1. Introduction

Reactions between liquid iron droplets and slag occur while liquid iron droplets are ejected from the metal bath into the slag phase in the smelting reduction of iron ore and oxygen steelmaking processes. Understanding the kinetic features of the FeO reduction or decarburisation related to the iron droplets, and the accompanying droplet swelling phenomenon are important for describing droplets’ behavior in slag. The swelling of an iron droplet in the slag will increase the reaction area and reduce the apparent density of the droplet to enhance its suspension in the slag. The droplet swelling in slag is therefore desired for high speed reduction or high speed decarburisation in the processes.

The metal droplet formation, droplet size distribution, and residence time of droplets in slag phase for steelmaking processes have been investigated in many studies.\(^1\)–\(^6\) The swelling of an iron droplet under oxidizing conditions was observed by Min and Fruehan,\(^7\) and Molloseau and Fruehan\(^8\) for an Fe–C droplet exposed to oxidizing slag. The phenomenon was also observed by Sun et al.\(^9\) and Gao et al.\(^10\) for an Fe–C droplet exposed to oxidizing gases. Kinetic studies on the iron droplet–slag reaction have been focused to carbon, silicon and manganese transfers between slag and the droplet.\(^7\)–\(^11\) In this work, the attempts were made to interpret the reaction kinetics and the swelling phenomena.

2. Interfacial Phenomena and Reaction Kinetics

2.1. CO Bubbles at the Interface

For a gas bubble presenting at the slag–metal interface as illustrated in Fig. 1(a), the contact angle between gas bubble and metal in slag media can be estimated by balancing the interfacial forces according to Young’s equation:

\[
\sigma_{gm} \cos \theta = \sigma_{sm} - \sigma_{gs}
\]

The contact angle was calculated to be \(~157^\circ\) using the interfacial tensions between slag and metal (~1200 mN/m),\(^12\) slag and gas (~380 mN/m),\(^13\) and metal and gas (~1550 mN/m).\(^14\) The high contact angle indicates poor “wetting” of gas bubble with metal in slag media. From X-ray fluoroscopy observations of an Fe–C in slag,\(^8\) the droplet appeared to be surrounded by a thin layer of slag less dense than the bulk slag during the reaction. This layer is considered to be numerous gas bubbles attached to the droplet as schematically presented in Fig. 1(b). These surface bubbles contact with the metal by limited area because of the poor “wetting”.

2.2. Elementary Steps

The mass transfer of elements and oxides in the vicinity of the droplet is a critical factor in the reaction kinetics. In this work, the attempts were made to interpret the reaction kinetics and the swelling phenomena.
of the interface between Fe–C droplet and the slag containing FeO are shown in Fig. 1(c). If there is an internal gas bubble presenting in the droplet, the mass transfers between the internal bubble and liquid iron are shown in Fig. 1(d).

The mass transfer and interfacial reaction processes at the slag–metal interface are summarized in Fig. 2. During the reaction, the migration of FeO from slag bulk to the slag side of the interface occurs by FeO diffusion in the slag phase, or Step 1. FeO arrived at the interface undergoes the reduction reaction (2) as represented by Step 2.

\[
\text{FeO} + \text{C} \rightarrow \text{CO} + \text{Fe} \tag{2}
\]

The oxygen produced from the reaction (2) reacts with carbon at the interface by the reaction (3), which is represented by Step 3.

\[
\text{C} + \text{O} \rightarrow \text{CO} \tag{3}
\]

Reaction (3) removes oxygen at the interface supplied from Step 2 and carbon supplied from Step 4, the latter is the carbon diffusion from droplet bulk to the interface. The removal of oxygen from the interface by reaction (3) keeps the interfacial oxygen at a lower level to sustain further oxygen supply from Steps 1 and 2. Reaction (3) produces CO gas bubbles at the interface, and these bubbles leave the interface by Step 5 to complete decarburisation process. If oxygen supplying rate by Step 2 is higher than the oxygen removal rate by Step 3, the extra oxygen will diffuse into the droplet by Step 6. As will be discussed later, the oxygen diffusion into the droplet bulk, or oxygen absorption by the droplet, may trigger the reaction (3) to occur within the droplet as seen in Fig. 1(d). This internal reaction produces CO bubbles within the liquid to swell the droplet.

If silicon presents in the droplet, desiliconisation occurs at the interface by reaction (4).

\[
\text{Si} + \text{O} \rightarrow \text{SiO}_2 \tag{4}
\]

This reaction, or Step 7, also removes oxygen supplied from Step 2, and silicon supplied from Step 8, the latter is the silicon diffusion from the droplet bulk to the metal side of the interface. Silica, the product of reaction (4), leaves the interface by the diffusion into slag bulk, which is represented by Step 9. If other elements such as manganese, titanium or phosphorus present in the droplet, their involvements in the kinetics can be treated in a similar manner as that for silicon.

2.3. Limiting Steps

These 9 steps proceed simultaneously during the reaction, those steps that process much faster than the others can be excluded from controlling steps. Diffusions of FeO and SiO₂ in slag, i.e., Steps 1 and 9, were found to be slow in many studies and they usually control or partly control the overall reaction rates. The reaction (2), i.e., FeO reduction at slag–metal interface, was found fast at high temperatures. The influence of the rate of reaction (2), or Step 2, on the overall reaction rates can be excluded and it is able to assume that the equilibrium of reaction (2) is maintained at the interface. The Step 3, i.e., decarburisation reaction (3) at slag–metal interface, has been found slow and is involved in controlling the overall decarburisation rate. The CO bubble detachment from the interface, Step 5, is considered as a part of reaction (3) and will be dealt together with Step 3 in the subsequent model development. The mass transfers of carbon, oxygen and silicon, i.e., Steps 4, 6 and 8, were found to be slow in slag–metal reactions and are involved in the kinetics of the reactions. The fast interfacial desiliconisation enables the equilibrium of the reaction (4) or Step 7 to be assumed at the slag–metal interface.

In summary, for an iron droplet containing carbon and silicon reacting with slag containing FeO, Steps 2 and 7 indicated by asterisk in Fig. 2 can be excluded from rate limiting steps while other steps may affect the overall reaction rates and should be included in the kinetic model.
3. Kinetic Model

3.1. Reaction between Droplet and Slag

3.1.1. Rate Equations

On the basis of the above discussions, a kinetic model was developed by including diffusion in the slag (Steps 1 and 9), diffusion in the metal (Steps 4, 6 and 8), the reaction (3) at the interface (Steps 3 and 5), and the equilibria of reactions (2) and (4) at the interface (Steps 2 and 7).

The composition of the slag bulk is uniform, and a composition profile exists only at the vicinity of the slag side of the interface where FeO is removed by Step 2 and SiO2 is produced by Step 7. This enables the boundary layer theory to be applied for describing the mass transfers of FeO and SiO2 in the slag as,

\[ J_{FeO} = \frac{A_k p_{FeO}}{100 M_{FeO}} \{ (FeO)^n - (FeO) \} \] 
\[ J_{SiO_2} = \frac{A_k p_{SiO}}{100 M_{SiO}} \{ (SiO_2)^n - (SiO_2) \} \]

where the superscript \( r_0 \) denotes the position at \( r_0 \) of the droplet and \( (\cdot)^n \) means the concentrations at the slag side of the interface between slag and metal, and the materials fluxes are in the direction of the droplet radius.

Within the vicinity of the metal side of the interface, mass transfer rates of carbon, oxygen and silicon in the direction of the droplet radius are given by Fick's first law as,

\[ J_C = \frac{AD_C p_m}{100 M_C dr} \{ [C]^{n-dr} - [C]^n \} \] 
\[ J_O = \frac{AD_O p_m}{100 M_O dr} \{ [O]^{n-dr} - [O]^n \} \] 
\[ J_{Si} = \frac{AD_{Si} p_m}{100 M_{Si} dr} \{ [Si]^{n-dr} - [Si]^n \} \]

where the superscript \( r_0 \) denotes the position at \( r_0 \) of the droplet and \( [\cdot]^{n-dr} \) means the concentrations at the metal side of the interface between slag and metal, and the reactions at a distance of \( dr \) from the interface. As will be presented later, the concentrations at \( r_0 - dr \) is calculated from the concentration profiles in the droplet by Eqs. (5)–(7). It should be noted that the Fick's first law is applicable only when \( dr \ll r_0 \).

The driving force for the reaction (3) at the interface is the deviation from the equilibrium state, and the rate of the reaction is given by,

\[ J_t = \frac{A_k}{100 M_C} \left\{ a_0^n a_C^n \frac{P_C^n}{K_C} \right\} \] 

\[ \frac{-J_t + J_C = 0}{-J_{FeO} + J_O - 2J_{SiO_2} - J_{C} = 0} \] 
\[ J_{Si} = -J_{SiO_2} = 0 \]

3.1.3. Equilibrium Relations

The equilibrium constants, \( K_{Fe}, K_{C}, \) and \( K_{Si} \) for the reactions (2), (3) and (4), respectively, are given by,

\[ K_{Fe} = \left( \frac{a_{FeO}}{a_O a_{Fe}^{1/2}} \right)_{FeO} \] 
\[ K_{C} = \left( \frac{P_O}{a_C a_O} \right)_{FeO} \] 
\[ K_{Si} = \left( \frac{a_{SiO}}{a_{Si}} \right)_{SiO} \]

Note that the reactions (2) and (4) reach equilibria at the interface as discussed in Sec. 2.3., and their equilibrium constants may be represented by the interfaceal values as seen in relations (14) and (16).

3.1.4. Activities

In slag, the activities of FeO and SiO2 at the interface are determined from Raoult's law with the activity coefficients,

\[ a_{FeO}^{FeO} = \gamma_{FeO}^{FeO} \] 
\[ a_{SiO_2}^{SiO_2} = \gamma_{SiO_2}^{SiO_2} \]

Where the molar fractions of FeO and SiO2 at the slag side of the interface are calculated for CaO–SiO2–MgO–FeO slag of a composition in weight percent by,

\[ X_{FeO}^s = \frac{(FeO)^s}{M_{FeO}} \sum \] 
\[ X_{SiO_2}^s = \frac{(SiO_2)^s}{M_{SiO_2}} \sum \]

Where

\[ \sum = \frac{(FeO)^s}{M_{FeO}} + \frac{(SiO_2)^s}{M_{SiO_2}} + \frac{(CaO)^s}{M_{CaO}} + \frac{(MgO)^s}{M_{MgO}} \]

In metal, the activities of carbon, oxygen or silicon at the metal side of the interface are determined by activity coefficients and composition at the metal side of the interface according to Henrian law,

\[ a_{FeO}^{FeO} = f_C^{FeO} [C]^n \] 
\[ a_{SiO_2}^{SiO_2} = f_O^{SiO_2} [O]^n \] 
\[ a_{SiO_2}^{SiO_2} = f_{SiO_2}^{SiO_2} [Si]^n \]

The activity coefficients are obtained by interaction parameters and metal composition at the interface,

\[ \log f_C^{FeO} = c_1^C [C]^n + \sum c_i^{FeO} [O]^n + e_i^{FeO} [S]^m + e_i^{FeO} [Si]^n \]
\[ \log f_O^{SiO_2} = c_1^O [O]^n + \sum e_i^{SiO_2} [S]^m + e_i^{SiO_2} [Si]^n \]
\[ \log f_{SiO_2}^{SiO_2} = e_i^{SiO_2} [C]^n + e_i^{SiO_2} [O]^n + e_i^{SiO_2} [S]^m + e_i^{SiO_2} [Si]^n \]
3.1.5. Determination of Interfacial Quantities

CO pressure at slag–metal interface is assumed to be equal to the ambient pressure. The activity of iron at surface is assumed to be unit for an iron based droplet. Since CaO, MgO and sulphur are not directly involved in the reaction when slag contains relatively higher FeO, their concentrations at the interface are approximately equal to their bulk counterparts. We therefore have $P_{\text{CO}}^*=P^*$, $a_{\text{Fe}}^*=1$, $(\text{CaO})^*=(\text{CaO})$, $(\text{MgO})^*=(\text{MgO})$ and $[S]^*=[S]$.

There are 15 interfacial values are needed to be determined for the reaction rate calculation. These values are, $a_{\text{FeO}}$, $a_{\text{MgO}}$, $X_{\text{EO}}$, $X_{\text{SiO}}$, $(\text{FeO})^*$, $(\text{SiO}_2)^*$, $a_{\text{Fe}}^*$, $a_{\text{Si}}^*$, $[\text{C}]^*$, $[\text{O}]^*$, $[\text{Si}]^*$, $f_{\text{C}}$, $f_{\text{O}}$, and $f_{\text{Si}}$. These unknown values can be solved by 15 relations. These relations are three stoichiometrical relations (11)–(13), two equilibrium relations (14) and (16), and ten activity relations (17)–(26). With the solution of these interfacial values, the rate of each element transport across slag–metal interface can be determined from Eqs. (5)–(10).

3.1.6. Mass Transfers in the Droplet

The concentration profiles of drop, oxygen and silicon in the droplet are obtained by applying Fick’s second diffusion law to a sphere as,

$$
\frac{\partial \text{[C]}}{\partial t} = D_{\text{C}} \left( 2 \frac{\partial \text{[C]}}{\partial r} + \frac{\partial^2 \text{[C]}}{\partial r^2} \right) \quad \text{(27)}
$$

$$
\frac{\partial \text{[O]}}{\partial t} = D_{\text{O}} \left( 2 \frac{\partial \text{[O]}}{\partial r} + \frac{\partial^2 \text{[O]}}{\partial r^2} \right) \quad \text{(28)}
$$

$$
\frac{\partial \text{[Si]}}{\partial t} = D_{\text{Si}} \left( 2 \frac{\partial \text{[Si]}}{\partial r} + \frac{\partial^2 \text{[Si]}}{\partial r^2} \right) \quad \text{(29)}
$$

Using the concentrations of carbon, oxygen and silicon obtained at the interface in the previous section, $[\text{C}]^*$, $[\text{O}]^*$ and $[\text{Si}]^*$, as the boundary conditions, and assuming that composition of the droplet is uniform before the reaction, the concentrations of carbon, oxygen and silicon in the droplet as a function of the distance from the droplet surface and reaction time can be determined by Eqs. (27)–(29). The concentrations at $r_0-\Delta r$ obtained from these calculations were used in Eqs. (7)–(9) for the determination of diffusion rates in Steps 4, 6 and 8.

3.1.7. Thermodynamic and Kinetic Parameters

To determine the reaction rates, relevant thermodynamic and kinetic parameters must be provided. The equilibrium constants and interaction parameters were used to calculate $K_{\text{FeO}}$, $K_{\text{C}}$ and $K_{\text{Si}}$, and the activity coefficients for metal components. The activities of FeO, SiO$_2$ in CaO–SiO$_2$–MgO–FeO slag were used to estimate the activity coefficients in the slag. The mass transfer coefficient in slag was selected for $k_t$ and $k_a$, respectively. Diffusivities of carbon, oxygen, and silicon in molten iron as a function of the temperature were used for $D_{\text{C}}$, $D_{\text{O}}$ and $D_{\text{Si}}$, respectively.

3.2. Stable CO Bubble in Metal

Once carbon and oxygen in the droplet become higher than the equilibrium as presented in relation (15), reaction (3) may occur in the droplet. When an instantaneous CO cluster is larger than a critical bubble size, the cluster will spontaneously grow into a bubble within the droplet. The critical bubble radius, $r_g$, is estimated from a stable gas bubble in a liquid. For a stable CO bubble in liquid iron, the sum of the pressure due to the surface tension and the ambient pressure must be balanced by CO pressure through,

$$
P_{\text{CO}}=2\sigma_{\text{gm}}/r_g+P^* \quad \text{(30)}$$

where $P_{\text{CO}}$ is the CO pressure in equilibrium with carbon and oxygen in the liquid iron.

Under a given ambient pressure, $P^*$, the critical bubble radius is determined by $P_{\text{CO}}$ and $\sigma_{\text{gm}}$. $P_{\text{CO}}$ and $\sigma_{\text{gm}}$ are in turn determined by the metal composition and temperature according to Eq. (15) and the surface tensions for Fe–O–S melts, respectively.

$r_g$ calculated from Eq. (30) is the minimum stable CO bubble size in liquid. The nucleation of an internal CO bubble or the occurrence of the internal reaction is likely to occur when $r_g$ is small because the higher possibility of CO clusters having a size over a smaller $r_g$. Any factors that reduce $r_g$ would increase the possibility of CO nucleation, and therefore the swelling of the droplet.

If there are inclusions in the droplet, and these inclusions have a good wetting with gas than with slag, the inclusions may provide sites for heterogeneous nucleation of CO bubble. The heterogeneous nucleation is much easier than that of homogeneous since for the same critical bubble radius, critical bubble volume for heterogeneous nucleation is smaller than that of homogeneous nucleation. Hence, the present of such inclusions will increase the possibility of CO nucleation, and therefore, promote the swelling of the droplet.

4. Results and Discussions

4.1. Reaction Rates

The model calculation was carried out for an 1 gram iron droplet containing 2.91 mass% carbon, 0.002 mass% oxygen and 0.011 mass% sulphur reacting with CaO–MgO–SiO$_2$–FeO slag of CaO/SiO$_2$ ratio of 1.2, 12 mass% MgO and 3–20 mass% FeO at 1 713 K. Except the oxygen content in the droplet was estimated from the equilibrium of reaction (3), these conditions are the same conditions as those used in the experimental observations. The calculated decarburisation rates during initial 60 s are compared with observed rates in Table 1. The calculations agree with the observed ones when FeO is less than 5 mass%. Beyond 10 mass% FeO, the calculations are much slower than the observed. This is because the droplet swelled for slag containing more than 10 mass% FeO and the reaction model did not include the increased surface area, the internal decarburisation, and enhanced mass transfer rate in a swollen droplet.

The variations of CO generation, FeO reduction and oxygen absorption rates during the reaction with slag of 5 mass% FeO are shown in Fig. 3(a). The reaction rates de-
creases with reaction time as reactions approaching the equilibria. Surprisingly, less than one half of oxygen brought by the reduction of FeO to the interface is removed by decarburisation, and the rest is absorbed by the droplet. Carbon content, silicon content, CO pressure, and critical bubble size are largely distributed quantities in the droplet. Their average values may be obtained from the integration across the droplet radius. For example, the average content of carbon in the droplet is obtained by,

\[ [C]_{av} = \int_{0}^{r} \frac{2r^2}{r_0^3} [C] dr \]  

As shown in Figs. 3(b) and 3(c), average carbon decreases and average oxygen increases with increasing reaction time; the average CO pressure increases and average critical bubble radius decreases. The droplet may not swell immediate after contacting the slag; a time period may be required for building up an over-saturation level for the CO bubble nucleation within the droplet. This corresponds qualitatively to the observations \(^8\) that there were a time period required for the occurrence of droplet emulsion, and the period length depends on the conditions.

**Table 1. CO generation rates during 60 s.**

| FeO, mass% | 3 | 5  | 10 | 15  | 20 |
|-----------|---|----|----|-----|----|
| Droplet swelling \(^8\) | no | no | yes | yes | yes |
| \( J_{CO} \times 10^6 \), mol/s, observed \(^8\) | 3.3–3.3 | 3.0–3.0 | 30–1 | 180–1 | 200–1.1 |
| \( J_{CO} \times 10^6 \), mol/s, calculated\(^*\) | 1.73–1.70 | 3.01–2.89 | 6.13–5.55 | 9.17–7.71 | 12.2–9.40 |

\(^*\)calculated \( J_{CO} \) during stagnant decarburisation period were not included.

**Fig. 3.** Rates of CO generation, oxygen absorption and FeO reduction (a), average concentrations of carbon (b) and oxygen (c), average CO pressure (d), and average critical bubble radius (e) during the reaction of a 1 g droplet (2.91 mass% carbon) reacting with slag (5 mass% FeO) at 1 713 K.

**Fig. 4.** Profiles of carbon concentration (a), oxygen concentration (b), CO pressure (c), surface tension (d) and critical bubble radius (e) at 1, 10, 50 and 200 s into reaction between a 1 g droplet (2.91 mass% carbon) and slag (5 mass% FeO) at 1 713 K.

### 4.2. Profiles in the Droplet

Figures 4(a)–4(e) show the concentrations of carbon and oxygen, CO pressure in equilibrium with the local metal composition, surface tension and critical bubble size in a Fe–C droplet as functions of radius in the droplet and reaction times (1, 10, 50 and 200 s) with the slag containing 5mass% FeO at 1 713 K. After 1 s of the reaction, carbon decreases in the vicinity of the droplet surface while remains unchanged at the droplet center. As shown by the curves for 10, 50 and 200 s of the reaction time, carbon concentration in the droplet decreases. Oxygen at the interface is kept high and diffuses into the droplet to create a profile in an opposite direction to that of carbon. The CO pressure in the droplet increases with the reaction time. When CO pressure in the droplet becomes greater than the ambient pressure (1 atm), the metal is said to be supersaturated, and the internal reaction (3) becomes possible. The surface tension decreases with time due to the increased oxygen. Critical bubble size near the surface is lower than those at the cen-
ter, and decreases with time. This indicates that CO bubble nucleation is easier at the regions close to the droplet surface or after longer reaction time. This corresponds well with X-ray observations\(^7\) of a swollen droplet in slag where the droplet center was found denser than the circumferential region of the droplet.

### 4.3. Influence of Variables on the Reaction Rates and the Droplet Swelling

Several reaction cases were modeled within 60 s reaction using conditions given in Table 2. The reference case is an 1 gram droplet containing 2.9 mass\% carbon, 0.002 mass\% oxygen and 0.01 mass\% sulphur reacting with a slag of 5 mass\% FeO at 1713 K. The influence of a variable was examined by varying the variable as shown in Table 2 while keeping other variables same as the reference case. The influence of FeO in slag, carbon and silicon contents in the droplet, droplet size, and ambient pressure were investigated. The average decarburisation rate during 60 s of reaction \(J_C\), reaction time required for removing 1\% of total carbon in the droplet \(t_{1\%}\), reaction time required for reaching max decarburisation rate \(t_{\text{max}}\), and average critical bubble size \(r_{B\text{av}}\) after 10 s reaction were investigated and will be discussed in the following sections.

#### 4.3.1. Decarburisation Rate

The decarburisation rate, \(J_C\), obtained with varying variables is compared with the reference case in Table 2. Decarburisation rate increases with increasing FeO in slag, carbon in the droplet and the size of droplet, but the rate decreases with increasing silicon in the droplet and ambient pressure. It is noted that the decarburisation rate in unit surface area of the droplet also increases with increasing the droplet size. On the qualitative base, these tendencies are in agreement with observations on the influences of FeO in slag, carbon and silicon contents in the droplet, droplet size, and ambient pressure were investigated. The average decarburisation rate during 60 s of reaction \(J_C\), reaction time required for removing 1\% of total carbon in the droplet \(t_{1\%}\), reaction time required for reaching max decarburisation rate \(t_{\text{max}}\), and average critical bubble size \(r_{B\text{av}}\) after 10 s reaction were investigated and will be discussed in the following sections.

#### 4.3.2. Retardation Period

A slow decarburisation rate at initial several seconds of slag–metal reaction was observed.\(^7,8,34\) This period is referred as the retardation periods. Figure 5 shows the reaction or mass transfer rates versus logarithm of reaction time for reference case (Ref), and cases of high carbon (4.5 mass\%), high silicon (0.5 mass\%) and low FeO (3 mass\%). The decarburisation starts slowly and progressively increases in the first several seconds for all cases as shown in Fig. 5(a). This is because the Steps 6 or 7 removes oxygen from the interface and less oxygen can be provided for decarburisation in this period. Since initial oxygen is lower in the metal, the oxygen diffusion (Step 6) is initially faster as shown in Fig. 5(b), which keeps interfacial oxygen at a low level to retard decarburisation. When silicon is present in the droplet, the desiliconisation keeps oxygen even lower. The decarburisation will not increase markedly until the desiliconisation becomes relaxed. As seen from curves of [Si]=0.5 mass\% in Figs. 5(a) and 5(b), both decarburisation and oxygen absorption rates increase markedly only after vigorous desiliconisation as shown in Fig. 5(d). A retardation period of decarburisation may be quantified by using the reaction time required for removing 1\% of total carbon in the droplet. The retardation periods, \(t_{1\%}\), for all cases are listed in Table 2 as well as shown by triangle marks in Fig. 5(a). Comparing with the reference case (Ref), the conditions that favor a longer retardation period of decarburisation was found to be higher carbon and higher silicon in the droplet, higher pressure, and lower FeO in the slag.

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**Table 2.** Average decarburisation rate during 60 s of reaction \(J_C\), reaction times required for 1\% decarburization \(t_{1\%}\) and reaching max decarburisation rate \(t_{\text{max}}\), and average critical bubble size \(r_{B\text{av}}\) in the droplet after 10 s reaction in slag.

| Rate, time and \(r_{B\text{av}}\) | Reference* | FeO=3 mass\% | C=4.5 mass\% | Si=0.5 mass\% | \(H_2\)=0.5 g | \(P=5\) atm |
|------------------------------|-------------|---------------|---------------|---------------|---------------|-------------|
| \(J_C\), mol/s × 10\(^{10}\)  | 2.95        | 1.72          | 3.42          | 2.12          | 1.73          | 2.11        |
| \(t_{1\%}\), s            | 8           | 14            | 11            | 18            | 7             | 10          |
| \(t_{\text{max}}\), s     | 5           | 7             | 17            | 105           | 5             | 6           |
| \(r_{B\text{av}}\), cm × 10\(^{10}\) | 1.31        | 2.38          | 1.16          | 1.94          | 1.08          | ∞           |

* Reference case: 1 gram droplet containing 2.9 mass\% carbon, 0.002 mass\% oxygen and 0.01 mass\% sulfur, slag of CaO:SiO\(_2\)=1.2 and containing 12 mass\% MgO and 5 mass\% FeO at 1713K.

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**Fig. 5.** Changes of CO generation (a), oxygen absorption (b), FeO reduction or diffusion (c, d), and silicon transfer (d) rates with time. The conditions for reference case (Ref) are a 1 g droplet (2.91 mass\% carbon, 0 mass\% silicon) reacting with slag (5 mass\% FeO) at 1713 K. Triangles and circles indicate reaction times required for 1\% decarburisation \(t_{1\%}\) and for reaching max decarburisation rate \(t_{\text{max}}\) or the maximum rates, respectively.
As reaction time further increases, the depletion of carbon in the droplet will bring the decarburisation rate down until the equilibrium of the reaction is achieved. There exists a maximum decarburisation rate during reaction. The maximum decarburisation rates are marked by the circles in Fig. 5(a). When silicon presents in the droplet, a maximum oxygen absorption rate as marked by a circle in Fig. 5(b) occurs for the similar reason. The time periods required for decarburisation reaching the maximum rate are also listed in Table 2. When the droplet contains higher carbon and silicon, slag contains lower FeO, longer time period is required for the decarburisation to reach the maximum rate.

4.3.3. FeO Reduction Rate

The reduction rate of FeO, which is equal to FeO diffusion rate, as presented in Figs. 5(c) and 5(d), increases with carbon and silicon contents in the droplet because increased oxygen removal rate by decarburisation and desiliconisation. The reduction rate of FeO increases with increasing FeO in slag because the driving force for the FeO diffusion is increased.

4.3.4. Influence of Variables on Droplet Swelling

Average critical bubble size \(r_{g,av}\) after 10 s reaction are given in Table 2. By comparing \(r_{g,av}\) of the reference case with those of other cases, the conditions that promote swelling of the droplet can be identified because of high probability of the bubble nucleation in the liquid if \(r_{g,av}\) is small. The swelling of the droplet would be promoted when the droplet contains higher carbon or lower silicon, slag contains higher FeO, the droplet is smaller, or ambient pressure is lower.

Figure 6 shows the effect of FeO on the profiles in the droplet after 10 s into reaction. Carbon decreases and oxygen increases with increasing FeO in slag. The increase of CO pressure with increasing FeO is the net result of the decrease of carbon, which decreases \(P_{CO}\), and the increase of oxygen, which increases \(P_{CO}\). The surface tension is decreased by the increased oxygen. The critical bubble radius decreases as FeO increases which mean the nucleation of gas bubbles in the droplet becomes easier. This prediction corresponds well with the observation that the swelling of the droplet is favored for high FeO in the slag.

The effect of initial carbon was examined by comparing profiles for the droplet of 2.9 mass\% carbon (solid lines) with those of 4.5 mass\% carbon (dash-dot lines) at 10 s of the reaction in Fig. 7. There is no large difference for oxygen between two cases. CO pressure increases and surface tension decreases with increasing initial carbon in the droplet. The critical bubble radius is smaller for high carbon droplet, and therefore, the bubble nucleation or the
droplet swelling would be easier for higher carbon metal.

The droplet contains 0.5 mass% silicon (dotted lines) was compared with that without silicon (solid lines) in Fig. 7. Due to the suppressions of both decarburisation and oxygen absorption by desiliconisation, oxygen and \( P_{\text{CO}} \) are lower for the droplet containing silicon. There is no significant change in the surface tension by adding silicon to the metal. The larger critical bubble radius for the silicon bearing droplet indicates that silicon will suppress droplet swelling.

The profiles in the droplet obtained for 5 atm (dashed lines) were compared with those at 1 atm (solid lines) in Fig. 8. Carbon is higher under 5 atm than that under 1 atm because the decarburisation is suppressed by the increased pressure. The changes in profiles of oxygen, \( P_{\text{CO}} \) and surface tension are insignificant. Critical bubble radius increases with increasing ambient pressure, and becomes infinity when ambient pressure is greater than \( P_{\text{CO}} \). Therefore, the nucleation of CO bubble within the liquid would be difficult under higher pressure.

Profiles for 0.5 g droplet at 10 (dash-dot lines) and 50 (solid lines) s were compared with those for 1 g droplet (dashed lines for 10 s and dotted lines for 50 s) in Fig. 9 as well as in Fig. 10 with a normalised radius. The profiles of two droplets are similar at 10 s. When the reaction time increases to 50 s, lower carbon, higher oxygen, higher \( P_{\text{CO}} \), lower surface tension and lower critical bubble radius profiles were obtained for smaller droplet than those for larger droplet. This indicates the nucleation of CO bubble within the liquid would be easier for a smaller droplet. As mentioned previously in Sec. 4.3.1, the decarburisation rate does not follow the expectation from proportionality between the rate and surface area. This is because the mass transfers occur throughout the droplet body which does not solely depend on the surface area. A contrast example was the observation by Sun and Pehlke35,36) for an internal stirred droplet exposed to oxidizing gases, where the decarburisation was limited by the processes in the vicinity of the droplet surface, and the rate was found to be simply in proportion to the surface area of the droplet.

5. Conclusions

The reaction of a Fe–C droplet with FeO bearing slag was analyzed with aid of a kinetic model that includes thermodynamic relations, diffusions in the slag and the droplet, and the interfacial reactions. The droplet swelling was interpreted by incorporating the kinetics of CO bubble nucleation within metal phase in the kinetic model. Decarburisation rate increases with increasing (FeO), [C] and droplet size, and decreases with increasing [Si] and pressure. The decarburisation rate in unit surface area of the droplet also increases with increasing the droplet size. Initial slow de-
Carburisation occurs due to oxygen absorption and desiliconisation, and this stagnancy is more significant when contents of carbon and silicon in metal, ambient pressure and droplet size are increased or when (FeO) is decreased. The swelling of the droplet occurs due to the CO bubble generation within the droplet. The conditions that promote swelling of the droplet during the reaction are higher carbon or lower silicon in the droplet, the smaller droplet size, lower ambient pressure and higher FeO in slag.

Nomenclature

- $A$: Surface area of the droplet (cm$^2$)
- $a_i$: Activities of carbon, oxygen or silicon, in metal or activities of FeO or SiO$_2$ in slag
- $D_i$: Diffusivity of carbon, oxygen and silicon in molten iron (cm$^2$ s$^{-1}$)
- $e_i'$: Interaction parameter between carbon, oxygen, sulphur and silicon in metal
- $f_i$: Activity coefficient of carbon, oxygen and silicon in metal
- $K_i$: Equilibrium constant
- $k_s$: Mass transfer coefficient in slag in gas (cm s$^{-1}$)
- $k_c$: Decarburisation rate constant at surface (mol cm$^{-2}$ s$^{-1}$)
- $M_i$: Molecular weight of elements and oxide (g mol$^{-1}$)
- [i]: Content of carbon, oxygen or silicon in metal (mass%)
- (i): Content of oxides in slag (mass%)
- $J_i$: Materials fluxes in the radius direction of the droplet (mol s$^{-1}$)
- $P_{CO}$: CO pressure (Pa)
- $P^o$: Ambient pressure (Pa)
- $r$ and $r_0$: Radius in droplet and that of the droplet (cm)
- $r_g$: Critical nucleation radius of gas bubble in liquid (cm)
- $r_{g,av}$: Average critical nucleation radius of gas bubble in liquid (cm)
- $T$: Temperature (K)
- $t$: Reaction time (s)
- $X_i$: Molar fraction of FeO or SiO$_2$ in slag
- $Y_i$: Activity coefficient of FeO or SiO$_2$ in slag
- $p_s$: Density of slag (g cm$^{-3}$)
- $r_m$: Density of metal (g cm$^{-3}$)
- $s$: Interfacial tension (mN m$^{-1}$)
- $q$: Contact angle
- $\theta$: Contact angle

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