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Comprehensive Model of Oxygen Steelmaking
Part 3: Decarburization in Impact Zone

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The development of a global model for oxygen steelmaking and its validation against industrial data has been reported in Part 1 of this paper. Part 2 of this paper explained the model development of decarburization reaction of emulsified droplets and discussed the effects of bloating behaviour of metal droplets on the overall kinetics of the process. Part 3 of this paper focused on the development of one sub-model on the decarburization reaction in the impact zone and critically evaluates the important process variables affecting the decarburization kinetics. Decarburization rates in the impact zone were calculated using the semi-empirical relationships developed from experimental results. Based on previous experimental studies, both diffusion through the gas phase and surface control were considered in the rate calculations. The model was validated against experimental data from Belton and Sain, and against plant data from Cicutti et al. The model developed was consistent with the experimental and plant data, and provided a reasonable basis for predicting the decarburization of iron in the impact region of oxygen steelmaking. The model predicted that the decarburization rate is sensitive to the partial pressure of carbon dioxide and oxygen in the impact zone. As the partial pressure of oxygen decreased from 26.4 to 16.2 kPa, the model predicted that the reaction rate of decarburization via oxygen decreased from 202 to 134 kg/min. The model predicted that approx. 40% of decarburization takes place in the impact zone during the main blow.

KEY WORDS: impact area; gas diffusion; sulphur; decarburization rate; oxygen steelmaking.

1. Introduction

The reactions between metal bath and oxidizing gasses are of particular interest, as a significant proportion of carbon removal in oxygen steelmaking occurs in the impact zone in the reactor.1–3) Unfortunately, distinguishing between decarburization from the emulsion and impact zones is very difficult from the analysis of plant data, where only the overall rate of carbon removal is measured. Also, duplicating the conditions of oxygen steelmaking experimentally is challenging. We have attempted to address this issue by developing models for both the emulsion zone (Part 24)) and the impact zone (current paper), combining them together into a global model and comparing them to the best plant data available (Part 15) of this series of papers).

It is known, as oxygen from a top blown lance reaches the surface of the liquid bath, it reacts with carbon dissolved in the metal at the impact zone and forms a mixture of CO and CO2 gases. Subsequently, dissolved carbon also reacts with CO2 simultaneously at this region. The following equations can be used to represent the decarburization reactions at the impact area;6)

\[ \text{C} + \text{CO}_2(\text{g}) = 2\text{CO}(\text{g}) \quad \text{(1)} \]

\[ \text{C} + \text{1}/2\text{O}_2(\text{g}) = \text{CO}_2(\text{g}) \quad \text{(2)} \]

The possible reaction mechanisms for above reactions are given below;

(i) mass transfer of oxygen in the gas phase
(ii) mass transfer of carbon monoxide in the gas phase
(iii) mass transfer of carbon dioxide in the gas phase
(iv) mass transfer of carbon through metal phase
(v) chemical reaction between carbon and oxygen at the interface

Steps (i),(ii) and (vi) apply to decarburization reaction with O2 whereas (ii), (iv) and (vi) apply to decarburization reaction with CO2. It has been postulated that the mass transfer of carbon can be neglected at high carbon concentrations (above critical carbon content).1,6–11) The adsorption of oxygen can also be neglected since oxygen has been present already in the liquid iron.

There have been numerous studies1,3,7–35) on the mechanism and kinetics of decarburization of Fe–C melts with oxidizing gasses using levitation and crucible techniques. The conclusion from the previous studies is in general agreement, i.e. gaseous diffusion controls the decarburization rate down to the critical level of carbon, where carbon diffusion in liquid phase becomes rate-limiting step. However, some investigators have suggested that sulphur has a determining effect on the reaction rate and chemical reaction at the interface also controls the reaction rate.30)

Previous experimental studies, both in laboratory and...
industrial scales, have improved the fundamental understanding of decarburization reactions via gases during the blowing process.\(^{1,3,7–35}\) Despite this, there is only limited industrial data\(^{36}\) that consider carbon removal rates via direct oxidation at the impact zone due to the complexity of the process. This study is an attempt to use the theoretical findings from the experimental studies to the full-scale operating conditions.

2. Model Development

2.1. Rate-Determining Step

In this study, it has been considered that decarburization reactions in the impact zone varying over a large range of temperature and fluid flow conditions, with constant concentration of sulphur. In the case of decarburization reaction via CO–CO\(_2\), Nomura and Mori\(^{17}\) and Fruehan and Martonik\(^{20}\) proposed that the effect of sulphur (sulphur concentration below 0.3 mass\%) is relatively small on the reaction rate in CO–CO\(_2\) gas mixture at high carbon concentrations. Goto et al.\(^{34}\) reported that sulphur has no significant effect on the reaction kinetics. Later, Sain and Belton\(^{7}\) and Mannion and Fruehan\(^{11}\) studied the decarburization kinetics under high gas flow rates and found that CO\(_2\) dissociation on the surface controls the reaction rate. Lee and Rae\(^{8,9}\) proposed that the surface active elements, the gas flow rate, partial pressure of oxygen and gas composition have an influence on the decarburization rate. However, the rate of decarburization is markedly controlled by the composition of gas since it determines the amount of oxygen transferred to the system.

A model based on a mixed control kinetics including gas phase mass transfer and chemical kinetics from the study by Sain and Belton\(^{7,30}\) was used since it has been established that sulphur has a retarding effect on the kinetics of decarburization reaction\(^{1,26,30,37}\) and this effect was included in our model to investigate the reaction rates of decarburization under various operational conditions. The rate equation for decarburization reaction via CO\(_2\) can be written by: \(^{7,30}\)

\[
\frac{-dW_C}{dt} = 100m_{wC}Ak_j\rho_{w}P_{CO2} \quad \text{---------- (3)}
\]

where

\[
k_j = \frac{1}{RT_j / k_g + 1 / k_s} \quad \text{---------- (4)}
\]

Here \(k_g\), \(k_j\) and \(k_a\) are the gas phase mass transfer coefficient, chemical rate constant and apparent rate constant, respectively. \(P_{CO2}\) is the partial pressure of CO\(_2\) in the gas mixture, \(R\) is the gas constant, \(T_j\) is the average gas temperature, \(A\) is the surface area and \(m_{wC}\) is molecular weight of carbon.

In this study, it was assumed that the rate of decarburization via oxygen was controlled by mass transfer in the gas phase since it has been established that gas diffusion has the predominant effect on the reaction kinetics and surface active elements play no important role on the reaction mechanism.\(^{10,11,16,21}\) The rate equation can be written using: \(^{10,16}\)

\[
\frac{-dW_C}{dt} = 200m_{wC}Ak_s \ln \left(1 + P_{O2}^b\right) \quad \text{---------- (5)}
\]

Equations (3) and (5) were applied to determine the reaction rates during the blowing period in which the carbon content was high. Below a critical value, carbon diffusion in the liquid phase was considered to control reaction rates of decarburization. The rate equation was represented in mass unit by the Eq. (6):\(^{7}\)

\[
\frac{dW_C}{dt} = k_{ma}\frac{\rho_m}{100} \left(\text{mass}\% \text{ } C_b - \text{mass}\% \text{ } C_{eq}\right) \quad \text{---------- (6)}
\]

where \(\rho_m\) is density of liquid iron and \(k_{ma}\) is mass transfer coefficient of carbon in liquid iron. The equilibrium carbon content, \(C_{eq}\) was small and neglected in the calculations.

Most of the experimental studies focused on the decarburization reaction of Fe–C melts or droplets and the effect of sulphur on the reaction mechanism was investigated. There are few studies\(^{12,23,24,27}\) that consider the influences of other refining reactions on the decarburization kinetics. Robertson and Jenkins\(^{12}\) observed the behaviour of levitated droplets containing C, Si and Al with oxygen. They found that the silicate layers formed initially on Fe–C–Si droplets which retard the decarburization reaction. Similar behaviour was observed by Sun and Pehlke.\(^{24}\) They studied the kinetics of simultaneous oxidation of carbon, silicon, manganese and sulphur in a liquid metal droplet by oxygen and/or carbon dioxide in nitrogen gas at 1 873 to 1 993 K. They observed that there was a delay in silicon and manganese oxidation reactions at high temperatures and high carbon contents since the decarburization reaction consumed the most of the oxygen supplied to the system. In the case of low carbon content (<0.4 mass\%), the simultaneous oxidation of manganese and silicon was observed in their experiments. The effects of other refining reaction was ignored in this study, it is difficult to relate the experimental results from droplets to the impact region of an oxygen steelmaking vessel and it is reasonable to assume that any silicate layer formed would quickly dissolve into the slag. Further experimental work is required to clarify the impact of other elements on decarburization in the impact region.

2.2. Calculation of Rate Constants

There are few studies\(^{7,38–42}\) for impinging gas jets at metal surface. Based on the study of Rao and Trass,\(^{30}\) Sain and Belton\(^{7}\) suggested a mass transfer correlation for the case of an impinging jet onto a liquid surface.

\[
Sh = 0.026 Re^{1.06} Sc^{0.53} \left(\frac{z’}{d_c}\right)^{-0.09} \quad \text{---------- (7)}
\]

This correlation is valid for \((z’/d)\leq 6.5, (x/d)\leq 4.5\), which is not the case in the industrial configuration. Lohe\(^{41}\) also suggested mass transfer correlations for gas side mass transfer, which were given based on the range of Reynolds number;

\[
Sh = 1.41 Re^{0.51} Sc^{0.33}, \quad 2 \cdot 10^2 \leq Re \leq 3 \cdot 10^4 \quad \text{---------- (8)}
\]

\[
Sh = 0.41 Re^{0.75} Sc^{0.53}, \quad 3 \cdot 10^4 \leq Re \leq 2 \cdot 10^5 \quad \text{---------- (9)}
\]

Here \(Sh = k_{d\rho D}/D\), \(Re = ur\rho/\mu\) and \(Sc = \mu/(\rho D)\). \(d_c\) refers to nozzle throat diameter and \(z’\) denotes the distance of the nozzle from the surface of the liquid. Interdiffusivity, viscosity and density of fluid (gas) at \(T_j\) are \(D\), \(\mu\) and \(\rho\),
respectively. Sc varies between 0.1 and 2 for gases. These correlations were used for the range of the Reynolds number applicable in this study. The velocity of the gas to be inserted into the Reynolds number was designated as the mean velocity of the gas. In this study, it was assumed to be equal to the impact velocity from the oxygen lance. The impact velocity was related with jet centreline velocity at the metal surface which can be obtained using:

\[ U_G = \eta U_f \]  

\( \eta \) is a constant and its value is 0.44721. The jet centreline velocity can be obtained by the equation for the dynamic impact pressure of the jet at the metal surface.6)

Variable \( r_0 \) is the radius of inducted surface area. In this study, the variable \( r_0 \) was considered to be the radius of individual penetration area. There are few proposed correlations \( 46-50 \) to calculate the diameter of the penetration area. The correlation developed by Koria and Lange \( 50 \) was used to estimate the diameter of the cavity since this study was based on an experimental study at steelmaking temperatures and the penetration correlations have been widely used by many researchers.

According to the Sain and Belton \( 7,30 \) and Nagasaka and Fruhman, \( 51 \) the rate constant of dissociative adsorption of CO\(_2\) for liquid \( \varphi \) iron by CO–CO\(_2\) was:

\[ k_f = \frac{k_f}{1 + K_f \gamma_f (\text{mass} \% \text{S})} + k_k \]  

where \( k_f, k_k, K_f \) and \( \gamma_f \) refer to chemical rate constant for pure iron, residual rate at high sulphur contents, adsorption coefficient of sulphur and activity coefficient of sulphur in liquid iron, respectively. In the study of Nagasaka et al., \( 51 \) the standard state for sulphur activity was taken as 1 mass% in carbon-saturated liquid iron. Rate constants \( k_f, k_k \) and adsorption coefficient, \( K_f \) can be calculated as a function of temperature using, \( 30,51 \)

\[ \log k_f = -\frac{5080}{T} - 0.21 \]  

\[ \log k_k = -\frac{5600}{T} - 1.75 \]  

\[ \log K_f = -\frac{3600}{T} + 0.57 \]

In the case of mass transfer in the metal phase, mass transfer coefficient was related to the stirring intensity in the metal bath. Several researchers \( 32-35 \) proposed a correlation between mass transfer coefficient to gas flow rate for gas stirred liquid-liquid systems. Accordingly, the mass transfer coefficient can be found using, \( 5,56 \)

\[ k_m = \beta \left( \frac{D_c}{F_G} \right)^{\frac{1}{2}} \]  

where \( \beta \) is constant and equal to 500 m\(^{-0.5}\) \( F_G \) is the volumetric gas flow rate (m\(^3\)/s) and it was assumed that inert gas blowing from the bottom of the furnace influences the mixing in the bath. \( D_c \) is the diffusion coefficient of carbon. The relationship between diffusivity in liquids and temperature and viscosity of liquid by the Stokes-Einstein and Eyring equations was used to calculate the diffusivities of carbon in liquid iron for various temperatures.

### 2.3. Calculation of Partial Pressure

Determination of partial pressure of oxidizing gasses such as CO\(_2\) and O\(_2\) is crucial since it governs the amount of gas delivered to the system in order to achieve the reactions at the impact zone. The partial pressure of the gasses in the system varies as a function of reaction rates and gas composition. In this model, it was assumed that the amount of gas blown (O\(_2\) and Ar–N\(_2\)) and 10% of the total amount of gas generated from the decarburization reaction were available in the impact zone for the given time interval. The gas generation by the decarburization reaction via emulsion was not included in the calculations of partial pressure of gasses in the impact zone. In this model, the total amount of gas available in the system was calculated by the summation of the gasses (O\(_2\), CO, CO\(_2\) and Ar–N\(_2\)) in unit of mol over time interval. The partial pressure of CO\(_2\) and O\(_2\) were calculated from the molar rates of Ar–N\(_2\) (\( N_A \)), O\(_2\) (\( N_{O_2} \)), CO (\( N_{CO} \)) and CO\(_2\) (\( N_{CO_2} \)) and the average pressure in the bath (\( P \)):

\[ P_{CO_2} = \frac{N_{CO_2}}{N_{CO_2} + N_{CO} + N_A + N_{O_2}} \times P \]  

\[ P_{O_2} = \frac{N_{O_2}}{N_{CO_2} + N_{CO} + N_A + N_{O_2}} \times P \]

### 2.4. Calculation of Gas Temperature

The average gas film temperature \( T_f \) is the mean of the temperature of the bulk gas and the temperature at the gas-liquid interface. The film temperature was estimated using, \( 16 \)

\[ T_f (K) = \frac{T_s + T_b}{2} \]

In this study, the temperature of CO\(_2\), \( T_s \) was assumed to equal to the temperature of impact zone whereas the temperature of O\(_2\) was assumed to be 25°C. Koch et al. \( 57,58 \) suggested that the temperature at impact area increases very rapidly in the early blow and it stays at a maximum level during the active decarburization period. Towards the end of the blow, the impact area temperature disappears. According to these studies, the impact temperature was assumed to be 2000°C until 4 min after the start of the blow, followed by an increase to 2500°C until 14 min after the blow and decreasing to the bath temperature towards the end of the blow in this study.

All dimensionless groups, thereby mass transfer coefficients of CO\(_2\) and O\(_2\) were calculated at the film temperatures of the related gasses as per previous studies \( 3,16 \) whereas rate constants \( k_f, k_k \) and adsorption coefficient, \( K_f \) were calculated at the impact temperature of the process.

### 2.5. Calculation of Impact Area

It is known that an increase in impact area increases the reaction rates significantly. \( 26,59 \) Zaghib et al. \( 26 \) studied experimentally the effects of bath surface area on the kinetics of decarburization reaction of Fe–C melts using crucible tech-

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nique at 1 450°C. They found that a decrease in the bath area lowers the reaction rates. They observed that the reaction also takes place outer of the penetration area. However, there is limited knowledge on the reaction area at the impact zone due to the difficulties in measurements and visualization at high temperatures.

In this study, the penetration area was assumed to be the reaction area for carbon removal reaction. The multi-head lance creates individual cavities on the liquid bath. The total impact area of a jet can be calculated by summation of individual areas for multi-head lance. The shape of the cavity was assumed to be paraboloid. The individual impact area was calculated using:

\[
\text{Area} = \int_0^r \int_0^{dh} \left[ 1 + \left( \frac{dh}{dr} \right)^2 \right] dr \quad \text{...... (19)}
\]

where \( h \) is paraboloid height and \( r \) is paraboloid radius. The height equals to the penetration depth which can be calculated using Korea and Lange’s relationship. Similarly, the radius equals to half of the penetration diameter and was also taken from Koria and Lange. There would be also the change in the cavity throughout the blow due to the surface waves. But this effect was ignored based on the findings from a study of Cheslak et al., since the cavity oscillations did not affect the final result of their observations.

2.6. Calculation of Critical Carbon Content

It might be hard to accept a constant value for the critical carbon content. However, it is well known that the value is below 1. Different proposals were made to determine the critical carbon content of liquid iron. Goldstein and Fruehan defined the critical carbon content as the carbon content where the decarburization reaction rate during the main blow equal to the decarburization rate at the end of the blow. In this study, this approach was used and it was assumed that the point where the total rate of decarburization via gases is equal to the decarburization rate controlled by mass transfer of carbon in the liquid iron represents the critical carbon content. This value was obtained using:

\[
\frac{24A}{\rho V} k_w \ln \left( 1 + \frac{P_{O_2}}{P_{O_2}} \right) + \frac{2A}{\rho V} k_w P_{CO} \quad \text{...... (20)}
\]

2.7. Calculation of Physical Properties of Gas

2.7.1. Viscosity

The viscosity of gases can be estimated using the Lennard-Jones parameter. The relationship is given:

\[
\mu_g = 266.93 \times 10^{-7} \frac{\sqrt{m_{w_g} \times T_f}}{d_c^2 \times \Omega} \quad \text{...... (21)}
\]

where molecular weight of gas is expressed as \( m_{w_g} \) and collision diameter of gas is as \( d_c \). The collision integral can be approximated by:

\[
\Omega_{AB} = 1.147 \left( \frac{kT}{\varepsilon} \right)^{0.145} + \left( \frac{kT}{\varepsilon} + 0.5 \right)^2 \quad \text{...... (22)}
\]

In this equation, a molecule’s kinetic energy was represented by \( kT \) while the potential energy of two colliding molecules was represented by \( \varepsilon \), which was Lennard-Jones potential well depth. The characteristic parameters, \( \varepsilon/k \) and \( d_c \) for \( O_2 \), \( CO \) and \( CO_2 \) were taken from the literature and are given in Table 1.

### Table 1. Characteristic parameter of gases

<table>
<thead>
<tr>
<th>Characteristic Parameters</th>
<th>( O_2 )</th>
<th>( CO )</th>
<th>( CO_2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \varepsilon/k )</td>
<td>113.2</td>
<td>110.3</td>
<td>190</td>
</tr>
<tr>
<td>( d_c (10^{-10}\text{m}) )</td>
<td>3.433</td>
<td>3.59</td>
<td>3.996</td>
</tr>
</tbody>
</table>

The Chapman-Enskog theory was applied to predict the interdiffusivity of gases as a function of temperature. The relationship is:

\[
D_{AB} = \frac{0.0018583 T_f^{1/2} \left( \frac{1}{mw_A} + \frac{1}{mw_B} \right)}{P \times d_{AB}^2 \times \Omega_{D,AB}} \quad \text{...... (23)}
\]

where the collision diameter between gases can be found using:

\[
d_{AB} = 0.5 \left( d_{A} + d_{B} \right) \quad \text{........... (24)}
\]

The collision integral for mixtures can be approximated by Cloutman using:

\[
\Omega_{D,AB} = \left( \frac{kT}{\varepsilon_{AB}} \right)^{0.145} + \left( \frac{kT}{\varepsilon_{AB}} + 0.5 \right)^2 \quad \text{...... (25)}
\]

\[
\left( \frac{\varepsilon_{AB}}{k} \right) = \left( \frac{\varepsilon_A}{k} + \frac{\varepsilon_B}{k} \right)^{0.5} \quad \text{........... (26)}
\]

2.8. Formulation of the Model

The sequence of calculation procedure for decarburization in impact zone model is shown in Fig. 1. The data from hot metal composition such as carbon and sulphur, oxygen flow rate, lance height and bottom gas flow rate with time were taken from the related sub-models. The penetration area was calculated as a function of lance dynamics and gas flow rates. The values from bath temperature sub-model were used to calculate the gas film temperatures since the physical properties of gases, thereby dimensionless groups such as \( Re, Sc \) were calculated as a function of gas film temperature. Mass transfer rates of \( CO_2 \) and \( O_2 \) and the rate constant for \( CO_2 \) reaction were estimated using Eqs. (7), (8), (9) and (11) to calculate individual decarburization rates via oxygen and carbon dioxide.

2.9. Model Validation

Model predictions for overall rate constants of \( CO_2 \) including both gas diffusion and chemical kinetics are given as a function of sulphur concentration in Fig. 2. The predictions were compared with the experimental data of Sain and
Belton. Experiments were carried out with a lance height of 2–3 mm and a flow rate of 0.02–0.03 m$^3$/min. CO$_2$ gas was blown with Ar and N$_2$ gases onto the liquid iron bath between 1160 and 1600°C. The solid lines represent the model predictions whereas the points were related to the experimental data by Sain and Belton.

The model results are consistent with the experimental data reported by the previous researchers. The mass transfer coefficient values are much higher (approximately 6 × 10$^{-4}$ mol/m$^2$.Pa.s for 0.2–0.1 mass% sulphur and 10 l/min gas flow rate at 1600°C) than the chemical kinetics constants (3.7–5.5 × 10$^{-5}$ mol/m$^2$.Pa.s) that the decarburization reaction is controlled by the dissociative adsorption of CO$_2$. Accordingly, the reaction rate is influenced by the temperature of the bath and sulphur concentration of liquid iron. As the temperature of the bath increases, the rate increases.

3. Results and Discussion

3.1. Rate Constants

The model was further studied using the industrial conditions reported by Cicutti et al., i.e. top-blown oxygen with a flow rate of 620 Nm$^3$/min onto the 200 ton oxygen steel-making vessel. The inert gas was ejected through the bottom of the furnace to increase the mixing properties of the liquid bath. The lance design and some other process conditions were provided in Part 1 of this series of papers. The bath sulphur concentration was assumed to be constant with a value of 0.015 mass% during the blow. Figure 2 shows the predictions for the rate constants for CO$_2$ during the blow. The mass transfer coefficient, $k_g$, was influenced by gas velocity, penetration profile and physical properties of the gasses, simultaneously and the values of $k_g/RT_f$ varied between 25–35 × 10$^{-5}$ mol/m$^2$.s.Pa (270–378 m/min). The predicted values for the mass transfer coefficient of oxygen in the gas phase were between 560 and 670 m/min. An increase in gas velocity and impact area radius causes an increase in the gas-metal transfer coefficients. In practice, these properties vary dynamically and have an important impact on the overall kinetics of the system.

The reaction rate constant, $k_t$ is only a function of impact temperature since sulphur concentration is assumed to remain constant during the blow. Consequently, as the impact zone temperature decreases, the reaction rate constant also decreases. As can be seen in Fig. 3, the values for gas diffusion and chemical reaction constants are close to each other 2 min after the start of blowing. This suggests that the decarburization of liquid iron via CO$_2$ at the impact zone is influenced by both mixed controlled at high carbon concentration under the various operating conditions such as temperature, gas flow rate and lance height.

3.2. Impact Area

The impact area was calculated as a function of penetration depth and diameter. Figure 4 shows the predicted individual impact area as a function of lance dynamics. As the lance height decreases, the penetration depth increases and the radius of the penetration decreases. At lower lance heights, the penetration would be deeper with lower penetration radius; therefore, penetration area would decrease. These findings are in agreement with those by Koria and Lange. The individual penetration area ranges between 2.6 and 2.3 m$^2$ using Eq. (21). In this study, it is assumed that the interaction between the separate jets does not occur. This assumption is valid for jets with an inclination angle higher than 8°. Consequently, the total impact area is predicted to vary from 13 to 15.5 m$^2$. 

Fig. 1. Algorithm of the decarburization at impact zone model.

Fig. 2. Rate constant of CO$_2$ as a function of sulphur concentration calculated at different temperatures using the data of Sain and Belton. Closed circles are for experimental data, solid lines are for model results.

Fig. 3. The variations in rate constants for CO$_2$ throughout the blow.
3.3. Decarburization via O₂

The reaction rates of decarburization via oxygen were calculated using Eq. (5) as a function of partial pressure of oxygen, impact area and mass transfer of oxygen in the gas boundary layer given in Fig. 5. The reaction rates varied from 180 to 220 kg/min. The reaction rate increases the impact area or mass transfer constant or partial pressure of oxygen increases. The reaction rates increases throughout the blow except the periods when the partial pressure of oxygen drops significantly. It was found that, if the partial pressure decreases from 26.4 to 16.2 kPa, the decarburization rate decreased from 202 to 134 kg/min. This suggests that partial pressure of oxygen has a decisive impact on the decarburization rates. However, it should be noted that these parameters have relative importance on the kinetics of decarburization reactions via oxygen gaseous in a dynamic oxygen steelmaking process.

3.4. Decarburization via CO₂

Figure 6 gives the estimated decarburization rates via CO₂ as a function of partial pressure of CO₂, apparent rate constant and impact area for the region above the critical carbon content. The rates of decarburization were calculated for CO₂ using Eq. (3). The reaction rate dropped much less rapidly with the partial pressure of CO₂ with compared to rates of decarburization via oxygen. For example, the increase in partial pressure of CO₂ increases the reaction rate slightly with time in the early part of the blow as the impact area and rate constant remain constant.

A similar pattern emerges in the comparison of the estimated rate constant and decarburization rate in Fig. 6. This implies that apparent rate constant is relatively more important on the kinetics of the decarburization reaction via CO₂.

3.5. Effect of Bottom Stirring

In industrial practice, inert gas was blown through the bottom of the furnace at a flow rate of 150 Nm³/h. This flow rate was increased to 500 Nm³/h in the last two min of the process. Below critical carbon content, metal-phase mass transfer controlled the refining rates. It is known that bottom stirring increases the transfer rates in the bath significantly. Fig. 7 shows the predictions of decarburization rate as a function of gas flow rate, carbon concentration and mass transfer coefficient of carbon in liquid iron. As can be seen,
Decarburization rate is dependent on carbon concentration and decreases as the carbon content decreases towards the end of the blow. However, it is also shown that as the bottom gas flow rate is increased to 500 Nm\(^3\)/h, the value of mass transfer coefficient and decarburization rate increase.

### 3.6. Decarburization Rate in Impact Zone

In the study by Cicutti et al.,\(^6\)\(^7\) the instantaneous decarburization rates were not provided and the overall reaction rates were calculated based on the predictions on the carbon content of liquid iron. The direct comparison of reaction rates in the impact zone is difficult. In this study, the decarburization rate in the impact zone was estimated by subtraction of calculated decarburization rates in emulsion phase from the overall decarburization rates estimated from the study of Cicutti et al.\(^6\)\(^7\) In the model, total decarburization rate in the impact zone was obtained by the summation of two different decarburization rates via O\(_2\) and CO\(_2\) above the critical carbon content.

Figure 8 compares the decarburization rate via O\(_2\) and CO\(_2\) in the impact zone during the blow predicted by the model and estimated from the study of Cicutti et al.\(^6\)\(^7\) As seen from the figure, the reaction rate of carbon was divided into two distinct regions according to the critical carbon content of the liquid bath. The critical carbon content was obtained using Eq. (22). However, the values obtained from the numerical calculations were higher than the real case. (above 1 mass%) This is most likely due to the low mass transfer coefficients. In this calculation, an empirical relationship was used to calculate the mass transfer coefficient. Further study is required for better understanding of mixing in the liquid bath under the industrial conditions. Therefore, the authors used a fixed value of 0.5 for the values of critical carbon content for simplicity. In region 1 (above the critical carbon content), the rate of carbon oxidation is independent of carbon concentration but subjected to the fluid flow and partial pressure of gasses. The decarburization reaction via oxygen is controlled by gas diffusion and has the major role on the overall kinetics of the reaction at the impact zone. In the case of CO\(_2\), both chemical reaction and gas diffusion limit the reaction kinetics.

In region 2, the decarburization rate decreases rapidly below the critical carbon content of the liquid iron (mass% C<1). The reaction rate is assumed to be controlled by the carbon diffusion in the liquid metal using Eq. (6). The reaction rate is strong function of mass transfer coefficient and carbon concentration of liquid bath. As the carbon concentration decreases, the reaction rate decreases simultaneously.

The estimated decarburization rates from the study of Cicutti et al. were represented by the points and the model results were represented by the line in Fig. 8. Some differences were observed in the figure. Nevertheless, as discussed in Part 1 of this series of papers, the evolution of carbon content of liquid iron is in good agreement with the industrial data. This means that combining the two models provides a reasonable estimate of the overall decarburization rate and therefore the model for the impact zone provides a reasonable basis for predicting the decarburization rate in the impact region.

### 4. Conclusion

A kinetic model involving decarburization reactions with O\(_2\) and CO\(_2\) that provides a quantitative understanding on how different operational parameters affect the decarburization rates at the impact zone under full-scale operating conditions has been developed. The model simulations were applied to given top and bottom gas flow rates on full scale operating conditions. The results from the model indicate the followings:

- In region 1 (above critical C content), higher decarburization rates were predicted when O\(_2\) was used as oxidizing gas instead of CO\(_2\). Partial pressure of oxygen has a marked affect on the decarburization kinetics via O\(_2\).
- The predicted rate constants showed that sulphur has a retarding effect on the decarburization reaction via CO\(_2\).
- In region 2 (below critical C content), decarburization rates decreased as the carbon content is decreased towards the end of the blow. The increase in bottom stirring from 150 to 500 Nm\(^3\)/min increases the transfer rates significantly.

The impact zone model was validated against the estimated values derived from the study of Cicutti et al. since it is very difficult to measure the decarburization rates at the impact zone individually and distinguish the gas production (CO and CO\(_2\)) from the off-gas analysis. Based on the model predictions, it can be approximated that 40% of decarburization takes place in the impact zone during the main blow. Limited data exist for real systems, and further investigations are needed to refine these correlations and further establish their integrity and validity.

**REFERENCES**