RECOVERY OF IRON AND PHOSPHORUS REMOVAL FROM GARA DJEBILET IRON ORE (ALGERIA)

Introduction. The metallic iron recovery from refractory iron ores, as a substitute to easy-to-process iron ore resources, is a major global issue to deal with the fast increase in need for iron and relieve the threat of raw material shortage in steelmaking industry [1]. Nowadays, the extraction of metallic iron via the BF remains the most dominant route in steelmaking industry. However, it is neither cost-effective nor feasible for high-phosphorus oolitic iron ores as almost elemental P is absorbed by hot liquid metallic iron owing to the high temperature of the procedure [2]. Phosphorus is a detrimental impurity in ferrous metallurgy, since it increases the brittleness and decreases the ductility and low-temperature toughness of steel products [3]. Several bench-scale tests have been conducted on promoting the beneficiation of high-phosphorus iron ores across the world. Available routes include hydrometallurgical and pyrometallurgical processes [4]. Froth flotation, gravity separation, magnetic separation, chemical leaching, and bioleaching are the beneficiation of high-phosphorus oolitic iron ores as almost elemental P is absorbed by hot liquid metallic iron owing to the high temperature of the procedure [2]. Phosphorus is a detrimental impurity in ferrous metallurgy, since it increases the brittleness and decreases the ductility and low-temperature toughness of steel products [3]. Several bench-scale tests have been conducted on promoting the beneficiation of high-phosphorus iron ores across the world. Available routes include hydrometallurgical and pyrometallurgical processes [4]. Froth flotation, gravity separation, magnetic separation, chemical leaching, and bioleaching are the main hydrometallurgical processes. The pyrometallurgical processes such as magnetisation roasting, microwave heating, gas-based reduction, and coal-based DR are promising routes for the production of DRI. The direct reduction (DR) prior to the electric arc furnace (EAF) is an alternative route to the standard BF process. It has been proven satisfactory for metallic iron recovery from fines, pellets, sinter and lump by converting them into DRI using several materials as reductant at reduction temperatures below the melting point of iron (Fe) [5]. This study focuses on the recovery of metallic iron and the removal of phosphorus from mixed pellets sourced from Gara Djebilet deposit via the procedure of coal-based DR followed by WLIMS. The effects of temperature, time periods and sodium sulfate dosage on phosphorus removal, metallisation degree and iron recovery rate were tried and optimised. Furthermore, phase changes in iron oxides and the distributing features of phosphorus in both reduced and magnetic materials were investigated as well.

Findings. The appropriate addition of sodium sulfate improves the Fe-P separation during the coal-based DR of Gara Djebilet mixed pellets.

Originality. Using additives of CaO and sodium sulfate during the coal-based DR-magnetic separation of mixed pellets sourced from Gara Djebilet deposit.

Practical value. The results reveal that a final direct reduced powder (DRI) assaying 96 wt% Fe and 0.16 wt% P at a recovery rate of 97.72% was obtained when the ore-coal-CaO mixed pellets were reduced in the presence of 5 wt% Na2SO4 at 1250 °C for 30 min. Thus, the coal-based DR could be used as an alternative to the blast furnace (BF) route in the steelmaking industry from refractory iron ores.

Keywords: Gara Djebilet, high-phosphorus oolitic iron ore, coal-based direct reduction, sodium sulfate, magnetic separation

Purpose. This research aims to promote the assay of iron and reduce the phosphorus grade of the final DRI.

Methodology. A high-phosphorus oolitic iron ore from Gara Djebilet deposit underwent the procedure of coal-based direct reduction (coal-based DR) followed by wet low-intensity magnetic separation (WLIMS). The effects of temperature, periods of time and Na2SO4 dosage on phosphorus removal, metallisation degree and iron recovery rate were tried and optimised. Furthermore, phase changes in iron oxides and the distributing features of phosphorus in both reduced and magnetic materials were investigated as well.

Materials. The ironstone samples used in this research were gathered from the Gara Djebilet deposit in Tindouf province of Algeria. The chemical and mineralogical compositions of the raw ore sample are presented in Table 1 and Fig. 1 respectively.

Practical value. The results show that the total iron (TFe) grade is up to 53.2 wt% and exists in the form of hematite, magnetite, goethite and maghemite. However, the assay of phosphorus amounts to 0.8 wt% and occurred as fluorapatite. SiO2 exists as quartz and Al2O3 distributes in chamosite. Fig. 2 presents photomicrographs of the raw ore observed under the ore microscope (Nikon Eclipse LV100POL) and the cathodoluminescence microscopy (CITEL MK5-2 CLM). It is clear from Fig. 2 that the oolitic texture is made of ooids having the shape of ovals ranging from 150 µm to 800 µm in size. The non-detrital facies (FOND) of non-magnetite ore (Fig. 2, a) and non-magnetitic ore (Fig. 2, b) correspond to scattered ooids in a quartz-poor groundmass. In the cemented facies (FOC) of non-magnetic ore (Fig. 2, c), the chamositic spastoliths and ooids with chamositic nucleus and swapping hematitic-chamositic layers are bounded together by a hematitic matrix. Apatite exists as prismatic crystals in both ooids and groundmass. The groundmass apatite is light-green luminescing, while the detrital apatite in grains and oolite laminae exhibits bright-yellow green emission (Fig. 2, d).

Methods. The raw ore and coal were crushed and screened separately to 100 % less than 1 mm by means of a laboratory jaw crusher. Then, iron ore samples were blended with coal and CaO according to particular proportions by rigorously stirring during 30 minutes. The quantity of excess coal in the mixture was determined according to C/O molar ration of 2.

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F. A. Mansour1, orcid.org/0000-0002-6694-2725, M. Ould-Hamou1, orcid.org/0000-0002-8770-5323, A. Merchichi1, orcid.org/0000-0001-8136-601X, O. Güven2, orcid.org/0000-0002-5267-9153

1 – Laboratoire de Génie Minier, Département Génie Minier, École Nationale Polytechnique, El Harrach Algiers, Algeria, e-mail: farid_aghilasse.mansour@g.enp.edu.dz
2 – Department of Mining Engineering, Faculty of Engineering, Adana AlparslanTürk Science and Technology University, Sarıçam Adana, Turkey

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1 – Laboratoire de Génie Minier, Département Génie Minier, École Nationale Polytechnique, El Harrach Algiers, Algeria, e-mail: farid_aghilasse.mansour@g.enp.edu.dz

2 – Department of Mining Engineering, Faculty of Engineering, Adana Alparslan Türk Science and Technology University, Sarıçam Adana, Turkey
Table 1

Chemical composition of raw iron ore sample (wt%)

| Component | Value |
|-----------|-------|
| TFe       | 53.20 |
| SiO₂      | 0.719 |
| Al₂O₃     | 0.596 |
| Fe₃O₄     | 63.17 |
| FeO       | 11.55 |
| CaO       | 02.36 |
| MgO       | 0.42  |
| MnO       | 0.23  |
| TiO₂      | 0.13  |
| Loss on ignition (LOL) | 7.15 |

Table 2

Results of proximate analysis of coal

| Component | Value |
|-----------|-------|
| TC        | 69.06 |
| FC₆₃      | 49.54 |
| V₆₃       | 45.28 |
| A₆₃       | 6.18  |
| M₆₃       | 14.95 |
| S         | 0.66  |

Fig. 1. X-ray diffraction pattern of Gara Djebilet oolitic iron ore sample

Fig. 2. Photomicrograph collages of ironstone samples:

- CH – chamosite; GO – goethite; HM – hematite; MH – maghemite; MG – magnetite

Results and discussions.

The effect of reduction temperature.

The prepared pellets were sintered at temperature values of 1000, 1100, 1200 and 1300 °C respectively. While the reducing period was fixed at 30 min and no sodium sulfate was added to the mixtures. The data revealed that, the temperature varied from 1000 to 1300 °C, η, $\beta_{Fe}$ increased sharply from 74.52 to 92.36 % and from 73.23 to 84.81 wt% respectively. Moreover, ε exhibited a significant improvement from 86.35 to 94.36 % (Fig. 3, a).

This is due to the fact that increasing further the reduction temperature improves the reduction of iron oxides, destroys the oolitic structure and promotes the iron grains growth [6]. However, there was a dramatic increase in $\beta_{Fe}$ in the whole (i.e. the molar ratio of fixed carbon in the coal to reducible oxygen in the iron oxides was 2). CaO was added to attain the predetermined binary basicity of 1 (i.e. the mass ratio of CaO to SiO₂ was 1). Each run requires 20 g iron ore sample, 13 g coal and 1.6 g CaO. An appropriate amount of water (12 wt%) was added to the prepared mixtures and subsequently pressed into cylindrical pellets using a steel die with a size of 30 mm diameter and a thickness of 8 mm with the aid of a hydraulic press, and thereafter dried in drying oven at 105 °C for 4 hours. The reduction trials were performed in a high-temperature electric resistance muffle furnace (MoS-B 170/8 Protherm Furnace). First, the pellets were charged into heat-resistance graphite crucibles, and a layer of coal (about 1 mm thick and 2 grams weight) was spread over the top surface of the pellets to assure the reducing atmosphere within the crucibles. Then, the mixtures were reduced under the designated experimental conditions in the furnace which had already been heated to the various desired reduction temperatures. Meanwhile, purge gases were not utilized during the roasting and the tests were run at atmospheric conditions. After reduction, the charged crucibles were withdrawn from the furnace and cooled down to the room temperature under an air-isolated environment. Once the mixtures in the crucibles cooled, a part of reduced mixed pellets were polished for SEM-EDS analyses, and the remainder were milled to 80 % less than 74 μm using a vibratory disc mill.

Then, a random sample weighing 20 g deducted from each of the milled products and slurried to 20 wt% solids by weight, was divided into concentrate and tailing fractions through a Davis magnetic tube (KHD Humboldt Wedag International AG, Germany) at a magnetic field intensity of 110 kAm⁻¹; after that, the magnetic products were filtered and dried in a vacuum drying oven at 105 ± 5 °C for 24 hours. The evaluation indices of test results were assessed by the iron metallisation degree η (%) and the iron recovery rate ε (%) of DRI powder. η is calculated according to the following formula

$$\eta = \frac{\delta_{Fe}}{a_{Fe}} \times 100$$

where $\delta_{Fe}$ is the metallic iron content of the reduced material (wt%); $\alpha_{Fe}$ is the total iron content of reduced material (wt%).

Whereas, ε is calculated according to the following formula

$$\varepsilon = \frac{m_{Fe}}{m_{Fe}} \times 100$$

where $m_1$, $m_2$ are the dry weight of feed and concentrate materials respectively (g); $\beta_{Fe}$ is the total iron content of magnetic material (wt%).

The P content and Fe grade of both reduced and magnetic products were assayed using chemical analyses.

The mineralogical phases present in both reduced and magnetic materials (under selected roasting conditions) were identified by powder X-ray diffraction (XRD). The morphology and microstructure of reduced samples were observed using scanning electron microscopes equipped with an X-ray energy dispersive spectrometer. Micrographs were taken and point EDS analyses were used to recognise local compositions.
range of tested reduction temperatures to peak 1.23 wt% at 1300 °C (Fig. 3, b). This increase could be attributed to the reduction of fluorapatite to generate gaseous phosphorus (P$_2$) which, due to its high reactivity, melted into metallic iron rather than emitting to the atmosphere.

The effect of reduction time. A set of experiments were conducted at reduction periods of 15, 30, 45 and 60 min. The reducing temperature was set at 1250 °C and the experiments were run in the absence of sodium sulfate. Fig. 4 indicates that there was a marked rise in $\varepsilon$ and $b_{Fe}$ to peak 96.21 % and 88.88 wt% respectively at 30 min, but then fell significantly over the next periods of time to approximately 92 % and 81 wt% respectively.

Likewise, $\eta$ dropped sharply as the reduction periods exceed 30 min from 94.15 to 87.17 %. In effect, an additional extension in the periods of reduction can lead to a weakening of the reducing atmosphere due to the exhaustion of the reductant at this high temperature. Thus, the engendered metallic iron would be subjected to re-oxidation, leading to a decrease in both $\eta$ and $\varepsilon$ [6]. In addition, a further increase in reduction time, results in a dramatic rise in $b_P$, and after the maximum reduction period (60 min), it was as high as 1.43 wt%. This is due to the fact that more fluorapatite in the raw ore is reduced to P, which migrates into the metallic phase as the reduction period is extended.

The effect of sodium sulfate dosage. The prepared mixtures were mixed thoroughly with a proper amount of water (12 wt%) in the presence of various quantities of sodium sulfate.

Then, the mixtures were pressed into pellets and reduced at 1250 °C for 30 min, after having been dried in a vacuum oven at 105 °C for 4 hours. The results are presented in Fig. 5. The data show that both $\eta$ and $\varepsilon$ first increased sharply from 95.72 and 95.11 % to 98.35 and 97.72 % respectively as the Na$_2$SO$_4$ dosage varied from 2.5 to 5 wt%, then declined obviously when the Na$_2$SO$_4$ dosage increased further up to 10 wt%. Meanwhile, a marked rise is observed in $b_{Fe}$ to peak 96 at 5 wt% Na$_2$SO$_4$. However, a further increase in the Na$_2$SO$_4$ dosage results in a slight decrease in $b_{Fe}$ to reach 90.04 wt%. Indeed, owing to the increase in Na$_2$SO$_4$ dosage, more FeS would be produced during the reduction process leading to a diminution
in $\beta_{f}$ [6]. Conversely, increasing the Na$_2$SO$_4$ dosage from 2.5 to 5 wt%, results in a sharp decrease in $\beta_{f}$ from 0.18 to 0.16 wt%.

However, a slight increase is observed when the Na$_2$SO$_4$ dosage is beyond 5 wt% (Fig. 5, b).

**Phase constitution of reduced pellets.** In order to understand the aforementioned experimental results, selected reduced pellets were subjected to mineralogical analyses, and their XRD pattern is shown in Fig. 6. The results reveal that the peaks associated with Fe$_3$O$_4$ vanished completely due to its reduction to metallic iron stepwisely in the order of Fe$_3$O$_4$ → Fe$_2$O$_3$ → Fe → FeO → Fe through gaseous intermediate of CO and CO$_2$ resulting in fixed carbon gasification. In the absence of sodium sulfate, the SiO$_2$ diffractions became less intense, and those of chamosite fade away. Meanwhile, the peaks associated with anorthite appeared after the reduction process, and became more intense with increasing the temperature (Fig. 6, c). This is attributed to the fact that a reaction between Ca$_2$(PO$_4$)$_3$F, SiO$_2$, CaO and chamosite took place during the reduction process. Moreover, iron appeared most often in the form of metallic state which is easily recovered by WLIMS.

However, diffraction peaks associated with intermediate products like wustite and minor iron minerals such as fayalite (Fe$_5$SiO$_4$) and hercynite (FeAl$_2$O$_4$) appeared in the XRD pattern of reduced samples, suggesting that some FeO take part in reaction with SiO$_2$ and chamosite during the reduction process, despite the consumption of some amount of these minerals in advance by CaO. Therefore, $\eta$ is relatively low, and the reaction activity of FeO and thereby, $\eta$ gets obviously enhanced. Although most of iron in the reduced powder exists as metallic iron, but some non-magnetic iron minerals are also present in the reduced powder like wustite (FeO), as well as a small amount of troilite (FeS) which is highly improbable to transform into Fe or FeO in the CO reducing atmosphere, since FeS could likewise be formed from FeO [7]. Furthermore, the unreacted fluorapatite turns into tailings along with non-magnetic materials. Therefore, $\epsilon$ and the Fe-P separation get obviously improved.

**Morphology and microstructure features of the reduced pellets.** A sample reduced at 1250 °C for 60 min was investigated by SEM-EDS analysis (SEM-EDS, JEOL JSM-5410, Japan). The SEM image and the results of EDS measurements at several spots of both metallic and slag phases from the selected sample are presented in Fig. 7. After reduction, the iron oxides were reduced to metallic iron particles which clump together causing the destruction of the entire oolitic structure (Fig. 7, a). In the absence of sodium sulfate, the metallic iron particles had not combined to each other completely, and connected yet with gangue minerals. The results of point analysis show that Si, Ca, Al, O are the main elements of spot 1 from the grey slag matrix phase and it may be (Mg, Fe)$_2$SO$_4$ and CaAl$_2$SO$_4$. Meanwhile, the Fe content was higher than 13 wt%, suggesting that many ultrafine iron crystallites had not grown and still associated closely with gangue minerals. Besides, the P content at point 2 amounted to 2.03 wt%, which indicates that most of fluorapatite in the mixture was reduced and the new-born phosphorus melted into the metallic iron grains to generate phosphorus iron FeP. This is due to the fact that, at these roasting conditions, more FeO will react with gangue minerals to produce low-melting point materials due to the poor reducing atmosphere supplied (the wide band proving the presence of glass phase materials was spotted between 10 and 40° in the XRD pattern provided in Fig. 6, c) [8]. Since the existence of liquid phase is contributing to the spread of phosphorus-bearing particles to those of carbon, the reduction rate of fluorapatite (Ca$_5$(PO$_4$)$_3$F) is improved and thereby more elemental P melted into metallic iron [9]. As the physical beneficization ways are unable to remove the elemental P in metallic iron, high-phosphorus content DRI is acquired by WLIMS. The sample produced under optimal reduction parameters (reduced at 1250 °C for 30 min with adding 3 wt% Na$_2$SO$_4$) was investigated by SEM-EDS analysis (SEM-EDS, JEOL JSM-7610FPlus, Japan). The SEM photomicrograph and the outcomes of EDS measurements at diverse spots of both metallic and slag phases from the selected sample are presented in Fig. 8. As revealed in Fig. 8, a, most metallic iron grains are connected, presenting a net-like microstructure, and some small gangue particles (dark grey) are enveloped in the mesh. By adding sodium sulfate, the metallic iron grains (in bright white) aggregated together and grew in sizes mostly over 75 μm. Furthermore, the metallic phase was apart from the gangue phase with clear boundaries, which improves the liberation of metallic iron from slag matrix and thereby, a DRI with high metallic iron grade is acquired.

Fig. 8, c reports that Si, Ca, Na, Al, O, P are the main elements of spot 1 from the slag phase and it may be anorthite (Ca$_2$Al$_2$SiO$_6$), nepheline (NaAlSiO$_4$), CaNaPO$_4$ and the unreduced part of fluorapatite. Meanwhile, there was found less amount of Fe, suggesting that iron grains and gangue minerals were less intergrown. Therefore, it is easy to attain a clean separation of iron from gangue by subsequent milling and WLIMS. The results of point analysis of spot 2 (Fig. 8, d) show that elemental phosphorus has not been spotted in the metallic iron phase (in bright white). This is because the addition of Al$_2$O$_3$, already consumed by CaO and Na$_2$SO$_4$, despite that the reduction temperature is over 1313 K [7]. In addition, Na$_2$SO$_4$ had reacted with SiO$_2$ and Al-bearing minerals in the raw ore, to generate NaAlSiO$_4$ and anorthite (CaAl$_2$SiO$_4$), in preference to wustite (FeO), leading to a marked rise in the reducing reaction activity of FeO and thereby, $\eta$ gets obviously enhanced. Although most of iron in the reduced powder exists as metallic iron, but some non-magnetic iron minerals are also present in the reduced powder like wustite (FeO), as well as a small amount of troilite (FeS) which is highly improbable to transform into Fe or FeO in the CO reducing atmosphere, since FeS could likewise be formed from FeO [7]. Furthermore, the unreacted fluorapatite turns into tailings along with non-magnetic materials. Therefore, $\epsilon$ and the Fe-P separation get obviously improved.

Fig. 6. XRD patterns of raw ore (a), pellet reduced at 1200 °C during 30 min (b), pellet reduced at 1250 °C during 60 min (c), and pellet reduced in the existence of 10 wt% Na$_2$SO$_4$ at 1250 °C during 30 min (d).
sodium sulfate hinders the reduction of Ca₅(PO₄)₃F by the production of a liquid phase which accelerates the interaction between SiO₂, Al₂O₃, CaO in the raw ore. However, there was found a little quantity of S in the metallic iron phase, and it may be troilite (FeS), which reacts with Fe during the reduction process to generate a low-melting point Fe-FeS eutectic system. The generation of the Fe-FeS liquid phase facilitates the mass transfer of metal ions and accelerates the metallic iron grains combination.

Distribution of phosphorus in the microstructure of DRI. In order to study the distribution of phosphorus in the DRI, the magnetic concentrate of the mixed pellet reduced at 1250 °C for 30 min in the absence of sodium sulfate was investigated by SEM-EDS analysis (FEI Quanta 250 ESEM, The Netherlands). Fig. 9, a shows that phosphorus exhibits a reticular distribution in DRI, and its content is high where elemental carbon distributes vigorously (Table 3) [10]. The affinity of phosphorus to metallic iron is quite associated with carbonisation in the metallic phase and the particular texture of oolitic iron ore. Indeed, the ore is made up of numerous layers, and the iron oxides layers are joined with those of gangue minerals [11]. Therefore, previous to reduction of fluorapatite, the new generated metallic iron would envelop the unreduced fluorapatite layers. Once the fluorapatite is reduced, P₂O₅ permeates readily into DRI, already wrapped in the Ca₃(PO₄)₂ surface and the heterogeneous melting of the iron phase led to the reticular distribution [10, 11]. By contrast, the reduction of fluorapatite becomes difficult in the existence of sodium sulfate, due to the consumption of SiO₂ and Al₂O₃ before their reaction with the P-containing minerals during the reduction process.

Characterization of magnetic concentrates. The magnetic concentrate obtained from reduction under optimal reduction parameters (reduced at 1250 °C for 30 min with adding 5 wt% Na₂SO₄) followed by grinding and WLIMS, was investigated using SEM-EDS analysis (Environmental FEI Quanta 250 SEM, The Netherlands) and the result is presented in Fig. 10. It is clear that the metallic iron grains occupied almost all the picture of the magnetic material (Fig. 10, a).
iron oxides are completely disseminated and closely related both optical and cathodoluminescence microscopy reveals that field were investigated. The characterization of the raw ore by DR of iron ore–coal–CaO mixed pellets from the Gara Djebilet cy of improving the Fe–P separation through the addition of a could be recovered under optimized roasting conditions.

Conclusion. According to the EDS quantitative analysis data presented in Table 4, metallic iron is the main element detected in the magnetic material of the pellet reduced at 1250 °C for 30 min in the presence of 5 wt% Na₂SO₄ during solid-state reductive roasting of oolitic hematite ore. These results demonstrate that the coal-based DR could be used as an alternative to the BF for the steel production from Gara Djebilet oolitic iron ore.

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Table 3
EDS quantitative analyses data of the selected spot from the magnetic material of the pellet reduced at 1250 °C for 30 min in the absence of sodium sulfate

| Element | % mass | % atomic |
|---------|--------|----------|
| C K     | 3.92   | 11.88    |
| O K     | 2.78   | 7.92     |
| F K     | 0.92   | 2.20     |
| Al K    | 0.65   | 1.10     |
| Si K    | 1.59   | 2.58     |
| P K     | 1.18   | 1.71     |
| Ca K    | 0.33   | 0.38     |
| Fe K    | 88.62  | 72.22    |

Table 4
EDS quantitative analysis data of the selected spot from the magnetic material of the pellet reduced at 1250 °C during 30 min in the presence of 5 wt% sodium sulfate

| Element | % mass | % atomic |
|---------|--------|----------|
| C K     | 1.94   | 8.10     |
| O K     | 1.42   | 4.25     |
| Al K    | 0.30   | 0.56     |
| Si K    | 0.55   | 0.84     |
| P K     | 0.16   | 0.25     |
| S K     | 0.18   | 0.28     |
| Fe K    | 95.44  | 85.72    |

with gangue minerals. Moreover, the phosphorus content is as high as 0.8 wt%, which makes it not suitable for steel industry.

When the mixed pellets were reduced at various high temperatures, without any additives, part of gaseous P₂, generated by the reduction of fluorapatite, readily penetrates into the metallic iron rather than emitting to the atmosphere, resulting in a high phosphorus content of DRI. By contrast, a final DRI with 96 wt% Fe and 0.16 wt% P can be acquired at e of 97.72 by adding 5 wt% Na₂SO₄ to the mixed pellets when reduced at 1250 °C for 30 min. Furthermore, optical microscopy and SEM analyses of reduced pellets show that the oolitic structure is destroyed and metallic iron particles gathered together and grew significantly in sizes mostly over 75 µm when re-duced in the existence of sodium sulfate.

Fig. 10. Photomicrograph of magnetic concentrate from the pel-let reduced at 1250 °C during 30 min in the presence of 5 wt% Na₂SO₄ (a) and EDS analysis of the selected spot (b)
Мета. Дослідження щодо зниження вмісту фосфору в залізі, отриманого методом прямого відновлення із залізної руди.

Методика. Оолітова залізна руда з високим вмістом фосфору з родовища Гара Джебілет пройшла процес прямої відновлення на основі вугілля (ПВ на основі вугілля) з подальшою мокрою магнітною сепарацією в полі низької інтенсивності (WLIMS). Були досліджені та оптимізовані впливи температури, часу обробки й концентрації Na₂SO₄ на видалення фосфору, ступені металізації та швидкість вилучення заліза. Також були досліджені фазові перетворення оксидів заліза та особливості розподілу фосфору як у відновлених, так і в магнітних матеріалах.

Результати. Відповідне додавання сульфату натрію покращує розділення Fe-P під час прямоого відновлення окатишів на вугільній основі родовища Гара Джебілет.

Наукова новизна. Використання добавки CaO і сульфату натрію під час прямоого відновлення окатишів на основі вугілля родовища Гара Джебілет і подальшої магнітної сепарації.

Практична значимість. Результати показали, що остаточний продукт, отриманий прямою відновленням, містить 96 % мас. Fe і 0,16 % мас. P. Ступінь вилучення заліза склала 97,72 % коли гранули з суміші руда-вугілля-CaO відновлювалися з 5 % мас. Na₂SO₄ за температури 1250 °C протягом 30 хв. Таким чином, ПВ на основі вугілля можна використовувати як альтернативний метод отримання металу з міцних залізних руд методу отримання металу в доменній печі (ДП) у сталеплавильній промисловості.

Ключові слова: Гара Джебілет, оолітова залізна руда з високим вмістом фосфору, пряме відновлення окатишів на основі вугілля, сульфат натрію, магнітна сепарація

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