Selective Removal of Zinc from BOF Sludge by Leaching with Mixtures of Ammonia and Ammonium Carbonate

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Abstract

The zinc content of basic oxygen furnace (BOF) sludges is too high for direct recycling into the blast furnace via the sinter plant, as excessive zinc concentrations are detrimental for the refractory lining of the blast furnace. However, by partial and selective removal of zinc from the BOF sludge, the residual sludge can be used as a secondary iron resource in the blast furnace. In this paper, BOF sludge was leached with aqueous ammonia, aqueous solutions of ammonium salts (chloride, carbonate, and sulfate), and aqueous mixtures of ammonia and ammonium salt. The mixtures of ammonia and ammonium salt could leach more zinc with respect to either the aqueous ammonia or the aqueous ammonium salt solution. The ammonia–ammonium carbonate (AAC) mixture was selected as the most suitable lixiviant due to the high zinc leaching efficiency in combination with a high selectivity towards iron; furthermore, this combination does not introduce unwanted chloride or sulfate impurities in the residue. The leaching process was optimized in terms of the liquid-to-solid ratio, total ammonia concentration, ammonium:ammonia molar ratio, temperature, and leaching time. The co-dissolved iron was precipitated as a hydroxide after oxidation of ferrous to ferric ions by an air stream, without co-precipitation of zinc, while the dissolved zinc could be easily recovered as zinc sulfide by precipitation with ammonium sulfide. The (almost) closed-loop process was successfully up-scaled from 10 mL to 1 L scale.

Graphical Abstract

Keywords Ammoniacal leaching · Ammonium salt leaching · Basic oxygen furnace sludge · Selective removal of zinc · Zinc recovery

Introduction

The average generation of basic oxygen furnace (BOF) dusts and sludges is approximately 22 kg/tonne of crude steel, with a minimum of 10 kg/tonne of crude steel and a
maximum of 40 kg/tonne of crude steel [1]. Due to improvements in the gas cleaning systems, more material is being collected rather than exiting into the atmosphere via exhaust stacks. This leads to slightly increasing production rates of BOF dusts and sludges over the years. Furthermore, generation of BOF dusts and sludges is expanding due to changes in process conditions, i.e., blowing rates, slag practices, bath additions, bath agitations.

BOF sludge predominantly consists of iron particles ejected from the BOF, which then partially oxidize within the gas cleaning system [1]. A study of the distribution of zinc in BOF off-gases of two ArcelorMittal steelmaking plants showed that the dust collected close to the BOF vessels contains much less zinc than the dust collected downstream in the off-gas cleaning system. The primary dust contains such a low zinc content, but rich in iron, that it has a good quality compared to virgin iron ores. Hence, it can be used as a secondary iron resource [2]. The average composition of the BOF sludge is about 60–75% iron, 0.3–1.6% of zinc, 1–2% carbon, and 1% lead [1]. A mineralogical study of BOF sludge from ArcelorMittal Monlevade (Brazil) showed that the following mineralogical phases were present both in the coarse and the fine fraction: wustite (FeO), magnetite (Fe₃O₄), metallic iron (α-Fe), lepidocrocite (γ-FeOOH), calcite (CaCO₃), and portlandite (Ca(OH)₂) [3].

The zinc distribution and zinc speciation in a zinc-rich (3.4% Zn) BOF sludge by micro-XRD and micro-XANES highlighted that the main zinc phases were franklinite (ZnFe₂O₄) and smithsonite (ZnCO₃) [5, 6]. The composition and phase distribution of BOF sludges are showing a large variability, depending on the production plant and specific campaigns.

Due to the high iron and metallic iron content of BOF dusts and sludges, their recycling and recovery is of utmost importance. However, the high concentration of zinc and lead in these residues inhibits the internal recycling via the sinter plant/blast furnace (BF) route. When the zinc content is too high for recycling via the sinter plant/BF route, some steel plants blend materials and produce briquettes/pellets, which are then charged back into the BOF vessels [1]. Another part of high-zinc BOF dust and sludge is temporarily stockpiled, and a significant part is recovered via the cement industry. There is a decreasing trend in landfilling of these materials, due to increasingly stringent environmental legislation and an increasing focus on waste minimization.

The high moisture content of BOF sludge is a major obstacle in its recycling [7].

Different leaching technologies have been reported for the recovery of zinc from BOF sludge. These routes comprise leaching with mineral acids [8–12], leaching with organic acids [13, 14], alkali leaching [3, 15], and ammoniacal leaching [16–18]. The pros and cons of these routes are discussed below, while Table 1 briefly summarizes the main conclusions and challenges.

Although sulfuric acid can leach high amounts of zinc, the high iron co-dissolution represents a main disadvantage. A process for removing zinc from BOF sludge by leaching with spent pickling liquor (HCl) was reported, where iron co-dissolution can be reduced by keeping the pH at 4–5 [12]. Organic acids have also been investigated for the recovery of zinc from BOF sludge [13, 14]. The use of organic acids is justified because they are biodegradable and environmentally benign, which implies that the generation of harmful secondary wastes can be avoided. Furthermore, leaching with organic acids is, in general, more selective than leaching with inorganic acids. The reported leaching tests aimed to maximize zinc recovery while minimizing iron co-dissolution. Butyric acid was found to be the most efficient lixiviant, with 49.7% Zn removal and 2.5% Fe leached. None of the organic acids could dissolve franklinite, which is a key phase in the BOF sludges.

Concurrently, alkaline leaching of BOF sludge with 5 M NaOH solution was found to be very selective for the removal of zinc over iron [3]. However, also in this case, the refractory franklinite phase was not dissolved. The zinc

| Table 1 Summarized analysis of the technologies for the recovery of zinc from BOF sludge |
|---------------------------------|---------------------------------------------------------------|
| Mineral acid leaching            | High zinc leaching efficiency but low selectivity of zinc over iron |
| Organic acid leaching            | Lower zinc leaching efficiency and higher selectivity compared to mineral acids |
| Alkali leaching                  | Very selective removal of zinc over iron |
| Ammoniacal leaching              | Very selective removal of zinc over iron |

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recovery was only between 40 and 60%. However, franklinite could be decomposed by thermal treatment with NaOH ($T = 450 \, ^\circ C$, $t = 1–5 \, h$, NaOH/sludge ratio $= 0.75$), and subsequently leached with 5 M NaOH resulting in 94% of zinc removal. However, the main disadvantage of the latter method is the large reagent consumption, which is detrimental for the cost-effectiveness of this treatment.

Although ammoniacal leaching has been widely explored for the valorization of by-products from the steelmaking industry [17, 19–26], it has barely been applied to BOF sludges. Only one study investigated the use of aqueous ammonia or aqueous solutions of ammonium salts for leaching of BOF sludge [18]. A recent patent describes the ammoniacal leaching of zinc from BF and BOF sludges, but a roasting pretreatment step was required to get high zinc dissolution [27]. In the present study, the leaching of BOF sludge using aqueous ammonia, an aqueous solution of ammonium salt, and also an aqueous mixture of ammonia and ammonium salt was investigated, in view of finding a cost-effective BOF sludge treatment methodology. The ammonia–ammonium carbonate (AAC) mixture was found to be the most suitable combination. The precipitation of the co-dissolved iron, and the recovery of the dissolved zinc from the pregnant leach solution was also studied. The present work proposes a closed-loop process for the recovery of zinc from BOF sludge using AAC leaching.

**Experimental**

**Chemicals**

Hydrochloric acid, HCl, (> 37%), ammonium sulfate, (NH$_4$)$_2$SO$_4$, (> 99.5%), and ammonia, NH$_3$, (25%) were purchased from VWR (Fontenay-sous-Bois, France). Nitric acid, HNO$_3$, (> 65%) and ammonium chloride, NH$_4$Cl, (> 99.8%) were obtained from Chem-Lab NV (Zedelgem, Belgium). Ammonium carbonate, (NH$_4$)$_2$CO$_3$, (99.5%) and ammonium sulfide solution, (NH$_4$)$_2$S, 20 wt% in water, were purchased from Sigma-Aldrich (Overijse, Belgium). A silicon solution in isopropanol was purchased from SERVA Electrophoresis GmbH (Heidelberg, Germany). Water was always of ultrapure quality, deionized to a resistivity of 18.2 MΩ cm with a Millipore ultrapure water system. All chemicals were used as received without any further purification. The BOF sludges were kindly provided by the ArcelorMittal steel plant in Bremen (Germany).

**Instrumentation**

The material was ground and sieved using a mortar grinder (Fritsch, Pulverisette 2) and a vibratory sieve shaker (Fritsch, Analysette 3). The metal content of the different samples was measured using a benchtop Total Reflection X-ray Fluorescence (TXRF) spectrometer (Bruker, S2 Picofox). X-ray powder diffraction (XRD) was used for the phase identification of the crystalline fraction (Bruker D2 Phaser). Experimental parameters for XRD analysis were $2\theta = 20^\circ–90^\circ$, radiation = CuK$_\alpha$, acceleration voltage = 40 kV, acceleration current = 40 mA, a step size of 0.020$^\circ$, and a counting time of 1 s per step, spin mode. The X’Pert HighScore software was used to analyze the collected data by comparison with the ICDD (International Centre for Diffraction Data). Electron probe microanalysis (EPMA) was performed on a JEOL JXA-8530FTMS-300 apparatus, operated at an acceleration voltage of 15 kV and an electrical current of $5 \times 10^{-8}$ A was used to characterize the BOF sludge. Thermoshakers from ThermoFisher Scientific were used to mix the vials of the leaching experiments. A Heraeus D-6450 oven was employed to dry the solid samples. For the large-scale experiments, a 1 L glass reactor (HiTec Zang, Herzogenrath, Germany) was used.

**Procedure**

The received samples were dried in a ventilated oven at 80 $^\circ$C for 48 h, milled and sieved (250 μm pore size mesh). The composition of the BOF sludge was determined by total digestion of the solid sample, followed by TXRF analysis. For the digestion, 5 mL of $aqua regia$ was added to a glass vial containing 50 mg of dried, milled, and sieved sludge. After 10 min, the lid was placed on the vial but it was not tightly closed to avoid pressure build-up due to the production of fumes. The mixture was left to react for 24 h. Afterwards, the metal content of the digested samples was measured via TXRF. Throughout the entire work, the sample preparation for the TXRF measurements was as follows: the sample was diluted in a nitric acid solution (2 vol%) and internal standards (ISs) were added (gadolinium and gallium). The dilution factor was chosen so that the final concentration was lower than 100 mg L$^{-1}$. Silicone solution SERVA (30 μL) was added on the carrier surface and dried for 5 min at 60 $^\circ$C in a hot air oven. A small droplet (5 μL) of the diluted sample was added onto the hydrophobized carrier. Then, the carrier was dried in a hot air oven for 30 min at 60 $^\circ$C. More experimental details about the TXRF analysis procedure can be found in the literature [28].
For the leaching experiments, 1 g of sample was added to a glass vial, the lixiviant was added, and the vial was closed and mixed using the Thermo shakers. After leaching, the pregnant leach solution and the residue were separated by syringe filtration (PET, 0.45 μm). Immediately after the filtration, the required amount of sample for the TXRF analysis was pipetted and placed in a vial containing the required amount of HNO₃ (2 vol%) for the dilution. In the case of multiple contact leaching experiments, the separation of the PLS was performed by vacuum filtration. The solid residue was dried in the oven for characterization before the next leaching experiment, and the composition of the PLS was analyzed. All the experiments were done in triplicate, and each sample within the triplicate was measured once. The standard deviations were calculated based on the measured values of this triplicate.

The precipitation of iron from the filtered PLS via both spontaneous oxidation of ferrous to ferric ions and forced oxidation by an air stream was investigated. For the spontaneous oxidation experiments, the filtered PLS was left in an open vial, while for the forced oxidation, air was bubbled through the samples. At the specified time intervals, a small aliquot of the sample was taken and analyzed via TXRF.

For the zinc recovery experiments, the required amount of ammonium sulfide was added to the filtered iron-free PLS and stirred at room temperature. Then the mixture was centrifuged and filtered. The zinc content of the remaining liquid was analyzed via TXRF. The precipitate was washed with MilliQ water and dried at 60 °C to determine its dry mass. The precipitate was digested in hydrochloric acid, while the metal content was measured by TXRF.

**Results and Discussion**

**Characterization of BOF Sludge**

The BOF sludges used in this work were collected before the filter press. The materials were dried, ground, and sieved, and only the fraction smaller than 250 μm was used (Fig. S1). The chemical composition was determined via digestion in aqua regia, followed by TXRF analysis. Two samples of BOF sludge were used in this work, namely BOF1 and BOF2. Both materials were from the same steel plant, but collected at different times. The composition of BOF1 was 55.37 ± 1.85 wt% iron and 6.73 ± 0.10 wt% zinc and that of BOF2 was 52.38 ± 1.76 wt% iron and 6.04 ± 0.20 wt% zinc. The mineralogical analysis of the powdered sample showed that the main phases present in the sludge were carbon (C), free iron (Fe), wüstsite (FeO), and calcite (CaCO₃), along with magnetite (Fe₃O₄) and/or franklinite (ZnFe₂O₄), and zincite (ZnO), (Fig. 1). This is in agreement with what has been previously reported in the literature [3–6]. The differentiation between magnetite and franklinite phases via XRD analysis is difficult due to the overlapping of their low-intensity characteristic diffraction peaks throughout the pattern, and because franklinite and magnetite are end members of solid solution series. Hence, the sample was also analyzed for elemental mapping of zinc, iron, and oxygen with EPMA-WDS. Backscattered electron (BSE) images of the powder along with respective elemental mappings are provided in Fig. 2 and in the supporting information (SI) (Figs. S2, S3, S4, and S5). The regions dominated by varying shades of turquoise and green are rich in iron indicating metallic iron or its oxide(s) (i.e., magnetite and wüstite). Zinc-rich regions are represented by red and pink indicating zinc oxide or zinc present in a solid solution between wüstite and zincite, (Fe,Zn)O. Free zinc cannot be present under BOF conditions. Blue regions represent oxides of metals in the sample other than zinc and iron. Finally, gray and yellow regions show the regions where zinc and iron are concentrated, indicating franklinite sites. Based on this analysis, it can be concluded that zinc is mainly distributed between its simple oxide and the hard-to-dissolve, refractory magnetite–franklinite solid solution phase.

**Selection of Lixiviant**

In the BOF sludge, both the zinc and (most of) the iron are present in the divalent state, as wüstite, zincite or a solid solution between wüstite and zincite. The ammoniacal leaching of zinc is thermodynamically feasible because of the favorable stability of the tetrammine zinc(II) complex [Zn(NH₃)₄]²⁺, as demonstrated by the predominance diagrams, and the high value of the stability constant [29]. The solubility of Fe(II) in aqueous ammonia is very low, although the hexaammine iron(II) complex [Fe(NH₃)₆]²⁺ can be formed under certain conditions [30]. When the ferrous ion Fe(II) in solution is oxidized to the ferric ion Fe(III), it precipitates as Fe(OH)₃, because the hexaammine iron(III) [Fe(NH₃)₆]³⁺ is not stable.
When BOF1 sludge was leached with aqueous ammonia ([NH₃] = 13.6 M, T = 60 °C; t = 3 h; liquid-to-solid ratio, L/S = 3 mL/g), only a small fraction of the zinc was dissolved (10.3%) with minor co-dissolution of iron (< 0.1%). The obtained solubility of zinc is much lower than what would be expected based on the stability of the [Zn(NH₃)₄]²⁺ complex. Unexpectedly low or negligible ZnO solubility has been previously reported in aqueous ammonia and NaOH solutions [3, 18, 19]. A possible reason for this behavior could be the high pH, at which the zinc complexes are not favorable, or that zinc could precipitate as Zn(OH)₂ [29]. A similar behavior has been reported for the ammoniacal leaching of chrysocolla, and it was suggested that free ammonia is not the active species in the ammoniacal leaching of copper (which is also based on the formation of tetraammine complexes), but rather the ammonium ion [31]. However, a more plausible reason for the limited leaching of zinc in ammonia solution is the fact that a counter anion is required for the charge balance with the [Zn(NH₃)₄]²⁺ complex and that the only anion present in ammonia solution is the hydroxide ion, which forms a poorly soluble complex with [Zn(NH₃)₄]²⁺.

The use of aqueous solutions of ammonium salts or aqueous mixtures of ammonia–ammonium salts increases the zinc leaching efficiency, due to the presence of anions stabilizing the [Zn(NH₃)₄]²⁺ complex (apart from OH⁻), as well as the lower pH associated to the presence of ammonium salts. BOF1 was leached with aqueous solutions of NH₄Cl, (NH₄)₂CO₃, (NH₄)₂SO₄, and {NH₄Cl + NH₃}, {(NH₄)₂CO₃ + NH₃}, and {(NH₄)₂SO₄ + NH₃} mixtures. For comparison purposes, the total {ammonium + ammonia} concentration ([NH₄⁺ + NH₃]) was kept constant at 3 M, and for the mixtures the ammonium:ammonia molar ratio, (NH₄⁺:NH₃), was kept at 1:1. The obtained results are present in Fig. 3.

Figure 3a shows that higher leaching efficiencies of zinc can be obtained with aqueous solutions of ammonium salts instead of aqueous ammonia. The leaching efficiency of zinc increased in the order (NH₄)₂CO₃ < (NH₄)₂SO₄ < NH₄Cl. The higher zinc leaching efficiency with NH₄Cl compared to (NH₄)₂CO₃ has been previously reported for BOF sludges [18]. Thermodynamic studies showed that the tetraammine zinc(II) complex is more stable than any of the chloro complexes, and that the role of the chloride ion is to stabilize the complex [29, 32]. The same has been reported for the carbonate system [17]. Therefore, the differences in the zinc solubility are attributed to the capacity of the anion salt to stabilize the tetraammine zinc(II) complex. An increase in the iron leaching efficiency was also noticed for the ammonia–ammonium salt system. It has previously been reported that the solubility of iron in the ferrous oxidation state can be surprisingly large in concentrated aqueous ammoniacal solutions containing an anion. In order to solubilize iron, ammonia needs to be added in excess with respect to the solute and a sufficient amount of conjugate ions need to be present to combine with a divalent cationic complex. These conditions are necessary to form the ammonia ligands. Otherwise, the ammonia is replaced by water and the iron becomes insoluble [30].

The addition of an excess of ammonia to the ammonium salt increased the leaching efficiency of zinc in all the cases, especially for (NH₄)₂CO₃. The zinc leaching efficiency of the ammonia–ammonium mixture increased in the order (NH₄)₂SO₄ < (NH₄)₂CO₃ < NH₄Cl. The excess of ammonia favors the zinc solubility due to the formation of soluble ammine complexes. The addition of NH₃ reduced the leaching efficiency for iron (i.e., increased the selectivity) in the (NH₄)₂CO₃ and the (NH₄)₂SO₄ systems, but not in the NH₄Cl system. The leaching efficiency for zinc is higher than that for iron in all the systems. However, the concentration of zinc in the PLS was not always higher due to the large difference in concentration of the initial residue (Fe = 55.0%, Zn = 6.7%). Figure 3 shows the composition of fresh PLS,
where the selectivity of the studied systems can be easily assessed. For the systems containing (NH₄)₂SO₄, the concentration of iron in the PLS was higher than that of zinc. Aqueous solutions of NH₄Cl are selective for zinc, but the selectivity decreased upon addition of an excess of ammonia. The systems containing (NH₄)₂CO₃ are the most selective towards iron. Based on the zinc leaching efficiency, both the {(NH₄)₂CO₃ + NH₃} and the {NH₄Cl + NH₃} mixtures are promising. However, the {(NH₄)₂CO₃ + NH₃} system is preferred due to the larger selectivity towards iron and the absence of corrosive chloride ions. The presence of chloride ions in the solid leaching residue could damage the blast furnace.

The filtered PLS obtained from the leaching of BOF sludge with ammonium salts or ammonia–ammonium salts mixtures was not stable over time; a brown precipitate appeared after the filtration of PLS. This precipitation corresponds to the formation of Fe(OH)₃ due to oxidation of the Fe(II) ion, dissolved as [Fe(NH₃)₄]²⁺, to the Fe(III) ion by contact with air. The precipitation might have started during the leaching step, but it continues after the filtration until all the dissolved iron has been precipitated as Fe(OH)₃. This behavior was not observed when leaching with aqueous ammonia due to the lower solubility of Fe(II) in the form of Fe(OH)₂.

The filtered PLSs were kept in a closed vial for 24 h (Fig. S6), and the composition was measured and compared to the composition of the PLS immediately after filtration. Figure 3 shows that the concentration of zinc in the PLS remained constant over time, while the concentration of iron had largely decreased. This phenomenon can be exploited to increase the selectivity for zinc, as will be shown later in this study. However, unless otherwise stated, the leaching efficiencies and compositions of the PLS have been calculated from the composition of the fresh PLS.

**Optimization of the Leaching Process**

AAC leaching was optimized in terms of liquid-to-solid ratio (Fig. 4, Table S1), NH₄⁺:NH₃ molar ratio (Fig. 4, Table S2), and total [NH₄⁺ + NH₃] concentration (Fig. 4, Table S3). Due to the limited availability of BOF1, the optimization was performed using a different BOF sludge sample (BOF2). The compositions of both residues were similar (see characterization section) and the leaching behavior was found to be identical (Fig. S7). The leaching efficiency for both zinc and iron increased while enhancing the L/S ratio (Fig. 4).
Table S1 shows that an increase in the L/S ratio decreased the selectivity against iron. Since the primary objective of this work was to obtain a solid residue with low zinc content to be fed into the blast furnace, high leaching efficiencies are preferred.

AAC system could leach more zinc than either aqueous ammonia or an aqueous solution of ammonium carbonate separately (Fig. 3). The effect of the NH₄⁺:NH₃ molar ratio on the leaching efficiency of zinc was investigated (Fig. 4b, Table S2), and an optimum NH₄⁺:NH₃ molar ratio of 1:1 was identified when leaching with [NH₄⁺ + NH₃] at 3 M. Further increase of ammonia or ammonium carbonate ratio decreased the zinc leaching efficiency. This is consistent with CO₃²⁻:NH₃ molar ratio of 1:4 required for the formation of the soluble [Zn(NH₃)₄]CO₃ species. This ratio has been reported as optimal in other studies, but, in general, there is disagreement on which value is optimal [17]. In another paper, the effect of the NH₄⁺:NH₃ ratio was reported to be negligible [20]. The latter used high total ammonia concentrations. In additional experiments performed with our BOF material, we have confirmed that at higher total ammonia concentrations the effect of the NH₄⁺:NH₃ ratio is negligible in the 2:1 to 1:2 region. Based on the literature data and our experimental results, it can be concluded that the optimal NH₄⁺:NH₃ is highly dependent on the total amount of ammonia and the mean solid-to-liquid ratio.

Figure 4 shows that increasing the total ammonia concentration increased the leaching efficiency for both zinc and iron. In this work, a total ammonia concentration of 4 M was selected as the optimal value because further increase in the concentration did not significantly enhance the leaching efficiency for zinc, but it increased the leaching efficiency for iron. At this concentration, the effect of the NH₄⁺:NH₃ ratio was negligible, and a ratio of 1:2 was selected for the further experiments.

The leaching time and temperature were simultaneously optimized for BOF2 (Fig. 5). Independently of the leaching temperature, the leaching efficiency of zinc increased with the leaching time until ≈ 24 h of leaching, after which the leaching of zinc decreased with time. This behavior has been previously reported in the literature; it was attributed to the precipitation of zinc as franklinite [18, 33]. Increasing the temperature enhanced the leaching efficiency for zinc. The
leaching efficiency for iron initially increased with the leaching time, but after certain time, it decreased. The time at which the maximum iron leaching efficiency was obtained, depended on the leaching temperature. The higher the temperature, the faster the maximum was reached. This behavior is attributed to the precipitation of iron in the form of Fe(OH)₃. Based on the obtained results, a leaching time of 6 h and a temperature of 60 °C can be considered as optimal, but longer leaching times at lower temperatures can produce the same results.

In an effort to reduce the amount of lixiviant, the performance of a multiple contact leaching was studied. At each step, the solid was contacted with fresh lixiviant at $L/S = 2.5$. Figure 6 shows that after two contacts, i.e., corresponding to a total $L/S$ of 5, the leaching of zinc was negligible. The leaching efficiency for zinc at 30 °C was about 60% already after just one contact, and it did not increase after multiple leaching steps. Therefore, the combination of low $L/S$ and low temperatures is not suitable for the leaching of zinc (Fig. 6). In the multiple contact leaching at 60 °C, the large error bars can be attributed to issues during the sample handling (i.e., the reason why the composition of the leaching residue after the second contact could not be measured, Table 2). Since this experiment was sequential, a significant cumulative measurement error occurred, but from the average values a trend can still be observed.

The composition of the solid residue after each contact is shown in Table 2. The results corroborate that the zinc concentration did not further decrease after the second leaching contact, while the iron concentration did. The zinc content of the leaching residue obtained after two leaching steps at

| Contact number | $T = 30^\circ$C | | $T = 60^\circ$C |
| --- | --- | | --- |
| Fe (wt%) | Zn (wt%) | Fe (wt%) | Zn (wt%) |
| 0 | 55.7±1.8 | 5.5±0.5 | 55.7±1.8 | 5.5±0.5 |
| 1 | 50.7±1.2 | 2.9±0.1 | 47.7±2.1 | 2.6±0.1 |
| 2 | 50.0±1.3 | 2.7±0.1 | 48.1±1.1 | 1.8±0.2 |
| 3 | 48.2±0.3 | 2.6±0.1 | 43.3±1.5 | 1.8±0.1 |
| 4 | 48.6±1.2 | 2.8±0.1 | 43.3±1.5 | 1.8±0.1 |

The leaching conditions were $t = 3$ h; $L/S = 2.5$ (mL/g) each step; $\text{NH}_4^+:\text{NH}_3 = 1:2$; $[\text{NH}_4^+ + \text{NH}_3] = 4$ M

Figure 5 Effect of the leaching time on the leaching efficiency of BOF sludge at different temperatures of: a zinc and b iron. The leaching conditions were $L/S = 10$ (mL/g); $\text{NH}_4^+:\text{NH}_3 = 1:2$; $[\text{NH}_4^+ + \text{NH}_3] = 4$ M

Table 2 Composition of dried BOF sludge residue after leaching steps
60 °C corresponded to a zinc leaching efficiency of approximately 72%. A single contact leaching experiment under the same conditions but \( L/S = 5 \) showed a zinc leaching efficiency of 66 wt%. From the obtained results, it appears that multiple contact leaching does not significantly boost the zinc leaching efficiency, rendering a two-step process irrelevant.

**Upscaling of the Leaching Process**

The optimized leaching process was up-scaled from 10 mL to 1 L. The selected leaching conditions were \( t = 6 \) h; \( T = 60 \) °C; \( L/S = 10; \) \( \text{NH}_4^+ : \text{NH}_3 = 1:2; \) \( [\text{NH}_4^+ + \text{NH}_3] = 4 \) M. The upscaling experiment was performed using a second batch of BOF2 with composition 55.70 ± 1.81 wt% iron and 5.53 ± 0.47 wt% zinc. The results of the upscaling experiment showed comparable leaching yields for zinc (76 wt% instead of 81 wt%) and iron (0.7 wt% instead of 1.3 wt%). The leaching residue obtained after filtration (see Fig. S8) was dried and characterized. The zinc content of the BOF sludge was reduced from 5.5 to 1.5 wt%, and the iron content decreased from 55.7 to 42.3 wt%. The iron composition in the leaching residue was lower than what would be expected based on the leaching efficiency. The reason is that FeO from the initial residues was transformed into Fe(OH)\(_3\); therefore, the relative mass of the residue increased.

From the optimization and upscaling of the leaching process, it can be concluded that if the temperature/time and the total ammonia concentration are sufficiently high, a maximum zinc leaching efficiency of about 80% can be achieved. This leaching efficiency indicates the fraction of zinc present as ZnO or (Fe,Zn)O solid solution. The remaining zinc is present as a solid solution of magnetite and franklinite, and cannot be leached via AAC leaching. Therefore, the feasibility of the process to produce a solid residue that can be fed into the BF relies on the initial franklinite content of the BOF sludge.

**Purification of the PLS**

It has previously been shown in Fig. 3 that the iron in the solution precipitated as Fe(OH)\(_3\) by oxidation of the hexaammine iron(II) complex, \([\text{Fe(NH}_3)_6]^2+\). Therefore, the precipitation of iron from the filtered PLS was further studied as a function of time. The iron precipitation was investigated by leaving the filtered PLS in an open vial (spontaneous oxidation) and by bubbling air through the sample (forced oxidation). In both cases the concentration of iron in the PLS was measured as a function of time (Fig. 7). Although all the iron in solution could be precipitated without air addition, air bubbling clearly accelerated the process. In both cases, minor co-precipitation of zinc was observed. An XRD spectrum and an image of the obtained XRD precipitate are shown in Figs. S9 and S10, respectively. The precipitation of iron via air bubbling before filtration was also tested, and full precipitation of iron without zinc co-precipitation was achieved. The precipitation of iron after the leaching, instead of during the leaching, is preferred because it can be performed at lower temperature and avoids the evaporation of ammonia.

The recovery of the dissolved zinc from the PLS by precipitation as ZnS by addition of ammonium sulfide was investigated. A stoichiometric amount of ammonium sulfide could precipitate 95% of the zinc present in solution, while a 10% excess of ammonium sulfide could precipitate 100% of the zinc. The obtained precipitate was dissolved and iron could not be detected by the TXRF. Other zinc recovery methods have been reported in the literature and could be potentially applied to this process [17]. After precipitation of iron and zinc, the lixiviant can be re-used for leaching, but ammonia or ammonium carbonate make-up will be necessary to maintain the appropriate \( \text{NH}_4^+ : \text{NH}_3 \) ratio.

**Conceptual Flowsheet**

Figure 8 shows a conceptual flowsheet of the process proposed in this paper. The dried BOF sludge was leached with an AAC mixture. The optimal leaching conditions were identified as \( t = 6 \) h; \( T = 60 \) °C; \( L/S = 10; \) \( \text{NH}_4^+ : \text{NH}_3 = 1:2; \) \( [\text{NH}_4^+ + \text{NH}_3] = 4 \) M. This process could leach all the zinc, except that in the form of franklinite, while only a very small fraction of iron was co-dissolved. The co-dissolved iron could be precipitated by bubbling
air through the PLS. The air can be bubbled directly in the leaching reactor immediately after leaching. The undisolved solids, together with the precipitated iron can be filtered and recycled to the blast furnace. The iron-free PLS was contacted with an aqueous solution of ammonium sulfide for the recovery of the dissolved zinc as zinc sulfide. The obtained zinc sulfide can be sold as a by-product to the zinc smelting industry. The depleted lixiviant can be re-used in multiple leaching steps. A certain amount of ammonia or ammonium carbonate must be added to maintain the optimum AAC ratio.

**Conclusions**

The recovery of zinc from BOF sludge via ammoniacal leaching was investigated. Aqueous ammonium salt–ammonia mixtures were found to leach more zinc than either the aqueous ammonia or an aqueous ammonium salt solution. The ammonia–ammonium carbonate (AAC) mixture showed the best compromise between zinc leaching efficiency and zinc over iron leaching selectivity. The iron of the pregnant leach solution present as hexaammine iron(II) complexes was precipitated as Fe(OH)₃ by oxidation with air. The oxidation process could be accelerated by bubbling air through the pregnant leach solution, while the dissolved zinc could be recovered as zinc sulfide by the addition of ammonium sulfide. Finally, the (almost) closed-loop process was successfully up-scaled in a 1 L reactor.

Unfortunately, for this specific BOF sludge the final zinc concentration in the treated BOF sludge was still too high to allow its recyclability to the blast furnace, given the extremely strict zinc constraints enforced by ArcelorMittal Bremen. Nonetheless, the presented AAC process could be suitable for other BOF sludges with lower frankinite content and/or steel plants with less strict Zn-limits for internal recycling into the blast furnace.

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**Compliance with Ethical Standards**

**Conflict of interest** On behalf of all authors, the corresponding author states that there is no conflict of interest.

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