Feasibility of Using Coal Fly Ash for Mine Waste Containment

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ABSTRACT
This study investigates the feasibility of using coal fly ash and fly ash-bentonite mixtures as a barrier material for mine waste. The hydraulic conductivity of the coal fly ash was measured to be in the order of $2 \times 10^{-9}$ m/s when it was permeated with deionized water, and this value decreased significantly when the permeant was switched to acid mine drainage (AMD). Addition of bentonite to coal fly ash lowered the hydraulic conductivity during water permeation but no further significant change was observed upon switching the permeant to AMD. Chemical analyses on the effluent from the hydraulic conductivity tests indicated that heavy metals present in AMD were attenuated and were well below the leachate criteria set by the Ontario Government. X-ray diffraction and scanning electron microscopy analyses results of post permeation samples showed significant structural differences and formation of secondary minerals after AMD permeation. The results of this study suggest that the addition of 10% bentonite to coal fly ash reduced the hydraulic conductivity of the coal fly ash to less than $1 \times 10^{-9}$ m/s and improved the chemical compatibility for mine waste containment.

Key words: fly ash; bentonite; acid mine drainage; mine waste containment; hydraulic conductivity; saturation index.

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INTRODUCTION
Acid mine drainage (AMD), the leachate from oxidized mine waste rock and tailings, has been a major environmental challenge for the mining industry. AMD is formed when sulfide-rich mine waste oxidize in the presence of water and oxygen, and is characterized by low pH, high acidity, elevated heavy metal and sulphate concentrations. AMD has been known to cause severe impacts on surface and ground water systems. Current practices of prevention and mitigation measures include the use of water covers or underwater disposal, use of soil covers as oxygen and infiltration barriers, and treatment of acidic effluent by lime neutralization.

Solid contaminant barriers can provide efficient and relatively inexpensive means of containing mine wastes. Many different barrier materials, notably compacted soil liners and geotextiles, have been used for the containment of municipal solid wastes. Compacted soil liners are commonly used to control or restrict contaminant migration from the landfill into the environment. When suitable natural soils are not available for use as a compacted clay liner, bentonite may be added to a non-cohesive soil (e.g., silty sand) to achieve a liner with acceptable hydraulic conductivity (Daniel 1993). Bentonitic clay materials are preferred because of their low hydraulic conductivity and good adsorption or retention capacity (Daniel 1993). However, application of clay, clay-bentonite mixtures and polymer based synthetic liners may become extremely expensive because of the lack of suitable clay materials at or near the disposal site or because of high costs of synthetic liners (Cokca and Yilmaz 2004). Therefore, re-use of industrial waste products and by-products such as coal fly ash can be a viable alternative for barrier construction and an important step toward sustainability.

Owing to its pozzolanic and highly alkaline properties resulting from its high calcium content, fly ash from coal-fired power generating stations has the potential to serve as a hydraulic and contaminant barrier. Various applications of coal fly ash as hydraulic barriers have been studied by a number of researchers. Usmen et al. (1988) investigated the use of fly ash in landfill liners and landfill covers because of its low permeability characteristics. They reported permeabilities on the order of $7 \times 10^{-6}$ to $2 \times 10^{-7}$ m/sec for two Class C fly ashes. Edil et al. (1992) also reported the use of fly ash with or without sand, as a construction material for waste containment liners and impermeable covers. Bowders et al. (1987) evaluated the potential for using Harrison and Amos Class F fly ashes from West Virginia in hydraulic barriers. Varying amounts of lime or
cement were mixed with the fly ashes and test specimens were compacted at their optimum water content using the standard Proctor method. The hydraulic conductivity tests showed that unstabilized Harrison and Amos fly ashes had hydraulic conductivities of $7.2 \times 10^{-8}$ and $5 \times 10^{-7}$ m/s, respectively using distilled-deionized water as permeant. Hydraulic conductivity of Harrison ash dropped about an order of magnitude when 15% lime was added. Studies on the utilization of coal fly ash in combination with clay in landfill bottom liners were also performed by Nhan et al. (1996) and Cokca and Yilmaz (2004). Nhan et al. (1996) investigated the hydraulic conductivity of Lakeview Class F fly ash mixed with lime kiln dust and bentonite. They found that with the addition of lime dust and bentonite, coal fly ash liner has a hydraulic conductivity of $4.3 \pm 1.6 \times 10^{-8}$ m/s. They also found the addition reduced the dissolved metal concentration in the synthetic municipal solid waste leachate. Cokca and Yilmaz (2004) investigated the combination of rubber, bentonite and coal fly ash as a liner material. The composite material had a hydraulic conductivity of less than $10^{-9}$ m/s using water as permeant and the effluent leachate was found to be non-hazardous.

The permeants used in previous laboratory studies were mostly distilled water or synthetic alkaline solution such as municipal solid waste leachate. In the present study, the use of acidic permeant in the utilization of coal fly ash as a barrier for AMD containment was investigated. Further investigation of the enhancement of the barrier performance by addition of bentonite was also undertaken. The specific tasks performed in order to achieve the above objective are itemized as follows: (i) evaluation of the hydraulic conductivity of compacted Atikokan coal fly ash (referred to as AFA hereafter) and fly ash-bentonite mixture (referred to as BAF hereafter) permeated with distilled water and acid mine drainage; (ii) evaluation of the efficiency of AFA and BAF samples in neutralizing AMD and removing contaminants contained in the AMD; and (iii) determination of the optimum mix of the compacted fly ash-bentonite mixture in terms of the hydraulic conductivity and overall chemical compatibility.

MATERIALS AND EXPERIMENTAL METHODS

Materials Used

Characterization of fly ash and bentonite used
The fly ash used in this study was collected in the dry state from Atikokan Thermal Generating Station located 190 km west of Thunder Bay, Ontario, Canada. The physico-chemical properties and environmental evaluation of Atikokan coal fly ash for environmental applications are described in detail in Yeheyis et al. (2008). The major chemical composition of Atikokan fly ash calculated as major oxides are SiO$_2$, Al$_2$O$_3$, Fe$_2$O$_3$, CaO and Na$_2$O contributing 43.83%, 21.85%, 4.01%, 14.36% and 7.15% (w/w) of the total, respectively. The next most abundant components are MgO, K$_2$O, P$_2$O$_5$ and SO$_3$, contributing about 2.71%, 0.51%, 0.54% and 1.05% (w/w) each. It also consists of trace elements such as manganese, barium, zinc, lead, chromium, copper and vanadium. According to American Society for Testing and Materials Standard Specification for Coal Fly Ash and Raw or Calcined Natural Pozzolan for Use as a Mineral Admixture in Portland Cement Concrete (ASTM C 618), Atikokan fly ash is classified as a Class C fly ash. The specific surface area and specific gravity of the AFA were found to be 0.6357 m$^2$/g and 2.73, respectively. AFA contains 88% fines (< 0.075 mm) with predominantly silt-sizes (0.075 mm - 0.002 mm). The results obtained from Yeheyis et al. (2008) also demonstrated that Atikokan fly ash are non-toxic and can be re-utilized for various environmental applications.

In addition to pure fly ash, bentonite added fly ash samples were prepared and tested to study its engineering property as a potential mine waste barrier material. The bentonite used in this study was sodium montmorillonite from Wyoming, USA. The liquid limit (LL) of the bentonite was 500% and the plastic index (PI) was 400%. Its cation exchange capacity (CEC) measured by silver thiourea method (Chhabra et al. 1975) and KCl extraction (Barone et al. 1989) amounts to 76 meq/100g and the predominant minerals present in the bentonite were quartz, feldspar, calcite, montmorillonite and mica. Two fly ash-bentonite mixtures were tested in this study, containing 5% bentonite (5% BAF) and 10% bentonite (10% BAF), respectively. The mixtures were prepared on the basis of dry weight and proper care was taken to ensure homogeneity of the mixes. The fly ash-bentonite mixtures tested were chosen to represent the probable field range of bentonite contents.

**Acid mine drainage permeant**

The samples were first permeated with distilled water and then with AMD generated in the laboratory from reactive mine rocks and tailings. The AMD used in the study had a pH of
2.78 and contained many of the chemical constituents of natural AMD. It had high concentration of metals including Fe (738.9 mg/L), Mn (106.7 mg/L), Zn (880.5 mg/L), Cu (53.4 mg/L), Ni (89.8 mg/L), Cr (6.7 mg/L), Cd (4.5 mg/L), Pb (8.7 mg/L), As (1.4 mg/L), Si (25.3 mg/L), Ca (512.1 mg/L) and Al (83.2 mg/L). It also contained high sulphate concentration (10,100 mg/L) and electrical conductivity of 9.31 mS/cm (@ 25 °C).

**Experimental Methods**

**Hydraulic conductivity measurement**

In order to assess the suitability of AFA and BAF samples as barrier materials, laboratory hydraulic conductivity tests were carried out and the results were evaluated to see if they satisfy the general requirement of hydraulic conductivity for contaminant barriers (i.e. 10^{-9} m/s or less; Daniel 1993) and to assess the chemical compatibility with AMD. The tests were performed on fly ash with 0, 5 and 10% bentonite contents using a fixed-wall constant-flow-rate permeameter consisting of four permeameter cells that run at the same flow rate (Fernandez and Quigley 1985). The relative advantages of the fixed-wall permeameter have been discussed by Bowders et al. (1986) and Daniel et al. (1985). At the beginning of the test, the samples were compacted in the sampling cylinder (permeameter cell) in three layers with a small spring-loaded compactor. The samples were compacted at about 2% above optimum moisture content (2% wet of optimum) determined from the standard Proctor moisture-density relationships. Each test sample had a diameter of 5 cm and was trimmed to 2 cm by the rotation of a T-shaped trimming plate. A confining pressure of 32 kPa was applied to the samples by means of a calibrated stainless steel spring-loaded device, to prevent the swelling of the samples and eliminate sidewall leakage (Kashir and Yanful 2001). Subsequently, each sample specimen was subjected to a constant hydraulic flow that was applied via a motor driven piston press. The samples were first permeated with distilled water to obtain the reference hydraulic conductivity of the specimen simulating pre-weathering condition of mine waste. After the specimens were brought to equilibrium with distilled water, the AMD was introduced into the specimen. During the experiments, room temperature, sample thickness, flow rate, and influent pressure were continually monitored and recorded by automated data acquisition system. A fluid outlet at the bottom of the cell allowed effluent to be collected. The hydraulic conductivity, pore volume of flow, and hydraulic head were calculated from the imposed constant flow rate.
Effluent chemistry analysis

The hydraulic conductivity and its susceptibility to changes with time or exposure to chemicals are the major factors in the selection of clay for use in waste containment barriers (Mitchell and Madsen 1987). In order to evaluate chemical compatibility of the barrier samples, the effluent from each cell was collected periodically for chemical analysis during permeation. pH and sulphate were analyzed immediately after sample collection. The pH was measured with a combination electrode (Orion 410A) calibrated at pH 4, 7 and 10. The sulphate concentration was measured using high-performance liquid chromatography (HPLC), and metals in acidified filtrates were analyzed using inductively coupled plasma optical emission spectroscopy (ICP-OES).

Post-testing analysis

Following completion of hydraulic tests, the samples were extruded from the cells and sub-samples were analyzed for mineralogy, morphology, major oxides and trace metals in order to identify changes after AMD permeation. Mineralogical analyses were carried out on random powder samples by means of x-ray diffraction (XRD) analysis, using a D/max Rigaku diffractometer (40 kV, 160 mA) with cobalt Kα radiation. A scanning electron microscope (Hitachi S-4500 SEM) was used to observe changes in the morphology of the surface after permeation. Major and trace elements of the samples were analysed by using x-ray fluorescence (XRF) technique (Philips PW-1400).

RESULTS AND DISCUSSION

Hydraulic conductivity

The results of hydraulic conductivity tests on AFA and two BAF samples permeated first with distilled water and then with AMD are presented in Figure 1. The AMD was introduced after the hydraulic conductivity of the samples had been stabilized with distilled water, which occurred after approximately 5.25 pore volumes of water permeation. The experiment was run for a total of approximately 30 pore volumes (PVs) of water and AMD permeation.
The average hydraulic conductivity of AFA was initially $2.5 \times 10^{-9}$ m/s within the first 0.55 PVs of water permeation, increasing slightly to $5.8 \times 10^{-9}$ m/s between 0.65 PVs and 1.25 PVs. The hydraulic conductivity then more or less stabilized to an average value of $2 \times 10^{-9}$ m/s until the end of water permeation (the first 5.25 PVs). After switching the permeant to AMD, the hydraulic conductivity decreased steadily and became lower than $1 \times 10^{-9}$ m/s after 10.8 PVs, and further decreased sharply to less than $4 \times 10^{-11}$ m/s at the end of AMD permeation (30 PVs). This sharp decrease in the hydraulic conductivity of the AFA sample is due to permeant-fly ash interactions. The pozzolanic and self-cementing properties of Atikokan fly ash result in the formation of hydration products that could possibly block void spaces and reduce the interconnection between fly ash particles. The precipitation of new minerals as result of chemical interaction between AMD and fly ash could also contribute to clogging of the pores between the particles. These two processes could lead to a reduction in hydraulic conductivity. Since the hydraulic conductivity of the fly ash during the first 11 PVs was higher than $1 \times 10^{-9}$ m/s, the target regulatory hydraulic conductivity for compacted soil liners, one may conclude that the fly ash alone may not be suitable for use as a liner by itself.

As seen in Figure 1, the addition of bentonite decreased the hydraulic conductivity of the fly ash-bentonite mixtures during water permeation. The average hydraulic conductivity of 5% BAF was initially $2.2 \times 10^{-9}$ m/s during the first 0.55 PVs of water permeation, and followed a trend similar to the AFA by increasing slightly to $3.75 \times 10^{-9}$ m/s between 0.65 PVs and 1.25 PVs and then eventually stabilizing to an average value of $1.25 \times 10^{-9}$ m/s until the end of water permeation 5.25 PVs. The average initial hydraulic conductivity of 10% BAF was $2 \times 10^{-10}$ m/s, decreasing approximately one order of magnitude from that of AFA during water permeation. However, upon switching to AMD the hydraulic conductivity of the mixture samples remained fairly constant in contrast to the pure fly ash sample (AFA) which kept on decreasing sharply. This shows the existence of an opposing phenomenon during AMD permeation in the samples containing bentonite. This opposing phenomenon could be attributed to clay double-layer contraction due to the displacement of the resident bentonite pore water by the AMD permeant. This is consistent with the findings of Kashir and Yanful (2001), who observed an increase in the hydraulic conductivity of bentonite filter cake permeated with AMD. The hydraulic conductivity of the 10% BAF sample remained steady relative to that of the 5% BAF sample, indicating that
bentonite-AMD interactions were more pronounced on the sample with higher bentonite content. The AMD attack on pure fly ash was comparatively low.

The distance between the negatively charged clay particle surface and the centre of mass of the diffuse double layer, 1/κ, may be related to the thickness of the double layer (Mitchell 1993). For a nonsymmetrical electrolyte containing various ions, κ may be given by (Hunter 1993):

\[ \kappa^2 = \frac{(e^2 \sum n_i z_i^2)}{\varepsilon \kappa T} \]  

(1)

Where \( e \) is the charge of the electron \((1.60 \times 10^{-9} \text{ C})\); \( n_i \) is the number of ions of type \( i \) in the bulk solution per cubic metre; \( z_i \) is the valence of ion \( i \); \( k \) is Boltzmann’s constant \((1.38 \times 10^{-23} \text{ J/K})\); \( \varepsilon \) is the permittivity of the bulk solution \((\text{C}^2/\text{J} \cdot \text{m})\); and \( T \) is the absolute temperature \((\text{K})\).

Kashir and Yanful (2001) showed that the ionic strength, \( I \), of effluents from compacted clays permeated with AMD could be related to \( \kappa \) as follows:

\[ \kappa = 3.28 \sqrt{I} \text{ } (\text{nm}^{-1}) \]  

(2)

Where \( I \) is the ionic strength.

Equation (2) was used to estimate double layer thickness of AFA and BAF samples. The ionic strength of the effluents sampled during the hydraulic conductivity tests were computed using MINTEQA2 (described in the next section). The variation of the diffuse double layer thickness surrounding the fly ash and bentonite particles in each compacted AFA and BAF samples during AMD permeation is presented in Figure 2. When the samples were permeated with distilled water, the ionic strength decreased due to the flushing of dissolved salts from the samples resulting in an increase in double layer thickness. After switching the permeant to AMD (i.e., 5.25 PVs), the double layer thickness reduced significantly up to 10 PVs, but remained fairly constant thereafter. Although double layer reduction might have occurred in all samples, it was evidently greater in BAF samples compared to the reduction in AFA, resulting in an increase in hydraulic conductivity. The change in hydraulic conductivity that occurred as a result of double layer contraction was likely compensated by a decrease in hydraulic conductivity due to pore clogging, as indicated by the steady hydraulic conductivity values for 10% BAF after 10 PVs of permeation.
The 5% BAF sample also did not meet the minimum hydraulic conductivity (1 x 10^{-9} \text{ m/s}) criterion during the early permeation stages (i.e. the first 5.6 PVs), and may not be suitable for use as barrier for mine waste containment. Fly ash amended with 10% bentonite satisfied the regulatory hydraulic conductivity requirement (1 x 10^{-9} \text{ m/s}) throughout the test. From these results, one may conclude that, compacted fly ash containing 10% bentonite may be required to achieve acceptable level of hydraulic conductivity for mine waste containment. It is also possible that slightly lower bentonite contents (6-8%) may be acceptable.

**Effluent chemistry analysis**

**Effluent pH and Sulphate**

Figure 3 shows pH measurements as a function of effluent pore volumes. When the samples were permeated with distilled water, the initial effluent pH remained highly alkaline (greater than 12.0). The pH decreased slowly to approximately 11.5 after 5.25 PVs of distilled water had passed through the samples. Upon switching the permeant to AMD, the effluent pH decreased more quickly compared to distilled water permeation because of the strong acidity of AMD. Both AFA and BAF samples showed essentially similar pH behavior. However, during the first 10 PVs of permeation, the pH of effluents from the BAF samples was slightly higher than that of the AFA sample (i.e. pH of 10\% BAF > 5\% BAF > AFA). This trend completely reversed after 13 PVs (i.e. AFA > 5\% BAF > 10\% BAF) and then stabilized more or less thereafter. The effluent pH of the samples showed plateaus from PV = 8 to PV = 14 and from PV = 19 to PV = 23 indicating that the fly ash showed high buffering capacity at about pH of 10.8 and 9.1 respectively. The alkaline oxides and calcium carbonate present in fly ash and fly ash-bentonite samples provided this buffering capacity. Although the influent pH of AMD was strongly acidic (2.78) after 30 total PVs, the effluent pH remained alkaline at pH 8.9 (compared to 11.5 with distilled water). This high effluent pH is attributed mainly due to dissolution and hydrolysis of CaO and other major alkali contributing oxides (MgO, Na_2O, and K_2O) originally present in fly ash and fly ash-bentonite mixtures. This implies that in addition to its low hydraulic conductivity, alkalinity of fly ash has the capacity to offset and neutralize highly acidic AMD asserting its potential use for mine waste containment.
The sulphate concentration of the effluent from all samples was monitored since it is a dominant anion in mine drainage waters (Figure 4). The sulphate concentration dropped from approximately 3000 ppm to 110 ppm during water permeation. This was likely due to dissolution of sulphate salts and anions from fly ash and fly ash-bentonite samples. Upon switching the permeation to AMD, the sulphate started increasing and reached approximately 1900 mg/L and remained constant thereafter. Although the trend in effluent sulphate concentration during water and AMD permeation was more or less similar in all samples, the fly ash sample has lower sulphate concentration compared to BAF samples. Comparing BAF samples, the fly ash with 5% BAF also showed relatively lower sulphate concentration than 10% BAF during most of the test period.

**Effluent chemical composition and geochemical modeling**

In order to establish chemical compatibility of the barrier samples and determine the fate of the metals present in AMD, the effluent metal concentrations were analyzed and presented in Figure 5 (a) to (l). The data are plotted as absolute effluent concentration (in mg/L) versus pore volumes. The effluent concentrations were also compared to Ontario Ministry of Environment Regulation 558 (MOE Reg. 558) leachate criteria (MOE, 2000).

Geochemical modeling was also conducted to determine whether specific solid phases were in equilibrium with the permeant, as an indication of possible precipitation and solid phase control of the metal concentrations. The geochemical equilibrium model Visual MINTEQ version 2.50 (Gustavsson 2006), a Windows based version of MINTEQA2 (Allison et al. 1991), was used for saturation indices calculation. The saturation index (SI) for an aqueous solution with respect to a mineral indicates the thermodynamic tendency to precipitate or to dissolve certain phases. The saturation index is defined by:

$$ SI = \log \left( \frac{\text{IAP}}{K_{sp}} \right) $$

(3)

where $\text{IAP}$ is the ion activity product calculated from the water sample, and $K_{sp}$ is the theoretical solubility product, both adjusted to the temperature of the sample. A negative value of the saturation index indicates that the solution is undersaturated with respect to a particular solid phase and that the solid phase would tend to dissolve if present, whereas a positive value
indicates the tendency for a mineral to precipitate. A value close to zero suggests that the mineral is in equilibrium in the solution and may either precipitate or dissolve.

The input parameters were measured pH, temperature and concentrations of inorganic constituents of the effluent (As, Al, B, Ca, Cd, Cr, Cu, Fe, K, Mg, Mn, Na, Ni, Pb, Zn, and Si) and SO₄. The pH was fixed and solids were allowed to precipitate. The thermodynamic database provided by the code was used. Both AFA and BAF samples showed similar effluent chemistry and the difference was minimal. As a result, detail discussion on the effluent chemistry as well as geochemical modeling was presented on the 10% BAF sample only. Table 1 presents the calculated saturation indices (SI) for some minerals that can affect the concentration of heavy metals in the effluent.

The initial effluent concentration of aluminium from the hydraulic conductivity testing cells was relatively high (~ 140 mg/L) but reduced to approximately 60 mg/L during water permeation (within 5.25 PV), and further reduced to less than 0.05 mg/L at the end of AMD permeation (Figure 5 (a)). Geochemical simulation using vMINTEQ showed the effluent was supersaturated with respect to the aluminium bearing mineral, diaspore (α-AlO(OH)) during the test. The effluent was also supersaturated with respect to crystallized gibbsite (γ-Al(OH)₃) during the test, except between PVs = 6.9 to PVs = 14 when the effluent was slightly undersaturated. Similarly, effluent iron and copper concentrations decreased significantly compared to the corresponding influent AMD concentration as shown in Figure 5 (b) and Figure 5 (c), respectively. Geochemical modeling results indicated that the effluent was supersaturated with respect to the iron oxyhydroxides: hematite (α-Fe₂O₃), goethite (α-FeOOH) and ferrihydrite (Fe₅HO₈·4H₂O). The effluent also appeared to be supersaturated with respect to crystalline tenorite (CuO) throughout the test. The model also predicted that the effluent was at or near saturation with respect to Cu(OH)₂. Cu likely co-precipitated with Fe as cupric ferrite. The effluent was also supersaturated with respect to crystalline Ni(OH)₂ during the first 14 pore volumes of permeation, and was slightly undersaturated thereafter. The effluent concentrations of chromium and lead were significantly reduced and were far below the Reg. 558 leachate criterion of 5 mg/L for both metals. Geochemical modeling using vMINTEQ predicted precipitation of Cr(OH)₂ for the first
10.5 pore volumes (at pH values greater than 10.9). The effluent was also supersaturated with respect to Pb(OH)$_2$ through out the test period.

Effluent calcium concentration showed a distinctly different behavior (Figure 5 (L)). The effluent calcium concentration decreased and became relatively low (~ 10 mg/L) during water permeation. However, after switching the permeant to AMD, the concentration of Ca started to increase and reached 700 mg/L after 10 PVs, and then remained stable during the rest of the experiment. This trend is similar to that of SO$_4^{2-}$ concentration suggesting that both ions were mainly controlled by gypsum and ettringite solubility as predicted by vMINTEQ. The computed saturation indices indicated that both gypsum (CaSO$_4$.2H$_2$O) and ettringite (Ca$_6$Al$_2$(SO$_4$_3)(OH)$_12$.26H$_2$O) were at or near saturation state.

Given an influent AMD highly loaded with hazardous heavy metal ions, the effluent chemical analysis results showed significant decrease in effluent concentration. All regulated elements have concentrations below the MOE Reg. 558 leachate criteria. This significant reduction is mainly attributed to the precipitation of metals as metal hydroxides in high alkalinity environment obtained from the coal fly ash. The reduction in effluent concentration could also be due to the encapsulation of the metals by hydration products formed during fly ash-permeant interaction, as discussed in the next section of this paper. The effluent chemical analysis showed the fly ash and fly ash-bentonite mixture barriers are compatible and efficient in retarding heavy metals from AMD.

**Post-test solid samples analysis**

**Mineralogical analysis**

X-ray diffraction (XRD) patterns obtained from powder samples of AFA, 5% BAF and 10% BAF before and after permeation are presented in Figure 6, Figure 7 and Figure 8 respectively. The major crystalline minerals present in the pre-permeation AFA sample (Figure 6-a) are quartz (SiO$_2$), mullite (Al$_6$Si$_2$O$_{13}$) and anhydrite (CaSO$_4$). It also contained a significant amount of amorphous phase as shown by a broad diffraction hump between 20 =15° and 20 = 35°. The AFA sample after permeation showed significant structural alterations compared to the original fly ash (Figure 6-b). The XRD pattern of the post permeated AFA sample showed the presence
of new, strong peaks at d-spacings of 0.762 nm (2θ = 11.60°) and 0.306 nm (2θ = 29.07°), which correspond to that of gypsum (CaSO₄·2H₂O). Gypsum had formed as a result of the reaction between the calcium from fly ash and sulphate from AMD. Gypsum is normally stable at pH values lower than 10.5, and this is verified as shown from the vMINTEQ geochemical modeling results (Table 1). The other major mineral phase detected on the x-ray traces is ettringite (Ca₆Al₂(SO₄)₃(OH)₁₂·26H₂O). Ettringite is reported to exist as a stable phase in the pH range of 10.5 to 12.5 (Damidot and Glasser 1993). However, Mynenei et al. (1998) reported that ettringite may still be present down to a pH of 9.5, dissolving partially to gypsum and aluminium hydroxide. Although the final effluent pH from the columns were slightly less than 9.5, characteristic peaks of ettringite were identified at d-spacing of 0.966 nm (2θ = 9.14°) and 0.561 nm (2θ = 15.78°). Ettringite is formed through the reaction of calcium and alumina mainly from fly ash and sulphate ions present in AMD, in the alkaline environment created by the fly ash as indicated by the following reaction:

\[
6Ca^{2+} + 2Al(OH)_4^- + 3SO_4^{2-} + 4OH^- + 26H_2O \rightarrow Ca_6Al_2(SO_4)_3(OH)_{12}·26H_2O
\]

The anhydrite peak, which appeared as a weak trace at 0.348 nm (2θ = 25.56°) on the x-ray diffractogram for the unpermeated fly ash sample, disappeared after AMD permeation indicating conversion to ettringite and gypsum.

The x-ray diffraction patterns of the BAF samples after permeation shown in Figure 7-b and Figure 8-b are similar to the pattern of post permeation AFA, except for some changes in the XRD intensities of the major peak associated with the different phases. The ettringite peak intensity generally became relatively weaker and the gypsum peak intensity became relatively stronger with the increase in the bentonite content. This was probably due to the fact that lower pH in high bentonite content samples favored gypsum formation through the dissolution of ettringite.

Most precipitates predicted by vMINTEQ were not confirmed by XRD analysis, probably because they were present in amounts below the detection limits or were poorly crystallized.

**Scanning electron microscopy (SEM) analysis**

Scanning electron microscopy (SEM) was employed to investigate the morphological changes of the fly ash and fly ash-bentonite mixtures after AMD permeation. SEM photographs of natural
and AMD-permeated samples are presented in Figure 9 (a) to (e). It is observed that the original fly ash before AMD permeation showed smooth spherical texture. However, the SEM photograph of post-permeation solid samples (Figure 9 (b) to (e)) showed agglomeration and deposits on the surfaces of fly ash and bentonite particles. This change in surface morphology on post permeation samples is attributed to the formation of new precipitates and cementitious matrix due to the inherent pozzolanic and cementitious property of fly ash used. The formation of gypsum crystal on post permeated samples was seen clearly.

**Physico-chemical analysis**

The physico-chemical characteristics of post permeation samples are presented in Table 2. The changes in the major oxide compositions of post permeation BAF samples showed similar trend to the post permeation AFA (Table 2). Comparing the major oxides of samples before and after AMD permeation, the result showed a general trend of decrease in major oxides. The decrease in basic oxides (CaO, MgO, Na₂O and K₂O) in post permeated samples is attributed to their dissolution which is the main process contributing to the alkalinity of the fly ash. The basicity (CaO/SiO₂) of AFA fly ash decreased from 0.33 before permeation to 0.28, showing 18% decrease at the end of AMD permeation. Another noticeable change occurred in the sulfide content (SO₃), which showed a significant increase after permeation. It increased from 1.05% prior permeation to 1.24% after AMD permeation of AFA, an 18% increase. This increase is probably attributed to sulfide minerals precipitated during permeation. There was also a large difference in the loss on ignition (LOI) of fly ash prior to and following permeation with AMD. The LOI of post permeation AFA increased from 0.48% to 7.32%. This significant increase in LOI could be due losses of the newly formed sulphates, carbonates and other compounds. The concentration of most trace metals in all solid samples increased after permeation. Trace metals including zinc, cobalt, copper, nickel and manganese increased significantly compared to the concentrations in the original fly ash before permeation (Table 2). The alkaline environment generated by coal fly ash immobilized the migration of heavy metals from AMD. Consequently, the heavy metals were precipitated and/or sorbed to the surface of the specimen.
SUMMARY AND CONCLUSIONS

Laboratory experiments were performed to investigate the hydraulic conductivity of fly ash and fly ash-bentonite mixtures at different mix proportions and its chemical compatibility with mine waste (AMD). The following conclusions are drawn from the study.

1. The test results showed that pure Atikokan fly ash did not meet the hydraulic conductivity regulatory requirement for compacted soil liners (i.e. 1 x 10^-9 m/s) during the first 11 pore volumes. Hence the fly ash alone is not adequate for use in mine waste containment. The hydraulic conductivity of the fly ash-bentonite mixtures decreased with increasing bentonite content during distilled water permeation. However, the hydraulic conductivity of BAF samples remained the same after AMD permeation. This was due to the decrease in the hydraulic conductivity due to pozzolanic reactions of fly ash and chemical precipitation is offset by the increase due to reduction in the double layer thickness as a result of AMD attacking the fly ash and bentonite particles. Based on these laboratory results, 10% bentonite may be adequate for fly ash-bentonite mixture to attain an acceptable level of hydraulic conductivity for mine waste containment. However, field applications may depend on several factors including proper mixing of bentonite with fly ash and AMD chemistry.

2. Chemical analysis results showed fly ash provides alkalinity to the system and therefore is able to precipitate heavy metals from AMD. As a result the concentrations of most heavy metals in the effluent decreased significantly and were well below MOE Reg. 558 leachate criteria. This shows that the coal fly ash and fly ash-bentonite mixtures barriers are efficient in removing hazardous metals from AMD and Atikokan coal fly ash can be used for mine waste containment.

3. Geochemical modeling of the effluent chemistry, based on vMINTEQ, predicted precipitation of several potential secondary minerals during AMD fly ash-bentonite interaction. These precipitates possibly controlled the composition of the effluent and reduced the hydraulic conductivity of the samples studied.

4. Comparative study of pre and post permeation of AFA and BAF samples using XRD and SEM conclusively showed the formation of new minerals and precipitates such as ettringite and gypsum, which were not detected on the original fly ash before permeation. Gypsum was the predominant mineral identified in post permeated solid samples by both XRD and SEM.
ACKNOWLEDGEMENTS
The work described in this paper was funded by the Natural Sciences and Engineering Research Council of Canada (NSERC) Corporate Research and Development Project Grant (No. CRDPJ 319808-04), Goldcorp Musselwhite Mine, Ontario Power Generation-Atikokan generating station and Golder Associates Ltd.

REFERENCES


Figure Caption List

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Figure 2. Variation of Sample Double Layer-thickness with Pore Volume of Permeant.

Figure 3. Effluent pH of Fly Ash and Fly ash-Bentonite mixture Samples Permeated with Water and AMD.

Figure 4. Effluent Sulphate Concentration of Fly Ash and Fly ash-Bentonite Mixture Samples Permeated with Water and AMD.

Figure 5. Leachate Concentrations of Heavy metals and Trace elements from Fly ash and Fly ash-Bentonite samples. (a) Al, (b) Fe, (c) Cu, (d) Mn, (e) Ni, (f) Zn, (g) Cr, (h) Cd, (i) Pb, (j) As, (k) Si and (l) Ca

Figure 6. Comparison of X-ray Powder Diffraction Pattern Only for Low Angles (Less than 45°).
   (a) AFA Before permeation; (b) AFA after Permeation.

Figure 7. Comparison of X-ray Powder Diffraction Pattern Only for Low Angles (Less than 45°).
   (a) 5% BAF Before permeation; (b) 5% BAF after Permeation.

Figure 8. Comparison of X-ray Powder Diffraction Pattern Only for Low Angles (Less than 45°).
   (a) 10% BAF Before permeation; (b) 10% BAF after Permeation.

Figure 9. SEM Images of Samples before and after AMD Treatment. (a) AFA before AMD Treatment Showing Smooth Surfaces of Fly Ash, (b) AFA after AMD Treatment Showing Cementation; (c), (d) and (e) SEM images of AFA, 5% BAF and 10% BAF respectively after AMD Treatment Showing the Formation of Gypsum Crystals.
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Fig 5

(a) Al

(b) Fe

(c) Cu

(d) Mn
Figure 5. Leachate Concentrations of Heavy metals and Trace elements from Fly ash and Fly ash-Bentonite samples. (a) Al, (b) Fe, (c) Cu, (d) Mn, (e) Ni, (f) Zn, (g) Cr, (h) Cd, (i) Pb, (j) As, (k) Si and (l) Ca

AMD conc. = Influent AMD concentration used as permeant (mg/L)
MOE Reg. 558 = Ontario Ministry of Environment Regulation 558/00 leachate criteria
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Table 1. Summary of Saturation Indices of Minerals as Predicted by vMINTEQ for 10% BAF.

<table>
<thead>
<tr>
<th>Mineral name</th>
<th>1.25</th>
<th>2.8</th>
<th>4.35</th>
<th>6.94</th>
<th>10.49</th>
<th>13.98</th>
<th>18.43</th>
<th>21.69</th>
<th>25.33</th>
<th>28</th>
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<tbody>
<tr>
<td>Cr(OH)$_2$</td>
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<td>1.22</td>
<td>0.87</td>
<td>1.42</td>
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<td>-2.57</td>
<td>-2.39</td>
<td>-2.55</td>
<td>-2.59</td>
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<tr>
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<td>0.06</td>
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<td>-0.02</td>
<td>-0.03</td>
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<td>16.88</td>
<td>17.23</td>
<td>16.79</td>
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<td>Ferricydrite</td>
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<td>3.55</td>
<td>3.59</td>
<td>3.71</td>
<td>3.69</td>
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<td>Gibbsite (c)</td>
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<td>0.69</td>
<td>0.44</td>
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<td>Gypsum</td>
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<td>0.07</td>
<td>0.03</td>
<td>0.07</td>
<td>0.06</td>
<td>0.07</td>
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<td>Hematite</td>
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<td>1.61</td>
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Table 2. Physico-Chemical Characteristics of AFA and BAF Samples before and after Permeation.

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<th>Property</th>
<th>FFA Before</th>
<th>FFA After</th>
<th>5% BAF Before</th>
<th>5% BAF After</th>
<th>10% BAF Before</th>
<th>10% BAF After</th>
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<tbody>
<tr>
<td>Major oxides (wt. %)</td>
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<td>SiO$_2$</td>
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<tr>
<td>Fe$_2$O$_3$</td>
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<td>3.83</td>
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<td>3.79</td>
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<td>3.75</td>
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<td>TiO$_2$</td>
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<td>0.80</td>
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<td>K$_2$O</td>
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<td>MgO</td>
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<td>2.93</td>
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<td>MnO</td>
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<td>Na$_2$O</td>
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<td>P$_2$O$_5$</td>
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<td>L.O.I (%)</td>
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<td>Total</td>
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Minor and Traces (mg/kg)

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<th>FFA 5% BAF Before</th>
<th>FFA 5% BAF After</th>
<th>10% BAF Before</th>
<th>10% BAF After</th>
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<td>Manganese</td>
<td>121.00</td>
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<td>Nickel</td>
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<td>595.56</td>
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<td>548.13</td>
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<td>Lead</td>
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<td>38.19</td>
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<td>Cobalt</td>
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<td>Chromium</td>
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