Solid-state reduction kinetics and mechanism of pre-oxidized vanadium–titanium magnetite concentrate

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Abstract: The solid-state reduction kinetics of pre-oxidized vanadium–titanium magnetite concentrate was studied. The phase and microstructure of the reduction product were characterized by XRD, SEM and EDS methods, based on which the mechanism of the solid-state reduction was investigated. The results showed that using coal as reductant at 950–1100 °C, the solid-state reduction of the pre-oxidized vanadium–titanium magnetite concentrate was controlled by interface chemical reaction and the apparent activation energy was 67.719 kJ/mol. The mineral phase transformation during the reduction process can be described as follows: pre-oxidized vanadium–titanium magnetite concentrate → ulvospinel → ilmenite → FeTi2O5 → (Fe,Mn)Fe2O5·3M. M3O5-type (M can be Fe, Ti, Mg, Mn, etc) solid solutions would be formed during the reduction process of the pre-oxidized vanadium–titanium magnetite concentrate at 1050 °C for 60 min. The poor reducibility of iron in M3O5 solid solutions is the main reason to limit the reduction property of pre-oxidized vanadium–titanium magnetite concentrate.

Key words: vanadium–titanium magnetite; solid-state reduction; reduction kinetics; reduction process

1 Introduction

Panxi Area of China is endowed with large reserves of vanadium–titanium magnetite, and the reserves of vanadium and titanium account for 11.6% and 35.17% of the total reserves in the world, respectively [1–4]. At present, China and Russia adopt the BF-BOF process to use the ores, while South Africa and New Zealand adopt the pre-reduction–electric furnace process, both of which can only recover the iron and vanadium [5,6]. In recent years, several metallurgy processes have been proposed to treat the vanadium–titanium magnetite concentrate, such as the reduction–magnetic separation and sodium roasting–reduction–electric furnace [7,8]. Obviously, the solid-state reduction is the key of all the technologies discussed above. The vanadium–titanium magnetite concentrate has a complicated structure, which contains minerals such as titanite magnetite, ilmenite, sulfur-bearing magnetic pyrite, gangue as well as other minerals; vanadium exists in a form of isomorphism in the titanium magnetite. Therefore, the reduction of vanadium–titanium magnetite concentrate is more complicated than other common ores, requiring higher temperature and longer reduction time to achieve a relatively high metallization ratio [9]. Hence, it is important to investigate the reduction kinetics and mechanism of the vanadium–titanium magnetite concentrate.

In recent decades, a large number of researches have been conducted to study the utilization and reduction of vanadium–titanium magnetite concentrate [10–15]. It was reported that ilmenite can be observed when ulvospinel was reduced at 500–800 °C, and the reduction of vanadium–titanium magnetite concentrate was mainly diffusion-controlled below 1050 °C, while reaction-controlled over 1050 °C [16,17]. The influencing factors of the reduction were investigated by many scholars, and the results suggested that the increase of reduction temperature, time and carbon to oxygen molar ratio could promote vanadium–titanium magnetite concentrate’s reduction, and they also found that the iron-joined crystal was formed, the size of which increased at first and then diminished with the increase of reduction time and carbon to oxygen molar ratio [18,19]. In addition, the intensification of the reduction
process was also studied by scholars, which demonstrated that the addition of metal powder and sodium salt could greatly reduce the reduction temperature [20,21].

Most of the aforementioned studies mainly aimed at the gas reduction kinetics and the influencing factors of vanadium–titanium magnetite concentrate reduction. However, few researches have been conducted on the solid-state reduction kinetics and mechanism. The purpose of this work is to investigate the solid-state kinetics of pre-oxidized vanadium–titanium magnetite concentrate reduction with coal. The phase and the microstructure of the reduction product were characterized by XRD, SEM and EDS, based on which the mechanism of the solid-state reduction was studied.

2 Experimental

2.1 Materials

The vanadium–titanium magnetite concentrate (73% of concentrate less than 74 μm) used in this study was obtained from Panzhihua Iron & Steel Company (Sichuan province, China). The main chemical compositions and phases are shown in Table 1 and Fig. 1, respectively. The oxidized samples (TFe 50.76%; FeO 15.03%) were prepared by oxidizing vanadium–titanium magnetite concentrate, and the specific operation is described in section 2.2.2. The reduction coal was a soft coal obtained from Inner Mongolia, China (Fixed carbon 57.07%, volatile matter 33.83%, ash 4.52%, size <1 mm). The pelleting binder was a multifunctional composite binder developed by Central South University [22].

Table 1 Chemical composition of vanadium–titanium magnetite concentrate (mass fraction, %)

<table>
<thead>
<tr>
<th>TFe</th>
<th>FeO</th>
<th>V_2O_5</th>
<th>TiO_2</th>
<th>SiO_2</th>
<th>Al_2O_3</th>
<th>Fe_3O_4</th>
</tr>
</thead>
<tbody>
<tr>
<td>51.36</td>
<td>28.21</td>
<td>0.66</td>
<td>9.98</td>
<td>4.12</td>
<td>3.03</td>
<td>3.03</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CaO</th>
<th>MgO</th>
<th>MnO</th>
<th>S</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.12</td>
<td>1.66</td>
<td>0.15</td>
<td>0.14</td>
<td>0.018</td>
</tr>
</tbody>
</table>

2.2 Procedure

2.2.1 Preparation of pellets

Vanadium–titanium magnetite concentrate was thoroughly mixed with 1% binder, and then pelletized to d10–15 mm in a disc pelletizer (d1000 mm). The wet pellets were dried at the temperature of (105±5) °C for 4 h.

2.2.2 Preparation of pre-oxidized

The pre-oxidation was carried out in a horizontal resistance furnace (d600 mm) with an automatic temperature control system. When the furnace temperature reached 900 °C, the corundum crucible filled with pellet samples was located into the furnace to oxidize for 6 min, and then removed, cooled to the room temperature in the air. The sample obtained would be used as the reduction sample.

2.2.3 Reduction

The reduction was carried out in a vertical resistance furnace (d600 mm) with an automatic temperature control system. 10 g of pre-oxidized pellets and 20 g of coal were loaded in a heat-resistant steel retort (d50 mm×100 mm). When the furnace was heated to the required temperature, the heat-resistant steel retort was put into the furnace. After a period of time, power supply was cut off and the reduction was stopped; the resort was take out quickly and the samples were poured into a container with nitrogen to cool down to the room temperature. The contents of total iron, ferrous and metallic iron were analyzed to calculate the reduction degree of iron. The microstructure and the phase of the reduction products were characterized by X-ray diffraction (Cu Kα radiation, λ=1.54056 Å, D/Max2200, Rigaku, Japan) and scanning electron microscope (SEM, Quanta200, FEI, Holland) equipped with energy dispersive analysis of X-ray (EDAX). Figure 2 illustrates the reduction experimental apparatus.

Fig. 1 XRD pattern of vanadium–titanium magnetite concentrate

The reduction degree of iron is calculated by the following formula [23]:
where \( R \) is the reduction degree of iron; \( w(TFe) \) is the total iron content in the reduced sample; \( w(Fe) \) is the metallic iron content in the reduced sample; \( w(FeO) \) is the ferrous content in the reduced sample; \( w(TFe0) \) is the total iron content in the pre-oxidized sample; \( w(FeO0) \) is the ferrous content in the pre-oxidized sample.

3 Results and discussion

3.1 Solid-state reduction kinetics

The isothermal reduction result of the pre-oxidized vanadium–titanium magnetite concentrate with coal reduced at 950–1100 °C is shown in Fig. 3, and \( R \) is the reduction degree of iron.

![Fig. 3 Isothermal reduction curves of pre-oxidized vanadium–titanium magnetite concentrate](image)

Figure 3 shows that the reduction degree of the iron is greatly affected by reduction temperature when the vanadium–titanium magnetite concentrates are reduced at relatively low temperature. However, the increase of the reduction degree slowed down when the degree was close to 95%.

The unreacted shrinking core model was used to analyze the results and a graphic of the relationship between the reduction degree and the reduction time \( t \) was made, namely \( 1-(1-R)^{1/3} \) vs \( t \) and \( 1-2R/3-(1-R)^{2/3} \) vs \( t \). At the test temperature, there is a better linear relationship between \( 1-(1-R)^{1/3} \) and \( t \). Figure 4 shows that the solid-state reduction of the pre-oxidized vanadium–titanium magnetite concentrate with coal at 950–1100 °C is controlled by interface chemical reaction. From the slopes of the lines in Fig. 4, the speed constant \( k \) under different temperatures and the apparent activation energy can be calculated by adopting the Arrhenius equation, namely,

\[
k = k_0 \exp\left(-\frac{E}{RT}\right)
\]

Taking the logarithm of both sides in the above equation yields:

\[
\ln k = \ln k_0 - \frac{E}{RT}
\]

where \( E \) is the activation energy; \( k_0 \) is the coefficient; \( k \) is the reaction rate constant; \( R \) is the molar gas constant, \( 8.314 \times 10^{-3} \); \( T \) is the temperature.

![Fig. 4 Relationship between 1-(1-R)^{1/3} and t at 950–1100 °C](image)

Figure 5 shows the Arrhenius curve of reaction rate constant \( k \) and temperature. As shown in Fig. 5, \( \ln k \) has a linear relationship with \( 1/T \). From the slope \( (-E/R) \), it can be calculated that the apparent activation energy of the solid-state reduction of the pre-oxidized vanadium–titanium magnetite concentrate is 67.719 kJ/mol.

![Fig. 5 Arrhenius curve of reaction rate constant k and temperature](image)

3.2 Phase transformation during reduction

Phase transformation during reduction was investigated to find out the reaction mechanism of the
solid-state reduction. Samples were reduced with coal at a given temperature for 60 min, and their XRD patterns are shown in Fig. 6.

![Fig. 6](image-url)

**Fig. 6 Phase transformation of reduction products**

As shown in Fig. 6, the main phases of the pre-oxidized vanadium–titanium magnetite concentrate before reduction are Fe₉TiO₁₅ and Fe₂O₃. It should be noted that Ca, Mg, Al and other impurities were not detected by XRD. This is probably because the concentrations of these impurities are below the detection limit of the XRD technique.

When the reduction temperature was 950 °C, the phases of Fe₂O₃ and Fe₉TiO₁₅ disappeared and the main phases of the reduction products were Fe and Fe₂TiO₄, suggesting that the following reactions occurred:

Fe₂O₃+C=2 FeO + CO                          (3)
FeO+C=Fe + CO                              (4)
Fe₉TiO₁₅+11C=7Fe+Fe₂TiO₄+ 11CO              (5)

When the reduction temperature was 1000 °C, the diffractive peaks of Fe₂TiO₄ weakened and that of FeTiO₃ appeared, suggesting that the following reaction occurred:

Fe₂TiO₄+C=Fe+FeTiO₃+ CO                    (6)

When the reduction temperature was 1050 °C, the diffractive peaks of Fe₂TiO₄ disappeared, the specific diffractive peaks of FeTiO₃ weakened and that of Fe₄Ti₅O₁₈ appeared, suggesting that the following reaction occurred:

2FeTiO₃+C=Fe+Fe₂Ti₅O₁₈+CO                  (7)

According to the results of the isothermal reduction in Fig. 3, the reduction degree of iron is 94.41% after the pre-oxidized vanadium–titanium magnetite concentrate was reduced at 1050 °C for 60 min. Assuming that Fe₂TiO₃ was the only unreduced iron-bearing and Ti-bearing mineral in the end, and TiO₂ grade did not change after reduction, the reduction degree of iron could be calculated theoretically according to the definition, i.e., the loss of oxygen in the reduction.

The total oxygen in iron oxide: \( w(O₂) = \frac{15.03 \times (16/72) + 39.78 \times (48/160)}{100\%} = 16.91\% \);
the remaining oxygen in iron oxide: \( w(O₂_{\text{remaining}}) = \frac{9.98 \times (79.88 \times 2)}{100\%} = 1\% \). So, \( R = \frac{(16.91 - 1)}{16.91} = 94.08\% \).

Obviously, the actual reduction degree is approximately equal to the theoretical reduction degree, which means that iron oxide is almost reduced to Fe₄Ti₅O₁₈. GUO [23] reported that Fe₂TiO₃ can be further reduced according to the following reaction:

\[(3-n)Fe₂Ti₅O₁₈ + 5(1-n)C = 3(1-n)Fe + 2(FeₙTi₁₋₅n)Ti₂O₅ + 5(1-n)CO, 0 < n < 1 \] (8)

For a lack of XRD characteristic spectral line data of \((FeₙTi₁₋₅n)Ti₂O₅\), this phase can not be found in the XRD detection results.

According to the earlier research [24], in the stage of generating float by magnetite iron reduction, the new phase of Fe₂TiO₄ was generated, which was further reduced to FeTiO₃, and finally vanadium–titanium magnetite concentrate is reduced into Fe and (Fe,Mg)Ti₂O₅. The reduction process discussed in this work is consistent with the results obtained by the earlier researchers.

### 3.3 Microstructure of reduction products

Figure 7 shows the SEM and EDS analyses of the reduction products, which were obtained by reducing the pre-oxidized samples at 1050 °C for 60 min. As shown in Fig. 7, most of the metallic iron is located in the edges or cracks of the particles. From the result of EDS analysis (Spot 2), it can be inferred that part of iron oxide in the pre-oxidized vanadium–titanium magnetite concentrate was not reduced and this part of iron oxide usually combined with Ti, Mg, Al, V, Mn, etc. The atomic number ratio suggested that these substances existed in the form of M₃O₅-type (M can be Fe, Ti, Mg, Mn etc).

To sum up, M₃O₅-type solid solution will be formed in the reduction of pre-oxidized vanadium–titanium magnetite concentrate. GUO et al [2] demonstrated that Fe₂TiO₃ solid solution would be stable in the presence of MgO and MnO, and the iron ionic activity would be reduced. The vanadium–titanium magnetite concentrate used in this work has higher MgO content (1.66%), in which the iron in the M₃O₅ solid solution is hard to be completely reduced. Additionally, it is easy for the Fe₂TiO₃ and the low-valence-state titanic oxide to be mutually soluble to the un-reacted ilmenite, which will reduce the iron ionic activity in the newly-formed ilmenite. Therefore, the poor reducibility of the iron in M₃O₅ solid solution is the main reason to limit the reduction property of pre-oxidized vanadium–titanium magnetite concentrate.
4 Conclusions

1) The kinetics study indicated that, with coal reducing at 950–1100 °C, the solid-state reduction of the pre-oxidized vanadium–titanium magnetite concentrate was controlled by interface chemical reaction and the apparent activation energy was 67.719 kJ/mol.

2) The analyses on phases transformation of the reduced products suggested that, with coal reducing at 950–1050 °C, its reduction process was: pre-oxidized vanadium–titanium magnetite concentrate $\rightarrow$ ulvospinel $\rightarrow$ ilmenite $\rightarrow$ FeTi$_2$O$_5$ $\rightarrow$ (Fe$_n$Ti$_{1-n}$)Ti$_2$O$_5$.

3) SEM and EDS analyses of the reduced products show that Mn$_3$O$_5$-type (M can be Fe, Ti, Mg, Mn, etc) solid solutions would be formed in the reduced products when the pre-oxidized vanadium–titanium magnetite concentrate was reduced at 1050 °C for 60 min, and the poor reducibility of iron in Mn$_3$O$_5$ solid solution is the main reason to limit the reduction property of pre-oxidized vanadium–titanium magnetite concentrate.

References


预氧化钒钛磁铁精矿的固态还原动力学及其机理

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摘 要: 研究预氧化钒钛磁铁精矿固态还原反应的动力学, 采用 XRD、SEM 和 EDS 等手段研究还原产物的显微结构和物相变化, 在此基础上, 对其固态还原机理进行研究, 结果表明: 以煤为还原剂, 在还原温度为 950~1100 ℃时, 预氧化钒钛磁铁精矿的固态还原受界面化学反应控制, 反应的表观活化能为 67.719 kJ/mol; 预氧化钒钛磁铁精矿的还原历程可描述为: 预氧化钒钛磁铁精矿→钒铁晶石→钒铁矿→亚铁板钛矿(FeTi2O5)→(Fe3Ti2)nTi2O7。预氧化钒钛磁铁精矿在 1050 ℃还原 60 min 后, 还原产物中会形成 FeO 固溶体; 前述反应过程中, 存在于 M2O3 固溶体中的铁的难还原性是限制预氧化钒钛磁铁精矿还原的主要原因。

关键词: 钒钛磁铁矿; 固态还原; 还原动力学; 还原过程

(Edited by Hua YANG)