Comparison of butyric acid leaching behaviors of zinc from three basic oxygen steelmaking filter cakes

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Article

Comparison of Butyric Acid Leaching Behaviors of Zinc from Three Basic Oxygen Steelmaking Filter Cakes

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Abstract: The selective leaching of zinc from three different basic oxygen steelmaking (BOS) filter cakes by butyric acid was investigated to compare the leaching behaviors of zinc and further to establish the correlation of the zinc leaching performances and the chemical compositions. The effects of acid concentration and the acid to solid (L/S) stoichiometric ratio were studied, with different optimal leaching conditions obtained. BOS-1 showed the lowest leachability with only less than 10% of zinc removed by 0.5 M acid concentration and 90% of the L/S stoichiometric ratio in 10 h. The best zinc selectivity was achieved with BOS-2 at 51.2% of zinc leaching efficiency, with only 0.47% of iron loss under optimal conditions of 1.5 M acid concentration and a 70% stoichiometric ratio. BOS-3 showed the highest leaching of zinc but the optimal conditions depend on the priority consideration. Using 1.0 M acid and 90% stoichiometric ratio for 10 h, the leaching efficiency of zinc was 84.6% with 20% iron loss. The filter cakes and the leaching residues were characterized. The results indicate different zinc and iron leaching behaviors, which were probably related to the storage conditions, zinc containing phases and the leaching parameters.

Keywords: BOS filter cakes; butyric acid; selective leaching; leaching behaviors; zinc; iron

1. Introduction

In the iron and steel industry, over 30 million tonnes of dusts are generated each year [1,2]. Basic oxygen furnace steelmaking (BOS) represents about 57% of the annual steel production worldwide and electric arc furnace (EAF) accounts for 27%, hence more dusts will be produced and this increasing trend is likely to continue [3]. These dusts are categorized as hazardous metallurgical waste in light of the heavy metals contained [4]. BOS plays a key role in steel production where hot metallic iron from blast furnace (BF) and scrap steel is converted to steel [3,5]. The BOS dust contains metal oxides, thus zinc mostly exists as ZnO and ZnFe₂O₄ while iron is mainly found as Fe, FeO, Fe₂O₃, Fe₃O₄ and ZnFe₂O₄. The zinc content ranges from 0.5% to 8% with 50–80% of zinc as ZnO and the rest as ZnFe₂O₄. The iron content in the dust varies between 50% and 75% [6–9]. Considering its high iron content, BOS dust is potentially a secondary iron resource for recovery. However, direct recycling is hindered by the problematic zinc and therefore BOS dust has to be stored or disposed of in landfills [10–13].

It has been demonstrated that the composition of the steelmaking dust varies widely depending on the quality of the scrap used, the type of steel produced, and the operating and the aging conditions [14,15]. EAF dust has a much higher content of non-ferrous metal oxides than BOS dust. The content of carbon in
the BF dust is usually higher than that contained in the BOS dust while zinc content is correspondingly 2–6% lower. Also, BF dust is coarser than BOS dust due to the difference between the processes. It should be noted that the zinc content in EAF dust is the highest among these three steelmaking dusts.

The best way to solve this problem is to selectively remove zinc from such waste materials by using hydrometallurgy methods, which complies with the policy of sustainable development, helps to alleviate furnace damage, reduces the environmental load of stored waste and also reduces the costs of the replacement of raw materials. Considering the lack of selectivity owing to the large amount of iron dissolution produced using inorganic acids, organic acids are preferred due to having no secondary waste, being environmentally benign and showing indications of a potential bioleaching technology. After selective removal of zinc, the waste becomes an excellent iron-bearing material that can be used as a feed component to produce iron and steel [4,16]. Hence, it is feasible to selectively remove zinc from the BOS dusts.

It is presumed that zinc leaching from steelmaking dust is related to the fraction of Zn in the form of ZnO because ZnFe$_2$O$_4$ has a very stable spinel structure that is considerably refractory against leaching [17–20]. However, no specific relationship between the leaching efficiency of zinc and the ZnO fraction has been established. Steer and Griffiths investigated the leaching of zinc from BF slurry with 2.25 wt.% of Zn in the form of ZnO using 1 M prop-2-enoic acid, and 83.1% of zinc was removed [21]. Kelebek et al. found that the fine fraction of BOS sludge contained a greater proportion of ZnFe$_2$O$_4$ than the coarse fraction, and 81% of zinc removal was achieved for the coarse fraction by leaching with H$_2$SO$_4$ while only 29.2% removal was achieved for the fine fraction [7]. It was reported by Vereš et al. that BOS dust contained 9.37 wt.% zinc with 14.5% of the total zinc as ZnFe$_2$O$_4$ and the remaining portion as ZnO, and the zinc extraction was less than 50% even if the H$_2$SO$_4$ concentration was increased to 2.0 M at room temperature [22]. Using H$_2$SO$_4$ to leach zinc from BOS sludge with 2.74% of Zn, about 10% of zinc was removed at room temperature and low acid concentration (0.1 and 0.2 M) since only Zn in ZnO was leached out. However, Zn in ZnFe$_2$O$_4$ was further dissolved at higher temperatures and higher concentrations (1 M), giving increased zinc and iron leaching efficiencies over 50% [11]. So, the characteristics of the steelmaking dusts are closely related to the zinc leaching performance.

Butyric acid (CH$_3$CH$_2$CH$_2$COOH), as a 4-carbon short-chain fatty acid, is known to have many applications in the chemical, food, and pharmaceutical industries. It is partially dissociated weak acid, and the stabilization of CH$_3$CH$_2$CH$_2$COO$^-$ could potentially alter the extraction capability as follows.

$$\text{CH}_3\text{CH}_2\text{CH}_2\text{COOH} + \text{H}_2\text{O} = \text{CH}_3\text{CH}_2\text{CH}_2\text{COO}^- + \text{H}_3\text{O}^+$$ \hspace{1cm} (1)

In a previous study [23], butyric acid was shown to be highly efficient in selective leaching of zinc over iron from a BOS filter cake. However, each steelmaking dust is unique in terms of the chemical composition and herein it is necessary to study the leaching performances of different steelmaking wastes using butyric acid. This paper compares the leaching behaviors of zinc from three different BOS filter cakes using butyric acid to develop a better understanding of the relations to their chemical, physical, morphological and mineralogical characterization. This is vital to the design of alternative techniques for treating such BOS filter cakes. The optimal leaching conditions for maximum zinc removal and minimum iron removal from the three BOS filter cakes were determined by varying acid concentration and the acid to solid (L/S) stoichiometric ratio. The filter cakes and the corresponding leaching residues were characterized by X-ray fluorescence (XRF), X-ray diffraction (XRD), scanning electron microscopy with energy dispersive X-ray spectrometer (SEM/EDS), thermo-gravimetric/differential scanning calorimetry (TG/DSC).
2. Materials and Methods

2.1. Materials

The three different BOS filter cake wastes were collected from a steelwork in Australia, and the details are provided in Table 1. These samples were oven dried, crushed and sieved to 300–500 µm for leaching experiments.

Table 1. The information of the three different BOS filter cake samples.

| Sample | Storage Time | Pretreatment | Color       |
|--------|--------------|--------------|-------------|
| BOS-1  | Several years| Sintered in Stockpile | Red brown   |
| BOS-2  | Half a year  | Heated at 130 °C for 40 days | Shiny black |
| BOS-3  | Within one month | Freshly stored in fridge | Dull black  |

Butyric acid with 99% purity supplied by Sigma Aldrich, Australia, was used as a leaching reagent. Zinc and iron standard solutions were used for the calibration of inductively coupled plasma-optical emission spectrometry (ICP-OES 710, Agilent, Australia) analysis to determine the concentrations of zinc and iron. The deionized water used for dilution of different acids was purified using a water super-purification apparatus (Milli-Q, Millipore, North Ryde, Australia).

2.2. Leaching Experiments

The leaching experiments were performed in 250 mL conical flasks on a horizontally oscillating shaker (RM2, Ratek, Boronia, Australia) with 120 rpm to stir the samples for 10 h at ambient temperature. The effects of acid concentration and L/S stoichiometric ratio were investigated in the study. The acid concentrations tested are 0.5, 1.0, 1.5, and 2.0 M with the butyric acid solution fixed at 150 mL. At each acid concentration, the 10, 30, 50, 70 and 90% of L/S stoichiometric ratio [23] were investigated to express the solid weight added.

The sampling was taking under agitated conditions by removing 1.0 mL of leaching solution to be diluted to 10 mL using deionized water and filtered by 0.22-µm cellulose nitrate membrane using syringe. Further dilution was made by 2% nitric acid to appropriate Fe and Zn concentration ranges for the analysis by ICP-OES. The leaching efficiency was calculated from a mass balance as follows:

\[
\text{Leaching efficiency (\%)} = \frac{\text{Mass of M in leachate}}{\text{Mass of M in filter cake added into flask}} \times 100
\]

where M is either Zn or Fe.

The pH of the leaching solutions was measured. After leaching, the final solid residues were washed with deionized water, filtered, dried and weighed for further characterization.

2.3. Characterization of the Filter Cakes and Leaching Residues

The chemical compositions of the BOS filter cakes were determined by X-ray fluorescence (XRF, AMETEK SPECTRO XEPOS ED-XRF, Kleve, Germany). To avoid loss of zinc when it was melted with fluxing material, all the samples were first heated slowly to 600 °C in an oxygen stream to remove their carbonaceous matter. The mineralogical compositions of the original filter cakes and leaching residues were analyzed by X-ray diffraction (XRD, GBC MMA, Braeside, Australia). The XRD patterns for quantitative analysis were obtained at 35 kV and 28.5 mA with monochromated Cu-Kα X-ray radiation (λ = 1.5406 Å) from 10° to 142° at a scanning speed of 0.5°/min with a step size of 0.014°. The patterns for qualitative analysis were from 15° to 85° at a scanning speed of 1.5°/min with a step size of 0.05°. The morphologies were studied by scanning electron microscopy with energy dispersive X-ray spectrometer (SEM/EDS, JSM-6490LV, JEOL, Tokyo, Japan). The thermo-gravimetric (TG) and differential scanning calorimetry (DSC) analysis of the thermal behavior of the filter cake samples was
performed using the NETZSCH STA 449 F5 Jupiter (Selb, Germany) under air and argon atmospheres. Approximately 100 mg of a sample was loaded in an Al$_2$O$_3$ crucible on a pan of the microbalance, and scanned in the temperature range of 50–800 °C at a heating efficiency of 10 °C/min and a flow efficiency of 100 mL/min. An empty pan served as the reference.

3. Results and Discussion

3.1. Characterization of the BOS Filter Cakes

The chemical compositions of the three BOS filter cakes were listed in Table 2, and similar elements were observed. They mainly contained Fe and Zn together with small amounts of Ca, Mg and Si. Other metals exhibited very low concentrations below 1%. BOS-1 contained the lowest zinc level at 2.42 wt.%, followed by 6.53 wt.% in BOS-2, which is approximately half of that in BOS-3 at 13.77 wt.%.

Table 2. The chemical compositions of the three BOS filter cakes (wt.%) based on the oxide content.

| Sample | BOS-1 | BOS-2 | BOS-3 |
|--------|-------|-------|-------|
| Fe     | 56.0  | 56.4  | 60.0  |
| Zn     | 2.42  | 6.52  | 13.8  |
| Ca     | 4.04  | 2.86  | 3.02  |
| Mg     | 1.73  | 1.16  | 0.95  |
| Si     | 0.72  | 0.65  | 1.01  |
| Al     | 0.07  | 0.02  | 0.09  |
| Mn     | 0.83  | 0.67  | 0.61  |
| Ti     | 0.015 | 0.022 | 0.007 |
| K      | 0.031 | 0.029 | 0.020 |
| Pb     | 0.096 | 0.085 | 0.387 |
| Cr     | 0.053 | 0.022 | 0.035 |
| Ni     | 0.030 | 0.008 | 0.000 |
| V      | 0.039 | 0.012 | 0.013 |
| LOI    | 4.73  | 1.89  | 3.56  |

Table 2. The chemical compositions of the three BOS filter cakes (wt.%) based on the oxide content.

LOI-Loss on Ignition at 1050 °C.

Figure 1 presents the XRD patterns of the three BOS filter cakes. It can be found that zinc existed as ZnO and ZnFe$_2$O$_4$ in all the samples, but the contents of zinc in these two compounds varied widely. The quantitative analysis showed that the content of zinc in ZnO was significantly lower, equal to, while much higher than that in ZnFe$_2$O$_4$ for BOS-1, BOS-2, and BOS-3, respectively. For iron containing phases, FeO, Fe$_3$O$_4$ and Fe were detected in all these samples but Fe$_2$O$_3$ was only found in BOS-1. Fe$_3$O$_4$ and Fe$_2$O$_3$ were the predominant crystalline phases in BOS-1 while FeO and Fe were the major iron states in both BOS-2 and BOS-3, indicating that BOS-1 is more oxidized. Calcite was also confirmed in all these three samples.

Figure 1. XRD patterns of the three BOS filter cakes: (a) BOS-1; (b) BOS-2; (c) BOS-3.
Figure 2 presents the morphology of the three BOS filter cakes. All the samples consist of fine grains with various shapes and sizes, but the grain size of BOS-1 is obviously larger than those of BOS-2 and BOS-3. BOS-1 contains non-spherical grains and clusters of submicron grains, but the majority of the grains are >1 µm. EDS analysis confirmed the presence of metal oxides that were mainly iron oxides, followed by the oxides of Zn, Al, Ca and Mg. For BOS-2 and BOS-3, the individual grains are predominantly spherical at submicron level, and large grains >1 µm are rare. The fine grains agglomerated together to form a porous structure of the filter cakes which favors the access of acid to zinc in leaching.

![Figure 2. SEM micrographs of the three BOS filter cakes. (a) BOS-1; (b) BOS-2; (c) BOS-3.](image)

Figure 3 presents the TG-DSC curves of the three BOS filter cakes, which were obtained in the temperature range of 50–800 °C both in air and in argon with a gas flow rate of 100 mL/min. Overall, the TG patterns of BOS-2 and BOS-3 show similar trends of weight increase in air atmosphere up to 500 °C, while mass loss in argon atmosphere. However, BOS-1 mainly was subjected to a weight loss with increasing temperature in both atmospheres. In air, the weight loss of BOS-1 was 5.15%. For filter cakes BOS-2 and BOS-3, there was 4.4% and 5.11% of total weight gain, respectively. The
results indicated some oxidation and possible combustion reactions for both BOS-2 and BOS-3 with two strong exothermic peaks at about 340 °C and 430 °C. In argon atmosphere, the absence of free oxygen in the furnace led to no oxidation reactions in contrast to the analysis in air, and hence the mass loss became increasing monotonically as reflected by the TG curves. The weight loss recorded for BOS-1 was 5.96%, followed by 4.44% for BOS-2 and 3.36% for BOS-3.

Figure 3. TG-DSC curves of the three BOS filter cakes in air and argon atmospheres: (a) BOS-1, in air; (b) BOS-2, in air; (c) BOS-3, in air; (d) BOS-1, in argon; (e) BOS-2, in argon; (f) BOS-3, in argon.

Figure 4 presents the XRD patterns of the filter cakes subjected to the TG-DSC analysis (reacted samples). The results confirmed that the weight changes are due to oxidation and reduction of the iron and its oxides in the BOS filter cakes. In air, all the three filter cakes are nearly fully oxidized during heating to 800 °C, since Fe$_2$O$_3$ and ZnFe$_2$O$_4$ (both containing Fe$^{3+}$) are the dominant phases in the reacted samples as seen in Figure 4a–c. However, the weight gain resulted from BOS-1 was less than the weight loss caused by the dehydration and dissociation of the calcium carbonate. This led to the total mass loss in air for BOS-1. On the other hand, BOS-2 and BOS-3 are converted mainly to FeO phase in addition to metallic Fe in the argon atmosphere (Figure 4e,f). The slight difference of mass loss for BOS-1 in air and argon can be attributed to the oxidation of low contents of Fe, FeO and Fe$_3$O$_4$ to Fe$_2$O$_3$ as shown in Figure 4a,b [24].
The change pattern of the iron dissolution from BOS-2 was which seemed to be independent of the L/S stoichiometric ratio. It is inferred that acid concentration with an increase of the L/S stoichiometric ratio, and up to 30% of iron was dissolved at the ratio of 90%. A further increase in the acid concentration to 1.5 and 2.0 M caused the iron dissolution even below 1%. This behavior of iron with that of zinc, revealing the remarkable effect of the L/S stoichiometric ratio and the small impact of the acid concentration. At the low acid concentration of 0.5 M, the iron leaching efficiency increased significantly with an increase of the L/S stoichiometric ratio, and up to 30% of iron was dissolved at the ratio of 90%. Increasing the acid concentration to 1.0 M reduced the iron dissolution dramatically to less than 5%. Further increases did not show an obvious increase in zinc removal. Like BOS-3, acid concentration also had a slight effect on the leaching efficiency. It should be noted that the maximum zinc leaching efficiency was limited at about 55%, and no larger increase can be achieved by increasing acid concentration and L/S stoichiometric ratios. BOS-1 showed no increasing trend at all the above leaching conditions, indicating the poorest leaching performance of zinc in butyric acid solution.

The three BOS filter cakes were leached by butyric acid under the same leaching conditions and the corresponding zinc and iron leaching efficiencies were obtained. Figure 5 compares the leaching efficiency of zinc at different L/S stoichiometric ratios and acid concentrations. BOS-3 filter cake shows the highest efficiency, followed by BOS-2, while less than 10% of zinc was dissolved from BOS-1 at all the above leaching conditions. BOS-3 shows a strong dependence on the L/S stoichiometric ratio, and increasing the ratio caused considerable increase in the leaching efficiency of zinc at all the acid concentrations. However, the acid concentration did not have a significant impact on the zinc leaching efficiency. For BOS-2, the increase in the L/S stoichiometric ratio from 10% to 50% increased zinc removal considerably, but further increases did not show an obvious increase in zinc removal. Like BOS-3, acid concentration also had a slight effect on the leaching efficiency. It should be noted that the maximum zinc leaching efficiency was limited at about 55%, and no larger increase can be achieved by increasing acid concentration and L/S stoichiometric ratios. BOS-1 showed no increasing trend at all the leaching conditions, indicating the poorest leaching performance of zinc in butyric acid solution.

Figure 6 shows the iron dissolution from the three BOS filter cakes in relation to L/S stoichiometric ratio at various acid concentrations. Like zinc leaching from BOS-1, iron dissolution was less than 1% at all the leaching conditions tested. The change pattern of the iron dissolution from BOS-2 was complex. At the low acid concentration of 0.5 M, the iron leaching efficiency increased significantly with an increase of the L/S stoichiometric ratio, and up to 30% of iron was dissolved at the ratio of 90%. Increasing the acid concentration to 1.0 M reduced the iron dissolution dramatically to less than 5%. A further increase in the acid concentration to 1.5 and 2.0 M caused the iron dissolution even below 1%, which seemed to be independent of the L/S stoichiometric ratio. It is inferred that acid concentration played a crucial role in controlling the iron dissolution from BOS-2. BOS-3 shows the same leaching behavior of iron with that of zinc, revealing the remarkable effect of the L/S stoichiometric ratio and the small impact of the acid concentration.

It can be concluded that the leaching of BOS-3 filter cake reached the best outcome with butyric acid concentration at 1.0 M. Under this concentration, the optimal leaching conditions can be determined for different objectives to achieve as indicated by Supplementary Figure S1. To achieve the maximum zinc removal, a leaching for 10 h with L/S stoichiometric ratio at 90% can be adopted which can reach 85% zinc removal and 20% iron loss. However, a better selectivity of zinc can be attained by decreasing the ratio to 70% or reducing the leaching time to 2 h.
Figure 5. The leaching efficiency of Zn from three different BOS filter cakes at L/S stoichiometric ratios 10% to 90% and acid concentrations (a) 0.5 M, (b) 1.0 M, (c) 1.5 M and (d) 2.0 M.

Figure 6. The leaching efficiency of Fe from three different BOS filter cakes at L/S stoichiometric ratio 10% to 90% and acid concentration (a) 0.5 M, (b) 1.0 M, (c) 1.5 M and (d) 2.0 M.
Overall, the three BOS filter cakes showed various leaching behaviors of zinc and iron. BOS-1 showed a poor leachability of zinc and iron in butyric acid solutions, which is likely related to the self-sinter behavior resulting from long term stockpile of the waste, and some exothermic oxidation-sintering reactions probably occurred [25]. The obtained sample generally has stronger physical properties with a harder surface, higher density and strength, and a larger grain size in comparison with the fresh filter cake, which can make the acid leaching very difficult. Another reason is the higher content of zinc in ZnFe$_2$O$_4$ than that in ZnO as demonstrated by quantitative XRD analysis, and ZnFe$_2$O$_4$ was very stable and insoluble in traditional acid leaching process [17–20]. The best leaching selectivity of zinc over iron was achieved from BOS-2, which depends on the increasing acid concentration to form the hydrophobic butyric film adsorbed on Fe-containing phases hindering the leaching of Fe. It was reported that zinc exhibits amphoteric character and is soluble in a wide range of pH from 0 to 6, while iron dissolution mainly dependent on pH (see Supplementary Table S1). By increasing the acid concentration to form a hydrophobic protective film for the adsorption of butyric acid by iron ions rather than zinc ions, iron dissolution can be effectively limited. However, the maximum zinc leaching can only be reached without the unleachable zinc contained in ZnFe$_2$O$_4$. In comparison, BOS-3 tended to be most easily dissolved in terms of zinc and iron. This could be ascribed to the adequate storage of this fresh BOS filter cake to avoid the exothermic oxidation reaction in stockpiles causing a higher stability of the zinc phases contained in the leaching material, and the binding force was not strong enough to resist acid leaching. To conclude, the optimum leaching conditions for the selectively leaching of zinc over iron from the three BOS filter cakes, the resulted leaching efficiencies of zinc and iron, and the selectivity as the Zn/Fe ratio are summarized in Table 3 (BOS-2 Data from [23]).

### Table 3. The optimal conditions and the corresponding leaching efficiencies of the three BOS filter cakes.

| Sample | Zn Removal (%) | Fe Removal (%) | Zn/Fe Ratio (w/w) | Optimal Parameters |
|--------|----------------|----------------|-------------------|--------------------|
|        |                |                |                   | Acid Concentration (M) | L/S Stoichiometric Ratio (%) |
| BOS-1  | 8.4            | 1.2            | 0.8               | 0.5                | 90                      |
| BOS-2  | 51.2           | 0.47           | 12.6              | 1.5                | 70                      |
| BOS-3  | 84.6           | 20.0           | 0.5               | 1.0                | 90                      |

#### 3.3. Characterization of the Leached Residues

Figure 7 presents the XRD patterns of the leaching residues obtained under the optimal leaching conditions. No obvious change of the phases were observed for the residue of BOS-1 except for the disappearance of ZnO and some peaks of Fe, and this matched well with the corresponding lower zinc and iron leaching efficiencies. The same phases of Fe, FeO and Fe$_3$O$_4$ were still detected from the residue of BOS-2 due to the 0.47% iron dissolution. Apparently, the residue of BOS-3 showed remarkable changes of phases and the relative contents. For iron-containing phases, Fe was mostly dissolved while no obvious dissolution of FeO and Fe$_3$O$_4$ was observed. The disappearance of ZnO peaks from all the filter cakes implied that it was completely dissolved, or the remaining amount was below the limit of the detection. As expected, there was insignificant variation of ZnFe$_2$O$_4$ peaks.

Figure 8 presents the morphology of the corresponding leaching residues and there were some changes in contrast with the original samples. The shape and grain size of the BOS-1 residue are almost the same as before leaching. The EDS analysis also does not show much difference. This corresponds to the low leaching efficiencies of zinc and iron by butyric acid. For the residue of BOS-2, although the morphology generally unchanged after leaching, the overall zinc content appears to be decreased based on the EDS results. Considering that up to 50% of zinc was removed by leaching with very low loss of iron, it can be recognized that iron oxides forming the basic structure of the filter cake were not affected by the removal of zinc. For the residue of BOS-3, it seems that the fine particles were leached, leaving the residue with relatively coarse grains (comparable to BOS-1) and higher porosity, which is consistent with the high zinc and iron leaching efficiencies.
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Table 3. The optimal conditions and the corresponding leaching efficiencies of the three BOS filter cakes.

| Sample  | Zn Removal (%), Fe Removal (%), Zn/Fe Ratio (w/w), Optimal Parameters |
|---------|---------------------------------------------------------------|
| BOS-1   | 8.4, 1.2, 0.8, 0.5, 90                                        |
| BOS-2   | 51.2, 0.47, 12.6, 1.5, 70                                      |
| BOS-3   | 84.6, 20.0, 0.5, 1.0, 90                                       |

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Figure 7 presents the XRD patterns of the leaching residues obtained under the optimal leaching conditions. No obvious change of the phases were observed for the residue of BOS-1 except for the disappearance of ZnO and some peaks of Fe, and this matched well with the corresponding lower zinc and iron leaching efficiencies. The same phases of Fe, FeO and Fe3O4 were still detected from the residue of BOS-2 due to the 0.47% iron dissolution. Apparently, the residue of BOS-3 showed remarkable changes of phases and the relative contents. For iron-containing phases, Fe was mostly dissolved while no obvious dissolution of FeO and Fe3O4 was observed. The disappearance of ZnO peaks from all the filter cakes implied that it was completely dissolved, or the remaining amount was below the limit of the detection. As expected, there was insignificant variation of ZnFe2O4 peaks.

Figure 7. The XRD patterns of the corresponding leaching residues. (a) Residue of BOS-1; (b) Residue of BOS-2; (c) Residue of BOS-3.

Figure 8 presents the morphology of the corresponding leaching residues and there were some changes in contrast with the original samples. The shape and grain size of the BOS-1 residue are almost the same as before leaching. The EDS analysis also does not show much difference. This corresponds to the low leaching efficiencies of zinc and iron by butyric acid. For the residue of BOS-2, although the morphology generally unchanged after leaching, the overall zinc content appears to be decreased based on the EDS results. Considering that up to 50% of zinc was removed by leaching with very low loss of iron, it can be recognized that iron oxides forming the basic structure of the filter cake were not affected by the removal of zinc. For the residue of BOS-3, it seems that the fine particles were leached, leaving the residue with relatively coarse grains (comparable to BOS-1) and higher porosity, which is consistent with the high zinc and iron leaching efficiencies.

Figure 8. SEM micrographs of the corresponding leaching residues. (a) Residue of BOS-1; (b) Residue of BOS-2; (c) Residue of BOS-3.

4. Conclusions

Selective leaching of zinc over iron from three BOS filter cakes by butyric acid was investigated. The effects of acid concentration and L/S stoichiometric ratio were systematically examined. The following conclusions are obtained:

1. Among the three BOS filter cakes, BOS-3 showed the highest removal of zinc by butyric acid, while the best selectivity of zinc over iron was achieved with BOS-2. The BOS-1 filter cake showed the lowest leaching performance of zinc and iron.

2. The optimal leaching conditions for BOS-3 can be selected depending on the priority consideration. Considering both zinc removal and zinc selectivity, 90% L/S stoichiometric ratio and
4. Conclusions

Selective leaching of zinc over iron from three BOS filter cakes by butyric acid was investigated. The effects of acid concentration and L/S stoichiometric ratio were systematically examined. The following conclusions are obtained:

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2. The optimal leaching conditions for BOS-3 can be selected depending on the priority consideration. Considering both zinc removal and zinc selectivity, 90% L/S stoichiometric ratio and 1.0 M acid concentration for 10 h were chosen as the optimal conditions with 84.6% zinc removal and 20.0% iron loss.

3. BOS-1 probably have the self-sinter behavior resulting from long term stockpile with some exothermic oxidation reactions occurred, and those materials generally have stronger physical properties with harder surface, higher density and strength, and larger grain size which can make the acid leaching very difficult. The zinc leaching from BOS-2 was limited by the franklinite but can reach the maximum by increasing the acid concentration.

4. For the leaching behaviors of zinc and iron from the BOS filter cakes using butyric acid, mineral compositions played a more important role than the leaching parameters.

Supplementary Materials: The following are available online at http://www.mdpi.com/2075-4701/9/4/417/s1, Figure S1: Progress of leaching of (a) Zn and (b) Fe from BOS-3 using 1.0 M butyric acid with different L/S stoichiometric ratios, Table S1: The pH variations obtained from the leachates for the three BOS filter cakes under different leaching conditions.

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