<td>0.64</td>
<td>0.02</td>
<td>0.64</td>
<td>0.044</td>
<td>0.023</td>
</tr>
<tr>
<td>7</td>
<td>0.78</td>
<td>0.02</td>
<td>0.71</td>
<td>0.017</td>
<td>0.003</td>
</tr>
<tr>
<td>5</td>
<td>1.38</td>
<td>0.02</td>
<td>0.68</td>
<td>0.016</td>
<td>0.013</td>
</tr>
<tr>
<td></td>
<td>2.60</td>
<td>0.02</td>
<td>0.65</td>
<td>0.013</td>
<td>0.006</td>
</tr>
<tr>
<td>4</td>
<td>1.78</td>
<td>0.01</td>
<td>0.69</td>
<td>0.010</td>
<td>0.035</td>
</tr>
<tr>
<td></td>
<td>1.43</td>
<td>0.01</td>
<td>0.32</td>
<td>0.010</td>
<td>0.029</td>
</tr>
<tr>
<td>3</td>
<td>1.43</td>
<td>0.05</td>
<td>0.62</td>
<td>0.004</td>
<td>0.031</td>
</tr>
<tr>
<td></td>
<td>1.14</td>
<td>0.03</td>
<td>0.61</td>
<td>0.021</td>
<td>0.025</td>
</tr>
<tr>
<td></td>
<td>1.09</td>
<td>0.02</td>
<td>0.60</td>
<td>0.010</td>
<td>0.027</td>
</tr>
<tr>
<td></td>
<td>1.81</td>
<td>0.02</td>
<td>0.42</td>
<td>0.013</td>
<td>0.021</td>
</tr>
</tbody>
</table>

Table 4. Analytical results of slag sampled from top of the bath after the termination of injection. (%)

<table>
<thead>
<tr>
<th>Total Fe</th>
<th>Met. Fe</th>
<th>FeO</th>
<th>Fe$_3$O$_5$</th>
<th>CaO</th>
<th>SiO$_2$</th>
<th>Al$_2$O$_3$</th>
<th>MgO</th>
<th>MnO</th>
<th>P$_2$O$_5$</th>
<th>S</th>
</tr>
</thead>
<tbody>
<tr>
<td>24.83</td>
<td>18.23</td>
<td>1.66</td>
<td>7.59</td>
<td>37.19</td>
<td>1.34</td>
<td>13.82</td>
<td>13.05</td>
<td>0.68</td>
<td>5.06</td>
<td>0.75</td>
</tr>
</tbody>
</table>

Table 5. Analytical results of slags floating up in hot metal by EPMA. (%)

<table>
<thead>
<tr>
<th>Sample No.</th>
<th>Sampling position</th>
<th>FeO</th>
<th>CaO</th>
<th>SiO$_2$</th>
<th>Al$_2$O$_3$</th>
<th>MgO</th>
<th>P$_2$O$_5$</th>
<th>S</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10 mm above nozzle</td>
<td>30.00</td>
<td>5.74</td>
<td>5.01</td>
<td>0.47</td>
<td>0.73</td>
<td>0.98</td>
<td>0.15</td>
<td>43.08</td>
</tr>
<tr>
<td>2</td>
<td>&quot;</td>
<td>16.28</td>
<td>22.74</td>
<td>2.76</td>
<td>0.69</td>
<td>1.36</td>
<td>12.05</td>
<td>1.21</td>
<td>57.09</td>
</tr>
<tr>
<td>3</td>
<td>&quot;</td>
<td>16.66</td>
<td>31.11</td>
<td>25.01</td>
<td>0.57</td>
<td>2.02</td>
<td>8.13</td>
<td>1.89</td>
<td>85.39</td>
</tr>
<tr>
<td>4</td>
<td>80 mm above nozzle</td>
<td>6.87</td>
<td>43.85</td>
<td>2.42</td>
<td>0.77</td>
<td>0.27</td>
<td>22.22</td>
<td>0.26</td>
<td>76.66</td>
</tr>
<tr>
<td>5</td>
<td>&quot;</td>
<td>8.44</td>
<td>3.58</td>
<td>17.25</td>
<td>6.63</td>
<td>2.77</td>
<td>9.78</td>
<td>1.50</td>
<td>49.95</td>
</tr>
</tbody>
</table>
a single-crystal lime lump immersed for 60 s during oxygen blowing. This photograph clearly shows a layer indicating the beginning of CaO slagging (I) and a slag layer (II~IV). In this case, the thickness of the layer indicating the beginning of slagging and that of the slag layer were 30 µ and 150 µ, respectively.

Photograph 4 shows the electron scanning images of the layer indicating the beginning of slagging and the succeeding reaction layer taken by EPMA. The layer indicating the beginning of slagging shows that Fe and Mn are concentrated (see Fig. 5). However, the penetration of P into the layer was found to be very little. High concentrations of Ca, Fe, Si and P were observed in the slag layer.

Figure 5 shows the results of line scanning of the reaction layer by EPMA. It is possible to divide the reaction layer into four parts according to the features of the distribution of each element.

Layer I which is in contact with CaO shows high concentrations of Fe and Mn and low concentrations of P and Si. This is the layer indicating the beginning of slagging.

![Graph](image)

Fig. 4. Relationship between the sampling position of slag and the concentration of FeO.

![Diagram](image)

1: Layer slag formation initiating
II~IV: Slag layer

Photo. 3. Microstructure of the cross-section of lime single-crystal immersed in hot metal for 60 s.
of slagging. Layer II shows high concentrations of Fe and Mn and the slagging of CaO is considered to be under way in this layer. The penetration of P has taken place in this layer. Layer III has high concentrations of Fe and Mn and the slagging of CaO is considered to be under way in this layer. The penetration of P has taken place in this layer. Layer IV has very high concentrations of Fe and Mn, and Ca is also observed. Therefore, the composition of this layer is considered to be \( (\text{CaO-Fe(Mn)O}) \).

To identify the phosphorus containing phases of \( \text{CaO-SiO}_2-P_2O_5 \) mineralogically, concentrations of Ca, Si, Fe, Mn and P at each peak of P were read from Fig. 5 and the compositions of CaO, SiO_2, FeO, MnO and P_2O_5 were calculated. The phosphorus containing mineral phases were identified by molar composition ratios of CaO, SiO_2 and P_2O_5.

The phosphorus containing mineral phase of Layer III was identified as either nagelschmidtite \( (7\text{CaO}_2\text{SiO}_2\cdot\text{P}_2\text{O}_5) \) or silicocarnotite \( (5\text{CaO} \cdot \text{SiO}_2 \cdot \text{P}_2\text{O}_5) \).

To identify the phase in reaction Layers I to IV during dephosphorization at 1350 °C, compositions of the phosphorus-containing phase (P-phase) and the iron oxide-containing phase (F-phase) in each layer of the reaction layer were estimated from Fig. 5 and are shown in Table 6. Further, each phase was regarded as the pseudo-ternary system of \( \text{CaO}^-\text{(SiO}_2+\text{P}_2\text{O}_5)^-\text{(FeO+MnO)}^+ \) and the liquidus temperature \( (T_{\text{LL}}) \) and solidus temperature \( (T_{\text{SL}}) \) of each phase were calculated by assuming that the phase diagram of the \( \text{CaO-SiO}_2-\text{FeO} \) system can be applied to the pseudo-ternary system.

As a result, it was found that in Layer I, i.e., near the starting location of slagging, only some F-phases were detected. And the liquidus and solidus temperatures are calculated from their compositions. In Layer I, since the experimental temperature (1350 °C) is lower than \( T_{\text{SL}} \) or between \( T_{\text{LL}} \) and \( T_{\text{SL}} \), those F-phases are solid or solid and liquid mixtures. The P-phase in Layer II, in the same manner, is a solid phase, while the F-phase in this layer is close to a liquid phase. In Layer III, the P-phase is a solid phase and the F-phase is a solid and liquid mixture. In this layer, however, the proportion of the P-phase is high. Therefore, it is presumed that as a whole, the liquid phase exist at the grain boundaries.

Fig. 5. Concentration profiles of the reaction layer obtained by EPMA.

Fig. 6. Changes in the reaction layer thickness formed on single-crystal lime (after immersion for 60 s at 1350 °C).
boundaries of solid particles. In Layer IV, the P-phase is in a solid and liquid coexisting state, and the F-phase is high in proportion, suggesting the predominance of liquid phase as a whole.

Figure 6 shows the reaction layer thickness of single-crystal lime samples immersed for 60 s at the positions 20, 40 and 60 mm apart from the furnace bottom during the bottom blowing of oxygen into hot metal. The reaction layer thickness decreases with increasing distance from the furnace bottom, i.e., oxygen blowing nozzle.

Further, the effect of iron oxide addition was studied under the condition that the total amount of oxygen consumption was constant. Figures 7 and 8 show the results obtained when a part of oxygen was replaced by iron oxide. The reaction layer thickness and the concentration of phosphorus in the layer decrease more slightly in the case of iron oxide addition with increasing distance from the furnace bottom than the case without iron oxide injection. Therefore, it is so judged that the range of dephosphorization reaction is enlarged by the addition of iron oxide.

IV. Discussion

1. Slagging Process of Lime in Hot Metal

Photographs 3 and 4 clearly show a layer indicating the beginning of CaO slagging. As is apparent from Fig. 5, the concentrations of Fe and Mn are very

Table 6. Identification of the phase in reaction Layers I~IV during dephosphorization at 1350 °C.

<table>
<thead>
<tr>
<th>Position in reaction layer</th>
<th>Mineral phase</th>
<th>CaO (%)</th>
<th>SiO₂ (%)</th>
<th>FeO (%)</th>
<th>MnO (%)</th>
<th>P₂O₅ (%)</th>
<th>Temperature calculated from phase diagram</th>
<th>State of phase (at 1350 °C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>F-1</td>
<td>53.5</td>
<td>4.2</td>
<td>31.1</td>
<td>8.3</td>
<td>2.9</td>
<td>≈ 1 750, ≈ 1 570</td>
<td>solid</td>
</tr>
<tr>
<td></td>
<td>F-2</td>
<td>32.1</td>
<td>1.0</td>
<td>33.3</td>
<td>30.4</td>
<td>3.2</td>
<td>≈ 1 400, ≈ 1 210</td>
<td>solid</td>
</tr>
<tr>
<td>II</td>
<td>P</td>
<td>65.6</td>
<td>17.0</td>
<td>1.1</td>
<td>1.5</td>
<td>14.8</td>
<td>≈ 2 080, ≈ 1 960</td>
<td>solid</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>20.6</td>
<td>2.0</td>
<td>41.8</td>
<td>34.5</td>
<td>1.1</td>
<td>≈ 1 380, ≈ 1 220</td>
<td>solid</td>
</tr>
<tr>
<td>III</td>
<td>P</td>
<td>59.1</td>
<td>6.2</td>
<td>0.5</td>
<td>1.5</td>
<td>32.7</td>
<td>≈ 1 810, ≈ 1 450</td>
<td>solid</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>50.3</td>
<td>6.1</td>
<td>15.4</td>
<td>8.9</td>
<td>21.0</td>
<td>≈ 1 830, ≈ 1 280</td>
<td>solid</td>
</tr>
<tr>
<td>IV</td>
<td>P</td>
<td>51.2</td>
<td>13.2</td>
<td>11.5</td>
<td>6.8</td>
<td>17.3</td>
<td>≈ 1 820, ≈ 1 250</td>
<td>solid+liquid</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>6.0</td>
<td>0.9</td>
<td>48.6</td>
<td>44.5</td>
<td>0</td>
<td>≈ 1 320, ≈ 1 200</td>
<td>liquid</td>
</tr>
</tbody>
</table>

Fig. 7. Relationship between the distance from oxygen blowing nozzle and the thickness of reaction layer of lime during iron oxide injection.

Fig. 8. Relationship between the distance from oxygen blowing nozzle and the average concentration of P₂O₅ in the reaction layer during iron oxide injection.

* P: Phosphorus oxide containing phase, F: Iron oxide containing phase
high in this layer and it can be said that slagging takes place owing to the formation of CaO–Fe(Mn),O by the reaction of Fe(Mn),O and CaO. If silicon is contained in hot metal, the slag is naturally CaO–Fe(Mn),O–SiO₂. An investigation was made in the initial stage of slagging, and the results obtained revealed that the concentrations of Fe, Mn and Si are all high. This fact proved that the reaction of CaO and P begins after the diffusion of Fe, Mn and Si into CaO. On the other hand, the peaks of P and Fe are always inverse outside the F-2 phase of Layer I, i.e., the part which is high in phosphorus is poor in iron, and on the contrary, the part which is high in iron is poor in phosphorus. From Table 6, this F-2 phase is considered to be a mixture of solid and liquid phases at the experimental temperature (1350 °C), and it might be thought that after the reaction with phosphorus, a different phase has precipitated from liquid slag in the cooling process.

In Layer II, the concentrations of Fe, Mn, Ca, Si and P are not uniform and the peak of each component indicates (CaO–SiO₂–P₂O₅)ₚ [P-phase] and (CaO–Fe(Mn),O–SiO₂–P₂O₅)ₚ [F-phase], where, Tₗₑ and Tₛ₂ of F-phase are about 1380 °C and 1200 °C, respectively. Therefore, slagging is considered to have proceeded considerable in this layer.

From these results, it is thought that CaO injected into hot metal slags to form (CaO–Fe(Mn),O), at first which reacts with phosphorus later.

2. Mechanism of Dephosphorization during Lime Injection by Oxygen

The dephosphorization mechanism of hot metal by CaO was reasoned as follows and showed in Fig. 9 on the basis of the results of the single-crystal lime immersion and lime injection tests. When CaO is injected into hot metal with oxygen, oxidation of Si, Mn, Fe and C takes place near the fire point of the blowing nozzle.

\[ \text{Si} + \text{O}_2 \rightarrow \text{SiO}_2 \] ..................................(1)

\[ \text{Mn} + \text{O}_2 \rightarrow \text{MnO} \] ..................................(2)

\[ \text{Fe} + \text{O}_2 \rightarrow \text{FeO} \] ..................................(3)

\[ \text{C} + \frac{1}{2}\text{O}_2 \rightarrow \text{CO} \] ..................................(4)

Equation (3) is not referred to here because CO gas does not directly participate in the dephosphorization reaction. (CaO–Fe(Mn),O)ₚ is then formed by the reactions of CaO with MnₙO and FeₚO.

\[ \text{CaO} + \text{Fe}_p \text{O} \rightarrow \text{[CaO–Fe(Mn),O]} \] ..................................(4)

The dephosphorization reaction begins at the interface between this multicomponent liquid and hot metal.

\[ \text{[CaO–Fe(Mn),O]} + \text{P} \rightarrow \text{[CaO–Fe(Mn),O–P}_2\text{O}_5] + \text{Fe}_p \] ..................................(5)

When Eq. (5) proceeds and the P₂O₅ concentration becomes high, solid phases such as silicocarnotite and nagelschmidtite precipitate since SiO₂ is included in the liquid slag.

\[ \text{[CaO–Fe(Mn),O–SiO₂–P}_2\text{O}_5] \rightarrow \text{[CaO–SiO₂–P}_2\text{O}_5] + \text{[CaO–Fe(Mn),O]} \] ..................................(6)

Liberated Feₚ in Eq. (5) reacts with carbon in hot metal. As a result, its melting point drops and it melts.

\[ \text{Fe}_p + \text{C} \rightarrow \text{[Fe–C]} \] ..................................(7)

With an increase in distance from the oxygen blowing nozzle, the concentrations of Fe₂O₅ and MnₙO at the CaO interface decrease because the blown oxygen is consumed by decarburization, etc., and Eq. (4) becomes hard to occur. Further, it is presumed that the reduction of the reaction product by carbon also takes place.

\[ \text{[CaO–Fe(Mn),O]} + \text{C} \rightarrow \text{Ca(Mn),O} + \text{[Fe–Mn–C]} \] ..................................(8)

\[ \text{[Fe–Mn–Cl]} + \text{C} \rightarrow \text{[Fe–Mn–C]} \] ..................................(9)

From the results of the single-crystal lime immersion and lime injection tests, it is conceived that dephosphorization during CaO injection with oxygen occurs mainly near the oxygen blowing nozzle.

This is because at a position apart from the nozzle, the reaction of Eq. (4) becomes hard to occur and the amount of [CaO–Fe₂O₅–MnₙOₚ] decreases as a result of Eq. (8).

V. Conclusion

An investigation was made on the mechanism of dephosphorization during CaO injection with oxygen by single-crystal lime immersion and lime injection tests, and the followings were made clear:

1. Slagging of CaO in hot metal takes place in the process of (CaO–Fe(Mn),O)ₚ formation through the reaction of CaO with Fe₂O₅ and MnₙO near the oxygen blowing nozzle.

2. The dephosphorization proceeds when [CaO–
Fe(Mn)O of high FeO concentration is present. When the P2O5 concentration in the reaction layer increases, SiO2 present in slag also participates in the reaction and the concentration of Fe(Mn)O decreases, solid phases of (CaO-SiO2-P2O5)n, such as nagelschmidtite and silicocarnotite, precipitate.

(3) These reactions proceed in a short period of time at a position near the oxygen blowing nozzle. The amount of oxygen supplied becomes insufficient at a position apart from the oxygen blowing nozzle. Therefore, the formation of \( \text{CaO-Fe(Mn)O}_t \) slows down and reduction of \( \text{CaO-Fe(Mn)O}_t \) by C takes place. Accordingly, the dephosphorization reaction becomes hard to occur. By the addition of iron oxide together with oxygen, the region where \( \text{CaO-Fe(Mn)O}_t \) reacts can be enlarged. In order to promote the dephosphorization reaction, it is necessary to form \( \text{CaO-Fe(Mn)O}_t \) of high FeO concentration at an early stage.

In this sense, it is important to blow oxygen and is effective to inject iron oxide together with oxygen in order to enlarge the region where \( \text{CaO-Fe(Mn)O}_t \) reacts.

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REFERENCES