Impact of Portland Cement Content on Alkali Activated Bottom Ash

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Abstract. This study explores the behavior of blended mortars of low reactive bottom ashes (BA) and ordinary Portland cement (OPC) in an alkaline solution. Mortar mixtures incorporating OPC with different replacement levels (0% to 30% mass) were studied. Isothermal conduction calorimetric analysis was studied and 28-and 60-day mechanical strength values were found. SEM images was used to identifies porosity structures at 28-day. Water absorption was also investigated. The results revealed different behaviors to low (OPC 2.5 and OPC 5) and moderated (OPC 10 and OPC 30) OPC content. Increase in percentage of Portland accelerated hydration kinetics. There is a second peak formation for moderated OPC content, associated with C-A-S-H gel formation. The partial replacement of bottom ash by OPC tend to reduce the absorption. There is a slower initial water absorption to low OPC content. This behavior is due the higher unreacted BA content, that works as a filler. Otherwise, the increase of water absorption for OPC content is due to the coexistence of Portland cement hydrates and alkali activated reactions.

Keywords: Blended Mortars, Alkali Activation, Porosity, Bottom Ash.

1 Introduction

Bottom ash (BA) represents 40% of ashes from the coal-fired thermoelectric power plants (Cheriaf et al., 1999), it is estimated 4.5 million tons of BA per year in Brazil (IEA, 2016). Unlike fly ash, the disposal of bottom ashes goes to settling basin. Studies have shown the use of bottom ashes as an alternative aluminosilicate source for alkali activations (Topçu et al., 2014, Tambara Júnior et al., 2018). However, its performance is worse than fly ash. This is due the less amorphous content, higher unburned coal and low reactive SiO₂ and Al₂O₃.

An alternative to improving alkali activation BA properties is to increase surface area (through grinding) and to reduce unburnt material, through calcination (Sathonsaowaphak et al., 2009). Currently in the literature hybrid or blended cements are used to obtain better responses of the material (Garcia-Lodeiro et al., 2016).

Durability remains a property requiring further studies on alkali activations. A problem reported about alkali activation materials is the higher sorptivity when compared with Portland cement. Some works (Collins and Sanjayan, 2000) affirm that this is due the large proportion of fine pores which results in micro-cracks in the matrix, increasing the water capillarity. To reduce the water sorptivity it is necessary add some material that densifies the sample (Rostami and Behgarnia, 2017).

In this article was investigate the influence of ordinary Portland cement (OPC) on the mechanical strength, kinetic of water absorption and pore structure (scanning electron microscope) of blended alkali activation mortar of BA/OPC. The reaction was monitored with isothermal conduction calorimetry to evaluate the heat flow variations in the pastes in the early time of reaction. The OPC was blended from 0%, 2.5%, 5%, 10% and 30%.
2 Experimental Procedure

2.1 Materials Characterization

A Mastersizer 2000 laser granulometer was used to obtain particle size distribution of OPC and BA (Figure 1) and the powder was dispersed in isopropanol. It is seen that the BA used in this study contained a finer particle size than BA. Where 86% of OPC and 99.9% of BA particle size measured under 45μm, in agreement with the size particle parameter used to improve ash reactivity (Fernández-Jiménez e Palomo 2003).

The BA and OPC oxide analysis and clinker phase compositions are presented in Table 1. To the cement the phase was estimated based on Bogue calculation method. To BA was performed a Rietveld refinement method (Rietveld, 1969) using X-ray diffraction test data. To determine the amorphous and crystalline content in the BA sample, the analytical reagent corundum (Al₂O₃) was added in a total of 10% by mass as an internal standard. The OPC is a type III and the BA is classified as ash type F (low calcium content).

The sand used in the mortar mixtures was a natural fine sand. It was obtained a relative specific density of 2600 kg/m³, water absorption of 1.6%, maximum grain size of 1.2 mm and minimum grain size of 0.15mm. The sand showed a fineness modulus of 1.83.

Table 1. Chemical composition.

<table>
<thead>
<tr>
<th>Oxide analysis (%)</th>
<th>OPC</th>
<th>BA</th>
<th>Phase composition results</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>19.36</td>
<td>40.82</td>
<td>C3S 60.05</td>
</tr>
<tr>
<td>CaO</td>
<td>63.71</td>
<td>1.73</td>
<td>C2S 10.22</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>5.10</td>
<td>37.46</td>
<td>C3A 8.41</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>3.02</td>
<td>5.71</td>
<td>C4AF 9.19</td>
</tr>
<tr>
<td>SO₃</td>
<td>2.84</td>
<td>0.29</td>
<td>CS 4.83</td>
</tr>
<tr>
<td>Others</td>
<td>2.17</td>
<td>7.32</td>
<td>free lime 1.25</td>
</tr>
<tr>
<td>LOI</td>
<td>3.80</td>
<td>0.07</td>
<td></td>
</tr>
</tbody>
</table>

*Loss of ignition

A combination of sodium hydroxide (NaOH) pellets, water distilled and sodium silicate solution (Na₂SiO₃) was used as an alkaline activator (SiO₂ = 26.5%; Na₂O = 10.6%; H₂O = 62.9%; and density of 1.39 g/cm³). Initially a 14M NaOH solution was prepared with distilled water, waiting up to room temperature. After that a ratio of 1:2 Na₂SiO₃ to NaOH solution were prepared. A superplasticizer additive based on polycarboxylate ether, with solids content of 0.22, was used to adjust the workability (flow value > 250 mm) required for self-leveling mortars.

To evaluate the effect of OPC content on the microstructure of blended alkali-activated mortars, it was prepared 5 mixtures. The sum of OPC and BA was considerate as binder. Hydrated mortars were prepared using the same liquid-to-binder (l/b) ratio of 0.55. The substitution in mass of BA for Portland cement was: 0% (i.e., only bottom ash), 2.5%, 5%, 10% and 30%. The binder and sand ratio were kept constant from 1:2 (in mass). The molar ratios were between 2.6-3.1 to Si/Al, 0.27-0.30 to Na₂O/SiO₂ and equal to 12.3 to H₂O/Na₂O.
2.2 Experimental Procedure

The mixing method consisted of an initial premix of the solutions (NaOH and Na₂SiO₃) until room temperature. After that, a dry mix powder of cement, bottom ash and sand of each proportion was executed. The solid/liquid mixing was performed in a planetary mixer, the first 2 minutes at 150 rpm and one last minute at 300 rpm. Then the mortars were added to the molds and cured for 24h at 80°C.

Compressive strength was performed in cubic molds (4cm x 4cm), according to Brazilian Standard NBR 13279. To each mixture 6 samples were tested at 28 and 60 days of reaction. Calorimetric tests were performing to determine the heat evolution and the total heat through a conduction isotherm calorimeter TAM Air-Thermometric. The pastes were manually mixed for 3 min with the same w/b ratio than mortars and then sealed. The samples were immediately placed in the calorimeter, under the same conditions as the mortars (80°C ± 1°C).

To stop the reaction at the test ages for microstructural analysis, the samples were kept in an oven at 40°C until constancy of mass. Water absorption was evaluated over time for 24 hours through a method of Mariotte’s bottle developed of laboratory ValoRes-UFSC. To analyze the microstructure images, the samples were carbon-coated and mounted on electron microscope slides for study under a JEOL JSM-6390SL (15.0kV) with energy-dispersive X-ray spectroscopy (EDS) performed at Central Laboratory of Electron Microscopy (LCME-UFSC).

3 Results and Discussion

3.1 Isothermal Calorimetry and Compressive Strength

The effect of OPC content on the rate of heat release and total heat release during alkali activation of the BA is shown in Figure 2. For all the samples is seen a faster reaction at 80°C, due to this it was not possible to detect the first peak associated aluminosilicate dissolution, as observed in literature (Palomo et al., 1999; Garcia-Lodeiro et al., 2013). For the alkali
activation only with BA (OPC0) it is seen the lowest heat release and total heat. This is due its insufficient dissolution phases that start the formation of N-A-S-H gel.

Two different behaviors were observed according to the OPC content on the hybrid cements. Results are presented as low OPC content (OPC2.5 and OPC5) and moderate OPC content (OPC10 and OPC30). Even in small quantities, increasing OPC content accelerates the activation reaction. To low OPC content (Figure 2a) it is noticed the formation of a single peak in heat evolution. The increment of OPC content resulted in a higher heat release and total heat, this indicates an increase of reaction degree. OPC2.5 and OPC5 showed a total heat at 6 h of 59.79 J/g and 77.73 J/g, respectively.

Figure 2b shows heat flow curve and total heat for the alkali activation with moderate OPC content. It is observed a more intense and rapid first peak compared with low OPC content, followed by a short dormant period (occurs up to 15 minutes of activation). OPC10 and OPC30 presented a total heat at 6 h of 148.48 J/g and 126.76 J/g, respectively.

The quick first peak formation may be associated with an accelerated Si and Al dissolution and N-A-S-H formation. Also, a secondary peak starts after 15 minutes of activation. After the first gel formation the pH of pore solution decreases by which the calcium provided from OPC begins to react and form a (N,C)-A-S-H gel type (García-Lodeiro et al., 2016).

OPC30 showed the highest total heat up to 1h of activation, however, OPC10 overcame the total heat in higher period. Martinez-Ramirez and Palomo (2001) showed that OPC hydration is retarded when alkali concentrations increases. This suggests that is greater amount of inner anhydrous cement to OPC30 than OPC10.

Compressive strength was evaluated to later ages of 28 d and 60 days (Figure 3). OPC0 showed no strength development over the time of 60 days. This result shows that the exclusive alkali activation of BA results in a very slow evolution of reaction. The gain of compressive strength from 28 d to 60 d of OPC2.5, OPC5 correspond to 8.8%, 12.8% to low OPC content, a result in agreement with heat flow calorimetric analysis. To moderate OPC content the compressive strength for 28 d and 60 d increases 19.9% and 11.6% to OPC10 and OPC30, respectively. There is observed an optimum OPC content of 10% to blended mortar, increasing the bottom ash reaction.
3.2 Water Absorption

Water transportation of mortars is an important property that mainly influences its durability. Figure 4 shows water absorption over a 24 h of experiment to low and moderate OPC mortars at 1 d and 28 d. The water absorption of the blended mortars reported in this work were obtained in the final rage of 2.27%-10.01% and 1.57%-4.30% for 1 d and 28 d, respectively. At 1 d of hydration, it is observed higher absorption of activation with increased OPC dosage.

At 28 d was observed to all samples a reduction of water absorption. Compared with 1 d of hydration, OPC30, OPC10, OPC5 and OPC2.5 presented a water absorption reduction of 57.04%, 81.17%, 68.59% and 30.84%, respectively. Water absorption reduction is greater for moderate OPC mortars. It was observed that OPC10 showed the lowest water absorption to 28 days. The authors associate this behavior to the pore filling due best compatibility of last formation of (N,C)-A-S-H gels observed through microscopy analysis (Tambara Júnior et al., 2018).

![Figure 3. Compressive strength to 28 d and 60 d.](image)

![Figure 4. Water absorption of low OPC moderate OPC content at 1 day (a) and 28 days (b) of reaction.](image)
3.3 SEM

As Figure 5a shows, the bottom ash after treatment presented predominance of irregular particles with few spherical particles, which difficult the activation. Figure 5b shows the OPC0 at 28 days of activation, where there is a larger presence of unreacted BA and insufficient gel formation, featuring in a higher porosity and low compressive strength (Figure 3).

![SEM images](image)

As visual criteria, analysis of pores size was evaluated through an image analysis software tool of SEM images. Figure 6 shows micrographs of low and moderated OPC samples at 28d. To OPC2.5 is the sample with the largest pore presence. It is observed a great presence of macropores 50 µm - 400 µm. To OPC5 was observed a reduction in the content of smaller macropores, with no significant difference to pores larger than 300 µm.

There is observed higher pore formation and unreacted bottom ash to low OPC content mortars. As calorimetry analysis shows, the improve of OPC content results in a higher energy release, also leading to denser N-A-S-H/C-A-S-H gels. Moderate OPC mortars resulted in a reduced presence of the larger porous size. OPC30 and OPC10 presented similar pore size range of 20 µm – 140 µm. Otherwise, it was observed the lower water absorption to the OPC10, that can be associated to a higher microporosity to OPC30 mortar.
4 Conclusions

This study evaluated the OPC content on the blended mortars of bottom ash and Portland cement. It is concluded that there is different behavior between low to moderate OPC replacement, but even in low content (2.5% and 5%) there is a good contribution to activate bottom ashes with low reactivity.

To all the samples, as the OPC replacement increased the mechanical strength and the water absorption increases. This improved performance may be due to a denser microstructure generated to C-A-S-H gel formation observed through microstructural and calorimetry analysis that showed a secondary peak to moderate OPC content.

Furthermore, the 10% substitution of OPC significantly improved the parameters of durability, associated with the higher strength development at later ages and lower water absorption, due to its suitable Ca\(^+\) content provided by OPC. The increase of compressive strength and water absorption of OPC30 may be associated with a reduction of macroporosity but higher microporosity.

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References