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To cite this version:


HAL Id: hal-01798131
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Submitted on 23 Jan 2019

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Physicochemical properties and heavy metals leachability of fly ash from coal-fired power plant

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\textbf{Abstract}

The physicochemical properties of fly ash from two kinds of coal-fired power plants were studied. Three aspects were examined: the micro-morphology, the mineral composition and the content of heavy metals. The results show that the fly ash from plants using a circulating fluidized bed are more irregular particles, while the particles from the plants using a pulverized coal-fired boiler are mainly spherical in shape. Quartz and mullite are the main crystalline phases in the ash. Clearly, both the technology and the coal used by a power plant can influence the mineral composition of the ash. The mineral composition of fly ash from a circulating fluidized bed is more complex than that from a pulverized coal-fired boiler. The quantity of elements found in the fly ash is greater than that found in the bottom ash for the same plant. Heavy metals are likely to be enriched in the fly ash. Heavy metal leachability was studied using two leaching methods. The results indicate that most of the heavy metals that leached during either batch leaching or column leaching experiments did not exceed the related maximum concentration standards. But Ni concentrations in the leachates from both batch and column tests exceed the standard. The highest excess rates in both tests were 572% and 497%, which levels might threaten the environment.

Keywords: Fly ash, Physicochemical property, Leachability, Heavy metal

1. Introduction

Coal fly ash is the by-product of coal-fired power plants that use flue-gas desulfurization technology. Ash consists of both fly ash and bottom ash, which constitute about 25% and 75%, respectively of the total ash [1–3]. Many studies have shown that most fly ash consists of silicates, along with unburnt carbon and/or char. The mineral composition is quartz and feldspar and the ash is rich in trace metals [4–7]. Mastering the mineral composition, morphology, heavy metals content and chemical composition of the fly ash is essential for developing its utilization.

Fly ash contains trace heavy metals. If the fly ash is piled outside and comes in contact with rain or other water, the heavy metals will leach from the fly ash and contaminate the soil, groundwater, or surface water. Leaching tests can help establish the leachability and mobility of the heavy metals from the fly ash and help predict the environmental impact associated with ash pond disposal techniques [8].

Skodras studied the chemical, leaching and toxicity characteristics of circulating fluidized bed combustion residues in Greece [9]. The results showed that toxic trace elements preferentially concentrated in the fly ash particles, since they presented a smaller particle size. Wang investigated the leaching behavior and potential environmental impact of the 11 potentially hazardous elements found in coal combustion residues [10]. A similar leaching behavior from both fly and bottom ash was observed. Souhail analyzed the leaching behavior of heavy metals from a mineral processing waste (MPW) [11]. Batch and column leaching tests were conducted on ground MPW, for different liquid/solid ratios and pH values, to determine the metal release mechanisms.

In this study, the physicochemical properties of fly ash were analyzed in three aspects: micromorphology, mineral composition and the content of heavy metals. Batch (equilibrium) and column (dynamic) leaching tests were conducted to study the leachability of the heavy metals in the fly ash.

2. Materials and methods

2.1. Samples

The fly ash was collected from various coal-fired plants in Xuzhou city and other plants in Shandong province. According to the boiler type and desulfurization technology, these samples could be divided into two types. The first type came from plants...
using a pulverized coal-fired boiler and limestone-gypsum wet flue gas desulfurization technology. The second group of samples came from plants using a circulating fluidized bed boiler, where a desulfurization agent is added during the progress of combustion.

2.2. Analytical methods

The micromorphology was observed using a Hitachi S-3500N Scanning Electron Microscopy/Energy Dispersive Spectroscopy (SEM/EDS) device. The mineral composition was measured with a D/Max-3B X-ray Diffraction Analyzer (XRD). Finally, Inductively Coupled Plasma Mass Spectrometry (ICP-MS) was used to analyze the concentrations of the elements.

2.3. Leaching tests

One representative sample was selected for the leaching tests. There were two leaching methods, a batch leaching test and a column leaching test. The batch leaching test is a simple test and provides qualitative predictive information on the leaching behavior of heavy metals. It may not adequately simulate long term and intermittent leaching conditions. However, the column leaching test monitors the heavy metal release over a long period and can identify temporal variations in the contaminant concentrations during the course of its transport through the fly ash [11,12].

In the batch leaching experiment, the samples were leached directly using deionized water. The mass of fly ash was 25 g, the liquid/solid (L/S) ratios were 5 and 10, and the oscillation frequency was set at 110 ± 10 r/min. The duration of mixing was 12 h, after which the samples were allowed to stand for 12 h before the leachates were collected. The column leaching test was done at four different pH values with separate samples being continuously leached. The liquid/solid ratio was 5. The leaching time was seven days, and the leachate was collected throughout the leaching time. For both methods, the leachate was filtered immediately through a 0.45 micron filter membrane and then analyzed by ICP-MS.

3. Results and discussion

3.1. Mineral composition

Coal contains many minerals which are composed of a variety of elements. The minerals change after the coal is fired and salic (mullite and quartz), ferruginous and calcareous minerals are produced [13–16].

Non-crystalline materials made up a large part of the mineral composition of the fly ash. Quartz and mullite were the main crystalline phases (point labeled B in Fig. 1), and other crystalline phases like illite, feldspar, calcspar, gypsum and hematite could also be found in the ash. Because of the high-temperature combustion in the pulverized coal-fired boiler, the quantity of the mullite in the ash from these plants to exceed that seen in plants using a circulating fluidized bed (Fig. 2a). Furthermore, the type of coal also influences the mineral composition of the ash. The circulating fluidized bed burns coal of low calorific value and adds the desulfurization agent during combustion, so the mineral composition of the fly ash from this boiler is complex and the number of species is very high (Fig. 2b).

All the fly-ash samples have oxides of silicon, aluminum and iron as the main chemical components of the ash. The sum of these components was more than 80% of the total. Silicon oxides were the main chemical component of the fly ash, and the content of aluminum and iron oxides were lower than that of silicon oxides. There were also other metal oxides present, such as CaO. The content of CaO was less than 10% in the fly ash from these power plants, so the fly ash was classified as a low calcium ash.

3.2. Heavy metals composition

In addition to the normal elements, some trace elements were also included in the ash. The quantity of these trace elements varied considerably with different power plant. Some levels were capable of producing a cumulative toxic effect that could negatively affect the environment [5,17,18]. It was important to detect the types and contents of these heavy metals. Table 1 lists the elemental composition of fly ash (XZHX-FA, XZHM-FA) and bottom ash (XZHX-B, XZHM-B) from two different plants (XZHX, using the pulverized coal-fired boiler; XZHM, using the circulating fluidized-bed boiler).

The content of trace elements was significantly different in the different power plant ash. For the studied elements, the content of Ti was the highest in the fly ash, ranging from 5795 to 7382 µg/g. The content of Cd was the lowest, ranging from 0.19 to 0.94 µg/g. The contents of the elements in the fly ash were higher than those found in the bottom ash.
in the bottom ash, even though the samples came from the same plant. The metallic elements are easily enriched in the fly ash. Even when the ash came from the same type of boiler, but from different plants, the contents of the fly ash and bottom ash were different. Pb, for example, the minimum value was 44.65 plants, the contents of the fly ash and bottom ash were different.

3.3. Leaching tests

Fly ash from the SDJN power plant was selected as the leaching sample in this study, because the flue-gas desulfurization and leaching effect is obvious in this power plant. This plant uses the circulating fluidized bed boiler.

![Fig. 3. Concentration of heavy metals in the leachates: batch leaching.](image1)

![Fig. 4. Leaching rates of heavy metals: batch leaching.](image2)

3.3.1. Batch leaching test

In the batch leaching test, the concentrations of Ni in the leachates were the highest, while the concentrations of Cd were the lowest (Fig. 3). When the L/S ratio was increased, most of the heavy metal concentrations decreased. The concentrations of Ni, Cr and Mo in the leachates each exceeded the national standard and had the potential to harm the environment.

However, the leaching rates of Ni, Mo and Cd were higher than the other elements, and the rate for Cd was the highest, showed that Cd leached more easily from the fly ash (Fig. 4).

3.3.2. Column leaching tests

The column leaching test was done to analyze the leaching over time. In these trials Ni and Zn concentrations exceeded 100 µg/L in the leachates, and Cr, Mo and Mn were between 40 and 100 µg/L. Co, Cu, Cd, Pb and V were all less than 20 µg/L. Three different kinds of leaching curve were observed for these elements, see Fig. 5. The first type was a downward trending curve where the leached metal content decreased slightly with time. The harm caused to the environment by these elements will gradually diminish. Ni and V have this type of behavior. The second type was an increasing trend where the leached concentration increased with time. Cr and Co are part of this group. The last type was a fluctuating curve where the elemental concentrations fluctuated with time. The other concerned heavy metals examined belong in this group.

Different pH values affect leaching. The results showed that the lower pH values of the leaching solutions, the higher metal contents in the leachate. The reason for this might be that an acidic solution accelerates the mineral dissolution in the liquids.

3.3.3. Contrasting the batch and column leaching tests

Because the cumulative L/S ratio in the column leaching test was 5, and because deionised water was used as the liquid medium in the batch leaching test, the batch leaching test with an L/S ratio of 5 is compared to the column leaching test using deionised water as the eluent.

The fly ash and deionised water were mixed completely and were in continuous contact during the batch leaching tests, while the fly ash was in gradual contact with the water with time during the column leaching tests, which is a dynamic process.

Comparing these two leaching tests, the results showed that the leaching rate was both higher and lower in the batch leaching.

![Fig. 5. Concentration of Ni, Cr and Pb in leachates: continuous leaching.](image3)
test, depending upon the particular element. This was also true in the column leaching test. The batch leaching test promoted the leaching of Ni, Cr, Mo, Co and V from the fly ash, while the column leaching test promoted the leaching of Mn, Zn, Cu, Pb and Cd (Fig. 6). In the batch leaching test, the fly ash and water contacted completely, and friction between solid particles and solid-water particles accelerated the dissolution of the heavy metals. In the column leaching test, elemental concentration gradients existed between the fly ash and the deionised water could affect the release of the heavy metals.

Since the column leaching test was a better simulation of fly ash contact with rain or other liquids in a natural state, this method was used as a reference. However, the batch leaching test was mainly used to evaluate the maximum leaching toxicity of the solid.

3.3.4. Effect of the leaching test on fly ash micromorphology

The fly ash can be classified as spherical particles and irregular particles, according to the SEM analysis of their shapes and micromorphology [13,19]. The sample from this plant which did the leaching tests in my study was more irregular particles before the experimental leaching treatment.

After the leaching tests, parts of the components leached from the fly ash into the leachates and the microstructure of the particles changed. The solid samples from the column leaching test were dried and then analyzed by SEM. The samples were different in microstructure compared to the fly ash before the leaching test.

Fig. 7 shows the particle micromorphology before the leaching test. The particle size was larger (Fig. 7a) and the surface was coarse with a number of attached crystal clusters were (Fig. 7b). The EDS map from point B suggested the chemical compositions were mainly C, O, Al, Si and Ca (Fig. 7c).

When the fly ash was leached by deionized water, the particle size shrank compared to the original particles (Fig. 8a). This is a result of water erosion of the particles and the chemical dissolution of the elements. The surface of the particles was still coarse, but the quantity of crystals attached to the surface decreased (Fig. 8b). The elemental composition of corresponding points did not change too much, but Fe and K were detected (Fig. 8c).

Comparing Figs. 7a and 9a, the particle sizes were clearly reduced after leaching with the acid solution. Had the acidity of the leaching fluid been stronger, the particles size would possibly have been even smaller. The main reason for this size reduction was the strong chemical decomposition of the fly ash in the acidic solution.

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**Fig. 6.** Leaching rates of heavy metals from fly ash for different leaching tests.

**Fig. 7.** SEM photomicrographs and EDS maps of fly ash before leaching.

**Fig. 8.** SEM photomicrographs and EDS maps of fly ash after leaching with deionized water.

**Fig. 9.** SEM photomicrographs and EDS maps of fly ash after leaching with nitric acid solution at pH = 2.
4. Conclusions

Fly ash from plants using the circulating fluidized bed contained more irregular particles. Because of the desulfurization technology used, the desulfurization product was mainly gypsum crystals in these samples. Samples from plants using a pulverized coal-fired boiler have more spherical particles. All samples contained more non-crystalline material in the ash, regardless of their origin. Crystalline phases like illite, feldspar, calc spar, gypsum and hematite could also be seen in the ash, along with the main minerals quartz and mullite. The heavy metals tested for all appeared in amounts above 1 μg/g, except for Cd. Ti had the highest content, which was above 7300 μg/g. Most of the heavy metals tested were enriched in the fly ash. Heavy metals were hard to leach from the fly ash, but the concentration of Ni exceeded the standard using both the leaching methods. This might constitute a threat to the environment.

Acknowledgments

Financial support for this work, provided by the Europe-Asia Link (No. CN/ASIA-LINK/010 94556) and State Scholarship Fund of China Scholarship Council (No. 2010642035), are gratefully acknowledged.

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