Effect of Additives on Arsenic, Boron and Selenium Leaching from Coal Fly Ash

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Academic Editor: M. Thaddeus Ityokumbul
Received: 4 April 2017; Accepted: 7 June 2017; Published: 10 June 2017

Abstract: The establishment of an inexpensive leaching control method to prevent the leaching of trace elements from fly ash is required for the utilization of large-scale fly ash as an embankment material. This study examined the effects of the additives on suppressing As, B, and Se leaching from coal fly ash using Ca(OH)$_2$, paper sludge ashes (PS Ash 3, PS Ash 4 and PS Ash 5), and filter cake (FC). PS Ash and FC are waste generated in the papermaking and lime industry processes and contain high levels of calcium. The treated fly ash H (FAH) and the resulting mixtures were subjected to a leaching test as per the Environmental Agency of Japan Notifications No. 13. The results indicate that the leaching concentrations of As, B, and Se could be greatly reduced in FAH with the highest effect given by Ca(OH)$_2$, followed by PS Ash 3 and PS Ash 5. Ca(OH)$_2$ greatly reduced both the leaching concentrations of As, B, and Se by about 91–100%, while PS Ash 3 reduced the As and B leaching concentrations by approximately 89–96% and 83–92%, respectively; and PS Ash 5 reduced the Se leaching concentration by about 87–96%. FC did not have any impact on As and B leaching, but reduced Se leaching by about 58–78%. A reason for the decrease in leaching concentrations of As, B, and Se may be the precipitation with calcium or the formation of ettringite. The presence of leached Ca and Na ions are key factors affecting the decrease of As, B, and Se leaching concentrations from fly ash. The utilization of PS Ash 3 and PS Ash 5 as inexpensive additives is a promising method to control the leaching of As, B, and Se into the environment.

Keywords: coal fly ash; leaching; calcium; arsenic; boron; selenium

1. Introduction

Coal-based power generation is one of the major sources of environmental pollution due to the discharge of large amounts of fly ash into the environment. After burning in a boiler, as the flue gas cools down, trace elements in coal such as As, B, Cr, Sb, and Se condense on the surface of the fly ash and form new stable compounds [1]. Approximately 41% of the production of fly ash worldwide is utilized in various applications, such as a substitute material for Portland cement, structural fills (usually for road construction), soil stabilization, as a mineral filler in asphaltic concrete and mine reclamation, and the rest is disposed in landfills [2]. The disposal of fly ash in the environment involves the interaction of fly ash particles with weathering and hydrological processes where rainfall causes trace elements in the fly ash to elute and contaminate the environment. The leaching of As, B, and Se from coal fly ash (CFA) is likely to occur as these elements tend to form hydrophilic oxides that are dissolved as oxyanion forms [3].

The beneficial reuse of fly ash as embankment material in road construction has great potential in minimizing the amount of disposed fly ash [4–6], which will reduce the disposal costs incurred by industry, reduce landfill requirements, minimize damage to natural resources caused by excavating...
earthen materials for construction, obtain added value from fly ash, and ultimately conserve production energy.

Among the trace elements found in coal fly ash, As, B, Cd, Cr, Hg, Pb, and Se are of the greatest concern as environmental hazards [7]. This study investigated the behavior of As, B, and Se as these elements have recently become a major problem in soil contamination in Japan. Long-term exposure of arsenic-contaminated materials to water may lead to various diseases such as conjunctivitis, hyperpigmentation, cardiovascular diseases, skin cancer, gangrene, and disturbances in the peripheral vascular and nervous systems [8]. Boron can cause nausea, vomiting, redness of the skin, diarrhea, and difficulty swallowing; also, in animals, acute excessive exposure to B may cause rapid respiration, eye inflammation, swelling of the paws, and may affect male reproductive organs [9]. Similarly, excessive Se intake may yield circulatory problems and loss of hair and fingernails in humans [10]. Due to the increased awareness of the environmental impact of fly ash, the leaching of trace elements including As, B and Se needs to be controlled before fly ash utilization. Additionally, the reuse of fly ash as embankment material needs to meet regulations on soil contamination; in Japan, the permissible limits for As, B and Se are 10 µg/L, 1 mg/L, and 10 µg/L, respectively.

Understanding the factors that control the leaching behavior of trace elements is critical in predicting the potential impacts of fly ash on the environment. Several works have been conducted on the leaching behavior of As, B, and Se from CFA. Jiao et al. [11] studied the leaching characteristics of As in fly ash and they found that the presence of Ca in fly ash plays an important role in the leaching behavior of As. Iwashita et al. [12] suggested that the leaching of B and Se may involve the trapping of B and Se species by the ettringite phase, leading to a decrease in leaching under alkaline conditions. Wang et al. [13] investigated the effect of pH, S/L ratio, calcium addition, and leaching time on the leaching behavior of As and Se from two major types of CFAs and found that the leaching of As and Se from CFA generally increased with increases in the S/L ratio and leaching time; also, adsorption/desorption played a major role in As and Se leaching from the CFA. Van der Hoek et al. [14,15] showed that the leaching of As and Se from acidic ashes could be described by sorption of iron oxide, while the leaching from the alkaline ashes appeared to be controlled by sorption in the alkaline calcium-phase. Our previous study investigated the leaching characteristics of As from six CFA samples, and described a decrease in the As leaching rate with an increase in CaO content in fly ash [16].

Overall, Ca content and the sorption process are known to play important roles in the release of As, B, and Se from CFA. Although there have been extensive studies to explain the effect of calcium on As, B, and Se leaching and adsorption in fly ash [11–19], the application of additives (which contain high levels of calcium) to suppress As, B, and Se release has been less well established. Furthermore, the utilization of paper sludge ash and filter cake—which are generated as waste in the papermaking and lime industry processes—as inexpensive additives to suppress As, B, and Se leaching from CFA has never been tested. Therefore, the aim of this study was to examine the effects of inexpensive additives on suppressing As, B, and Se leaching from CFA. For this purpose, an appropriate amount of paper sludge ash and filter cake were added to fly ash, and the resulting mixture was subjected to a leaching test. The pH of the mixture leachates and the relation of As, B, and Se leaching (with major coexisting ions including Ca, Na, K, and Mg in mixture leachates) are discussed. This new information is expected to help in controlling the release of As, B, and Se into the environment to aid in the development of sustainable fly ash management strategies.

2. Materials and Methods

2.1. Coal Fly Ash and Additives

A relatively high concentration of trace elements leaching (As 48.66 µg/L, B 5.39 mg/L, Se 86.9 µg/L, detected using ICP-AES) and low calcium content fly ash sample (2.05% of CaO, detected using X-ray fluorescence) named fly ash H (FAH) was collected from a Japanese coal fired power plant.
(600 MWe) for the leaching test. Paper sludge ash (PS Ash 3, PS Ash 4, and PS Ash 5) and filter cake (FC) (considered as suppressing materials) and pure Ca(OH)$_2$ were used in this experiment as the additives. Ca(OH)$_2$ was applied to compare the effect of calcium addition among pure calcium-material (Ca(OH)$_2$) and native calcium-material (PS Ash 3, PS Ash 4, PS Ash 5, and FC). Paper sludge ash is waste generated in the papermaking industry as a by-product of the de-inking and re-pulping of paper, while filter cake comes from the lime industry as a waste from the CaCO$_3$ manufacturing process.

The utilization of paper sludge ash and filter cake was considered due to the relatively high content of calcium they contain. The CaO content detected using X-ray fluorescence (XRF) in PS Ash was varied, ranging from 18.77 to 46.31%. FC had the highest CaO content at 59.18%. Table 1 lists the composition of inorganic elements in FAH and additives.

### Table 1. Composition of inorganic elements in fly ash and additives.

<table>
<thead>
<tr>
<th>Sample</th>
<th>FAH (Is)</th>
<th>PS Ash 3</th>
<th>PS Ash 4</th>
<th>PS Ash 5</th>
<th>FC</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO$_2$</td>
<td>59.25</td>
<td>31.47</td>
<td>44.21</td>
<td>42.36</td>
<td>23.31</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>25.63</td>
<td>12.40</td>
<td>22.23</td>
<td>19.80</td>
<td>13.87</td>
</tr>
<tr>
<td>TiO$_2$</td>
<td>1.99</td>
<td>0.38</td>
<td>2.56</td>
<td>2.11</td>
<td>0.06</td>
</tr>
<tr>
<td>Fe$_2$O$_3$</td>
<td>7.49</td>
<td>5.13</td>
<td>2.63</td>
<td>5.56</td>
<td>2.33</td>
</tr>
<tr>
<td>CaO</td>
<td>2.05</td>
<td>46.31</td>
<td>18.77</td>
<td>19.51</td>
<td>59.18</td>
</tr>
<tr>
<td>MgO</td>
<td>0.79</td>
<td>3.28</td>
<td>3.42</td>
<td>3.30</td>
<td>0.96</td>
</tr>
<tr>
<td>Na$_2$O</td>
<td>0.60</td>
<td>0.24</td>
<td>0.95</td>
<td>0.41</td>
<td>0.03</td>
</tr>
<tr>
<td>K$_2$O</td>
<td>1.56</td>
<td>0.20</td>
<td>2.09</td>
<td>1.99</td>
<td>0.25</td>
</tr>
<tr>
<td>P$_2$O$_5$</td>
<td>0.18</td>
<td>0.18</td>
<td>1.75</td>
<td>1.54</td>
<td>0.00</td>
</tr>
<tr>
<td>MnO</td>
<td>-</td>
<td>0.03</td>
<td>0.05</td>
<td>0.06</td>
<td>0.04</td>
</tr>
<tr>
<td>V$_2$O$_5$</td>
<td>0.03</td>
<td>0.01</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>SO$_3$</td>
<td>0.42</td>
<td>0.36</td>
<td>1.31</td>
<td>3.33</td>
<td>0.00</td>
</tr>
<tr>
<td>pH of the leachate</td>
<td>10.38</td>
<td>13.72</td>
<td>12.71</td>
<td>12.26</td>
<td>7.28</td>
</tr>
<tr>
<td>Leached Ca ion [mg/L]</td>
<td>10.38</td>
<td>13.72</td>
<td>12.71</td>
<td>12.26</td>
<td>7.28</td>
</tr>
</tbody>
</table>

1 Sample fly ash H, from the chamber 1 of electrostatic precipitator.

2.2. Fly Ash Treatment and Leaching Test

The additive was added to the FAH sample so as to give a Ca content of 5% and 10% (see Appendix A). The mixture was then moved to a bowl and distilled water at 25% addition ratio of the total mixture was added; the mixture was kneaded for one minute, and then scraped and kneaded for a further two minutes. The mixture was then stored in a sealed plastic bag for seven days, following which the mixture was air-dried and the leaching test was conducted.

The procedure of standard leaching tests for fly ash (Notification No. 13 by the Environmental Agency of Japan) was employed as the protocol for leaching tests in this work. The mixture of FAH-additive of 5 g was mixed with 50 mL distilled water, which accounted for a liquid to solid ratio (L/S) of 10, and was shaken for six hours at room temperature with a shaking speed of 200 rpm. The solid-liquid sample was separated by filtration using a membrane filter of 0.45 µm to obtain the filtrate.

2.3. Characterization of the Elements in CFA and Additives

The total concentrations of the major chemical compositions in FAH and additives were determined using a Wavelength Dispersive X-ray Fluorescence Spectrometer (WDXRF S8 TIGER, Bruker AXS, Yokohama, Japan). For XRF analysis of the fly ash and additive samples, a small amount (approximately 500 mg) was poured onto a polypropylene thin-film that was attached previously to a plastic O-ring sample cup with an outer diameter of 40 mm. The samples were analyzed using XRF, and the chemical compositions of samples were determined.

The identification of calcium compounds in the additives was determined qualitatively using thermogravimetric analysis (TG/DTA6300 SII EXSTAR 6000, Hitachi, Hong Kong, China) and X-ray diffraction (LabX XRD6100, Shimadzu, Kyoto, Japan). For thermogravimetric (TG) analysis, a sample
of 10–20 mg was heated with a measurement temperature from 30 to 1000 °C at a heating rate of 10 °C/min under a nitrogen atmosphere at a flow rate of 200 mL/min. For XRD analysis, the sample was irradiated with Cu Kα-radiation between 10 and 80° (2θ) with a counting angle at 0.02° and slit of 0.3 mm, under a scanning speed of 2°/min, at an acceleration voltage of 40 kV and current of 30 mA. The diffraction pattern was analyzed with the help of the software module “DDView and Sleve,” the phases were viewed and identified by applying the database PDF-2/Release 2013 RDB from the International Center for Diffraction Data (ICDD).

2.4. Chemical Analyses

The concentrations of As, B, and Se in the filtrate were carefully analyzed by ICP-AES (ULTIMA2, HORIBA Ltd, Tokyo, Japan). Cations such as Ca²⁺, Na⁺, K⁺, and Mg²⁺ were quantified using ion chromatographs IA-300 (DKK-TOA Corporation, Tokyo, Japan). The final pH of the leachate reflected the interaction of the leaching fluid (distilled water) with the buffering capacity of the FAH-additive mixture. The pH measurement was carried out by a pH/ion Meter D-53, HORIBA. After pH meter calibration at pH 4, 7, and 9 using buffer solutions, the pH measurement of samples was carried out.

3. Results

3.1. pH of the Leachates

A strong relationship between the pH of the leachate and CaO content has been recognized since this mineral elevates pH in the leachates [20–23]. Figure 1 shows the effect of including additives in the pH of the leachates, where it increased with the treated amount. However, each additive had a different way of elevating pH that was related to their properties. Of the four additives discussed (after FAH-Ca(OH)₂ mixture leachates), the FAH-PS Ash 3 mixture leachates showed the highest final pH over FAH-PS Ash 5, FAH-PS Ash 4 and FAH-FC mixture leachates for both 5% and 10% Ca content samples, as shown in Figure 1. This indicated that the relatively higher CaO content of PS Ash 3 added into fly ash tended to increase the pH of the mixtures leachates. This was consistent with the previous study in Reference [24], where the release of Ca from CaO minerals yielded Ca(OH)₂ in aqueous solutions, which is an oxide mineral that significantly contributes to alkalinity. Conversely, FC, which contained the highest CaO content, did not appear to have any impact on elevating the pH of the mixture leachates.

To probe the effects of FC addition into FAH, the leaching amount of calcium ions in the FAH-FC mixture leachates was examined (Figure 2). As predicted, the amount of Ca ions leached in the FAH-FC mixture leachates was lower than that of other additives. This result indicates that the low amount of leached Ca ion in the FC mixture leachates was not enough to elevate the pH of the leachates. This is relevant with the previous observation in Reference [12], that pH tended to rise when the Ca leaching amount was larger, where the main species of Ca such as CaO elevated the pH in the leachate. Overall, it could be concluded that an increase of CaO content in the additives caused an increase in the leachates pH, and that the leached Ca ions contributed to elevate the leachates pH.

Based on the above discussion, it is clear that the leached Ca ions in the mixture leachates seemingly play an important role in contributing to the alkalinity of the mixture, especially Ca(OH)₂, which possessed high solubility during the leaching test given that Ca(OH)₂ is composed of completely water-soluble Ca. As the amount of leached Ca ions yielded was varied due to the diverse solubilities of different calcium compounds in the additives, it was important to identify the types of calcium compounds included in the PS Ash 3, PS Ash 4, PS Ash 5, and FC to better understand the effect of the additives on the pH of the leachates. To clarify such a hypothesis, TG and XRD analyses were carried out. The results are shown in Figures 3 and 4.

TG was performed on Ca(OH)₂ and CaCO₃ of 99.9% purity to confirm that the two weight losses in the TG curve of additives were due to the thermal decomposition of Ca(OH)₂ and CaCO₃. Figure 3 compares the TG curves of each calcium compound and the additives. The TG analysis of Ca(OH)₂,
CaCO₃, and additives was conducted under a nitrogen atmosphere based on the methods described in References [25,26]. Based on previous study, Ca(OH)₂ was thermally decomposed at 330–460 °C into CaO [26], and CaCO₃ was thermally decomposed into CaO at around 700 °C [27]. As seen in Figure 3, the weight losses in PS Ash 3 at around 390 °C and 600 °C corresponded to the thermal decompositions of Ca(OH)₂, and CaCO₃. The weight losses in PS Ash 4 (around 600 °C), PS Ash 5 (around 590 °C), and FC (around 700 °C) corresponded to the thermal decomposition of CaCO₃. From these results, it was found that the Ca(OH)₂ and CaCO₃ in additives could be detected by TG. Since the decomposition temperature of CaO is above 1000 °C [25] and cannot be detected by TG, the analysis of calcium compounds in the additives was performed using the XRD method.

![Figure 1. Leachate pH values from fly ash H alone and fly ash H under five kinds of additives for 5% and 10% Ca content samples.](image1)

![Figure 2. Ca ions leached from fly ash H alone and fly ash H under five kinds of additives for 5% and 10% Ca content samples.](image2)

![Figure 3. Thermogravimetric curves showing the thermal decomposition of Ca(OH)₂, CaCO₃, and additives in a N₂ atmosphere.](image3)
Figure 3. Thermogravimetric curves showing the thermal decomposition of Ca(OH)\textsubscript{2}, CaCO\textsubscript{3}, and additives in a N\textsubscript{2} atmosphere.

Figure 4. XRD Patterns of several calcium compounds and additives.

Figure 4 shows the X-ray diffraction patterns of several calcium compounds and additives. All of the peaks in the additives were compared with the peaks in each calcium compound. The results showed that PS Ash 3 contained CaCO\textsubscript{3}, CaO, and Ca(OH)\textsubscript{2}; PS Ash 4 and PS Ash 5 contained small amounts of CaCO\textsubscript{3}, while FC contained the most amount of CaCO\textsubscript{3}. The XRD analysis results were consistent with the TG analysis results described above, and revealed that a relatively high content of CaCO\textsubscript{3} in the FC caused this additive to yield low amounts of leached Ca ions and produce a relatively low value of pH mixture leachates, which was considered to be due to CaCO\textsubscript{3} being an insoluble substance in pure water. Therefore, the contents of CaO and Ca(OH)\textsubscript{2} (as water soluble Ca) in PS Ash 3 caused this additive to yield high amounts of leached Ca ions and a higher value of pH mixture leachate than that of PS Ash 4, PS Ash 5, and FC. Regarding the higher amounts of leached Ca ions in PS Ash 5 than in PS Ash 3 (Figure 2), this could be attributed to the relatively high content of SO\textsubscript{3} (detected by XRF) as an acid compound in PS Ash 5 (see Table 2). This is consistent with the reported study by Killingley et al. [28], where the balance between the concentration of alkaline-earth element, Ca, in the ashes, and the proportion of potentially acid generating SO\textsubscript{3} influenced the initial pH and leached Ca ions of the ash-water system.

Table 2. Trace element leaching suppression effect of additives.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Ca (wt%)</th>
<th>As leach. conc. \textsuperscript{1} (µg/L)</th>
<th>L.I.R \textsuperscript{2} (%)</th>
<th>B leach. conc. (mg/L)</th>
<th>L.I.R (%)</th>
<th>Se leach. conc. (µg/L)</th>
<th>L.I.R (%)</th>
<th>Final pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>FA H (1s)</td>
<td>1.46</td>
<td>48.66</td>
<td>0.0</td>
<td>5.39</td>
<td>0.0</td>
<td>86.39</td>
<td>0.0</td>
<td>10.38</td>
</tr>
<tr>
<td>Ca(OH)\textsubscript{2}</td>
<td>5</td>
<td>1.26</td>
<td>97.4</td>
<td>0.10</td>
<td>98.1</td>
<td>7.15</td>
<td>91.7</td>
<td>12.86</td>
</tr>
<tr>
<td>PS Ash 3</td>
<td>5</td>
<td>5.12</td>
<td>89.5</td>
<td>0.87</td>
<td>83.9</td>
<td>15.00</td>
<td>82.6</td>
<td>11.58</td>
</tr>
<tr>
<td>PS Ash 4</td>
<td>5</td>
<td>27.71</td>
<td>43.1</td>
<td>2.83</td>
<td>47.5</td>
<td>26.04</td>
<td>69.9</td>
<td>10.46</td>
</tr>
<tr>
<td>PS Ash 5</td>
<td>5</td>
<td>12.06</td>
<td>75.2</td>
<td>1.48</td>
<td>72.6</td>
<td>10.60</td>
<td>87.7</td>
<td>10.63</td>
</tr>
<tr>
<td>FC</td>
<td>5</td>
<td>57.75</td>
<td>−18.7</td>
<td>5.55</td>
<td>−3.0</td>
<td>36.04</td>
<td>58.3</td>
<td>9.66</td>
</tr>
<tr>
<td>Ca(OH)\textsubscript{2}</td>
<td>10</td>
<td>0.00</td>
<td>100.0</td>
<td>0.10</td>
<td>98.2</td>
<td>6.16</td>
<td>92.9</td>
<td>12.98</td>
</tr>
<tr>
<td>PS Ash 3</td>
<td>10</td>
<td>1.84</td>
<td>96.2</td>
<td>0.41</td>
<td>92.4</td>
<td>8.57</td>
<td>90.1</td>
<td>11.94</td>
</tr>
<tr>
<td>PS Ash 4</td>
<td>10</td>
<td>5.66</td>
<td>88.4</td>
<td>2.96</td>
<td>45.1</td>
<td>5.54</td>
<td>93.6</td>
<td>10.62</td>
</tr>
<tr>
<td>PS Ash 5</td>
<td>10</td>
<td>6.71</td>
<td>86.2</td>
<td>1.06</td>
<td>80.3</td>
<td>2.80</td>
<td>96.8</td>
<td>10.51</td>
</tr>
<tr>
<td>FC</td>
<td>10</td>
<td>65.29</td>
<td>−34.2</td>
<td>5.34</td>
<td>1.0</td>
<td>19.19</td>
<td>77.8</td>
<td>9.23</td>
</tr>
</tbody>
</table>

\textsuperscript{1} Leaching concentration. \textsuperscript{2} Leaching inhibition rate.
3.2. Effect of Additives on Arsenic Leaching

The presence of Ca in CFA, pH, and the sorption process are known to play important roles in the leaching behavior of arsenic. Arsenic reacts with calcium to form a new stable compound, which is slightly soluble in water [11,29]. As the additives contained high native-calcium that could enrich the calcium content in CFA, the leaching of arsenic into the environment was expected to be reduced.

Figure 5a shows the arsenic leaching concentration for FAH alone, and for fly ash under five kinds of additives at a 5% and 10% Ca content. The arsenic leaching concentration of FAH is shown on the far left as a comparison standard in Table 2. As seen in Figure 5a, Ca(OH)$_2$ was very effective in suppressing arsenic leaching, and the leaching amount of FAH greatly reduced from 48.7 µg/L to 1.3 µg/L in the 5% Ca content sample. The leaching inhibition rate was 97.4% and 100% for the 5% and 10% Ca content samples, respectively. The value of the ‘leaching inhibition rate’ was defined as [(leaching concentration of element in FAH)−(leaching concentration of element in FAH-additives mixture)]/(leaching concentration of element in FAH)]. Comparing PS Ash 3, 4 and 5, it was seen that the arsenic suppressing effect was high in the order of PS Ash 3 > 5 > 4 for the 5% Ca content sample and PS Ash 3 > 4 > 5 for the 10% Ca content sample. In addition, the arsenic leaching concentration was reliably reduced in the Ca 10% sample compared to that in the Ca 5% sample. A plausible reason to describe the dramatic reduction of arsenic leaching concentration could be due to the relatively high Ca ions (Ca$^{2+}$) in Ca(OH)$_2$ and in the PS Ash mixture leachates that react with oxyanionic species of arsenic at an alkaline pH, which can form precipitates as a new stable compound or trap the oxyanionic species of arsenic through the ettringite phase formed under alkaline conditions, leading to a decrease in the leaching concentration of arsenic from FAH. This is consistent with previous studies that have shown that under a high Ca condition and pH > 11.5 (alkaline leachate), the dominating species, AsO$_4^{3–}$, formed a precipitate with Ca as a less soluble compound or trapped the arsenic species by the ettringite phase which prevented the leaching of arsenic [11,13]. Figure 6a presents the plots of Ca ion concentration and arsenic leaching concentration in the mixture leachates. The arsenic leaching concentration tended to decrease as the Ca ion concentration increased.

As discussed previously, there was an increase in pH when Ca was abundant in the leachates; Figure 7a shows the relationship between the leachate pH and the arsenic leaching concentration for all samples. Since Ca(OH)$_2$ is completely water-soluble Ca, the pH of Ca(OH)$_2$ was the highest, around 13. Overall, the pH and arsenic leaching concentration showed a linear relationship, where the higher the pH, the lower the arsenic leaching concentration, and arsenic leaching was suppressed as the pH of the leachate became 11.5 or higher. This agrees with the observation reported by Jiao et al. [11] and Wang et al. [13], who claimed that at a pH above 11, AsO$_4^{3–}$ (as the dominating species) forms precipitates with Ca, which provides a suppressing effect on the mobilization of arsenic. Figure 7a shows that arsenic leaching was suppressed in PS Ash 4 and 5 despite a pH lower than 11.5, therefore, it was considered that alkaline elements other than Ca also contribute to suppress arsenic leaching. Thus, the concentration of alkaline elements in the leachates (such as Na, K, and Mg) were measured and the relationships between them and arsenic leaching concentrations were examined. Figure 8a shows the relationship between the Na ion concentration and the arsenic leaching concentration in the mixture leachates. As seen in Figure 8a, the Na ion concentration showed a corresponding correlation with the arsenic leaching concentration, where the arsenic leaching concentration tended to decrease as the Na ion concentration increased. For K and Mg (as shown in Figures A1a and A2a), neither of them were clearly related to the arsenic leaching concentration. This result shows the reason why PS Ash 4 and 5 can be understood to suppress arsenic leaching as an effect of Na ions. The reason why the Na ion concentration as an alkaline element other than Ca influences the arsenic leaching concentration has been insufficiently explored, but warrants further investigation to be able to estimate further mixture compositions of additives for the more effective suppression of arsenic leaching.
reason to describe the dramatic reduction of arsenic leaching concentration could be due to the relatively high Ca ions (Ca\(^{2+}\)) in Ca(OH)\(_2\) and in the PS Ash mixture leachates that react with oxyanionic species of arsenic at an alkaline pH, which can form precipitates as a new stable compound or trap the oxyanionic species of arsenic through the ettringite phase formed under alkaline conditions, leading to a decrease in the leaching concentration of arsenic from FAH. This is consistent with previous studies that have shown that under a high Ca condition and pH > 11.5 (alkaline leachate), the dominating species, AsO\(_4^{3-}\), formed a precipitate with Ca as a less soluble compound or trapped the arsenic species by the ettringite phase which prevented the leaching of arsenic [11,13].

Figure 6a presents the plots of Ca ion concentration and arsenic leaching concentration in the mixture leachates. The arsenic leaching concentration tended to decrease as the Ca ion concentration increased.
Figure 5. Trace element leaching concentration for fly ash H alone, and fly ash H under five kinds of additives addition for 5% and 10% Ca content samples: (a) arsenic; (b) boron; and (c) selenium.

Figure 6. Relationship between the Ca ion leaching concentration and trace elements leaching concentration of five kinds of additives: (a) arsenic; (b) boron; and (c) selenium.

As discussed previously, there was an increase in pH when Ca was abundant in the leachates; Figure 7a shows the relationship between the leachate pH and the arsenic leaching concentration for all samples. Since Ca(OH)\(_2\) is completely water-soluble Ca, the pH of Ca(OH)\(_2\) was the highest, around 13. Overall, the pH and arsenic leaching concentration showed a linear relationship, where the higher the pH, the lower the arsenic leaching concentration, and arsenic leaching was suppressed as the pH of the leachate became 11.5 or higher. This agrees with the observation reported by Jiao et al. [11] and Wang et al. [13], who claimed that at a pH above 11, AsO\(_4^{3-}\) (as the dominating species) forms precipitates with Ca, which provides a suppressing effect on the mobilization of arsenic. Figure 7a shows that arsenic leaching was suppressed in PS Ash 4 and 5 despite a pH lower than 11.5, therefore, it was considered that alkaline elements other than Ca also contribute to suppress arsenic leaching. Thus, the concentration of alkaline elements in the leachates (such as Na, K, and Mg) were measured and the relationships between them and arsenic leaching concentrations were examined. Figure 8a shows the relationship between the Na ion concentration and the arsenic leaching concentration in the mixture leachates. As seen in Figure 8a, the Na ion concentration showed a corresponding correlation with the arsenic leaching concentration, where the arsenic leaching concentration tended to decrease as the Na ion concentration increased. For K and Mg (as shown in Figures A1a and A2a), neither of them were clearly related to the arsenic leaching concentration. This result shows the reason why PS Ash 4 and 5 can be understood to suppress arsenic leaching as an effect of Na ions. The reason why the Na ion concentration as an alkaline element other than Ca influences the arsenic leaching concentration has been insufficiently explored, but warrants further investigation to be able to estimate further mixture compositions of additives for the more effective suppression of arsenic leaching.
Figure 7. Relationship between the leachate pH and trace elements leaching concentration of five kinds of additives: (a) arsenic; (b) boron; and (c) selenium.
For the arsenic leaching suppression effect on FC, despite the highest Ca content, FC was found to increase rather than decrease arsenic leaching. This could be explained by the relatively low pH of the FAH-FC mixture leachates of around 9.2–9.7, which is not an effective pH value for the suppression of arsenic leaching. Furthermore, the Ca ion concentration was 83–101 mg/L, which is not an effective...
Ca ion concentration for the suppression of arsenic leaching due to the fact that the composition of FC is calcium carbonate based, which is a relatively stable substance.

3.3. Effect of Additives on Boron Leaching

Figure 5b shows the boron leaching concentration for fly ash H (FAH) alone, and for fly ash under five kinds of additives at 5% and 10% Ca content (as shown in Table 2). As shown in Figure 5b, like arsenic leaching suppression, Ca(OH)$_2$ was also very effective in suppressing boron leaching with a leaching inhibition rate of 98% for both 5% and 10% Ca content, respectively. This result indicated that As and B leaching can be simultaneously suppressed by the addition of Ca(OH)$_2$. A comparison of PS Ash 3, 4, and 5, showed that the boron suppressing effect to be high in the order of PS Ash 3 > 5 > 4 at both 5% and 10% Ca content samples. The dramatic reduction in the boron leaching concentration could be due to the relatively high level of Ca ions (Ca$^{2+}$) in Ca(OH)$_2$ and in the PS ash mixture leachates that react with oxyanionic species of boron at an alkaline pH, which can form new stable precipitates or trap the oxyanionic species of boron through the ettringite phase formed under alkaline conditions, leading to a decrease in the boron leaching concentration from FAH. This was consistent with a previous study that showed that with a high amount of Ca leaching and a pH >11 (alkaline leachate), the dominating species—in this case, borate—formed a precipitate with Ca that was a less soluble compound or trapped the boron species in the ettringite phase and prevented the leaching of boron [12,30,31]. Figure 6b presents the plots of the Ca ion concentration and boron leaching concentration in the mixture leachates. Like the arsenic results, the boron leaching concentration tended to decrease as the Ca ion concentration increased. Figure 7b shows the relationship between the leachate pH and boron leaching concentration for all samples. The leaching test was performed simultaneously with As and Se, the pH data were the same data as As and Se (Table 2). As with arsenic, the pH and boron leaching concentration also showed a linear relationship; the higher the pH, the lower the boron leaching concentration, and boron leaching was suppressed as the pH of the leachate became 11.5 or higher. This finding is consistent with Iwashita et al. [20], Hollis et al. [30], and Cetin et al. [31], who claimed that the B concentrations decreased with an increase in pH above 11, where large quantities of Ca minerals in the leachates may have caused the precipitation of B with Ca. In the case of PS Ash 5, it seems that the boron leaching concentration was suppressed despite a pH lower than 11, and it was suggested that alkaline elements other than Ca could also contribute to the suppression of boron leaching. Therefore, the concentration of alkaline elements in the leachate (such as Na, K, and Mg) were measured and the relationships between them and the boron leaching concentrations were examined. Figure 8b shows the relation between the Na ion concentration and boron leaching concentration in the mixture leachates. As shown in Figure 8b, it appeared that the Na ion concentration showed a corresponding correlation to the boron leaching concentration; the boron leaching concentration tended to decrease as the Na ion concentration increased. For K and Mg (as shown in Figures A1b and A2b), neither of them was clearly related to the boron leaching concentration. This result demonstrates why PS Ash 5 could suppress boron leaching as an effect by Na ions. The reason why the Na ion concentration influences the boron leaching concentration has been insufficiently investigated, but it warrants further study in order to estimate the mixture composition of additives for the more effective suppression of boron leaching.

For a boron leaching suppression effect of FC like that observed with arsenic, FC did not have an inhibitory effect on boron leaching. Overall, the mechanism to suppress the leaching of arsenic and boron is similar. Figure 9 shows the relationship between the arsenic leaching concentration and the boron leaching concentration plotted for all samples. In general, they have a good correlation, indicating that As and B can be suppressed simultaneously.
was increased in the order of PS Ash 5 > 3 > 4 for the 5% Ca content sample, and PS Ash 5 > 4 > 3 for the 10% Ca sample. In addition, it could be seen that the selenium suppression effect was increased in the order of PS Ash 5 > 3 > 4 for the 5% Ca content sample, and PS Ash 5 > 4 > 3 for the 10% Ca sample. Therefore, the inhibitory effect of Ca was high in the order of B > As > Se. When comparing PS Ash 3, 4, and 5, it could be seen that the selenium suppression effect by Ca(OH)$_2$ for As, B, and Se showed that the leaching inhibition rate when Ca content was 5% was 97.4%, 98.1%, and 91.7%, respectively. Therefore, the inhibitory effect of Ca was high in the order of B > As > Se. When comparing PS Ash 3, 4, and 5, it could be seen that the selenium suppression effect was increased in the order of PS Ash 5 > 3 > 4 for the 5% Ca content sample, and PS Ash 5 > 4 > 3 for the 10% Ca sample. In addition, it could be seen that the selenium leaching concentration was reliably reduced in the Ca 10% sample compared to the Ca 5% sample. The dramatic reduction of the selenium leaching concentration could be due to the relatively high level of Ca ions (Ca$^{2+}$) and Ca(OH)$_2$ in the PS ash mixture leachates that react with oxyanionic species of selenium at an alkaline pH, which formed precipitates as a new stable compound or trapped the oxyanionic species of selenium through the ettringite phase formed under alkaline conditions, thus leading to a decrease in the leaching concentration of selenium from FAH. This is consistent with previous studies that under high Ca conditions and a pH > 11 (alkaline leachate), the dominating species, SeO$_3^{2-}$, formed a precipitate with Ca as a less soluble compound or trapped the selenium species in the ettringite phase and prevented the leaching of selenium [13,32,33]. Figure 6c presents the plots of the Ca ion concentration and selenium leaching concentration in the mixture leachates. Like the results of arsenic and boron, the selenium leaching concentration tended to decrease as the Ca ion concentration increased.

Figure 7c shows the relationship between the leachate pH and the selenium leaching concentration for all samples. As with arsenic and boron, the pH and the selenium leaching concentration also showed a linear relationship; in general, the selenium leaching concentration decreased with increasing pH, and selenium leaching was suppressed as the pH of the leachate became 11 or higher. This finding was consistent with Izquierdo et al. [21], Morar et al. [32], Jankowski et al. [33], and Solen-Tishmack et al. [34], who claimed that the Se concentrations decreased with an increase in pH above 11 due to the substitution of Se in the structure of ettringite. Figure 5c shows that selenium leaching was suppressed in PS Ash 4 and 5. Although the pH was lower than 11, it is thought that alkaline elements other than Ca also contributed to the suppression of selenium leaching. Therefore, the concentration of alkaline elements in the leachate (such as Na, K, and Mg) were measured and the relationships between them and selenium leaching concentrations were examined. Figure 8c shows the relationship between the Na ion concentration and the selenium leaching concentration in the mixture leachates. As seen in Figure 8c, it seemed that the Na ion concentration showed a corresponding
correlation to the selenium leaching concentration, where the selenium leaching concentration tended to decrease as the Na ion concentration increased. For K and Mg (as shown in Figures A1c and A2c), neither of them were clearly related to the selenium leaching concentration. This result showed why PS Ash 4 and 5 could suppress selenium leaching as an effect of the Na ions. The reasons as to why the Na ion concentration is one of the alkaline elements other than Ca that influence the selenium leaching concentration has also been insufficiently considered, but is needs to be further understood in order to be able to estimate further mixture compositions of additives to more effectively suppress selenium leaching.

In contrast to arsenic and boron, FC showed some effect on selenium leaching suppression. The decrease in selenium leaching may have been due to the presence of Al$_2$O$_3$ and Fe$_2$O$_3$, as well as the Ca content and leached Ca ion content in FC, as reported by previous observations where Se was controlled by sorption reactions that occurred on the surface of metal oxides such as Al$_2$O$_3$ and Fe$_2$O$_3$ [35,36]. These oxides may provide additional surface area to which positively charged ions may attach, resulting in decreased Se concentration in aqueous solutions [37].

Figure 10 shows the relationship between the arsenic leaching concentration and selenium leaching concentration, and Figure 11 shows the relationship between the boron leaching concentration and selenium leaching concentration for all samples. In general, they had good correlation, indicating that As, B and Se could be suppressed at the same time.
3.5. Comprehensive Evaluation of Leaching Suppression Materials

As shown above, PS Ash had the effect of simultaneously suppressing the leaching of As, B, and Se, and it has become clear that it is very promising as a practical leaching suppression material. Although FC had some effect on Se leaching suppression, it did not work for As and B, and was found overall to be inappropriate as a leaching suppression material. Figure 12 plots the leaching inhibition rates of As, B, and Se by the PS Ash as radar charts for the Ca 5% and 10% samples. In these figures, it can be comprehensively evaluated that the one having the largest area had the effect of simultaneously suppressing As, B, Se. From this figure, PS Ash 3 was the most suitable material as a suppression material, followed by PS Ash 5 and PS Ash 4, respectively. For the three elements As, B, Se, the only elution suppression material satisfying the soil environmental standard was PS Ash 3, with a Ca = 10% setting.

![Radar chart showing leaching inhibition rates for PS Ash samples](image)

**Figure 12.** Trace elements (arsenic, boron, and selenium) leaching inhibition rate for (a) 5% Ca content; and (b) 10% Ca content samples.

4. Conclusions

This study investigated the effect of additives on the leaching characteristics of As, B, and Se in coal fly ash. We proposed Ca(OH)_2, Paper Sludge Ash 3, Paper Sludge Ash 4, Paper Sludge Ash 5, and filter cake as new additives (suppression materials); adjusted the Ca concentration in the finished mixture
to 5% and 10%; and verified the simultaneous leaching suppressing effect of As, B, and Se. The results revealed that all additives (other than filter cake) showed a simultaneous leaching suppression effect. However, only the leaching suppression effect of PS Ash 3 satisfied the soil environmental standard with Ca concentration of 10%. Other additives could meet the soil standards by increasing the Ca concentration to a higher level.

Acknowledgments: The financial support from Tohoku Electric Power Company is gratefully acknowledged. The authors would like to thank Hayakawa Yukio for his valuable contributions in improving the manuscript. The authors also thank Erda Rahmilaila Desfitri for her help in XRD analysis. Sincere thanks to the editor and reviewers for their detailed reviews that led to substantial improvements in this paper.

Author Contributions: Akihiro Takeyama and Shinji Kambara conceived and designed the experiments; Akihiro Takeyama, Farrah Fadhillah Hanum and Sri Hartuti performed the experiments; Akihiro Takeyama and Sri Hartuti analyzed the data; Shinji Kambara contributed reagents, materials, analysis tools; Sri Hartuti and Akihiro Takeyama wrote the paper.

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

Appendix A

To obtain the Ca content of the finished mixture at 5% or 10%, the amount of each additive and FAH mixed was calculated based on the CaO concentration in the ash (Table 1). For example, when preparing a Ca 5% sample with FAH ($x$ gram) and PS Ash 3 ($y$ gram) based on 100 g of the finished mixture, considering the molecular weight of CaO 56.078 and the molecular weight of Ca 40.078, $x + y = 100$, $2.05 \times (40.078/56.078) \times x + 46.31 \times (40.078/56.078) \times y = 5$, from the simultaneous equations, the amount of $x$ becomes 88.78 g and that of $y$ becomes 11.22 g.

![Figure A1. Cont.](image-url)
Figure A1. Relationship between the K ion leaching concentration and trace elements leaching concentration of five kinds of additives: (a) arsenic; (b) boron; and (c) selenium.
References


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