Intrinsic Reduction Kinetics Investigation on a Hematite Oxygen Carrier by CO in Chemical Looping Combustion

Mingze Su,*† ‡ Haibo Zhao,‡ Xin Tian,‡ Pengfei Zhang,‡ Baozhou Du, † and Zhigang Liu*‡

Abstract: In the present work, the intrinsic reduction kinetics of a chemical looping oxygen carrier, hematite, reduced by CO has been investigated. Initially, the pore structure features of the hematite OC were tested to provide the necessary parameters for the kinetics analysis. Then, temperature-programmed reduction experiments were conducted in a thermogravimetric analyzer. The results showed that the weight loss rate appears to be a bimodal distribution and the appropriate experimental temperature range for the intrinsic kinetics testing of the first reduction stage of hematite is 400–650 °C. Additionally, a series of experiments were further conducted in a batch fluidized bed system with different temperatures and different CO concentrations. After elimination of the internal and external diffusion limitations, the intrinsic activation energy and pre-exponential factor were determined as 74.48 kJ/mol and 1.2 × 1012 s⁻¹, respectively. The analysis of the internal and external diffusion effects was beneficial to understand the reaction characteristics of this kind of hematite oxygen carrier. As concluded, the shrinking core series model, the external diffusion control, and the small apparent activation energy obtained from many previous studies were compatible with this analysis and can be explained by this work. The research results attained in this study can also contribute to the development of a practical model for computational fluid dynamics simulation.

1. INTRODUCTION

Chemical looping combustion (CLC), which has the characteristic of CO₂ inherent separation, has been considered as a very promising fossil fuel utilization technology for addressing the global warming issue.¹,² Because CLC splits the conventional combustion process into two separate redox processes, the gas composition separation (CO₂ separation from flue gas) in conventional CO₂ capture techniques can be substituted by the gas–solid [flue gas–oxygen carrier (OC) particles] separations in CLC. In such a way, much higher energy conversion efficiency can be achieved as well as much lower cost for CO₂ capture.³,⁴ Basically, lattice oxygen is cyclically transferred to oxidize the fuel with the aid of the OC circulation in CLC. In general, the CLC reactor is composed of an air reactor (AR) and a fuel reactor (FR). Air and fuel enter the AR and FR separately and without direct contact. The schematic view of the CLC reactor is shown in Figure 1. The fossil fuel is oxidized via OCs with high oxygen potential (usually metal oxides, Me₃Oₓ) in FR and comes out as CO₂, H₂O, and OCs with low oxygen potential (MeₓOᵧ−₁) (see eq 1). High-purity CO₂ can be easily obtained by a simple condensation treatment to the flue gas of the FR.

\[
(2n + m)Me₃O_y + C_nH_{2m} \rightarrow (2n + m)Me₃O_{y−₁} + mH₂O + nCO₂
\]  

(1)

\[
(2n + m)Me₃O_{y−₁} + (n + m/2)O₂ \rightarrow (2n + m)Me₃O_y
\]  

(2)

\[
C_nH_{2m} + (n + m/2)O₂ \rightarrow mH₂O + nCO₂
\]  

(3)

Me₃O_y can be regenerated in AR (see eq 2). In this way, a loop can be formed as produced Me₃O_y is further transferred back to the FR. The overall reaction of the traditional combustion process (see eq 3) is the sum of eqs 1 and 2 indeed.

Fe₃O₄ is regarded as a kind of cheap and environmentally friendly OC material. Among all of the Fe-based OCs, iron ore has been widely adopted as OC in most pilot-scale CLC systems. The reaction mechanism and kinetics of Fe-based OCs have already been reported in some literature. Elgeassy et al.⁵ and Ettabirou et al.⁶ have studied the reduction kinetics of Fe₃O₄ by CO decades ago. However, the obtained kinetics, which was only limited to the metallurgical industry, might not be applicable to the CLC processes. On the basis of thermodynamic analysis, Wang et al.⁷ pointed out that the
active material, Fe₃O₄, would react with the inert support, Al₂O₃, to produce Fe₃Al₂O₆, which plays a negative role in the reactivity of the OC. Zafar et al. found that Fe₃O₄-based OC demonstrates high reactivity in the first reduction stage (Fe₂O₃ to Fe₃O₄) but typical low reactivity was observed in the second reduction stage (Fe₃O₄ to FeO). Abad et al. examined the apparent kinetics of the synthetic Cu-, Fe-, and Ni-based OCs reduced by CO, H₂, and CH₄ for the purpose of providing key kinetic parameters for reactor design and computational fluid dynamics (CFD) simulation. Obviously, the obtained small activation energy indicates that the diffusion effect had not been fully eliminated in their experiments. Chen et al. investigated the reduction kinetics of hematite in a micro-fluidized bed reactor. Unfortunately, the effect of the reducing gas concentration was not discussed in their study.

The heterogeneous gas–solid reaction is regarded as one of the most complex processes in nature. The overall reaction includes both the chemical reaction process and the mass/heat transfer processes; moreover, the solid material/surface is also difficult to be characterized. The intrinsic reaction kinetics and the internal/external mass transfer effects should be comprehensively considered when a kinetic equation is established. As reviewed in the literature, there is no agreement on the rate-controlling step for the reduction of OC, let alone the reported hensively considered when a kinetic equation is established. As for the OC, the internal/external mass transfer processes; moreover, the solid material/surface is also difficult to be characterized. The intrinsic reaction kinetics and the internal/external mass transfer effects should be comprehensively considered when a kinetic equation is established. As reviewed in the literature, there is no agreement on the rate-controlling step for the reduction of OC, let alone the reported hensively considered when a kinetic equation is established.

First, the reduction activities of the surfaces (1102) and (0001) of FeO were studied, and the best adsorption sites were identified. As concluded, the surface (1102) is more prone to reacting with CO than the surface (0001). Then, a possible reaction mechanism for the reduction of Fe-based OC by CO was proposed: Fe₂O₃ will be activated by one CO molecule to produce Fe₃O₄, and then the activated O molecule will be reduced by another CO molecule to produce CO₂. After that, they pointed out that the energy barrier of the second reduction stage is far larger than that of the first stage; thus, it is inadvisable to reduce the Fe₃O₄-based OC deeply in practical CLC application. Finally, they examined the effect of the inert carriers, such as ZrO₂, MgO, and Al₂O₃, on the reactivity of Fe₂O₃ with CO. They concluded that the inert carriers promote the adsorption of CO on Fe₂O₃ and the electron transfer from the inert carriers to Fe₂O₃, lengthens the Fe—O bond, which is beneficial to the reduction of Fe₂O₃. The energy barriers of the reactions were determined as 2.59, 0.99, 0.98, and 0.83 eV for the reduction of Fe₂O₃, Fe₃O₄/Al₂O₃, Fe₃O₄/MgO, and Fe₃O₄/ZrO₂, respectively.

Although the reduction of hematite OC has been widely studied, its reduction kinetics is still unclear (especially for the first reduction stage) and there exists a huge gap between experimental measurements and quantum chemistry calculations. In this work, the intrinsic kinetics of a kind of hematite OC reduced by CO was experimentally investigated. Initially, the Brunauer—Emmett—Teller (BET) surface area test was conducted to provide necessary parameters for theoretical analysis. Then, temperature-programmed reduction experiments were conducted in TGA to determine an appropriate temperature range for the intrinsic kinetics testing. Finally, a series of formal experiments were conducted in a batch fluidized bed reactor with different temperatures and different CO concentrations to analyze the intrinsic kinetics, and the effects of the internal and external diffusion factors were also discussed.

2. EXPERIMENTAL SECTION

The iron ore originated from the E’kou mine, China. The physical and chemical properties of this hematite OC can be considered to be stable in this study. The iron ore used in this work has been previously used for more than 300 h in a 5 kWth interconnected fluidized bed reactor. The OC has been fully oxidized before tests, and the elemental analysis [as determined by X-ray fluorescence (XRF, EDAX EAGLE III)] showed that the mass fraction of Fe₂O₃ is 66 wt %. The particle size in the diameter range of 180–200 μm by sieving were collected as the samples in this experiment. The granule density of the particles is 3472 kg/m³. Characterizations of this material, such as X-ray diffraction (XRD, EM-PTREAN) analysis and BET surface area (JW-BK132F) analysis, will be shown in the following section.

Under the heating rate of 15 °C/min and 15 and 50 vol % CO balanced by N₂ as the reducing gas, temperature-programmed reduction experiments were conducted in TGA (Netzsch STA409PC). The sample mass was 12.5 mg, and the final temperature for both tests was 970 °C. The weight loss peaks will be discussed later.

After that, experiments were further conducted in a batch fluidized bed reactor to investigate the intrinsic reduction kinetics of the first reduction stage of hematite OC. A total of 5 g of hematite was chosen as the active OC, and 15 g of silica sand was chosen as the inert bed material. The main reason for adding silica sand is to reduce the error coming from the sample loss. Reducing gas of 5, 7.5, and 15 vol % CO balanced by N₂ was introduced into the reactor with a flow rate of 1 L/min in the reduction period. The experiments were conducted at 400, 425, 450, 475, 555, 600, and 647 °C, separately. The choice of these
temperatures depends upon the analysis of the TGA results. The schematic view of the fluidized bed experimental rig is presented in Figure 2. The diameter of the reaction chamber is 33 mm, and the exhaust gas was detected by the online gas analyzer (Gasboard 3100P, Cubic Optoelectronics).

For the batch fluidized bed experiment, we considered N$_2$ as the characteristic fluidization agent. The lowest operation temperature, 400 °C (where the worst fluidization occurs), was considered as the reference temperature for analysis. The density and viscosity of N$_2$ at 0.0442 m/s give always fluidized in our experiments.

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3. RESULTS

3.1. BET Surface Area and Pore Size Distribution. The pore size distribution of the OC particle after redox cycles is shown in Figure 3. The most probable pore radius and the mean pore radius ($\bar{r}_p$) are 3.051 and 12.904 nm, respectively. The BET surface area, $S_B^{\text{BET}}$, is 1.387 m$^2$/g. Thus, the porosity of this material, $\varepsilon = \rho_B S_B^{\text{BET}}/\rho$, is 0.031. The surface area per volume of the solid material, $S_B^{\text{BET}}$, was attained according to the following correlation:

$$S_B^{\text{BET}} = \frac{4.86 \times 10^6 \mu^2 m^3}{C_\text{BET}}.$$
into the fuel reactor will be heated rapidly; thus, the first reduction stage can only happen under high-temperature conditions. Because the first reduction stage could happen at a typical low temperature, the reaction under high temperatures may be controlled by external mass transfer. This is also consistent with conclusions obtained from our previous research.18

In the batch fluidized bed experiments, CO was consumed when it flowed through the bed materials, which results in the fact that the outlet CO concentration is smaller than that of the inlet. Therefore, the CO concentration adopted in data processing is an important issue that should be appropriately addressed. Although most researchers (including us) take the logarithmic average (see section 3.3.1) as the characteristic value, this treatment may also introduce some errors to the kinetics analysis. For a better accuracy, consumed CO should be restricted to a relatively small amount; thus, a low reaction rate is required. Another aim of the TGA experiments is to seek the appropriate temperature range for the batch fluidized bed experiments. As mentioned above, the intrinsic reduction kinetics of the first reduction stage of hematite should not be tested under the temperature condition of a real CLC process, i.e., 900 °C or higher, and should be tested within the temperature range of 400 to 600 °C, because the intrinsic kinetics will be distorted as a result of the limitation of gas diffusion at higher temperatures. On the basis of this understanding, 400−650 °C was chosen as the temperature window for the batch fluidized bed experiments later. Lower temperatures are also beneficial to determine the CO volume fraction more accurately.

3.3. Batch Fluidized Bed Experiment. 3.3.1. Data Processing Method. The flow rate and volume fraction of the exhaust gas components were detected online in the batch fluidized bed experiments. Because no CO2 was detected during the OC oxidation period, it can be considered that no carbon deposition (no coking) occurred during the reduction period. The conversion (X) as a function of time can be calculated as

\[
X = \frac{\int_{t_0}^{t_f} F_{\text{ex}} Y_{\text{CO}_2} \, dt}{\int_{t_0}^{t_f} F_{\text{ex}} Y_{\text{CO}} \, dt}
\]

(4)

where \( F_{\text{ex}} \) is the flow rate of the exhaust gas and \( Y_{\text{CO}_2} \) represents the volume fraction of \( \text{CO}_2 \) in the product gas.

A typical profile for the product gas concentration detected by the gas analyzer is also shown in Figure 5. CO is considered as excessive for the intrinsic kinetics testing, without the influence of the aeration rate. As mentioned above, the logarithmic-averaged volume fraction was introduced as the characteristic CO volume fraction. The logarithmic average can be calculated as

\[
Y_{\text{ave}} = \frac{Y_{\text{in}} - Y_{\text{out}}}{\ln(Y_{\text{in}}/Y_{\text{out}})}
\]

(5)

where \( Y_{\text{in}} \) and \( Y_{\text{out}} \) represent the inlet and outlet volume fractions for the same gas component, respectively. The logarithmic-averaged volume fraction will be used in the formal...
temperature, and $R$ represents the universal gas constant, $T$ calculated according to the mass transfer correlation.\(^3\) To the particle surface. The maximum reaction rate can be reached, there exists a rising process, which may result from the combination effect of gas mixing in the reactor and gas diffusion to the particle surface. The maximum reaction rate can be regarded as the reaction rate when the solid conversion, $X$, was equal to 0. Therefore, the maximum values were chosen as the characteristic reaction rates for analysis. The Arrhenius-type apparent kinetics equation can be used to describe the conversion profile as follows:

$$\frac{dX}{dt} = k' \exp(-E/RT)f(X)g(Y_{CO})$$  \hspace{1cm} (6)

where $X$ represents the conversion of OC, $k'$ is the pre-exponential factor, $E$ is the apparent activation energy, $R$ represents the universal gas constant, $T$ is the reaction temperature, and $Y_{CO}$ represents the mole/volume fraction of the reducing gas. $f(X)$ is the reaction mechanism function in differential form, which reflects the functional relationship between the reaction rate, $dX/dt$, and the solid conversion, $X$. According to the scale investigated, $f(X)$ can be grouped into two categories: the particle function and the grain function. Within the particle function, the homogeneous reaction model is usually represented as $f(X) = 1 - X$, while the shrinking core model is usually represented as $f(X) = (1 - X)^{2/3}$. Generally speaking, the homogeneous reaction model corresponds to the situation that the overall reaction is not affected by the internal diffusion, while the shrinking core model is satisfying when the overall reaction is highly affected by the internal diffusion. $g(Y_{CO})$ represents the relationship between the reaction rate with the reducing gas concentration. The simplest form, $g(Y_{CO}) = Y_{CO}$, was applied in this study, although several researchers (e.g., Monson et al.\(^3\)) have reported that the power function correlation does not apply well over a wide range of gas concentrations.

### 3.3.3. Effect of the External Diffusion

To evaluate the effect of external diffusion, the reaction rate measured in the experiment was compared to the maximum rate that could be reached under the gas diffusion control, as shown in Figure 6. The mass transfer rate, $r_{\text{ext}}$ (mol g\(^{-1}\) s\(^{-1}\)), can be calculated as

$$r_{\text{ext}} = k_s S_m (C_B - C_i)$$  \hspace{1cm} (7)

where $S_m$ represents the bulk concentration of CO (i.e., 2.76 mol/m\(^3\) under the condition of 450 °C and volume fraction of 15 vol % CO) and $C_i$ is the concentration of CO on the external surface of a particle. To calculate the maximum reaction rate, $C_i$ is assumed to be 0. $S_m = 0.0091$ m\(^3\)/g is the nominal external surface area per unit mass (g) of the OC particle (here, the particles were considered to be absolutely spherical), $k_s$ is the external mass transfer coefficient and can be calculated according to the mass transfer correlation.\(^3\)

$$Sh = 2\varepsilon_i + 0.69Re_p^{1/2}Sc^{1/3}$$  \hspace{1cm} (8)

where $k_s$ is the intrinsic reaction rate, and $\eta$ represents the ratio of the reaction rate with diffusion resistance to that without.
Diffusion resistance. $\eta \rightarrow 1$ means that the heterogeneous reaction rate is relatively lower when compared to the gas diffusion in the pore. Hence, the concentration of the reducing gas in the pore was equal to that on the external surface of the particle. In this case, the heterogeneous reaction happens only on the external layer of the particle, because the reducing gas will be consumed before diffusing into the interior of the particle through the pore. In this case, the mechanism function of the overall reaction may correspond to the shrinking core model. $\eta$ can be calculated by iterating eq 9 and the following two equations:

$$\eta = \Phi \Phi \Phi - \left(\frac{1}{3}\Phi\right)^2(3\Phi \coth 3\Phi - 1)$$

$$\Phi = \left(\frac{d_p}{6}\right)\sqrt{k m_v / D_{eff}}$$

where $\Phi$ is the Thiele modulus, $m_v$ represents the solid mass per unit volume of carrier ($3.47 \times 10^8$ g/m$^3$), $D_{eff}$ is the effective diffusivity of CO within the pore, which can be calculated as $D_{eff} = \varepsilon^2 / (1/D_K + 1/D_O)$ (i.e., $5.46 \times 10^{-9}$ m$^2$/s under the condition of 450 °C), $D_K$ is the Knudsen diffusivity (i.e., $6.34 \times 10^{-8}$ m$^2$/s at 450 °C), and $\varepsilon = 0.031$ is the porosity of the OC particle, which has been gained in section 3.1. The calculated effectiveness factors when using 15 vol % CO as the reducing gas are shown in Table 1.

From Table 1, it can be seen that the effectiveness factors are far smaller than 1 for all tests, which illustrates that the gas–solid reaction happens on the external layer of the particles and the overall reactions can be described by the shrinking core model. On the other hand, the effectiveness factors decrease as the temperature increases, indicating that the reaction layer becomes thinner with the increase of the temperature. The gas–solid contact surface for the heterogeneous reaction becomes smaller under a higher temperature condition than that under a lower temperature condition. This change partially offsets the reactivity-induced improvement of the overall reaction rate, which results in a much lower value of the activation energy. The intrinsic reaction rate, $k_v$, which can be obtained in the above analysis simultaneously, is also shown in Table 1.

The tests with the same reaction temperature are taken as a group. $k_v$ as a function of the CO volume fraction at different temperatures is shown in Figure 7. It can be seen that the intrinsic reaction rates have good linearity with the volume fraction of the reducing gas; thus, the reaction order, $n$, can be concluded as 1. Different constants, $b$, were obtained for different temperatures. Arrhenius analysis [$\ln(b)$ as a function of $1/T$, also shown in Figure 7] gave an activation energy of 74.48 kJ/mol and a pre-exponential factor of $3.46 \times 10^6$.

Considering the relationship between $k_v$ and $k_v$, i.e., $k_v = k_v m_v$, the final version of the reaction rate equation can be expressed as

$$k_v = 1.2 \times 10^{12} \exp\left(-\frac{74480}{RT}\right)Y_{CO}$$

The activation energy of 74.48 kJ/mol corresponds to the energy barrier of 0.774 eV in the DFT simulation, which is almost consistent with the previous simulation results. $^{23-25}$ Interestingly, the pre-exponential factor of $1.2 \times 10^{12}$ s$^{-1}$ was close to the theoretical value of the vibrational frequency of the atom in the potential minimum, at a low temperature ($k_B T/h = 0.04 K$).

### Table 1. Relationships of $\eta$ and $k_v$ with the Experimental Temperatures (15 vol % CO)

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>400</th>
<th>425</th>
<th>450</th>
<th>475</th>
<th>555</th>
<th>600</th>
<th>647</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\eta$</td>
<td>0.00477</td>
<td>0.00375</td>
<td>0.00330</td>
<td>0.00283</td>
<td>0.00217</td>
<td>0.00188</td>
<td>0.00171</td>
</tr>
<tr>
<td>$k_v$</td>
<td>0.0659</td>
<td>0.1088</td>
<td>0.1441</td>
<td>0.1991</td>
<td>0.3616</td>
<td>0.4991</td>
<td>0.6192</td>
</tr>
</tbody>
</table>

**Figure 7.** Arrhenius plot for $k_v$.
1.5 \times 10^{13} \text{ s}^{-1}, where the Planck constant \( h = 6.626 \times 10^{-34} \text{ J s} \) and the Boltzmann constant \( k_B = 1.381 \times 10^{-23} \text{ J/K} \).

For validation, as described in section 3.3.1, the results with the same inlet concentration were analyzed in a group. The results of the Arrhenius analysis \( \ln(k_s) \) as a function of \( 1/T \) for each group is shown in Figure 8. The activation energies were determined as 60.40 and 60.07 kJ/mol, respectively, which are smaller than the activation energy obtained above (74.48 kJ/mol). It is because, although the inlet CO concentrations have the same value for the same group, the characteristic concentrations are not equal for each point in the same group because the outlet concentrations are different. Grouping experimental results according to the inlet concentration will bring errors to the analysis. From Figure 8, it can also be seen that the results at a higher temperature deviate from the extrapolation of the low-temperature experimental results. It is because the high-temperature reduction reaction is influenced by the external mass transfer.

Many researchers obtained relatively low activation energies in their studies, even as small as 19 kJ/mol.38 If fitting the apparent reaction rate, \( k_{\text{overall}} \) directly, as shown in Figure 9, a lower activation energy result (about 31.94 kJ/mol) can also be obtained in our study. From this perspective, the internal diffusion effect should be carefully treated for pursuing the intrinsic reaction kinetics of the OC reduction.

XRD analysis was conducted for the samples in the oxidation and reduction states, respectively. As shown in Figure 10, SiO\(_2\)

![Figure 8. Direct Arrhenius plot for \( k_s \).](image)

![Figure 9. Direct Arrhenius plot for \( k_{\text{overall}} \).](image)

![Figure 10. XRD patterns of the samples in the oxidation and reduction states.](image)

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The above results show that the intrinsic activation energy is 74.48 kJ/mol, which is larger than some results of other researchers.10,38 Low intrinsic activation energy (e.g., 19 kJ/mol38) is unconvincing, because a reaction system with such a small activation energy may be unstable at room temperature.

4. DISCUSSION

Obviously, the above analysis and results were compatible with previous conclusions, and this analysis can provide a systematical explanation for the previous conclusions. It can be seen that this hematite OC is nearly compact and relatively impermeable, which results in the fact that reducing gas can hardly permeate into the particles deeply, even under a low-temperature condition. Under higher temperature conditions, the overall reaction rate is limited by the external gas diffusion (see section 3.3.2); therefore, the concentration of the reducing gas around the external surface of particle is nearly equal to zero, making the reducing gas more difficult to permeate into the particles. The above two factors support the shrinking core series model (particle is reduced layer by layer from outside to inside). It should be noted that, although the shrinking core series model was supported by the analyses, that is not to say that the relationship between the reaction rate and the solid conversion \( (1 - X) \sim \) 2/3 definitely, because the solid conversion process may also be affected by the ash content in the condition of \( X \neq 0 \). The actual relationship should depend upon experiments in which the complete conversion process is conducted under the characteristic reaction conditions.

The above results show that the intrinsic activation energy is 74.48 kJ/mol, which is larger than some results of other researchers.10,38 Low intrinsic activation energy (e.g., 19 kJ/mol38) is unconvincing, because a reaction system with such a small activation energy may be unstable at room temperature.
In fact, the value of 19 kJ/mol was even close to the energy barrier of chemisorption. Although the DFT calculation has its limitation, all simulation results reported do not support the low activation energy, while some results agree well with the activation energy that we obtained.

There are three reasons to explain why many researchers get low activation energies using different analysis techniques. First, the experimental results were influenced more or less by the external mass transfer, because the experimental conditions were not so reasonable. As shown in this study, the external mass transfer factor will become remarkable under a high-temperature condition. It is well-known that the apparent activation energy was usually smaller than 20 kJ/mol when the reaction rate was controlled by the external mass transfer. Second, even if the external mass transfer limitation could be eliminated completely, the reduction reaction may also be controlled by the internal mass transfer, as shown in this study. The effectiveness factor for the internal diffusion decreases as the temperature increases, indicating that the reducing gas permeating into the particles becomes more difficult, which results in the decrease of the gas-solid contact surface. The decreased reaction surface partially offsets the reactivity-induced improvement of the overall reaction rate as the temperature, contributing to much lower sensitivity of the apparent reaction rate with the temperature than that of the intrinsic reaction rate, and the apparent activation energy is then smaller than the intrinsic activation energy. Finally, the kinetic compensation effect may also lead to a low activation energy. However, a low pre-exponential factor will certainly be obtained simultaneously, and the two lower values for the parameters make no physical sense. As well known in the quantum chemistry calculation, the reaction rate has a lower sensitivity to the pre-exponential factor than the activation energy. Therefore, to obtain the intrinsic kinetics in the fitting process, the pre-exponential factor should be restricted in the order of magnitude as its physical background. Only in this way, the comparison of the activation energy is meaningful.

Although the intrinsic reduction kinetics of the hematite OC by CO was obtained in this work, the mechanism of the reaction at the molecular level is still unknown. That is to say, the reaction coordinate is still unclear. However, the kinetics is highly dependent upon the reaction mechanism. Although some DFT simulation results have been reported, it is still not so confident to accept these mechanism results as a result of the inherent disadvantage of the simulation and the complexity of the reaction pathway. The aim of this work is to establish the relation between the DFT simulation results and the experimental results; this is only a preliminary step forward obtained now, and more work is required in the future.

Through analysis of the internal and external diffusion factors, it is clear that the overall reaction may be affected by the external mass transfer under a high temperature condition and the reaction process can be described by the shrinking core series model. Further work should be performed with comprehensive consideration of the intrinsic kinetics, reactive surface, and internal/external diffusion to build a CFD-simulation-applicable model. The basic idea is shown in Figure 11. This model will suit not only regime I but also regimes II and III.

5. CONCLUSION

In this work, the intrinsic kinetics of the first reduction stage of a hematite OC reduced by CO was carefully investigated. Initially, the BET surface area and pore size distribution were obtained to provide fundamental data for the kinetics analysis. After that, temperature-programmed reduction experiments were conducted in TGA. The results showed that the weight loss rate demonstrates a bimodal distribution. For the first reduction stage of this hematite OC, the appropriate experimental temperature range for the intrinsic kinetics testing was determined as 400–650 °C. Additionally, a series of formal experiments were further conducted in a batch fluidized bed rig with different temperatures and different CO concentrations. After elimination of the internal and external diffusion limitations, the intrinsic activation energy and pre-exponential factor were determined as 74.48 kJ/mol and 1.2 × 10^12 s⁻¹, respectively. The analysis of the internal and external diffusion factors is beneficial to understand the reaction characteristics of this kind of hematite OC. The external diffusion control and the smaller apparent activation energy attained in previous studies can be well-explained by the result obtained in this work. This study can also help to develop the applicable model for CFD simulation.

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