Recovering Iron from Iron Ore Tailings and Preparing Concrete Composite Admixtures

Chang Tang 1,2,3, Keqing Li 1,2,*, Wen Ni 1,2,* and Duncheng Fan 1,2

1 Key Laboratory of the Ministry of Education of China for High-Efficient Mining and Safety of Metal Mines, University of Science and Technology Beijing, Beijing 100083, China; b20160058@xs.ustb.edu.cn (C.T.); b20170035@xs.ustb.edu.cn (D.F.)
2 School of Civil and Resource Engineering, University of Science and Technology Beijing, Beijing 100083, China
3 Hebei Technology Research Center for Application of High Performance Concrete with Ultra-low Environment Load, Shahe 054100, China

* Correspondence: lkqing2003@163.com (K.L.); niwen@ces.ustb.edu.cn (W.N.)

Received: 15 March 2019; Accepted: 9 April 2019; Published: 15 April 2019

Abstract: Iron ore tailings (IOTs) are a form of solid waste produced during the beneficiation process of iron ore concentrate. In this paper, iron recovery from IOTs was studied at different points during a process involving pre-concentration followed by direct reduction and magnetic separation. Then, slag-tailing concrete composite admixtures were prepared from high-silica residues. Based on the analyses of the chemical composition and crystalline phases, a pre-concentration test was developed, and a pre-concentrated concentrate (PC) with an iron grade of 36.58 wt % and a total iron recovery of 83.86 wt % was obtained from a feed iron grade of 12.61 wt %. Furthermore, the influences of various parameters on iron recovery from PC through direct reduction and magnetic separation were investigated. The optimal parameters were found to be as follows: A roasting temperature of 1250 °C, a roasting time of 50 min, and a 17.5:7.5:12.5:100 ratio of bitumite/sodium carbonate/lime/PC. Under these conditions, the iron grade of the reduced iron powder was 92.30 wt %, and the iron recovery rate was 93.96 wt %. With respect to the original IOTs, the iron recovery was 78.79 wt %. Then, highly active slag-tailing concrete composite admixtures were prepared using the high-silica residues and S75 blast furnace slag powder. When the amount of high-silica residues replacing slag was 20%, the strength of cement mortar blocks at 7 days and 28 days was 33.11 MPa and 50 MPa, respectively, whereas the activity indices were 89 and 108, respectively. Meanwhile, the fluidity rate was appropriately 109. When the content of high-silica residues replacing slag was not more than 30%, the quality of mineral admixtures was not reduced. Last but not least, reusing the high-silica residues during iron recovery enabled the complete utilization of the IOTs.

Keywords: iron ore tailings; iron recovery; concrete composite admixtures; reuse

1. Introduction

Iron ore tailings (IOTs) are a form of solid waste produced during the beneficiation process of iron ore concentrate. Among all kinds of mining solid waste, IOTs are one of the most common solid wastes in the world due to their high output and low utilization ratio. In China, the generation of IOTs has increased rapidly due to recent growth in the iron and steel industries. According to some recent statistics, the output of iron tailings was around 500 million tons in 2015 [1]. Although there has already been a recent trend of gradual reduction in the amount of IOTs produced, the total accumulation of IOTs has exceeded 7.5 billion tons in China [2,3]. The enormous amount of IOTs deposited as waste incurs a high economic cost for waste management and also creates serious environmental
problems and security risks. However, some studies have found that waste IOTs have an iron grade of approximately 8 wt %–12 wt % on average, and occasionally as high as 27 wt % [4,5]. If these materials contain 10 wt % iron on average, 750 million tons of metallic iron is lost during the disposal of IOTs.

With the passage of time, the earth’s iron ore resources continue to decline. Due to this reason, IOTs might become valuable resources in the future. Currently, the processes for utilizing IOTs can be mainly divided into two categories, namely iron recovery from IOTs and the use of IOTs as raw materials. Recovery is a well-known process, and many recovery methods have been developed, including gravimetric separation, magnetic separation, and flotation separation [5–8]. However, these processes create new waste streams after the recovery of iron and are inefficient when using traditional mineral processing techniques. IOTs can also be used to make other products, such as ceramic tiles [9,10], engineered cementitious composites [11], IOT-filled polypropylene cement, and epoxy composites [12,13], and underground backfill mining materials [3]. The conversion of solid industrial waste into other valuable products has received considerable attention in recent years. However, in these processes, iron within the utilized IOTs is not recovered, thus wasting large amounts of iron resources.

Recently, a method involving direct reduction followed by magnetic separation has been used to recover iron from red mud, vanadium tailings, pyrite slag, copper slag, cyanide tailings, and IOTs [14–19]. This method has the advantage of high iron recovery, and is especially suitable for ore with a low iron grade. However, there is no suitable method for dealing with the secondary tailings after iron selection.

In this study, Qidashan IOTs were comprehensively utilized to generate zero-emission IOT waste. The IOTs were owned by the Anshan Iron Steel Group (AISG) in northeastern China and were discharged after the processing of iron minerals from Anshan magnetite quartzite, because iron recovery or reuse of these IOTs as raw materials for value-added products was not economically viable. Meanwhile, due to the low iron content of IOTs, the cost of recovering iron through direct reduction–magnetic separation is high. Consequently, an innovative technique called “pre-concentration followed by direct reduction and magnetic separation technology” was developed [20]. According to this method, pre-concentration is initially undertaken to obtain a pre-concentrated concentrate (PC) and high-silicon residue. The PC is used as a raw material for direct reduction after pre-concentration, and the resulting high-silicon residue from this process is used to prepare slag-tailing concrete composite admixtures that consist of high-silicon residues and blast furnace slag (BFS) discharged by the Ansteel Group Corporation during iron smelting. The reduced iron powder can be conveniently used for electric arc furnace steelmaking by the nearby Ansteel Group Corporation facility as a high-grade alternative raw material to steel scrap. The admixtures can be used to partially or fully replace cement to produce various construction concretes or filling materials for backfilling the mined-out area. Therefore, this technology can truly achieve full iron recovery from IOTs and approaches zero-tailing mining, thus resulting in substantial economic and social benefits. Notably, the amount of silicate residues remaining after direct reduction followed by magnetic separation can reach up to 20 wt % of the total IOTs, and therefore they are used as cement raw meal. In this paper, iron was recovered from IOTs using pre-concentration followed by direct reduction and magnetic separation, whereas slag-tailing concrete composite admixtures were subsequently prepared using the high-silicon residues separated from IOTs.

2. Materials and Methods

2.1. Raw Materials

2.1.1. Iron Ore Tailings

IOTs were sampled from a storage dam at the Qidashan iron ore dressing plant in Anshan, which is in the Liaoning province of China. The chemical composition of this material is presented in Table 1: The total iron content was approximately 12.61 wt %. The crystalline phases in the received IOTs are shown in Figure 1, and included quartz, hematite, and magnetite. After analyzing the iron phases, approximately 58.92 wt % of the total iron was in the form of magnetite or hematite (Table 2), whereas
37.31 wt % of the total iron was in the form of silicate minerals, such as chlorite and hornblende. It is worth noticing that the latter materials cannot be separated effectively using traditional magnetic separation methods.

### Table 1. Chemical composition of the iron ore tailings (IOTs).

<table>
<thead>
<tr>
<th>Components</th>
<th>TFe</th>
<th>SiO₂</th>
<th>CaO</th>
<th>MgO</th>
<th>Al₂O₃</th>
<th>Na₂O</th>
<th>K₂O</th>
<th>MnO</th>
<th>TiO₂</th>
<th>S</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Content/wt %</td>
<td>12.61</td>
<td>75.46</td>
<td>1.70</td>
<td>1.75</td>
<td>1.65</td>
<td>0.32</td>
<td>0.34</td>
<td>0.13</td>
<td>0.06</td>
<td>0.10</td>
<td>0.03</td>
</tr>
</tbody>
</table>

**Figure 1.** XRD pattern of the IOTs.

<table>
<thead>
<tr>
<th>Phases</th>
<th>Magnetic Iron</th>
<th>Hematite and Limonite</th>
<th>Iron Carbonate</th>
<th>Iron Silicate</th>
<th>Iron Sulfide</th>
<th>TFe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iron content/wt %</td>
<td>2.29</td>
<td>5.16</td>
<td>0.39</td>
<td>4.72</td>
<td>0.08</td>
<td>12.64</td>
</tr>
<tr>
<td>Fraction/wt %</td>
<td>18.12</td>
<td>40.82</td>
<td>3.09</td>
<td>37.34</td>
<td>0.63</td>
<td>100.00</td>
</tr>
</tbody>
</table>

2.1.2. Reducing Coal

The reducing coal used in this study was bitumite. Its composition is listed in Table 3. Carbon (45.16 wt %) was the major active reagent during the direct reduction process.

### Table 3. General analysis of the bitumite.

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>Moisture</th>
<th>Ash</th>
<th>Volatiles</th>
<th>Fixed Carbon</th>
</tr>
</thead>
<tbody>
<tr>
<td>Content/wt %</td>
<td>9.88</td>
<td>16.20</td>
<td>28.76</td>
<td>45.16</td>
</tr>
</tbody>
</table>

2.1.3. Additives

Analytical-grade lime (CaO) and sodium carbonate (Na₂CO₃) were used as additives in this study.

2.1.4. S75, Portland Cement PI42.5, and Standard Sand

S75 ground-granulated blast furnace slag powder was produced by the Anshan Iron Steel Group in northeastern China, and the specific surface area was 353 m²/kg. The cement used was Portland cement with a strength grade of 42.5 (complying with the Chinese National Standard GB 175-1999) and was produced by the Tangshan Jidong Cement Co., Ltd., in China. Standard sand was manufactured and packed by the Xiamen ISO Standard Sand Co., Ltd., in China.

2.2. Experimental Methods

The experimental workflow is shown in Figure 2. The workflow includes pre-concentration, direct reduction–magnetic separation, and final products. Among them, iron concentrate I and
iron concentrate II represent the magnetic products after two magnetic separations (0.11 T and 0.8 T), respectively.

Figure 2. Experimental flowsheet: (a) The process of pre-concentration; (b) the process of iron recovery; (c) final products.

For pre-concentration, the IOTs were first milled to a size such that 50 wt % of particles were smaller than 38 μm. They were then separated in a low-intensity magnetic separator (model: RK/CXG; Φ = 50 mm; magnetic intensity: 0.11 T, Hualian Mining Co., Ltd., Tianjin, China) followed by a high-intensity magnetic separator (model: RK/CSQ; Φ = 50 mm × 70 mm; magnetic intensity: 0.8 T, Bureau of geology and minerals 102, Xichang, China). The magnetic product was the pre-concentrated concentrate (PC), whereas the nonmagnetic product was the high-silicon residues that would be reused as concrete admixture.

For direct reduction–magnetic separation, a direct reduction was conducted in a muffle furnace (model: CD-1400, Oriental Electrical Instrument, Yuyao, China). Six sample groups were designed to investigate the effects of various parameters on iron recovery from IOTs, including roasting temperature, roasting time, the carbon-to-IOT ratio, as well as the ratio of the additives and the grinding fineness of the roasted product. The PC was mixed with bitumite and additives at different ratios. Afterwards, the mixtures were roasted at various temperatures and times, and the roasted products were air-cooled and wet-milled in a rod mill before being separated in a low-intensity magnetic separator with a magnetic field intensity of 0.24 T. The iron grade of the reduced iron powder was chemically analyzed, and the recovery of iron was calculated based on the mass balance during the magnetic separation process. The nonmagnetic reducing slag (the chemical composition is presented in Table 4) in the roasted products was used as cement raw meal to directly burn a cement clinker, considering that the chemical composition was similar to that of cement raw materials.
Table 4. Chemical composition of the reducing slag/wt %.

<table>
<thead>
<tr>
<th>Components</th>
<th>SiO$_2$</th>
<th>Fe$_2$O$_3$</th>
<th>FeO</th>
<th>Al$_2$O$_3$</th>
<th>CaO</th>
<th>MgO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Content</td>
<td>59.57</td>
<td>1.36</td>
<td>1.93</td>
<td>3.10</td>
<td>24.81</td>
<td>2.65</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Components</th>
<th>Na$_2$O</th>
<th>K$_2$O</th>
<th>MnO</th>
<th>S</th>
<th>P</th>
<th>Loss</th>
</tr>
</thead>
<tbody>
<tr>
<td>Content</td>
<td>0.97</td>
<td>0.52</td>
<td>0.10</td>
<td>0.10</td>
<td>0.02</td>
<td>4.27</td>
</tr>
</tbody>
</table>

The process of preparing the concrete mineral admixture using high-silicon slag was as follows: S75 was mixed with high-silica residue in different proportions, and the mixtures were milled in the same mill for 30 min, thus yielding slag-tailing composite admixtures for concrete with a ultrafine particle size, which was determined using a laser diffraction analyzer (model: LMS-30, Seishin Enterprise CO., Ltd., Osaka, Japan). Afterwards, the slag-tailing concrete composite admixtures (225 g), Portland cement (225 g), and water (225 g) were mixed together and stirred slowly for 30 s in a mixer (model: NJ-160A, Jianyi Instrument and Machinery Co., Ltd., Wuxi, China), thus forming a cement paste. Then, sand (1350 g) was added to the mixer. The mixture was stirred rapidly for 30 s and poured into prismatic molds (40 mm × 40 mm × 160 mm) on a vibration table running at a moderate vibration rate. All the samples were demolded after curing at 24 h under 90% humidity at 20 ± 3 °C in a standard curing box (model: SHBY-40B, Luda Machinery Instrument, Shaoxing China). The samples were dipped in saturated lime water for curing under the same conditions. The compressive and flexural strengths of the cured specimens were tested after aging for 3, 7, and 28 days.

In this article, iron grade means the total iron (TFe) content, as detected by the chemical analysis (EDTA titration using the ethylenediamine tetra acetic acid as complexing agent to determine the chemical composition). The iron phases were determined using an atomic absorption spectrophotometer (model: AA-6800, Shimadzu, Kyoto, Japan) and titration. The mineral phases of the IOTs and roasted products were investigated using X-ray diffraction (Ultima IV diffractometer with a copper target, 40 kV, 40 mA, scanning speed of 20°·min$^{-1}$, scanning angle of 10°–100°, model: MXP21VAHF, MAC Science Co., Ltd., Osaka, Japan).

3. Results and Discussion

3.1. Pre-Concentration

The pre-concentration process was investigated using a previously described procedure [21,22]. The optimized conditions for pre-concentration were obtained from a previous work [22], and they were as follows: A particle size such that 50 wt % of the particles were smaller than 38 µm, a low magnetic field intensity of 0.11 T, and a high magnetic induction of 0.8 T. Under these conditions, the productivities of PC, the iron grade, and total iron recovery were 28.82 wt %, 36.58 wt %, and 83.86 wt %, respectively. The pre-concentration products are presented in Table 5. The chemical composition of the products is given in Table 6.

Table 5. Pre-concentration products/wt %. PC: Pre-concentrated concentrate.

<table>
<thead>
<tr>
<th>Products</th>
<th>Productivity Rate</th>
<th>Iron Grade</th>
<th>Iron Recovery Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>PC</td>
<td>28.82</td>
<td>36.58</td>
<td>83.86</td>
</tr>
<tr>
<td>High-silicon residue</td>
<td>71.18</td>
<td>2.85</td>
<td>16.14</td>
</tr>
<tr>
<td>IOTs</td>
<td>100.00</td>
<td>12.57</td>
<td>100.00</td>
</tr>
</tbody>
</table>

Table 6. Chemical composition of the pre-concentration products/wt %.

<table>
<thead>
<tr>
<th>Products</th>
<th>SiO$_2$</th>
<th>Fe$_2$O$_3$</th>
<th>FeO</th>
<th>CaO</th>
<th>MgO</th>
<th>Al$_2$O$_3$</th>
<th>Na$_2$O</th>
<th>K$_2$O</th>
<th>MnO</th>
<th>TiO$_2$</th>
<th>S</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>PC *</td>
<td>41.71</td>
<td>42.83</td>
<td>8.18</td>
<td>1.82</td>
<td>1.69</td>
<td>1.82</td>
<td>0.41</td>
<td>0.27</td>
<td>0.14</td>
<td>0.072</td>
<td>0.1</td>
<td>0.023</td>
</tr>
<tr>
<td>High-silicon residue **</td>
<td>91.29</td>
<td>2.96</td>
<td>0.72</td>
<td>1.3</td>
<td>1.25</td>
<td>1.19</td>
<td>0.42</td>
<td>0.17</td>
<td>0.074</td>
<td>0.024</td>
<td>0.066</td>
<td>0.023</td>
</tr>
</tbody>
</table>

* The PC was composed of iron concentrate I and Iron concentrate II; ** high-silicon residue is the nonmagnetic component after the pre-concentration process.
3.2. Iron Recovery through Direct Reduction Followed by Magnetic Separation

In order to improve iron recovery, sodium carbonate (Na$_2$CO$_3$) and lime (CaO) were used as additives in the reduction process. The effects of the bituminous content, roasting time, roasting temperature, sodium carbonate content, lime content, and grinding fineness of the roasted products on iron grade and iron recovery were studied.

3.2.1. Effect of Bituminite Ratio on Iron Recovery

When studying the influence of bituminous content on iron grade and iron recovery (based on previous studies [23–27]), the optimized conditions of sodium carbonate content, lime content, roasting temperature, roasting time, and fineness of roasted products were determined. The specific parameters used in the experiments were as follows: A 100:10:5 mixing ratio for PC/lime/sodium carbonate, respectively; a roasting temperature of 1150 °C; and a roasting time of 60 min. The grinding fineness of the roasted product at a -74 μm ratio was 65%. The results are shown in Figure 3a.

![Figure 3a](image-url)

**Figure 3.** Effect of different influencing factors on iron recovery and iron grade: (a) Bituminous ratio; (b) roasting time; (c) roasting temperature; (d) sodium carbonate ratio; (e) lime ratio; (f) grinding fineness.
Figure 3a shows that the iron grade and recovery rate increased rapidly as the bitumite ratio increased from 12.5 wt % to 17.5 wt %, respectively. After reaching this value (17.5 wt %), both quantities decreased. The iron recovery rate increased from 85.63 wt % to 91.21 wt %, indicating that increasing the bitumite dose significantly enhanced the iron recovery rate. However, when the bitumite was in excess, the reduced metallic iron could not overtake the porous coal and other gangue minerals. The increase in size of the metallic iron particles was hindered, which reduced iron recovery [15]. During this test, the optimal bitumite content was found to be 17.5 wt %. Meanwhile, the increase in noncombustible substances in bituminous coal reduced the iron grade.

3.2.2. Effect of Roasting Time on Iron Recovery

The effect of roasting time on iron recovery was investigated under the following conditions: A 100:17.5:10:5 mixing ratio of PC/bitumite/lime/sodium carbonate, respectively; a roasting temperature of 1150 °C; and a grinding fineness the same as that reported in Section 3.2.1. The corresponding experimental results are shown in Figure 3b.

As shown in Figure 3b, the iron recovery rates peaked after roasting for approximately 50 min. As the roasting time increased from 35 to 50 min, the iron recovery of the reduced iron powder increased from 87.65 wt % to 92.23 wt % before increasing slowly from 50 to 80 min. Meanwhile, the iron grade maintained a stable trend. Consequently, the optimum roasting time was found to be 50 min for the following conditions: A 100:17.5:10:5 mixing ratio of PC/bitumite/lime/sodium carbonate, respectively; and a roasting temperature of 1150 °C.

3.2.3. Effect of Roasting Temperature on Iron Recovery

The mixtures consisting of PC, bitumite, lime, and sodium carbonate in the mixing ratio of 100:17.5:10:5, respectively, were roasted for 50 min at various temperatures. The grinding fineness was the same as that reported in Section 3.2.1. Figure 3c shows that increasing the roasting temperature up to 1250 °C induced a rapid increase in the iron grade from 90.8 wt % to 91.48 wt %. However, when the temperature exceeded 1250 °C, the rate of improvement decreased dramatically. When the roasting temperature was too low, the metallic iron particles remained small and closely associated with the gangue minerals. Therefore, the gangue mineral particles were easily mixed with the metallic iron during the magnetic separation process. It is worth noticing that further increasing the roasting temperature improved the growth of the metallic iron particles [28]. When the roasting temperature exceeded 1350 °C, however, the metallic iron particles became coarser, and the minerals softened or melted, causing difficulty during grinding [29]. Consequently, a roasting temperature of 1250 °C was recommended as the optimum temperature.

3.2.4. Effect of Sodium Carbonate Ratio on Iron Recovery

In order to optimize the sodium carbonate dosage, various sodium carbonate ratios were studied, while the other process parameters were kept constant. The conditions were maintained as follows: A 100:17.5:10 mixing ratio of PC, bitumite, and lime, and roasting at 1250 °C for 50 min. The grinding fineness was the same as that reported in Section 3.2.1. The corresponding results are presented in Figure 3d.

When the sodium carbonate ratio increased from 2.5 wt % to 7.5 wt %, the iron grade of the reduced iron powder increased from 81.55 wt % to 91.75 wt %, respectively, whereas the recovery of iron increased from 90.12 wt % to 92.97 wt %, respectively. When the sodium carbonate ratio exceeded 7.5 wt %, no obvious improvements were observed in the iron recovery. When sodium carbonate was present, it reacted with SiO₂ in the gangue minerals to form low-melting phases, such as Na₂O·SiO₂, 2Na₂O·SiO₂, and Na₂O·2SiO₂ (melting point of 1088 °C), during the direct reduction. Furthermore, the liquid phase was composed of Na₂O·SiO₂, 2Na₂O·SiO₂, and Na₂O·2SiO₂, which were formed locally [30]. When a liquid phase is present in roasted products, the diffusion coefficient increases, thus accelerating the diffusion and migration of iron in some cases and causing the metallic iron particles to
gather and grow rapidly [28,31,32]. Based upon these results, the optimum sodium carbonate content was found to be 7.5 wt %.

3.2.5. Effect of Lime Ratio on Iron Recovery

The effect of lime content on iron recovery was also studied. The samples consisted of PC, bitumite, and sodium carbonate in a ratio of 100:17.5:7.5, respectively, and they were roasted at 1250 °C for 50 min. The grinding fineness was the same as that reported in Section 3.2.1. The corresponding results are shown in Figure 3e. It can clearly be seen that increasing the ratio of lime from 5 wt % to 12.5 wt % increased the iron grade from 80.67 wt % to 89.62 wt %, whereas the corresponding iron recovery increased from 90.17 wt % to 93.64 wt %. When the lime addition ratio exceeded 12.5 wt %, both the iron grade and recovery rate decreased. Based upon the results, the optimum sodium carbonate content was chosen to be 12.5 wt %.

Lime is a common and low-cost reaction additive that is used to improve the reduction of iron ore. Because PC contains numerous SiO$_2$ groups, the poor reactivity of fayalite, which forms readily during the reaction between FeO and SiO$_2$, is expressed using Equation (1):

$$\text{FeO} + \text{SiO}_2 = \text{FeO-SiO}_2.$$  

(1)

When the lime is added, it reacts with SiO$_2$ instead of FeO in the gangues to form calcium silicate (CaO·SiO$_2$) during reduction roasting, and meanwhile bituminous coal participates in the reaction as a reducing agent. This reaction can be expressed using Equation (2):

$$\text{FeO-SiO}_2 + \text{CaO} + \text{C} = \text{Fe} + \text{CaO-SiO}_2 + \text{CO}↑.$$  

(2)

Consequently, more FeO can be reduced to metallic iron, enhancing the direct reduction process of iron oxide.

3.2.6. Effect of the Grinding Fineness of the Roasted Product on Iron Recovery

Complete mineral liberation is essential when magnetically separating metallic iron particles [15]. Consequently, the roasted product must be ground before magnetic separation. The effect of the grinding fineness of the roasted product was also studied under the following conditions: A roasting temperature of 1250 °C, a roasting time of 50 min, and a mixing ratio of 100:17.5:12.5:7.5 of PC, bitumite, lime, and sodium carbonate, respectively. The effect of the grinding fineness of the roasted product on iron recovery is shown in Figure 3f. The relationship between the grinding fineness and grinding time of the roasted product is presented in Table 7.

<table>
<thead>
<tr>
<th>Grinding Time/min</th>
<th>10</th>
<th>15</th>
<th>20</th>
<th>25</th>
<th>30</th>
<th>35</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grinding fineness (−74 µm)/wt %</td>
<td>63.9</td>
<td>66.25</td>
<td>69.05</td>
<td>74.35</td>
<td>76.21</td>
<td>78.54</td>
</tr>
</tbody>
</table>

Figure 3f shows that as the grinding time was prolonged and the grinding fineness increased, the iron grade of the reduced iron powder and the iron recovery initially increased and then stabilized. When 74.35 wt % of the particles were smaller than 74 µm, the metallic iron particles were almost completely liberated, generating an iron grade of 92.30 wt % and an iron recovery of 93.96 wt %. Generally, the finer the grinding fineness, the more difficult the recovery of magnetic fraction is. When the grinding fineness was −74 µm, the iron recovery was not improved relative to 66 µm, though the iron grade improved. Due to this reason, −74 µm was chosen as the optimum grinding fineness.
3.2.7. XRD Analysis of the Roasted Product

The roasted products were analyzed using XRD to explain the experimental results discussed above. The XRD pattern of the roasted product is shown in Figure 4.

![XRD pattern of the roasted product](image)

Figure 4. XRD pattern of the roasted product.

Figure 4 shows that the peaks for metallic iron appeared without evidence of any other iron compounds in the roasted product, indicating that the iron-bearing minerals were almost completely reduced to metallic iron. The impurities included calcium silicate, cristobalite, and some residual quartz, as well as amorphous forms of the roasted product. Under high temperature conditions, a liquid phase formed in large amounts during the reduction. This liquid phase accelerated the growth of metallic iron particles.

This study did not investigate the combined effects of sodium carbonate, lime, and other factors on iron recovery. The optimum parameters determined by the above single factor test were found to be a roasting temperature of 1250 °C; a roasting time of 50 min; a mixing ratio of 100:17.5:12.5:7.5 of PC, bitumite, lime, and sodium carbonate, respectively; and the proportion of −74 μm being more than 74.35 wt %. Under such conditions, the iron grade and iron recovery rate were found to be 92.30 wt % and 93.96 wt %, respectively. Specifically, 78.79 wt % of the iron could be recovered from Qidashan IOTs through pre-concentration followed by direct reduction and magnetic separation. The XRD pattern showed that the main phase in the roasted product was metallic iron.

3.3. Preparation of Slag-Tailing Concrete Composite Admixtures Using the High-Silica Residues

Since the main component of high-silica residues is SiO₂, which has no mineral activity, it is necessary to prepare concrete mineral admixtures using mechanical excitation. Therefore, a grinding test was carried out. The method for determining the optimized grinding time and method was based on the approach of Huang [33] and was as follows: First, S75 and high-silica residue were mixed according to the ratios given in Table 8, and the mixture was ground for 30 min. Then, a cement mortar test block was prepared using ground composite admixtures of cement, ISO sand, and water. The mixing proportions of the cement mortar are presented in Table 9. The specific surface area of each group mixture, fluidity, and the compressive strength of each group mortar were investigated. Group C-0 was made of pure cement and ISO sand as a blank group for a calculation of the activity index of the composite admixtures.
Table 8. Composite admixture proportions/wt %.

<table>
<thead>
<tr>
<th>Number</th>
<th>S 75</th>
<th>Tailing</th>
</tr>
</thead>
<tbody>
<tr>
<td>S-0</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>S-10</td>
<td>90</td>
<td>10</td>
</tr>
<tr>
<td>S-20</td>
<td>80</td>
<td>20</td>
</tr>
<tr>
<td>S-30</td>
<td>70</td>
<td>30</td>
</tr>
<tr>
<td>S-40</td>
<td>60</td>
<td>40</td>
</tr>
<tr>
<td>S-50</td>
<td>50</td>
<td>50</td>
</tr>
</tbody>
</table>

Table 9. Mixing proportion of cement mortar/g.

<table>
<thead>
<tr>
<th>Number</th>
<th>Cement</th>
<th>Composite Admixture</th>
<th>ISO Sand</th>
<th>Water</th>
</tr>
</thead>
<tbody>
<tr>
<td>C-0</td>
<td>450</td>
<td>0</td>
<td>1350</td>
<td>225</td>
</tr>
<tr>
<td>CS-0–CS50*</td>
<td>225</td>
<td>225</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

CS-0–CS50* represents cement mortar when the composite admixtures were numbered S-0, S-10, S-20, S-30, S-40, or S-50.

3.3.1. Chemical Composition and Mineral Phases of the High-Silica Residues

According to the data shown in Table 6, the SiO$_2$ content of the high-silica residues was 91.29 wt %. The XRD pattern of the high-silica residues is shown in Figure 5. The main crystalline phase was quartz, with small amounts of hematite and hornblende.

![XRD pattern of the high-silica residues](image)

3.3.2. Specific Surface Area of the Slag-Tailing Composite Admixture and the Working Performance of the Mortar

Figure 6a shows the specific surface area of each group slag-tailing composite admixture after grinding for 30 min. The specific surface area of each group from S-0 to S-50 increased sequentially, indicating that the grindability of the mixture improved with an increase in the content of high-silica residue. Meanwhile, the energy consumption of the grinding could be reduced in industrial production.

The fluidity and fluidity ratio of each group mortar are shown in Figure 6b. The results show that both the fluidity and fluidity ratio increased with an increase in the content of high-silica residue. This was because the content of the fine particles increased: Due to this, the gap between the mortars could be filled, and the amount of interstitial water was reduced. The calculation of the fluidity ratio revealed that the fluidity values of each group were comparable to the reference group (Group C-0). Within a certain range, the greater the fluidity ratio, the better the performance of the mortar was. However, the compressive strength and fluidity of the mortar, which are the main properties of mineral admixtures, should be considered simultaneously for determining the optimum proportion.
The compressive strength of 7 days and 28 days in the CS-20 group was the highest, reaching values of 33.11 MPa and 50 MPa, respectively. This is because high-silica residue, which contains large quantities of ultrafine particles with diameters of less than 5 μm, is an inert material comparable to blast furnace slag, whereas the compressive strength of each group first increased and then decreased. This was due to the reason that high-silica residues are inert materials comparable to blast furnace slag, whereas the activity index of mineral materials did not decrease when the amount of blast furnace slag replaced by tailings did not exceed 30%. Furthermore, the 7-day and 28-day activity indices of the S-20 group were the highest, reaching values of 60.70, 80.90, and 100.11%, respectively. However, the activity index of concrete mineral admixtures should be considered when high-silica residue is used to prepare slag-tailing composite admixtures. Figure 7 shows the compressive strength of mortars and the activity index of the slag-tailing admixture.

### 3.3.3. Compressive Strength and Activity Index of Mortar

The activity index is one of the main indicators for evaluating the quality of mineral admixtures, and their calculation method was based upon the compressive strength of each group compared to the blank group. Figure 7b shows the activity index of different ages of each group. It is clear that the activity index of mineral materials did not decrease when the amount of blast furnace slag replaced by tailings did not exceed 30%. Furthermore, the 7-day and 28-day activity indices of the S-20 group reached values of 89 and 108, respectively, which exceeded the activity index of the S-0 group without high-silica residues. The high activity of the S-20 group was primarily attributed to high-silica residue, which contained large quantities of ultrafine particles with diameters of less than 5 μm (the particle size distribution curve for the slag-tailing admixtures of S-20 is shown in Figure 8). These particles not only served as filling materials, but they also increased the compactness of the matrix,
but also as nucleation catalysts, which accelerated the cement hydration process \[34,35\]. When only the quality of the slag-tailing admixture was considered, the S-20 group came out as being the best among all. However, the S-30 group may be more suitable if the maximum use of high-silica residue is considered.

![Particle size distribution curve for the slag-tailing admixtures of CS-20.](image)

**Figure 8.** Particle size distribution curve for the slag-tailing admixtures of CS-20.

4. **Conclusions**

This paper demonstrated an innovative technical method called “pre-concentration followed by direct reduction and magnetic separation technology”, which was developed to completely utilize IOTs. This method recovers a large amount of iron while solving the problem of secondary storage of reducing slag and high-silica residues. Based on the obtained results, the following conclusions can be drawn:

1. Primary grinding was followed by low-intensity magnetic separation and subsequent high-intensity magnetic separation processes to concentrate the iron in IOTs. The PC was characterized by an iron grade of 36.58 wt % and a total iron recovery of 83.86 wt % when 50 wt % of the particles were smaller than 0.038 mm, and the magnetic separation was conducted with a low magnetic field intensity of 0.11 T and a high magnetic induction of 0.8 T. Furthermore, high-silicon residues containing 91.29 wt % SiO$_2$ were obtained;

2. The bitumite ratio, roasting temperature, roasting time, lime ratio, sodium carbonate ratio, and particle size of the roasted product were the six major factors that influenced iron recovery. Optimal reduction conditions were obtained as follows: A roasting temperature of 1250 °C, a roasting time of 50 min, and a mixing ratio of 17.5:7.5:12.5:100 ratio of bitumite/sodium carbonate/lime/PC, respectively. Under these conditions, the iron grade of the reduced iron powder was found to be 92.30 wt %, and the iron recovery rate was 93.96 wt %. Relative to the original IOTs, the iron recovery was 78.79 wt %;

3. When the amount of high-silica residues replacing blast furnace slag was 20%, the strength of the cement mortar was 33.11 MPa and 50 MPa at 7 days and 28 days, respectively, whereas the corresponding activity index was 89 and 108, respectively. Furthermore, the fluidity rate of the mortar was approximately 109. When the content of high-silica residues replacing blast furnace slag was not more than 30%, the quality of slag-tailing concrete composite admixtures was not reduced;

4. Iron was recovered from IOTs through pre-concentration followed by direct reduction and magnetic separation processes. Moreover, the high-silica residues from the pre-concentration process were reused in slag-tailing concrete composite admixtures. Combining these recycling processes can ensure complete utilization of IOTs;
5. Incorporating an extra pre-concentration step before the direct reduction circumvented the high costs of recovering small amounts of iron through direct reduction. Consequently, this innovative technology can be extended to other IOTs with low iron content, exhibiting high potential for many other applications.

Author Contributions: C.T. performed the experiment, wrote the manuscript, and analyzed the data with the help of D.F.; K.L. revised the manuscript; W.N. designed the experiment.

Funding: This research was funded by the Ministry of Science and Technology International Science and Technology Cooperation Program, grant number 2016YFE0130700.

Acknowledgments: The authors would also like to thank the Analytical and Testing Centre of the University of Science and Technology Beijing, China University of Geoscience, and the Xi’an Tianzhou Mining Science and Technology Development Co., Ltd., of China, which supplied the facilities necessary for data collection.

Conflicts of Interest: The authors declare no conflicts of interest.

References


22. Fan, D. Research on Pre-Concentration and Deep Reduction of Qidashan Iron Ore Tailings and the Comprehensive Utilization of Tailings; University of Science and Technology Beijing: Beijing, China, 2018.


25. Yu, W.; Sun, T.; Kou, J.; Wei, Y.; Xu, C.; Liu, Z. The function of Ca(OH)₂ and Na₂CO₃ as additive on the reduction of high-phosphorus oolitic hematite-coal mixed pellets. *ISIJ Int.* 2013, 53, 427–433. [CrossRef]


