The Function of Ca(OH)$_2$ and Na$_2$CO$_3$ as Additive on the Reduction of High-Phosphorus Oolitic Hematite-coal Mixed Pellets

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A process with coal-based direct reduction followed by magnetic separation is presented for recovering metallic iron from high-phosphorus oolitic hematite in this study. Ca(OH)$_2$ and Na$_2$CO$_3$ were used as additives in the reduction roasting. A Direct Reduction Iron (DRI) with 93.28 mass% Fe, and 0.07 mass% P can be obtained at a recovery percentage of 92.30 mass% under optimal conditions. The mechanisms of Ca(OH)$_2$ and Na$_2$CO$_3$ were investigated by XRD and SEM with EDS. It showed that fluorapatite was reduced to P smelt into metallic iron without additives while the hematite was reduced. The addition of Ca(OH)$_2$ can not only inhibit the reduction of fluorapatite but also promote the reduction of hematite. Na$_2$CO$_3$ can promote the separation of iron from slag, meanwhile it may also inhibit the reduction of fluorapatite at the presence of 15 mass% Ca(OH)$_2$. Under optimal conditions, phosphorus remained as fluorapatite in the slag and can be removed by grinding and magnetic separation.

KEY WORDS: high-phosphorus oolitic hematite; fluorapatite; Ca(OH)$_2$; Na$_2$CO$_3$; direct reduction roasting; metallic iron.

1. Introduction

The feed for iron and steel making must be upgraded through a series of physical separation processes including crushing, grinding and separation, because both blast furnace and direct reduction reactors cannot tolerate high volumes of slag. In addition, the content of harmful impurity such as phosphorus must be strictly controlled since too high phosphorus will bring about the welding problem of steel. However, there are many iron ores so fine-grained that it is impractical to grind them to a fine enough size to separate the iron oxides from the gangue, thus these iron ores are not utilized now. Along with the fast depletion of easy-to-process iron ores, exploitation of refractory iron ores becomes more and more important all over the world, especially in China. China imports hundreds of millions of tons of iron ore every year. At the same time, China has many iron ore deposits which are difficult to process. The Ningxiang type high-phosphorus oolitic hematite ore, which typically contains 30 mass% to 45 mass% Fe and 0.4 mass% ~ 1.1 mass% P, is one of the refractory iron ores. Since the unique internal structure of the ore that the iron oxides, fluorapatite and chamosite are so intimately intermixed that it is not practical to obtain qualified iron concentrate by conventional mineral processing methods.

Recent research shows that, it is feasible to recover metallic iron from refractory ore and other iron-containing materials by coal-based direct reduction - magnetic separation process. Some researches have been reported on coal-based direct reduction of high-phosphorus oolitic hematite. It was reported that the Direct Reduction Iron (DRI), assaying more than 90 mass% Fe, less than 0.1 mass% P with an iron recovery of about 80 mass%, can be obtained at the optimum conditions. These researches were conducted below 1 150°C, because higher temperature will lead to the P content of DRI increasing. The cause for this is that the fluorapatite in the raw ore will be reduced to elemental P which can be smelt into metal iron. However, because the reduction of hematite was not complete under low temperatures, iron lost into the tailing in the form of fayalite and hercynite. In order to promote the recovery of iron, 20 mass% ~ 30 mass% additives, sodium salt or calcium salt, must be added in the reduction roasting. Ca(OH)$_2$ and Na$_2$CO$_3$ are used as additives in the direct reduction process to increase the recovery of iron, their effects on promoting the reduction of iron oxides and improving the separation of iron from slag have been proved. However, their action mechanisms for dephosphorization are not completely clarified, especially at high temperature which fluorapatite can be reduced by coal.

In this study, the direct reduction of high-phosphorus oolitic hematite-coal composite pellets were conducted at 1 200°C, Ca(OH)$_2$ is used alone or in combination with Na$_2$CO$_3$ as additives in the process and their action mechanisms of dephosphorization were studied.

2. Experimental

2.1. Raw Materials

The high phosphorus iron ore sample used in this work is...
a typical Ningxiang high phosphorus oolitic hematite from Hubei Province, China. The results of chemical analysis of raw ore show that the iron content of raw ore is 43.58 mass%, the content of phosphorus is 0.83 mass%, and the contents of acid gangue including SiO₂ and Al₂O₃ are 17.10 mass% and 9.28 mass% respectively. The X-Ray Diffraction (XRD) analysis of the raw iron found that iron mainly occurs in the form of hematite, with impurities including quartz, chlorite and fluorapatite, as shown in Fig. 1. The photomicrograph of raw ore is shown in Fig. 2. It can be seen clearly from Fig. 2 that hematite together with chlorite and fluorapatite builds the concentric shell of oolites, the iron oxides and gangue minerals are intimately intergrowth.

The coal used as reductant in this study was obtained from Ningxia Province in China. The results of industrial analysis (air dry) of the coal were 11.77 mass% moisture, 19.90 mass% ash, 28.18 mass% volatiles, 51.92 mass% fixed carbon. The raw ore and coal were crushed to 100 mass% passing 1 mm. Ca(OH)₂ and Na₂CO₃ used as additives in the experiments were analytical reagent (AR) grade.

2.2. Experimental Procedure

The pellets were produced by the following procedures: first, 20 g iron ore, a certain amount of coal, additives and water were mixed together, and then the mixture was pressed to form pellets using an die with a size of 30 mm diameter.

The reduction roasting was carried out in a muffle furnace with a temperature control programmer. Pellets were put in a graphite crucible of 600 mm in diameter and 800 mm in height. The graphite crucible was put into the furnace and held for scheduled time when the temperature arrived at 1200°C. After that, the roasted pellets were taken out of the furnace and cooled to room temperature under ambient temperature.

Once the pellets in the crucible had cooled, the roasted pellets were crushed to ~2 mm and then ground to 64.80 mass% – 0.074 mm using wet grinding. The XCGS-73 low intensity magnetic separator with a magnetic field intensity of 1120 Oe was used to recover metallic iron from roasted pellets. The iron concentrates obtained from the first separation were ground and magnetic separated once again with the grinding fineness 76.95 mass% – 0.044 mm to produce high iron content DRI. The main evaluation index of test results were the iron content, P-content and the iron recovery of DRI. The iron recovery refers to the percentage of the total iron metal contained in the ore that is recovered into the DRI.

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\text{Iron recovery} = \frac{\text{Total weight of DRI} \times \text{iron content of DRI}}{\text{Total weight of raw iron} \times \text{iron content of raw iron}} \times 100\%
\]

2.3. Analysis and Characterization

The chemical analyses were conducted by China University of Geosciences (Beijing) analysis laboratory. Scanning Electron Microscope with Energy Dispersive Spectrum (Carl Zeiss EVO18) analyses were carried out on samples mounted in epoxy resin and polished. Phases present in the samples were identified by Rigaku DMAX-RB X-ray Diffract Meter (Cu target).

3. Results and Discussion

3.1. The Effect of Ca(OH)₂

The effect of Ca(OH)₂ dosage on the reduction of high-phosphorus oolitic hematite was investigated. The experiments were conducted at a condition of temperature 1200°C, roasted time 40 min and coal dosage 25 mass%. The dosages of coal and additives are expressed by a percentage that refers to their mass ratio to the raw ore. The experimental results are shown in Fig. 3.

The results in Fig. 3 show that (1) the content of phosphorus in the DRI decreases with increasing dosage of Ca(OH)₂. A DRI with 0.30 mass% phosphorus can be obtained without Ca(OH)₂. When the dosage of Ca(OH)₂ increases to 15 mass%, the content of phosphorus decreases to 0.14 mass%, and then the dephosphorization effect maintained stable; (2) The recovery of iron increases with increasing dosage of Ca(OH)₂, when Ca(OH)₂ dosage increases from 0 mass% to 15 mass% the iron recovery increases from 86.58 mass% to 92.13 mass%. But, when Ca(OH)₂ dosage further increases, the iron recovery maintain stable; (3) The addition of Ca(OH)₂ is not conducive to the grade of the DRI, especially when Ca(OH)₂ dosage is more than 10 mass%, the iron content falls sharply with the increase of the content of Ca(OH)₂.

So it can be concluded that Ca(OH)₂ dosage has a great...
influence on the reduction of hematite. An appropriate Ca(OH)₂ dosage can improve the recovery of iron and decrease the content of phosphorus in the DRI. Excessive Ca(OH)₂ fails to further increase the recovery of iron and decrease the phosphorus content of DRI, it can leads to reducing the iron content sharply. The optimal dosage of Ca(OH)₂ is 15 mass%.

3.2. The Effect of Na₂CO₃

The tests above proved that although the addition of Ca(OH)₂ can decrease the phosphorus content of DRI, the phosphorus content in the iron products is still 0.14 mass% even in the optimal Ca(OH)₂ dosage condition. On this basis, in order to obtained DRI with less than 0.1 mass% phosphorus, different dosages of Na₂CO₃ were added together with the same Ca(OH)₂ dosage of 15 mass% in the reduction roasting, the other experimental conditions (temperature, roasted time, coal dosage etc.) are same as the previous experiments. The result is presented in Fig. 4.

Figure 4 shows that the content of phosphorus in the DRI decreases from 0.14 mass% to 0.07 mass% and the iron content of the DRI increases from 91.14 mass% to 93.28 mass% with increasing dosage of Na₂CO₃ from 0 mass% to 3 mass%. In addition, the addition of Na₂CO₃ has no significant effect on the recovery of iron in this condition. A DRI with 0.07 mass% of phosphorus is obtained by adding 3 mass% Na₂CO₃ and 15 mass% Ca(OH)₂, while the grade of iron is 93.28 mass% with a recovery of 92.30 mass%. The result is superior to the results of previous reports.⁸⁻¹⁰

4. Characteristics of Roasted Pellets

In order to investigate the phase translation and the morphological changes of iron-bearing minerals and fluorapatite at different roasting conditions. SEM with EDS and XRD were used.

4.1. Without Any Additive

Figure 5(a) shows the XRD patterns of roasted pellet roasted at 1200°C without any additives. It can be seen from Fig. 5(a) that hematite is reduced to metallic iron and the diffraction peak of fluorapatite disappears, in addition, some anorthite and unreacted quartz were observed. This shows that when additive was not at presence, the reduction of fluorapatite can occur at temperature of 1200°C.

The chemical reaction of the reduction of phosphate can be summarized as

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\text{Ca}_{10}(\text{PO}_4)_6\text{F}_2 + 15\text{C} + 9\text{zSiO}_2 \rightarrow 3\text{P} + 15\text{CO} + 9\text{[CaO(SiO}_2\text{z}] + CaF}_2
\]

It is believed that the reaction occurs at above 1100°C and the limiting step appears to be diffusion of phosphorus-bearing species to a carbon particle where reaction occurs at the range from 1100°C to 1250°C. Silica plays an important role in promoting the reaction by providing a thermodynamic driving force and modifying the melting phenomena.¹²⁻¹⁵

Figure 6 shows SEM image and EDS of the roasted pellet without any additives. As shown in Fig. 6(a), hematite was reduced to metallic iron and metallic iron particles gathered so that the oolitic structure was completely destroyed. Figures 6(b) and 6(c) show the dispersion of Fe and P in the observed area, respectively. It can be seen that the majority of Fe overlaps with P. Moreover, the result of point analysis of the metallic iron (Point 1) shows that it contains a high content of P, which could be explained that some fluorapatite was reduced to P and the P smelt into metallic iron. Since the P in metallic iron cannot be removed by physical separation, DRI obtained by magnetic separation contains high phosphorus content.
4.2. The Roasted Pellet with 15 mass% Ca(OH)$_2$

X-ray diffraction pattern of the roasted pellet with 15 mass% Ca(OH)$_2$ is presented in Fig. 5(b), which shows that (1) the diffraction peak of metallic iron increases markedly with 15 mass% Ca(OH)$_2$ compared with the roasted pellets without additives (Fig. 5(a)). This is due to the fact that CaO, dissociated from Ca(OH)$_2$, can replace the FeO from fayalite, which increases the reducing reaction activity of FeO, and as a result, the reduction of iron oxide can be promoted.$^{16}$ (2) The diffraction peak of fluorapatite still exists, which proves that the reduction of fluorapatite is hindered by Ca(OH)$_2$; (3) The diffraction peak of quartz reduces significantly and gehlenite appears with 15 mass% Ca(OH)$_2$, this is attributed to that the CaO dissociated from Ca(OH)$_2$ reacts with quartz and Al-bearing mineral in the raw ore to form gehlenite. The reaction leads to the reduction of quartz content in the roasted pellets. On account of that quartz can promote the reduction of fluorapatite, the decrease of quartz content may not benefit for the reduction of fluorapatite.

Figures 7 and 8 show the SEM image and EDS results from a roasted pellet with 15 mass% Ca(OH)$_2$. It can be seen from Fig. 7(a), the melting phenomena is slighter and the the size of iron particles formed are smaller compared to that without Ca(OH)$_2$ (Fig. 6(a)), and oolitic structure in the raw
ore was not completely destroyed. It is because that Ca(OH)$_2$ decomposes into CaO and H$_2$O at high temperature, since CaO can promote the reduction of iron oxide, FeO content in the slag will reduce and less low melting point materials will form which in turn improve the melting point and viscosity of the slag. As a result, the diffusion of mass in the slag is limited, and thereby the growth and aggregation of metallic iron is hindered in the reduction roasting process. The liberation of iron particles from slag thus hindered by adding Ca(OH)$_2$, which may be the main reason why the iron content of DRI decrease with increasing the dosage of Ca(OH)$_2$.

The EDS mappings of Fe and P in the observed area are shown in Figs. 7(b) and 7(c), respectively. It is observed that the majority of Fe is not overlaps with P content, phosphorus is mainly concentrated in the oolite almost the same with that in the raw ore. In addition, the result of point analysis of the metallic iron (Point 2) shows that it does not contain P. Combined with the result of XRD pattern, it can be concluded that the reduction of fluorapatite is inhibited by adding Ca(OH)$_2$ which hinders the diffusion of phosphorus-bearing species to a carbon particle where reaction occurs. What’s more, the reduction of quartz content roasted pellet may be not favorable to the reduction of fluorapatite. So less P will generate and smelt into metallic iron. Thus the content of phosphorus in DRI decreases with increasing the dosage of Ca(OH)$_2$.

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The XRD pattern of roasted pellet with 15 mass% Ca(OH)$_2$ and 3 mass% Na$_2$CO$_3$ shows that the diffraction peak of quartz disappears, as shown in Fig. 5(c), this shows that most of the quartz take part in reaction in the present of Na$_2$CO$_3$. Meanwhile, the diffraction peak of fluorapatite still exists.

Figures 9 and 10 show SEM images and EDS results of roasted pellet with 15 mass% Ca(OH)$_2$ and 3 mass% Na$_2$CO$_3$. It can be seen from Fig. 9(a) that the size of iron particle is coarser than that in the roasted pellet without Na$_2$CO$_3$ (Fig. 7(a)), and the oolitic structure was destroyed. This is due to the fact that Na$_2$CO$_3$ reacts with SiO$_2$ to form some low melting point materials, these low melting point materials appear in liquid phase in the course of reduction which is beneficial for the metallic iron to diffuse and aggregate.17) The separation of iron particles from slag is thus improved by adding Na$_2$CO$_3$ and thereby the content of phosphorus in the DRI decreases and the iron content increases.

Figures 9(b) and 9(c) show that the majority of Fe is not overlaps with P in the observed area. The result of point analysis of the metallic iron particle shows that P has not been detected even in the metallic iron (Point 6) which close to the fluorapatite, as shown in Figures 10(a)–10(c). So, it can be inferred that the addition of Na$_2$CO$_3$ restrains the reduction of fluorapatite further at the presence of 15 mass% Ca(OH)$_2$. This may be explained from two aspects. On the one hand, Na$_2$CO$_3$ reacts with quartz. On the other hand, the presence of Na$_2$CO$_3$ leads to the generation of liquid phase which accelerate the quartz reacting with CaO and other components in raw ore. As the depletion of quartz, the reduction of fluorapatite will be restricted.

In summary, the presence of Na$_2$CO$_3$ not only creates a favorable condition for the liberation of metallic iron from the fluorapatite but also restrains the reduction of fluorapatite in the DRI, thereby increasing the iron content and decreasing the phosphorus content.
slag but also inhibits the reduction of fluorapatite at the presence of 15 mass% Ca(OH)$_2$. Under the optimal conditions, phosphorus remained as fluorapatite in the slag and can be removed by ground and magnetic separation.

4. Conclusions

From the present investigation, the following conclusions can be drawn:

(1) Without any additive and at the temperature of 1200°C, fraction of fluorapatite in the raw ore is reduced to P by carbon and the phosphorus entering into metallic iron particle results in the high phosphorus content of DRI.

(2) A DRI with 93.28 mass% Fe, and 0.07 mass% P can be obtained at a recovery of 92.30 mass% by adding 15 mass% Ca(OH)$_2$ and 3 mass% Na$_2$CO$_3$ to the raw ore.

(3) The presence of Ca(OH)$_2$ restrains the reduction of fluorapatite and promotes the reduction of hematite. The mechanism of Ca(OH)$_2$: restraining the reduction of fluorapatite can be explained from two aspects. On the one hand Ca(OH)$_2$: increases the melting point and viscosity of slag which in turn deteriorates the conditions of solid-solid reaction in the slag and thereby restrains the reduction of fluorapatite. On the other hand the addition of Ca(OH)$_2$: leads to

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Fig. 9. SEM images and EDS results of roasted pellet with 15 mass% Ca(OH)$_2$ and 3 mass% Na$_2$CO$_3$.

Fig. 10. SEM image and EDS results of roasted pellet with 15 mass% Ca(OH)$_2$ and 3 mass% Na$_2$CO$_3$. 
the consumption of quartz and thus restrains the reduction of fluorapatite.

(4) At the presence of 15 mass% Ca(OH)$_2$, the Na$_2$CO$_3$ exerts a positive effect for iron particle growth and aggregation which improve the conditions of separation of metallic iron and slag. At the same time, that Na$_2$CO$_3$ accelerates the consumption of quartz, which may restrain the reduction of fluorapatite.

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