Original Article

Solidification/Stabilization of Arsenic in Red Mud upon Addition of Fe (III) or Fe (III) and Al (III) Dissolved in H₂SO₄

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ABSTRACT

In the aluminum industry, as an alkaline waste, red mud is composed of Fe and Al. It is difficult to safely dispose certain types of some arsenic-containing red mud owing to hazardous arsenic leaching. Hence, the method of arsenic stabilization using Fe (III) or H₂SO₄ to reduce arsenic leaching from red mud was investigated. The results indicated that the addition of 3−5% Fe₂(SO₄)₃ or H₂SO₄ (ratio of H⁺ (mM)/red mud (g) ≈ 1.5) into red mud and maintaining the pH at 4 until dryness was achieved at room temperature, decreased arsenic solubility from 360 to 2−6 µg/L (< 10 µg/L: Environmental criteria in Japan) and maintained the same for several months. In the mechanism of arsenic stabilization, Fe³⁺ ions added or Fe³⁺ and Al³⁺ ions dissolved from red mud by adding H₂SO₄ reacted with arsenic and formed insoluble Fe–As and Al–As compounds. Through X-ray diffraction analysis, insoluble arsenic compounds including iron arsenate, iron hydrogen arsenate, aluminum arsenate, and aluminum arsenate hydrate were detected and considered as responsible for low arsenic leaching. Additionally, the results suggested that HCl was not appropriate for stabilizing arsenic in red mud because the high amounts of Cl⁻ negatively influence the formation of insoluble arsenic compounds.

Keywords: solidification/stabilization of arsenic, red mud, Fe (III) Al ions, H₂SO₄

INTRODUCTION

Red mud as a bauxite residue is produced in alumina plants by the Bayer process in which Al-containing minerals are dissolved in hot NaOH solution to extract Al. Globally, approximately 120 million t of red mud are annually produced, and the global total discharged amount exceeded more than 3.5 billion t in 2014 [1]. Typically, the chemical properties of red mud include high alkalinity (pH = 9.7 − 12.8), elevated salinity, and elevated concentrations of Al and Fe.

Red mud is a naturally occurring heterogeneous material that is mainly composed of one or several aluminum hydroxide minerals, including primarily gibbsite [Al(OH)₃], boehmite [γ-AlO(OH)], and diaspore [α-AlO(OH)]. Except these aluminum hydroxide minerals (Average wt.%: 26 − 60), most red mud is also rich in oxides of Fe (Average wt.%: 10 − 35), Ti (Average wt.%: 2 − 4), Si (Average wt.%: 4 − 8), and some special elements such as Zn, Ni, and V [2−4]. With the exception of inherent high alkalinity, the leakage risk of toxic elements from red mud could also limit its reclamation and application, such as potentially toxic metals and metalloids: As, Cr, Ni, Pb, and Mo [5, 6]. In Minas Gerais State, Brazil, the red mud had an arsenic content of 19.3 mg/kg [7]. In Ajka, Hungary, red mud samples contained elevated concentrations (150 mg/kg) of potentially toxic arsenic (As). And in aqueous solution, Ajka red mud had dissolved As concentrations of 4 − 6 mg/L [8−10]. Many studies examine As due to its potential harmful effect on human health. Permanent As intake leads to chronic intoxication, and prolonged As exposure can damage the central nervous system, liver, and skin and results in the appearance of diverse types of cancers such as hyperkeratosis, lung, and skin cancer [11]. A wide range of studies related to As removal were conducted to reduce the risk that underwater As contamination poses to human health. Currently, several treatment methods such as adsorption, ion exchange, reverse osmosis, nanofiltration, coprecipitation, membrane distillation, and biological procedures are increasingly used for the removal of As [12−15]. Among these researches, the sorption and coprecipitation of As by Fe (II / III) compounds were widely conducted. The
main mechanisms of these include the adsorption of arsenate via surface complex on iron oxides (hematite goethite) and ferrihydrite mainly as an inner-sphere bidentate complex [16–19] and the formation of insoluble Fe-As compounds, ferric arsenate, scorodite and poorly crystalline ferric arsenate [20–22]. Additionally, Al ions, hydroxide and oxide of aluminum also showed good behaviors in adsorbing arsenic as well as Fe [23, 24].

The aims of the study involved decreasing the high alkalinity, reducing As leaching concentration from red mud (less than 10 ppb, satisfying Environment Agency Notification No.18 in Japan) and making this red mud safely usable. Given the effective As stabilization by Fe and to decrease the high pH of red mud, Fe ions were added into red mud to control As leaching concentration. And given that the main components of red mud were Fe and Al, inorganic acid was used to release Fe and Al ions and decrease the pH of red mud. In the experiments, additive amounts of an Fe reagent and inorganic acids, pH variation of the disposed red mud, and the long-term effect of As stabilization were investigated to optimize the reaction conditions. Moreover, the study also examined the mechanism of solidification/stabilization of As in red mud with the addition of Fe (III) or inorganic acids.

MATERIALS AND METHODS

Materials and analytical instruments

As-containing red mud was supplied by an aluminum company. Several million tons of the red mud were in the final disposal site for several decades. A Ferric Sulfate solution (assay as Fe$_2$(SO$_4$)$_3$: 64%) was supplied by JUJO Synthetic Chemistry Labo, Tokyo Japan. Additionally, FeSO$_4$·7H$_2$O (assay as FeSO$_4$: 55%) was provided by Wako Pure Chemical Industries, Ltd., Osaka Japan. It was considered that ferrous in solution was oxidized easily in air, so to avoid the influence of oxidization, FeSO$_4$ was used as solid. The proportion of additive amounts used in subsequent experiments, namely Ferric Sulfate solution and FeSO$_4$·7H$_2$O, were calculated as Fe$_2$(SO$_4$)$_3$ and FeSO$_4$. For example, the use of 5% Fe$_2$(SO$_4$)$_3$ or FeSO$_4$ to dispose 1.0 g red mud indicated that the actual additive amount corresponds to 0.078 mL Ferric Sulfate solution or 0.091 g FeSO$_4$·7H$_2$O. All chemicals were of analytical grade and used without further purification. Deionized (DI) water was used for all the experiments. All glassware were cleaned by soaking in 5% HNO$_3$ and rinsed thrice with DI water prior to use.

The concentration of As in the solution was determined using an atomic absorption spectrophotometer (AA7000, Shimadzu Corporation, Kyoto, Japan) and a hydride generator (Shimadzu Corporation). The detection limit of the instrument corresponded to 0.1 µg/L. A solution containing 0.4% NaBH$_4$ and 0.5% NaOH was mixed with 15% HCl to generate hydride. Prior to the analyses, a sample solution was mixed with a solution (20% potassium iodide) and (10% concentrated hydrochloric acid) at volume proportion of 8:1:1 for the reduction from As (V) to As (III). Finally, As (III) in solution was reacted with a hydride to form a hydride of As, which was directly determined by AAS.

Solid samples were characterized using an Ultima IV X-ray diffractometer (Rigaku Corporation, Tokyo, Japan) that was equipped with a copper target and a D/tek Ultra detector. The equipment was operated at 40 kV and 40 mA by step scanning from 10° to 80° in increments of 0.02° 2θ. Integrated X-ray powder diffraction software was used in the analysis. The shapes of the surface of the solid samples were analyzed on a SU8000 (Hitachi Corporation, Tokyo, Japan) scanning electron microscope. All the samples were imaged at 5 kV. Compositions of red mud were determined using a SU8000 (Hitachi Corporation) energy dispersive X-ray spectrometer at 20 kV.

Composition of red mud

Red mud was dried at 70°C for 3 h to calculate the moisture content and dried red mud was sieved (with an approximate particle size corresponding to 0.149 mm) and analyzed by X-ray diffraction (XRD) and Energy Dispersive X-ray (EDX) to characterize the mineralogical characteristics and composition, respectively.

Arsenic content test

Dried red mud (7.50 g) was mixed with 250 mL of 1 M HCl and shaken for 2 h using a shaker (SA 300, Yamato, Tokyo, Japan). The leachate was filtered using a 0.45-µm membrane, and the As concentration was determined by atomic absorption spectrophotometer (AAS). The As content in red mud was calculated based on the As concentration [25].

Arsenic leaching test

Red mud sample (10.0 g) was mixed with 100 mL pure water and shaken for 6 h. After a centrifugal separation, the leachate was filtered using a 0.45-µm membrane, and the pH value was measured. Arsenic concentration in the leachate was analyzed using hydride generator and atomic absorption spectrophotometer (HG-AAS), and this was considered as corresponding to the As leaching concentration from red mud [26].
Stabilization of arsenic in red mud by Fe (II/III) ions

**Stabilization test**

Red mud was mixed with Ferric Sulfate solution or FeSO\(_4\cdot7\)H\(_2\)O by iron spoon with hand, and the pH of the mixture was measured by the pH test paper. The red mud was placed at room temperature for 24 h to dry, and the samples were analyzed by Scanning Electron Microscopy (SEM). Arsenic concentration and pH of leachate was determined in the leaching test. In a few experiments, cement (cement number: Pacific S-GS211 marked as C in later statement) and pieces of old newspapers (0.1 − 0.3 cm\(^2\); marked as P in later statement) were also added into red mud with Fe reagents to enhance the strength of the red mud.

**Simulation test**

In order to investigate the stable As compounds potentially generated in the above operations, 5 mL As\(^{5+}\) solution (Na\(_2\)HAsO\(_4\)·7H\(_2\)O 0.15 g) and 5 mL Fe\(^{3+}\) solution (Fe\(_2\)(SO\(_4\))\(_3\) 1.92 g) were mixed, and the pH was adjusted to 4 using NaOH solution. The solution (Fe\(^{3+}\):As\(^{5+}\) = 20:1) was placed in room temperature for 6 h, and filtration was performed. The resultant product was dried at 70°C for 12 h and examined through XRD analysis.

Stabilization of arsenic in red mud by inorganic acids (H\(_2\)SO\(_4\), HCl)

**pH adjustment test**

In order to verify the relationship between pH of leachate and As adsorption by red mud, 2.0 g red mud and 20 mL pure water were mixed, and the pH of the solution was changed from 3 to 10 by using HCl. These samples were tumbled in a shaker for 6 h. Arsenic leaching concentration and the pH of the leachate was determined.

**Stabilization Test**

Red mud was mixed with a certain amount of H\(_2\)SO\(_4\) or HCl and the pH of the mixture was then measured. The red mud was placed at room temperature for 24 h to dry, and the samples were analyzed by SEM. Arsenic concentration and the pH of the leachate was determined in the leaching test.

**Simulation Test**

To examine the stable As compounds potentially generated in the forementioned operations in which H\(_2\)SO\(_4\) was added, 5 mL As\(^{5+}\) solution (Na\(_2\)HAsO\(_4\)·7H\(_2\)O 0.15 g), 5 mL Fe\(^{3+}\), and Al\(^{3+}\) solution (Fe\(_2\)(SO\(_4\))\(_3\) 0.96 g; Al\(_2\)(SO\(_4\))\(_3\) 0.82 g; Na\(_2\)SO\(_4\) 0.2272 g) were mixed and the pH was adjusted to 4 using NaOH solution. The solution (SO\(_4^{2-}\):Fe\(^{3+}\):Al\(^{3+}\):As\(^{5+}\) = 33:10:10:1) was mixed for 20 min and filtration was performed. Furthermore, in the simulation test in which HCl was added, 5 mL As\(^{5+}\) solution (Na\(_2\)HAsO\(_4\)·7H\(_2\)O 0.15 g), 5 mL Fe\(^{3+}\), and Al\(^{3+}\) solution (FeCl\(_3\)·6H\(_2\)O 1.29 g; AlCl\(_3\)·6H\(_2\)O 1.16 g; NaCl 0.19 g) were mixed and the pH was adjusted to 4 using NaOH solution. The solution (Cl\(^-\):Fe\(^{3+}\):Al\(^{3+}\):As\(^{5+}\) = 66:10:10:1) was mixed for 20 min and filtration was performed. The two types of resultants were dried at 70°C for 12 h and examined through XRD analysis. In the experiment, SO\(_4^{2-}\) and Cl\(^-\) concentrations were calculated based on the actual added amounts of H\(_2\)SO\(_4\) and HCl in the red mud.

**Long-term stability test**

The red mud disposed by 3% and 5% Fe\(_2\)(SO\(_4\))\(_3\) and inorganic acids (H\(_2\)SO\(_4\), HCl) with a ratio of H\(^+\) (mM)/red mud (g) corresponding to 1.5 mM/g were exposed to air for durations corresponding to 15, 30, 60 and 90 d. Arsenic leaching concentrations were determined in this period. After disposal, the samples were analyzed using SEM to compare the changes in surface structure prior to disposal.

**RESULTS AND DISCUSSION**

**Composition of Red Mud**

The moisture content of the red mud was determined as 20%, and the pH corresponded to 12. The proportion of elements is listed in **Table 1**. Specifically, Fe, Al, and O occupied a large proportion in the red mud. The XRD analysis (not shown in the study) indicated that the main components of the red mud corresponded to hematite (Fe\(_{1.67}\)H\(_{0.99}\)O\(_3\)), aluminum oxide (Al\(_2\)O\(_3\)), and a large amount of amorphous substances that were mainly composed of iron oxide hydroxide (FeOOH) and aluminum hydroxide oxide

<table>
<thead>
<tr>
<th>Elements</th>
<th>Concentration of atoms [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>O</td>
<td>65.29</td>
</tr>
<tr>
<td>Fe</td>
<td>9.82</td>
</tr>
<tr>
<td>Al</td>
<td>9.18</td>
</tr>
<tr>
<td>Na</td>
<td>7.66</td>
</tr>
<tr>
<td>Si</td>
<td>5.56</td>
</tr>
<tr>
<td>Ti</td>
<td>1.33</td>
</tr>
<tr>
<td>Ca</td>
<td>0.9</td>
</tr>
<tr>
<td>Cl</td>
<td>0.14</td>
</tr>
<tr>
<td>S</td>
<td>0.12</td>
</tr>
<tr>
<td>As</td>
<td>0.01</td>
</tr>
</tbody>
</table>

**Table 1** Elemental composition of the red mud.
(AlO(OH)) at the specific high pH. The total content of As in red mud approximately corresponded to 27.5 mg/kg. In the leaching test, the concentration of As leaching from red mud approximately corresponded to 360 ppb, which significantly exceeded the upper limit (10 µg/L) described in Japan Environment Agency Notification 18. The As (V) in the high pH condition mainly existed as AsO$_3^{2-}$ and AsO$_4^{3-}$ species, and As (III) may occur as both AsO$_2^{2-}$ and AsO$_3^{3-}$ species [27]. Soluble As compounds, namely iron arsenic (FeAs), svenekite (Ca(H$_2$AsO$_4$)$_2$), and calcium arsenide (CaAs), were also detected in red mud by XRD analysis. Svenekite is a soluble compound that is reported in extant studies [28], and iron arsenic and calcium arsenide are unstable in water. The forementioned compounds are considered as the main source of As leaching.

### Stabilization of Arsenic in Red Mud by adding Fe (II/III) ions

Red mud was disposed by Fe (II/III) reagents as described above. Arsenic leaching concentrations are listed in Table 2. In this case, As leaching decreased to approximately 4 − 7 ppb (Table 2, No.4,5,7,8) with the addition of Fe (II/III) ions, and this was less than the upper limit (10 µg/L) of environmental criteria in Japan with respect to As leaching concentration. When red mud was mixed with Fe reagents, the pH of the mixture approximately corresponded to 4 − 5. Additionally, the pH of leachate changed from 10 to approximately 6 − 8. There were various reasons for the stabilization of As with the addition of Fe (II/III) ions. First, the added Fe (II/III) ions reacted with As to form some insoluble Fe−As compounds such as poorly crystallized ferric arsenate. Furthermore, Fe$_{1.67}$H$_{0.99}$O$_3$, Al$_2$O$_3$, FeOOH, and AlO(OH) in the red mud were unable to adequately adsorb As in an alkali condition with respect to the surfaces of adsorbents prior to the addition of Fe (II/III) ions. The As ions were both negative and coulombic repulsions between the negatively charged ion and negatively charged surface significantly decreased the arsenate adsorption rate [29]. The added ferric and ferrous ions consumed OH$^-$ and generated ferric/ferrous hydroxide and ferrihydrite. This decreased the pH of the red mud and promoted the adsorption of As by the red mud itself. Moreover, the newly generated ferric/ferrous hydroxide and ferrihydrite fixed more As by coprecipitation.

### Table 2 Arsenic leaching concentrations at the stabilization test with the addition of Fe (II/III) ions.

<table>
<thead>
<tr>
<th>NO.</th>
<th>Matrices Composition</th>
<th>Arsenic leaching [ppb]</th>
<th>pH of leachate [-]</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Red mud</td>
<td>360.0</td>
<td>10.0</td>
</tr>
<tr>
<td>2</td>
<td>Red mud + 3.0% Fe(NO$_3$)$_3$</td>
<td>7.5</td>
<td>7.5</td>
</tr>
<tr>
<td>3</td>
<td>Red mud + 1.5% Fe$_2$(SO$_4$)$_3$</td>
<td>16.2</td>
<td>7.7</td>
</tr>
<tr>
<td>4</td>
<td>Red mud + 3.0% Fe$_2$(SO$_4$)$_3$</td>
<td>7.4</td>
<td>7.2</td>
</tr>
<tr>
<td>5</td>
<td>Red mud + 5.0% Fe$_2$(SO$_4$)$_3$</td>
<td>4.2</td>
<td>6.6</td>
</tr>
<tr>
<td>6</td>
<td>Red mud + 1.5% FeSO$_4$</td>
<td>14.0</td>
<td>8.2</td>
</tr>
<tr>
<td>7</td>
<td>Red mud + 3.0% FeSO$_4$</td>
<td>6.8</td>
<td>7.0</td>
</tr>
<tr>
<td>8</td>
<td>Red mud + 5.0% FeSO$_4$</td>
<td>6.2</td>
<td>6.5</td>
</tr>
<tr>
<td>9</td>
<td>Red mud + 3.0% C + 3% P</td>
<td>14.2</td>
<td>11.4</td>
</tr>
<tr>
<td>10</td>
<td>Red mud + 1.5% Fe$_2$(SO$_4$)$_3$ + 3% C + 3% P</td>
<td>10.4</td>
<td>8.8</td>
</tr>
<tr>
<td>11</td>
<td>Red mud + 3.0% Fe$_2$(SO$_4$)$_3$ + 3% C + 3% P</td>
<td>6.2</td>
<td>8.5</td>
</tr>
<tr>
<td>12</td>
<td>Red mud + 5.0% Fe$_2$(SO$_4$)$_3$ + 3% C + 3% P</td>
<td>6.8</td>
<td>8.1</td>
</tr>
<tr>
<td>13</td>
<td>Red mud + 1.5% FeSO$_4$ + 3% C + 3% P</td>
<td>28.7</td>
<td>8.6</td>
</tr>
<tr>
<td>14</td>
<td>Red mud + 3.0% FeSO$_4$ + 3% C + 3% P</td>
<td>12.3</td>
<td>8.5</td>
</tr>
<tr>
<td>15</td>
<td>Red mud + 5.0% FeSO$_4$ + 3% C + 3% P</td>
<td>11.7</td>
<td>8.2</td>
</tr>
</tbody>
</table>

C: Cement  P: Old newspapers (0.1 − 0.3 cm$^2$)
to stabilize As. However, in an alkali condition, most $\text{Fe}^{2+}$ quickly changed into $\text{Fe(OH)}_2$, and only a low amount of $\text{Fe}^{2+}$ reacted with As. Therefore, it is better to use $\text{Fe}^{3+}$ than $\text{Fe}^{2+}$ to dispose the red mud in this case. In air, a few white or transparent crystals also appeared in the surface of red mud, which is discussed in a later section.

**Stabilization of Arsenic in the Red Mud by Inorganic Acids**

The results confirmed that high alkali condition did not contribute to the adsorption of As on oxide or hydroxide of Fe/Al that largely existed in the red mud. The results of the pH adjustment test are shown in Fig. 1. The variation trend of As leaching suggested that As adsorption by red mud itself was highly pH dependent. In both low and high pH conditions, As was not adequately adsorbed by the red mud. However, at a leachate pH near 5, the As concentration decreased to 10 ppb, and this suggested that oxide or hydroxide of Fe/Al in red mud exhibited a strong adsorption capacity of As in a weakly acidic condition.

As shown in Fig. 1, slight changes in the leachate pH significantly changed the As concentration, and this indicated that the adsorption was not so stable. Despite this, it was more advantageous to clearly maintain the pH of the system as weakly acidic or neutral to reduce As leaching. When the pH decreased, a low amount of Fe and Al ions were also released from the red mud, and this is considered as helpful in the stabilization of As.

Furthermore, $\text{H}_2\text{SO}_4$ and $\text{HCl}$ were added into the red mud to decrease the pH of red mud to weakly acidic or neutral and release additional Fe and Al ions to stabilize the arsenic (Table 3). The results revealed that excess acids could increase the dissolution of As. When the ratio of $\text{H}^+$ (mM) and red mud (g) approximately corresponded to 1.5 mM/g (Fig. 2), the pH of mixture was maintained at 4, and arsenic leaching was approximately limited to 1.6 ppb (by $\text{H}_2\text{SO}_4$) and 2.5 ppb (by $\text{HCl}$) with the pH of the leachate approximately corresponding to 6.1 – 6.3. This constitutes significantly low As leaching when compared with the upper limit (10 µg/L) of the environmental criteria in Japan with respect to As leaching concentration.

The As leaching with the addition of $\text{H}_2\text{SO}_4$ or $\text{HCl}$ ($\text{H}^+$ (mM) and red mud (g): 1.5 mM/g) was significantly lower than that with the addition of Fe (III) ions. This was attributed to the addition of inorganic acids that cause a significant amount of Fe and Al ions to be released from the red mud, and the As in red mud could react with Fe/Al ions in the process of mixing and placing in air to form insoluble compounds at a pH of 4. Although the weakly acidic condition was advantageous for As adsorption, it did not reduce As leaching to the forementioned extremely low level. Thus, the formation of insoluble arsenic compounds is the main reason why As was successfully stabilized. The use of inorganic acids to dispose of As-containing red mud is an effective and stable method to stabilize As. Additionally, with respect to the viewpoint of the best stabilization effect, the addition of inorganic acid is a better proposition than the addition of Fe.

**Table 3** Arsenic leaching concentration from the red mud with the addition of inorganic acids.

<table>
<thead>
<tr>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>2.0 g Red mud + HCl</td>
<td>2.00</td>
<td>0.50</td>
<td>16.9</td>
<td>7.6</td>
</tr>
<tr>
<td></td>
<td>2.00</td>
<td>1.00</td>
<td>9.2</td>
<td>6.3</td>
</tr>
<tr>
<td></td>
<td>2.00</td>
<td>1.50</td>
<td>2.5</td>
<td>6.1</td>
</tr>
<tr>
<td></td>
<td>2.00</td>
<td>2.00</td>
<td>7.1</td>
<td>4.7</td>
</tr>
<tr>
<td>2.0 g Red mud + $\text{H}_2\text{SO}_4$</td>
<td>0.50</td>
<td>1.50</td>
<td>13.2</td>
<td>6.9</td>
</tr>
<tr>
<td></td>
<td>1.00</td>
<td>1.50</td>
<td>1.6</td>
<td>6.3</td>
</tr>
<tr>
<td></td>
<td>1.50</td>
<td>1.50</td>
<td>28.5</td>
<td>4.2</td>
</tr>
<tr>
<td></td>
<td>2.00</td>
<td>1.50</td>
<td>49.2</td>
<td>4.0</td>
</tr>
</tbody>
</table>
Additionally, 500 g red mud was tested at different ratios of the content of H⁺ (mM) and red mud (g) with H₂SO₄. The results are listed in Table 4. In a manner similar to the results shown in Fig. 2, when the ratio of H⁺ (mM) and red mud (g) is close to 1.5 mM/g, then leaching concentrations of As are low (3 − 5 ppb), and the pH of leachates approximately corresponded to 5 − 6.

The alkali substance remaining in red mud influences the formation of insoluble arsenic compound and increases As leaching, and thus it is necessary to ensure that the ratio of H⁺ (mM) and red mud (g) exceeds 1.2 mM/g. A few carbonates were present in the red mud, and therefore the addition of acids created a significant amount of CO₂ gas. In mixing the operation, red mud became ropy and was difficult to mix. After 24 h, several holes appeared on the surface of the red mud, and a few transparent threadlike or white crystals separated out in the surface of red mud. This was different from H₂SO₄ because only small crystals were created when HCl was added. This crystal was the same as those that appeared when Fe₂(SO₄)₃ was added to the red mud. The crystals were analyzed through SEM, EDX, and XRD (not shown here). The results revealed that crystals almost corresponded to disodium sulfate (Na₂SO₄), which was considered relative to the Fe₂(SO₄)₃ or H₂SO₄ that was added.

In the SEM images (Fig. 3), large amounts of amorphous substances are evidently observed in the original dried red mud. On account of the high pH condition, it was speculated that a small amount of amorphous substance observed in the original red mud corresponded to hydroxides of Fe/Al or ferricydrite. With respect to the addition of Fe (III) ions, the pH of red mud changed to neutral, and the amount of amorphous substance was relatively low (as shown in B of Fig. 3). As shown in C and D of Fig. 3, the surface shapes are the same as those in the original dried red mud. Although the pH condition changed when acids were added, the amorphous substances remained as the main visible substances in the SEM images.

### Stable arsenic compounds potentially generated in the red mud by the addition of Fe (III) ions or inorganic acids

The XRD analysis results of resultants in the simulation test are shown in Figs. 4–6. In the condition of Fe³⁺:As⁵⁺ = 20:1 (Fig. 4), with respect to the hydroxide of Fe, the main content corresponded to a high proportion in the resultants because of the Fe hydrolysis at pH 4. Conversely, several insoluble Fe-As compounds were also detected in this resultant and included pharmacosiderite (Fe₄(AsO₄)₃(OH)₄·5H₂O), iron arsenate (Fe₂As₄O₁₂), karibibite (Fe₂As₄O₉), and iron hydrogen arsenate (FeHASO₄). Pharmacosiderite is insoluble and important in many environments to control the mobilization of arsenic.

![Fig. 2 Arsenic leaching concentration at different ratio of H⁺ [mM]/red mud [g].](image)

<table>
<thead>
<tr>
<th>Red mud [g]</th>
<th>H₂SO₄ [M]</th>
<th>Volume [mL]</th>
<th>Total of H⁺ [mM]</th>
<th>Ratio of H⁺[mM]/red mud [g]</th>
<th>pH of mixture [-]</th>
<th>As arsenic leaching [ppb]</th>
<th>pH of leachate [-]</th>
</tr>
</thead>
<tbody>
<tr>
<td>500</td>
<td>1.20</td>
<td>120</td>
<td>288</td>
<td>0.58</td>
<td>5</td>
<td>35.4</td>
<td>7.0</td>
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<td>1.50</td>
<td>360</td>
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<td>1.80</td>
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<td>2.00</td>
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ity of As [30]. Additionally, FeHAsO$_4$ exhibits a very low solubility that is even lower than that of scorodite [31]. The formation of karibibite was also advantageous to As stabilization as reported in previous studies [22]. Other stable compounds, such as unnamed mineral (FeAsO$_4$) and diiron tris (arsenato) arsenate (Fe$_2$(As(AsO$_4$)$_3$)), were also generated in this resultant and possessed a low solubility that was the same as iron arsenate [32]. In the red mud, the As and Fe (III) ions potentially formed the Fe-As insoluble compounds, and this was considered responsible for the stabilization of As.

In the condition of SO$_4^{2-}$:Fe$^{3+}$:Al$^{3+}$:As$^{5+}$ = 33:10:10:1 (Fig. 5), the main contents corresponded to hydroxide and oxides of Fe and Al. A few new insoluble Al-As compounds also generated in addition to the formation of diiron tris (arsenato) arsenate, karibibite, and iron hydrogen arsenate. As reported in previous studies, aluminum arsenate (Al(AsO$_4$)) is a stable arsenic compound easily generated in the presence of Al$^{3+}$ [33]. Furthermore, the newly generated compound also included aluminum arsenate hydrate (AlAs$_3$O$_9$·5H$_2$O), which corresponds to an amorphous aluminum arsenate. There is a paucity of studies on AlAs$_3$O$_9$·5H$_2$O although studies examined analogs including AlAsO$_4$·3.5H$_2$O and AlASO$_4$·2H$_2$O with low solubility albeit higher than that of scorodite (FeAsO$_4$·2H$_2$O) [34]. When H$_2$SO$_4$ was added into the red mud, Fe$^{3+}$ and Al$^{3+}$ ions released from red mud could potentially react with As and form the Fe-As/Al-As insoluble compounds, and this contributed to the stabilization of As in the red mud.

In the condition of Cl$^-$:Fe$^{3+}$:Al$^{3+}$:As$^{5+}$ = 66:10:10:1 (Fig. 6), the main substances included iron oxide hydroxide chloride (FeO(OH)Cl$_{0.227}$), aluminum oxide (Al$_2$O$_3$), and iron oxide (Fe$_3$O$_4$). Several amorphous substances appeared with respect to the existence of high amounts of Cl$^-$. This is different from the As compounds that appear above since the
new compound, parascorodite (FeAsO$_4$·2H$_2$O), mansfieldite (AlAsO$_4$·2H$_2$O), and aluminum arsenic (AlAs), were detected in the resultant. Parascorodite was metastable with respect to scorodite, which possessed the same chemical formula [35]. However, aluminum arsenic was unstable in water [36]. Mansfieldite was more reactive when compared with scorodite, and it was not a satisfactory carrier for the fixation of As in the environment [37]. In contrast, given the high amounts of amorphous substances, crystallinity of the As compound was not good and only a few insoluble As compounds generated. It was possible that when HCl was added into the red mud in the presence of many Cl$^-$, the Fe$^{3+}$ and Al$^{3+}$ ions released from red mud were surrounded by Cl$^-$ to form coordination compounds such as [FeCl$_4$]$^{-}$ and [AlCl$_4$]$^{-}$ [38,39]. This potentially influenced the stabilization of As. Hence, it is preferable to use H$_2$SO$_4$ as opposed to HCl to dispose the red mud from a safety viewpoint.

From the XRD analysis, both the added Fe (III) ions and inorganic acids generate insoluble As compounds in the red mud. In the long-term stability test, after exposure to the air for 7, 30, 60, and 90 d, As leaching concentrations from the red mud disposed by Fe (III) ions and inorganic acids are shown in Figs. 7 and 8.

The results indicated that As leaching was limited at a suf-
sufficient low level that approximately corresponded to 4 – 6 ppb by 3 – 5% Fe\(_2\)(SO\(_4\))\(_3\) and 2 – 5 ppb by H\(_2\)SO\(_4\) and HCl in the long-term stability test, and this is significantly lower than the environmental criteria (10 µg/L).

It was confirmed that the effect of As stabilization in the above operation was stable and that the red mud disposed was safe for recycling. Additionally, after mixing with inorganic acids, the concentration of Fe and Al ions released immediately from red mud were about 1.16 and 1.06 ppm (leaching solution determined by AAS). And after the 90-d exposure in the air, the concentration of Fe and Al ions released from red mud were about 0.56 and 0.22 ppm (leaching solution determined by AAS). This decrease suggested that Fe and Al ions released from red mud at the beginning underwent coprecipitation that could adsorb As or formed the insoluble Fe-As and Al-As compounds in the exposure. With the addition of acids, the use of Fe and Al ions released from the red mud to stabilize As was better than the addition of Fe (III) ions from the viewpoint of the effect of As stabilization. Due to the negative influence of Cl\(^-\) as discussed above, the addition of H\(_2\)SO\(_4\) and HCl was more stable. Furthermore, acid addition was also suitable for a mixing operation in industrial treatment. However, with respect to the lost carbonate, red mud experienced a mass decrease and generated a high amount of CO\(_2\).

![Image 1](example.com/image1.png)

**Fig. 6** Insoluble arsenic compound generated upon adding HCl.

![Image 2](example.com/image2.png)

**Fig. 7** Stability of arsenic in red mud disposed by Fe (III) ions.

![Image 3](example.com/image3.png)

**Fig. 8** Stability of arsenic in red mud disposed by H\(_2\)SO\(_4\) and HCl.
CONCLUSIONS

This study involved investigating methods to stabilize As in red mud by the addition of Fe (III) ions, H$_2$SO$_4$, and HCl. The addition of 3 − 5% Fe$_2$(SO$_4$)$_3$ or the use of inorganic acids (ratio of H$^+$ (nM)/red mud (g) ≈ 1.5) to release Fe (III) and Al (III) ions in red mud controls As solubility at approximately 2 − 6 µg/L for several months, and this satisfies the requirement of the Environment Agency Notification 18 (less than 10 µg/L) in Japan. Additionally, the pH of red mud also decreased to neutral, and this could be used as roadbed material or as other materials safely. The main reason for As stabilization in the red mud corresponded to the formation of insoluble Fe-As and Al-As compounds at pH 4. In this study, it was a new viewpoint to add Fe which was the main substance of red mud to stabilize As. Though it was very different from other stabilization of waste, it has a high effect on As stabilization. The XRD analysis indicated that the generated stable As compounds included pharmacosiderite, iron arsenate, iron hydrogen arsenate, aluminum arsenate, and aluminum arsenate hydrate, which were considered responsible for the low As leaching. Moreover, the results revealed that the presence of high amounts of Cl$^-$ negatively influenced the formation of insoluble As compounds, and thus the use of HCl was not an appropriate method to dispose the red mud. As opposed to Fe (III) ions, the use of H$_2$SO$_4$ is a better method to stabilize As in the red mud from the viewpoint of a long-term As stabilization effect.

REFERENCES


