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Analysis of the Transition State of the Carbon and Iron Oxide Mixture Activated by Mechanical Milling

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In previous study, the extremely high reaction rate obtained by the mechanical milling has been reported. It was found that the phenomena were quite complicated and the many factors were affected on the reactivity. Among these factors, the crystal state of hematite and atmosphere (air and argon) during the milling were quite important ones.

Milling of hematite and graphite mixture was carried out both in the air and in the argon. Rate constants for the reactions were obtained by kinetic analysis. The enthalpy $\Delta H^*$ and entropy $\Delta S^*$ of activation were estimated on the basis of transition state theory. The interpretation of $\Delta S^*$ and $\Delta H^*$ was performed to explain the effect of milling on the reactivity related to the milling conditions.

The temperature of peak of reaction curve decreased with increasing milling time. The reactivity of the sample milled in the argon atmosphere showed higher reactivity than that milled in the air. The enthalpy $\Delta H^*$ and entropy $\Delta S^*$ of activation were obtained from experimental rate constants. The variation of $\Delta H^*$ and $\Delta S^*$ with milling time showed opposite tendency between the sample milled in the air and in the argon. It was considered that the active complex having loose bonding would form during milling in the air atmosphere. On the other hand, the active complex having a structure closed to the product (metallic iron) might form.

KEY WORDS: mechanical milling; active complex; entropy of activation; enthalpy of activation; transition state; iron oxide; graphite.

1. Introduction

In Japan, about 100 million tons iron are produced every year. The energy for the production of iron is significantly large and about 12% to the total energy consumption in Japan. Most of pig iron is made by Blast Furnace in Japan. In the view points of the productivity and the energy efficiency for the reactor of iron production, blast furnace has highest ones. However, such a blast furnace has further possibility for the increase of the efficiency in accordance with the high reaction rate and the low reaction temperature, because the theoretical utilization of CO gas and input energy has some allowance for the improvement.

In the Project of ‘the innovational ironmaking reaction in new blast furnace at half energy consumption and minimum environmental influences’ (from 1999 to 2004, Chaired by Prof. Ishii, Hokkaido University), many researchers in the ironmaking field have performed their investigations, in which they were classified into 4 groups: (1) Study on the high rate reduction and gasification, (2) Study on the optimization of the structure and composition of iron ore and coal, (3) Study on the lowering of the temperature of melting of metallic iron and slag, (4) Study on the establishment of process image and the quantification of influence of new ironmaking process).

In present study, the authors were belonging to the research group (1) on the high reduction rate and clarification of the mechanism of coupling reaction. We have conducted the study on the coupling phenomenon in previous paper.1)

Recently, it was reported that the iron ore–carbon composite pellet has a high reaction rate and a low reaction temperature. To make clarify the mechanism of the reaction in the iron ore–carbon composite, it could obtain a key to a breakthrough for the higher reaction rate and the lower reaction temperature.

From the results of previous study,2,3 in our laboratory, it was found that the complicated phenomena existed in the system depending on the location and the distance between iron oxide and graphite.

From the contact to the separate situation, there will be many states of the contact between carbon and iron oxide. We classify those three levels as follows.

I: A separate situation between iron oxide and carbon (Fig. 1)

II: A weak contact situation between iron oxide and carbon (Fig. 2)

III: A strong contact situation between iron oxide and carbon introduced by mechanical milling (Fig. 3)

• In the case of I, it is a separate situation between iron oxide and carbon. The direct reduction by the solid carbon never occur, apparent direct reduction, that is, the indirect reduction by CO gas and carbon gasification reaction will be dominant.

• In the case of II, it could be explained as a weak contact
situation, where not only the atomic level roughness but also the macroscopic level one always exist (Fig. 2), because the two reactants (iron oxide and carbon) are just mixed without any pressure. Even if the iron oxide and carbon are contacting at the beginning of reaction, the interface will soon separate, in which the coupling phenomenon (Case I) will be dominant.

It could be considered that the ultimate contact would be the distance that the both reactant atoms were closed to the position where the reaction easily occurred (activated complex). However, until this distance, the variety of distance would exist according to the preparing condition. Actually, the contact situation between iron oxide and carbon would be different from the way of mixing.

In addition to the microscopic feature, the macroscopic feature is important and will affect the apparent reaction rate strongly. Most important nature would be the separation of the both reactants during reaction, which means that the reaction mechanism would be changed in the course of reaction. From these considerations, the reaction rate would show the stable value and could be changed strongly in accordance with the condition of the contact. The mechanical milling is one of the method which can introduce the variety of contact situation into the carbon and iron oxide mixture from the various milling conditions, that is, many kind of activated complex can be formed.

- In the case of III, a strong contact situation between iron oxide and carbon introduced by mechanical milling is illustrated in Fig. 3. In previous study, 1) the milling of hematite and graphite mixture in long time (around 100 h) could give a quite high reaction rate and low reaction temperature. In such a situation, the nano-reactor was constructed in the sample. Especially, the structure of nano-reactor was important and took the situation that the nano-size hematite intruding into the graphite particle as shown in Fig. 3. By the nano-reactor having this structure, the reaction reduction could continue until the reaction completion. And contact mode between carbon and iron oxide is very strong and it could be explain as the activated state, which means the activated complex.

In this study, to clarify the mechanism of the reaction of iron oxide and carbon mixture made by the mechanical milling, the analysis of rate constants obtained in previous study was performed. 1) From the view point of the formation of activated complex introduced by mechanical milling, the enthalpy and entropy of activation were derived and the discussion was carried out on the structure of the activated complex. Furthermore, the changing of reaction mechanism was explained in accordance with the difference of milling atmosphere.

2. Experimental

The details of experiment were shown in previous paper. 1) Here, outline of the experiment is briefly shown. Several kind of hematite samples were prepare (details was shown in below) and high purity graphite with high density (1.77 g/cm³) and low ash content (<100 ppm) were used as starting materials. Reactivity of the milled sample was studied by TG-DTA (sample weight: 20 mg, resolution: 1 μg). The crucible was made of platinum and 4 mm in inner diameter and 3 mm in depth. The mixture of hematite and graphite with a fixed composition (19.84 mass% C in hematite, which corresponds to C/O = 1.1) was subjected to ball milling and the milling time was varied from 6 to 100 h. The milling experiment was performed using a planetary ball mill. A ceramics (Alumina) vessel of 500 cm³ volume, charged with 10 alumina balls of a nominal diameter of 20 mm, was used as the milling medium in all the experiments. The rotating speed of the vessel was maintained constant at 200 rpm for revolution. TG-DTA experiments were conducted under an Ar atmosphere with a constant flow rate of 50 Ncc/min. Samples were heating up with a constant heating rate of 10 K/min from room temperature up to 1373 K and kept at this temperature for 1800 s.

2.1. Difference of Sample as a Starting Material and Milling Condition

Since the difference of the particle morphology and crys-
tal structure of sample as a starting material affect strongly on the result of milling, several kinds of hematite was prepared. Even a reagent grade hematite (1st grade (>99%)) and special grade (>99.99%) have a different type of particle morphology as shown in Fig. 4. Moreover, the sample sintered at 1273 K, 24 h has shown the different grain size between 1st grade and special grade hematite (Figs. 4(c) and 4(f)). A crystalline size of 1st grade reagent sample sintered at 1273 K, 24 h was larger than that of special grade, and the range of crystalline size for 1st grade was from 1 to 5 μm and special grade hematite was from 1 to 2 μm. In the present experiment, only one kind of graphite was used for making clear the effect of hematite condition.

The atmosphere during milling was also very important factor affecting on the reactivity of the milled sample. In the case of air atmosphere, when the hematite crystal was broken during milling, the active new surface will react with oxygen quickly. If there was no oxygen in the atmosphere (argon atmosphere), the active surface will react with the active carbon (the milled carbon became also active), which meant the direct reduction between iron oxide and carbon (the detail mechanism was discussed below). Actually, we detected the CO gas evolution and the formation of FeO. The details will be published in the same journal. Therefore, the inert (argon) atmosphere and the air atmosphere were selected for the milling of different type of hematite.

3. Results and Discussions

3.1. Classification of Phenomena

As mentioned above, crystal conditions (reagent or sintered hematite) and milling atmosphere (argon and air) are very important factors. The experiments were carried out with samples prepared by those combinations. Figure 5 shows the results of the reactivity measurement experiment of the sample milled under the different conditions. Figure 5(a) shows the comparison of wustite reduction (W–Fe) curves (RTG means the rate of reaction which were obtained by the differential of TG curve (weight loss curve)), in which the reagent hematite (1st grade) was used without any treatment and the atmosphere during milling was air. The temperature at the peak of RTG curve decreased with the increasing milling time. Furthermore, the maximum value of the curve increased with the milling time.

On the other hand, when the hematite (1st grade) was sintered at 1273 K for 24 h (Fig. 5(b)), only 6 h milling could attain almost the same reaction rate (RTG was about 1.2%/s) as the 100 h milling (RTG=1.0%/s, Fig. 5(a)) for the reagent grade hematite under an air atmosphere. The peak temperature of the reaction curve decreased furthermore from 1277 K (6 h) to 983 K (100 h). From these results, it was found that the hematite sintered at 1273 K showed higher reactivity after milling. The sintered sample was also milled in air atmosphere (the results were not shown in present paper), the reactivity was higher than that of reagent grade hematite, but slightly lower than that of sintered hematite milled under argon.

Figure 6 shows the comparison of RTG curves of H–M, and M–W reduction curves between the reagent hematite in air and the sintered hematite in argon, which both the samples were milled in 100 h. It was quite interesting that the H–M and M–W reduction curves were disappear in the sintered hematite (Fig. 6(b)). The H–M reduction never showed the peak and the easy monotonous slope until 11.5% of reduction degree was observed in every sintered samples. This tendency of reduction curve means almost
constant value of the rate constant and is close to the zero order reaction. It might be resulted from a disordered hematite structure by the milling. It would suggest that the sintered hematite was easier to break than the reagent hematite.

As it was considered that the crystalline size after milling affected strongly on the reactivity of sample, X-ray diffraction (XRD) was carried out and crystalline size was estimated using Scherrer's equation. Here, we should note that the particle size and the crystalline size were not always equal. If the original crystalline size was large and equal to particle size, the decreasing mode between the particle and the crystal would be same. However, as shown in Figs. 4(b) and 4(c), the particle was constructed with smaller crystal (polycrystal), the way of decreasing would be different from the kind of reagent grade hematite. On the other hand, the sintered hematite has larger particle size from 1 to 2 \( \mu m \) for special grade hematite and 1 to 5 \( \mu m \) for 1st grade hematite. Since the crystalline size obtained by XRD is different from the particle size, it should be taken into account the difference of particle crystallinity.

Figure 7 shows the variations of crystalline size of sintered hematite according to the milling condition. Figure 8 shows the comparison of crystalline size of sintered hematite in mixture under argon atmosphere with the one under air atmosphere. In the case of argon atmosphere, the crystalline size decreased with the increasing milling time. From the result of gas analysis, it was found that the CO gas evolution occurred during the milling under the argon atmosphere, which meant the reduction reaction occurred during milling. This result also suggested that the re-
oxidation of hematite during milling under an air atmosphere might occur, simultaneously. Furthermore, this phenomenon (reduction–reoxidation) would related to the non-changed crystalline size under the air. These complicated phenomena during milling were summarized in Fig. 9. It was important to recognize the necessary condition that the milling of hematite and carbon should carry out in the mixture. When the milling was carried out separately and mixed together after milling, the reactivity was very low,1) and did not reach the completion of reaction. So that it was considered that the meaning of milling in mixture would be the utilization of the active surface generated at the instance of broken.5) If there was no oxygen gas in the atmosphere, the utilization of the active surface generated at the instance of broken would be small, the amount of the active complex would become small content of the complex would proceed forward.

The reaction rate constant expressed by the Arrhenius equation, for the milling in air and for the milling in argon, respectively. Subscripts ‘H’, ‘M’ and ‘W’ mean the HM, MW and WF reductions, respectively. S (-) means the specific contact area between oxide and carbon (S=Sp/Sp, Sp; contacting area of the particles (cm²), S0; total surface area of non-milling sample (cm²)) and k (1/s) is the reaction rate constant expressed by the Arrhenius equation, Eq. (5).

\[
k_f = A_f \exp \left(\frac{-E_f}{RT}\right), \quad (Y = H, M \text{ and } W) \quad (5)
\]

Where \(A_f\) (1/s) means a frequency factor, \(E_f\) (J/mol) means an activation energy, \(R\) (J/mol K) is the gas constant and \(T\) (K) is the absolute temperature. At first, the exact values of contact area \(S\) between the oxide and the carbon was treated as an unknown value, \(k\) (1/s) was introduced and Eq. (6) was used in the previous analysis instead of Eq. (5).\(^{1)}\)

\[
k'_f = S_f A_f \exp \left(\frac{-E_f}{RT}\right) = A'_f \exp \left(\frac{-E_f}{RT}\right), \quad (Y = H, M \text{ and } W) \quad (6)
\]

The parameter fitting to determine \(A'_f\) and \(E_f\) was carried out and RTG curves were recalculated.\(^{1)}\) The obtained rate constants for each experiments were summarized in Table I(a).

The surface area of the hematite particle that was increasing with the milling time was estimated in present study. The relationships among the milling time, the surface area (cm²) and average particle size (µm) were shown in Fig. 10, which were calculated by the following data: Density of hematite: \(\rho_H = 5.0 \text{ g/cm}^3\), FeO + C = 20 mg, FeO = 16.032 mg, Total volume of hematite \(V_y = 3.2064 \times 10^{-3} \text{ cm}^3\). The results were summarized in Tables 2 and 3, for the milling in air and for the milling in argon, respectively. The average particle size was measured based on SEM observation and the oxidation in low temperature at 550°C for separation of hematite particle from carbon mixture was adopted. According to milling, the average of par-
The particle size in argon atmosphere decreased rapidly and smaller than that of milling in air. Although the difference at 100 h was not so large, the particle size from 24 to 48 h showed a large difference between the milling in the air and in the argon. Corresponding to the smaller particle size, the total surface area of the sample milled in argon atmosphere was larger. Since the starting size of particle was the same, the magnification of total surface area in the argon milling at 100 h was largest ($S_e/S_o = 2.508$, at 100 h).

### 3.2.2. Kinetic Analysis on the Sample Milled in Argon Atmosphere: Two Steps Reduction

Based on Fig. 6(b), the reaction curves could be classified into two periods, kinetic analysis have performed with the following equations. The hematite reduction HM and the magnetite to metallic iron reduction, M–Fe would proceed as consecutive reaction, however, as the point of the transition was 11.5%, both reaction could be treated as the single reaction.

$$\frac{dM}{dt} = k_\text{H} \cdot x_\text{H}$$

$$\frac{dM}{dt} = k_\text{M} \cdot x_\text{M}$$

Obtained rate constants were summarized in Table 4, in which the term of the surface area were already eliminated using $S$ shown in Table 3. In the present study, since it was considered that the milling conditions (time, atmosphere, crystal state of samples) were resulted in the different state of product which had completely different reactivity, the rate constants summarized in Tables 1 and 2 could be different. The reasons are showing in the following section.

---

**Table 1.** Rate constants obtained by increasing milling time. (a) including the surface area, (b) except the term of surface area.

| Milling time (h) | (a) $k_\text{H}(1/s) = A_\text{H} \exp(-E_\text{H}/RT)$ | (b) $k_\text{H}(1/s) = A_\text{H} \exp(-E_\text{H}/RT)$ |
|------------------|-----------------------------------------------|-----------------------------------------------|
| 6 1x10⁶ exp | $\frac{-166,100}{RT}$ | $\frac{-3,570,000}{RT}$ |
| 24 200 exp | $\frac{-103,800}{RT}$ | $\frac{-265,800}{RT}$ |
| 72 500 exp | $\frac{-229,900}{RT}$ | $\frac{-6,300}{RT}$ |
| 100 1x10⁶ exp | $\frac{-122,900}{RT}$ | $\frac{-2,700}{RT}$ |

**Table 2.** Average particle size and total surface area of hematite. (sintered hematite, milling in air, $V_i = 3.2064 \times 10^{-3}$ cm³)

| Milling time (h) | Average particle radius $r$ (µm) | Volume of particle, $V_e$ (cm³) | Number of particle, $N_p=V_e/r^3$ | Total surface area, $S_e=4\pi r^2$ | $S_e/S_o$ |
|------------------|----------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------|
| 0 100 | 4.19x10⁻³ | 765 | 9.62 | 10 | 100 |
| 6 10 | 4.19x10⁻³ | 765 | 9.62 | 10 | 100 |
| 24 5 | 5.23x10⁻⁸ | 6.12x10⁸ | 19.2 | 20 | 20 |
| 72 1 | 4.19x10⁻³ | 765 | 9.62 | 100 | 100 |
| 100 0.1 | 4.19x10⁻³ | 765 | 9.62 | 1000 | 1000 |

**Table 3.** Average particle size and total surface area of hematite. (sintered hematite, milling in argon, $V_i = 3.2064 \times 10^{-3}$ cm³)

| Milling time (h) | Average particle radius $r$ (µm) | Volume of particle, $V_e$ (cm³) | Number of particle, $N_p=V_e/r^3$ | Total surface area, $S_e=4\pi r^2$ | $S_e/S_o$ |
|------------------|----------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------|
| 0 100 | 4.19x10⁻³ | 765 | 9.62 | 10 | 100 |
| 6 1 | 4.19x10⁻³ | 765 | 9.62 | 10 | 100 |
| 24 0.1 | 9.04x10⁻⁸ | 3.54x10⁸ | 1600 | 1664 | 1664 |
| 100 0.04 | 2.68x10⁻³ | 1.20x10⁸ | 2410 | 2508 | 2508 |

**Table 4.** Rate constants for the sample that sintered at 1 273 K for 24 h and milled in argon atmosphere.

| Milling time (h) | (a) $k_\text{H}(1/s) = A_\text{H} \exp(-E_\text{H}/RT)$, $E_\text{H}$ (J/mol) |
|------------------|-----------------------------------------------|
| 6 1.0x10⁻⁷ exp | $\frac{-50,000}{RT}$ |
| 24 1.0x10⁻⁷ exp | $\frac{-50,000}{RT}$ |
| 48 6.0x10⁻⁷ exp | $\frac{-50,000}{RT}$ |
| 100 2.4x10⁻⁷ exp | $\frac{-10,700}{RT}$ |

Hence, the kinetic analysis was carried out with the following equations.

$$RTG_H = -\frac{dM}{dt} = S_H \cdot k_H \cdot x_H = k_H \cdot x_H$$

$$RTG_M = -\frac{dM}{dt} = S_M \cdot k_M \cdot x_M = k_M \cdot x_M$$

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**Fig. 10.** Variation of average particle size and surface area with milling time.

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through the introduction of the active complexes.
In the case of argon atmosphere (as shown in Fig. 6(b)),
the reduction curve of HM never showed the peak and
showed almost constant value, which meant that the rate
constant of HM reduction should be almost the same (very
low temperature dependence).

3.3. Activated Complex Introduced by Mechanical Milling

3.3.1. Sample Milled in Air
It was found that the simple reaction of hematite with
carbon was changed by the mechanical milling, which
might be introduced the activated state into the mixture.
The interpretation of the meaning of the obtained rate
constants could be performed through the transition state
theory.6,7) According to the theory, Gibbs free energy of
activation, \( \Delta G^* \) can be expressed by Eq. (10).

\[
\Delta G^* = \Delta H^* - \Delta S^* \cdot T 
\]  

(10)

Where \( \Delta H^* \) and \( \Delta S^* \) are enthalpy of activation and entropy
of activation, respectively. The rate constant of the activation
can be expressed by Eq. (11).

\[
k = \kappa \left( \frac{k_B T}{h} \right) \exp \left( -\frac{\Delta G^*}{RT} \right) 
\]  

(11)

Where \( k_B \) is Boltzmann constant, \( \kappa \) is transmission coefficient
and \( h \) is planck constant. Substitution of Eq. (10) into
Eq. (11) can be obtained Eq. (12).

\[
k = \kappa' \left( \frac{k_B T}{h} \right) \exp \left( -\frac{\Delta H^* + \Delta S^* T}{RT} \right) 
\]  

(12)

On the other hand, the experimental rate constant obtained
in previous study6) is Eq. (13).

\[
k_1 (1/s) = A_y \exp \left( -\frac{E_y}{RT} \right) 
\]  

(13)

where \( E_y \) is the activation energy. The relationship between \( \Delta H^* \) and \( E_y \) can be expressed by
Eq. (14).6,7)

\[
E_y = \Delta H^* + (1 - \Delta n^*) \cdot RT 
\]  

(14)

where \( \Delta n^* \) means the difference of moles between the reac-
tant(s) and the activated complex, which is depending on
the reaction equation including the kind of activated com-
plex. Although overall reaction can be expressed by Eq. (15),

\[
\text{Fe}_2\text{O}_3 + 3\text{C} \rightarrow 2\text{Fe} + 3\text{CO} 
\]  

(15)

the actual reactions are considered as following consecutive
reactions for the sample milled in air atmosphere.

\[
\begin{align*}
3\text{Fe}_2\text{O}_3 + \text{C} & \rightarrow 3\text{FeO} + \text{CO} + \text{CO}_2 \\
\text{Fe}_2\text{O}_3 + \text{C} & \rightarrow 2\text{FeO} + \text{CO} \\
\text{FeO} + \text{C} & \rightarrow \text{Fe} + \text{CO} \\
\end{align*}
\]  

(16-18)

These reactions were described as a removal of the one
mole oxygen. The activated complexes included in Eq. (16)
could be considered as follows;

\[
\begin{align*}
3\text{Fe}_2\text{O}_3 + \text{C} & \rightarrow (3\text{Fe}_2\text{O}_3\text{C})^* + (2\text{Fe}_2\text{O}_4) \cdot \text{CO}^* \\
& \rightarrow (2\text{Fe}_2\text{O}_4) \cdot \text{CO}^* + 2\text{Fe}_2\text{O}_4 + \text{CO} \\
& \rightarrow 2\text{Fe}_2\text{O}_4 + \text{CO} 
\end{align*}
\]  

(19)

where \( i_2 > i_3 \) will be reasonable, because the formation and
the desorption of CO gas could be the rate controlling step
as mentioned above. Then Eq. (19) can be simplified as
follows.

\[
\begin{align*}
3\text{Fe}_2\text{O}_3 + \text{C} & \rightarrow (2\text{Fe}_2\text{O}_4 \cdot \text{CO})^* \rightarrow 2\text{Fe}_2\text{O}_4 + \text{CO} \\
& \rightarrow 2\text{Fe}_2\text{O}_4 + \text{CO} 
\end{align*}
\]  

(20)

Normally, the transition Fe 3O4* → FeO3 will be very fast
and it is no need to take into account. However in this
study, it was assumed that the activated complex could be
utilized for the next reaction, especially in the long time
milling, the contact between the iron oxide and carbon
maintained always until the reaction complete. So that fol-
lowing two cases could be considered.

\[
\begin{align*}
3\text{Fe}_2\text{O}_3 + \text{C} & \rightarrow (2\text{Fe}_2\text{O}_4 \cdot \text{CO})^* \rightarrow 2\text{Fe}_2\text{O}_4 + \text{CO} \\
& \rightarrow 2\text{Fe}_2\text{O}_4 + \text{CO} 
\end{align*}
\]  

(21)

3Fe2O3+C \rightarrow (2Fe2O4·CO)* \rightarrow 2Fe2O4+CO ... (22)

Using Eq. (21) or (22), \( \Delta n^* \) can be determined.

\[
\Delta n^* = 1 - 4 = -3 
\]  

(23)

\[
\Delta H_{\text{II}}^* = E_{\text{II}}^* + \Delta n^* \cdot RT = E_{\text{II}} - 4 \cdot RT 
\]  

(24)

Similarly, magnetite and wustite reduction were presented.

\[
\begin{align*}
\text{Fe}_3\text{O}_4 + \text{C} & \rightarrow (3\text{FeO} \cdot \text{CO})^* \rightarrow 3\text{FeO} + \text{CO} \\
& \rightarrow 3\text{FeO} + \text{CO} \\
\text{FeO} + \text{C} & \rightarrow (\text{Fe} \cdot \text{CO})^* \rightarrow \text{Fe} + \text{CO} \\
& \rightarrow \text{Fe} + \text{CO} 
\end{align*}
\]  

(25-26)

For convenience, notations, H, M, W and Fe are used for
Fe3O4, Fe2O3, FeO and metallic iron, respectively. The asterik
‘*’ means the activated state of the molecules and atom, Fe2O4*, FeO*, Fe* and complexes (2Fe2O4·CO)*,
(3FeO·CO)*, (Fe·CO)*.

Equation (20), Eq. (25) and Eq. (26) are rewritten as follows.

\[
\begin{align*}
3\text{Fe}_2\text{O}_3 + \text{C} & \rightarrow (2\text{Fe}_2\text{O}_4 \cdot \text{CO})^* \rightarrow (2\text{Fe}_2\text{O}_4 \cdot \text{CO})^* \rightarrow (2\text{Fe}_2\text{O}_4 \cdot \text{CO})^* \\
\text{M} + \text{C} & \rightarrow (3\text{W} \cdot \text{CO})^* \rightarrow (3\text{W} \cdot \text{CO})^* \\
\text{W} + \text{C} & \rightarrow (\text{Fe} \cdot \text{CO})^* \rightarrow (\text{Fe} \cdot \text{CO})^* 
\end{align*}
\]  

(20′-26′)

The enthalpy of activation for Eq. (25) and Eq. (26) can be
calculated by Eq. (27) and Eq. (28), respectively.

\[
\Delta H_{\text{M}}^* = E_{\text{M}} - 2 \cdot RT 
\]  

(27)

\[
\Delta H_{\text{II}}^* = E_{\text{II}} - 2 \cdot RT 
\]  

(28)

Moreover, Frequency factor \( A_y \) and the entropy of activa-
tion are expressed by Eq. (29).
Where $A_k = \kappa \left( \frac{k_B T}{h} \right) \exp \left( \frac{\Delta S^*}{R} \right) \quad \text{(29)}$

Using experimental $A_k$ and $E_0$ shown in Table 1(b), which were already eliminated the effect of surface area of hematite, the enthalpy $\Delta H^*$ and the entropy $\Delta S^*$ of activation were evaluated and summarized in Table 5. The temperatures for the enthalpy and entropy were adopted the peak temperatures of the respective reaction curves, which were obtained in previous study.\(^{6,7}\) In this study, the value of $\kappa$ in Eq. (29) was assumed as unity.

(1) Feature of Hematite (H–M) and Magnetite (M–Fe) Reduction

The inherent image of the transition state can be understood through the entropy of activation, $\Delta S^*$. The negative value of $\Delta S^*$ means the vanishing of the freedom for the vibrations of the rotation and the translation from reactant to complex.\(^{7}\) The H–M and M–W reduction are corresponding to this situation. Especially in present experiment, the 6 h milling time and the longer ones (24 h, 72 h and 100 h) showed different situation. It was considered that the activated complex did not constructed completely in the case of 6 h milling, which could be understood by the change of activation energy (the value of activation energy decreased from 6 to 24 h). $\Delta S^*$ of H–M reduction from 24 to 100 h almost the same (约-240 J/mol·K), however, it decreased from 6 h (约-179 J/mol·K) to 24 h (约-236 J/mol·K) which mean the complex, (2M·CO)* lost the freedom of some vibration from 6 h milling to 24 h milling.

The M–W reduction showed almost the same trend as the H–M reduction. These explanations on the nature of the activated complex would also discussed through the partition function as follows:

Equation (11) can be rewritten by using the partition function.\(^{6,7}\)

$$k = \kappa \left( \frac{k_B T}{h} \right) \exp \left( \frac{\Delta S^*}{R} \right) \quad \text{(30)}$$

Where $F_{MCO}^*, F_H^*$, and $F_C^*$ are the partition function for the complex, hematite and carbon.

If the reaction was simplified to $A^+BC \rightarrow A^+ \cdot \cdot \cdot B \cdot \cdot \cdot C \rightarrow AB+C$, each partition functions can be expressed by Eqs. (31) and (32).\(^{6,7}\)

\begin{align*}
F^* &= g^* \left( \frac{2m_n k_B T}{h^3} \right)^{3/2} \frac{8\pi^2 m_n k_B T}{\sigma_A h^2} \prod \left( 1 - \exp \left( \frac{-h v_n}{k_B T} \right) \right)^{-1} \quad \text{(31)}
\end{align*}

$$F_A \cdot F_{BC} = \left[ g_1 \left( \frac{2m_n k_B T}{h^3} \right)^{3/2} \frac{8\pi^2 m_n k_B T}{\sigma_A h^2} \right] \times \left( 1 - \exp \left( \frac{-h v_n}{k_B T} \right) \right)^{-1} \quad \text{(32)}$$

The term corresponding to the entropy of activation, $F^*/(F_A \cdot F_{BC})$ would decrease with the decrease of $F^*$ because of losing of some vibration mode, $\Pi (1 - \exp(-h v_n/k_B T))^{-1}$, which would correspond to the change of $\Delta S^*$ from 6 to 24 h.

Moreover, as this term related to the temperature dependence, the activation energy $E_A$ would also decrease with the term of $\Pi (1 - \exp(-h v_n/k_B T))^{-1}$, which is equivalent to the high-temperature reaction.

(2) Feature of Wustite (W–Fe) Reduction

On the other hand, wustite (W–Fe) reduction showed entirely different behavior. The values of the entropy of activation, $\Delta S^*$ was positive and quite large value (Table 5). Furthermore, the enthalpy of activation, $\Delta H^*$ was extremely high according to the milling time. The positive value of $\Delta S^*$ means that the activated complex has larger entropy than that of reactant. Moreover, a loose bonded complex has larger entropy than that of rigid bonded complex.\(^{3,7}\)

From this point of view, the positive and extremely high value of $\Delta S^*$ meant that the complex of (Fe·CO)* having a loose bond, which corresponded to the high reaction rate.

3.3.2. Sample Milled in Argon

Similarly to the analysis of activation complex on the sample milled in the air, the rate constants obtained from the reaction of sample milled in the argon atmosphere were analyzed as follows.

Hematite reaction was the same as Eq. (2). However, Magnetite reduction M–Fe can be expressed as Eq. (33).

$$\frac{1}{4} \text{Fe}_2\text{O}_4 + \text{C} = \frac{3}{4} \text{Fe} + \text{CO} \quad \text{.........(33)}$$

And active complex can be expressed by Eq. (34).

$$\frac{1}{4} \text{Fe}_2\text{O}_4 + \text{C} \rightarrow \frac{3}{4} \text{Fe} + \text{CO} \quad \text{.........(34)}$$

The difference of mole between the complex and the reactants is

$$\Delta n^* = 1 - \frac{5}{4} = -\frac{1}{4} \quad \text{.........(35)}$$

Then the enthalpy of activation can be calculated by Eq. (36).

$$\Delta H^*_M = E_M + (\Delta n^* - 1) \cdot RT = E_M - \frac{5}{4} \cdot RT \quad \text{.........(36)}$$

The enthalpy ($\Delta H^*$) and entropy ($\Delta S^*$) of activation were...
estimated by Eqs. (36) and (29), respectively. The obtained values are shown in Table 6 and plotted against the milling time in Fig. 11 (W–Fe reduction in air milling, M–Fe reduction in argon milling). It was found that $D_{H}^{*}$ and $D_{S}^{*}$ varied same way with milling time. When $D_{S}^{*}$ increased (decreased) with milling time, $D_{H}^{*}$ also increased (decreased). However, the way of variation between the milling in the air and in the argon was opposite, which meant that completely different feature (transition state) of product were assigned by milling between two conditions. As mentioned above, the positive and extremely high value of $D_{S}^{*}$ meant the complex of (Fe · CO)$^{*}$ having a loose bond owing to the milling in air. Contrary to the milling in air, the sample milled in argon atmosphere gave the decreasing tendency of the entropy of activation, $D_{S}^{*}$. In the case of milling in air, the active complex having a loose bond was generated during milling. On the other hand, in argon atmosphere, it would be the vanishing process of some vibration mode for the bondings. So that the milling in argon makes the active complex having relatively tight bond close to the state of the product. The complex would have a different structure from the reactant loosing some vibration mode. In other words, the complex made in argon atmosphere would have a structure similarly to the product one. Therefore, the reactivity of milling sample was extremely higher than that in the air.

Gibbs free energy of activation, $D_{G}^{*}$ was evaluated at 1 000 and 1 273 K using $D_{H}^{*}$ and $D_{S}^{*}$ obtained, as shown in Fig. 12. $D_{G}^{*}$ for milled in the air has extremely high value in longer milling time and it decreased with increasing temperature from 1 000 to 1 273 K. Corresponding to this, $D_{G}^{*}$ from the sample milled in the argon atmosphere decreased with the increasing milling time. Moreover, it decreased largely from 1 000 to 1 273 K, which meant the reaction needed the higher energy to proceed in the lower temperature.

4. Conclusions

Milling of hematite and graphite mixture was carried out both in the air and in the argon. Rate constants for the reactions were obtained by kinetic analysis. The enthalpy $D_{H}^{*}$ and entropy $D_{S}^{*}$ of activation were estimated on the basis of transition state theory. The interpretation of $D_{S}^{*}$ and $D_{H}^{*}$ was performed to explain the effect of milling on the reactivity related to the milling conditions. The results obtained are as follows.

(1) The temperature of peak of reaction curve decreased with increasing milling time. The reactivity of the sample milled in the argon atmosphere showed higher reactivity than that milled in the air.

(2) The reaction consisted of three steps (HM, MW and WFe reduction) for the sample milled in the air, while there were two reaction steps (HM and MFe reduction) existed in the reaction of sample milled in the argon.

(3) From the results of XRD, the crystalline size in the sample milled in air showed almost constant size, while the crystalline size milled in argon decreased. Taking into account of the CO gas evolution during milling, it was considered the reduction reaction occurred. Especially, reoxidation should occur during the milling in the air atmosphere. The difference of crystal state after milling should result in the quite different reactivity through the active complex.

(4) The enthalpy $D_{H}^{*}$ and entropy $D_{S}^{*}$ of activation were obtained from experimental rate constants on the basis estimated by Eqs. (36) and (29), respectively. The obtained values are shown in Table 6 and plotted against the milling time in Fig. 11 (W–Fe reduction in air milling, M–Fe reduction in argon milling). It was found that $D_{H}^{*}$ and $D_{S}^{*}$ varied same way with milling time. When $D_{S}^{*}$ increased (decreased) with milling time, $D_{H}^{*}$ also increased (decreased). However, the way of variation between the milling in the air and in the argon was opposite, which meant that completely different feature (transition state) of product were assigned by milling between two conditions. As mentioned above, the positive and extremely high value of $D_{S}^{*}$ meant the complex of (Fe · CO)$^{*}$ having a loose bond owing to the milling in air. Contrary to the milling in air, the sample milled in argon atmosphere gave the decreasing tendency of the entropy of activation, $D_{S}^{*}$. In the case of milling in air, the active complex having a loose bond was generated during milling. On the other hand, in argon atmosphere, it would be the vanishing process of some vibration mode for the bondings. So that the milling in argon makes the active complex having relatively tight bond close to the state of the product. The complex would have a different structure from the reactant loosing some vibration mode. In other words, the complex made in argon atmosphere would have a structure similarly to the product one. Therefore, the reactivity of milling sample was extremely higher than that in the air.

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of transition state theory. The variation of $\Delta H^*$ and $\Delta S^*$ with milling time showed opposite tendency between the sample milled in the air and in the argon.

(5) It was considered that the active complex having loose bonding would form during milling in the air atmosphere. On the other hand, the active complex having a structure closed to the product (metallic iron) might form.

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REFERENCES

1) J. Vahdati Khaki, Y. Kashiwaya, K. Ishii and H. Suzuki: *ISIJ Int.*, 42 (2002), 13.
2) Y. Kashiwaya, M. Kanbe and K. Ishii: *ISIJ Int.*, 41 (2001), 818.
3) Y. Kashiwaya and K. Ishii: *ISIJ Int.*, 31 (1991), 440.
4) Y. Kashiwaya, H. Suzuki and K. Ishii: *ISIJ Int.*, 44 (2004), 1970.
5) Y. Kashiwaya, R. Suzuki and K. Ishii: *ISIJ Int.*, 44 (2004), 1975.
6) S. Glasstone, K. J. Laidler and H. Eyring: The Theory of Rate Processes, McGraw-Hill, New York and London, (1941), 146.
7) W. J. Moore: Physical Chemistry 4th Ed., Tokyo Kagaku Dojin, Tokyo, (1978), 387.