Non-structural Performance of Pastes and Mortars Fabricated from Alkali Activated Electric Arc Furnace Slag

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Abstract

With the implementation of legislation on waste management in the European Union, which sets objectives in terms of promoting the re-use and recycling of products, the industry and the academic community have been making additional efforts in order to develop solutions that aim to use industrial by-products as raw materials, more specifically, in the study of alternative binders to Portland cement. In this work, an extensive experimental campaign is carried out to characterise, from the non-structural point of view, a set of alkali activated mixtures fabricated with electric arc furnace slag (EAFS) and phosphate sludge (PS) as the precursor, and two different activators (a commercial solution composed by sodium hydroxide and sodium silicate, and a solution used to clean the sand moulds from the aluminium processing industry). Results reveal that the mixtures based on EAFS present some handicaps in terms of shrinkage and workability, and that these properties are influenced by the nature of the activator and the presence of the PS.

Keywords: Electric arc furnace slag, Phosphate sludge, Alkali activated mortars, Non-structural behaviour

List of Abbreviations:

CS – solution used to clean the sand moulds from the Portuguese aluminium processing industry; EAFS – electric arc furnace slag; FS – flexural strength tests; HS – solution containing commercial sodium hydroxide and sodium silicate; LVDT – linear variable differential transformer; PS – phosphate sludge; PSD – particle size distribution; UCS – uniaxial compressive strength tests

List of Notations

- $C_m$ is the coefficient of water absorption for mortars other than renovation mortars
- $C_r$ is the coefficient of water absorption for renovation mortars
- $f$ is the factor of molarity
- $m$ is the mass of the sample
- $M_0$ is the dry mass of the specimen
- $M_1$ is the mass of the specimen after soaking for 10 minutes
- $M_2$ is the mass of the specimen after soaking for 90 minutes
- $M_3$ is the mass of the specimen after soaking for 24 hours
- $m_d$ is the dry weight of the specimen
- $m_h$ is the immersed weight of the specimen
- $m_s$ is the saturated weight of the specimen
- $p_0$ is the open porosity of the specimen
Introduction

Today, more than ever, the protection of the environment is a priority issue and the industry is being compelled to adopt proper and responsible behaviours, which includes the exploration, until the exhaustion of all possibilities, of using industrial by-products that show some potential as raw materials for specific applications. Within the framework of the global vector known as waste valorisation, the construction industry may play a key role regarding the aforementioned outcome, by promoting the development of more environmentally friendly products. Furthermore, the timescale until these solutions are implemented in the market depends, largely, on the reliability of the results presented by the academic community. Electric arc furnace slag (EAFS), which is an industrial by-product of the steel-making process, has been the subject of extensive research aiming at the application of this material in the construction industry, usually in the role of the aggregate [1-4]. Concerning its use as an aggregate (i.e. in substitution of natural, more traditional aggregates, for the production of concrete), several studies demonstrated that the mechanical performance of such concretes is suitable, when compared with ordinary concretes [5-7]. However, in order to limit the increase in density, partial replacements only should be considered [8]. Fewer studies have focused on the use of EAFS as a part of the binder, mostly based on its essentially crystalline nature and on its relatively high iron content [9]. In short, although the EAFS is preferentially used as an aggregate, some interesting results were attained, in terms of mechanical performance, when the EAFS, in powder form, was used in cementitious blends, usually incorporating Portland cement [10-12].

The procedure known as alkaline activation is being developed, in the last two decades, to produce binders from either industrial by-products or natural materials (e.g. kaolin). There are numerous materials that have been used in a precursor role, like fly ash (usually with a low calcium content) or blast furnace slag (with a high content of calcium, such as the EAFS) [13-16]. The earlier development of appreciable mechanical strengths is one of the most meaningful features of the alkaline cements based on blast furnace slag and other high-calcium precursors. However, these high-calcium AA binders face some important setbacks, such as shrinkage or rapid hardening [17]. As a valuable source of aluminium, silicon and calcium, EAFS is an industrial by-product that, apparently, meets several of the required conditions to be used as a precursor in alkali activated blends, and current research is now targeting its functional optimization. Even though the volume of research-based knowledge regarding alkali activated materials is constantly growing [18,19], information on its non-structural behaviour is clearly less than that available concerning its structural behaviour, which acts as a hindrance when potential industrial partners need confidence and security to move forward. By analyzing a set of non-structural properties, this work aims to be a contribution to a more consolidated knowledge about this kind of alkali activated pastes and mortars and, therefore, to assist the decision-making regarding the applications that can be developed. The research presented was preceded by a preliminary campaign, which tested the mechanical performance (through uniaxial compressive strength tests) of a wide spectrum of alkali activated pastes, composed by electric arc furnace slag (EAFS) and phosphate sludge (PS). Based on the results of this preliminary phase, which are not presented, the two most performing combinations were selected and used in the fabrication of mortars. Silica sand was used, in weight percentages of 10%, 20% and 30%, by weight of the paste, and the resulting mortars were submitted to flexural and compressive strength tests. Four mixtures (two pastes and two mortars) were then submitted to an extensive experimental campaign that included the assessment of the following properties: bulk density, porosity, shrinkage, water absorption, water-soluble chloride content and several fluidity related tests. Considering the lack of standards for alkali activated materials, tests were carried out based on the recommendations of the existing standards for cement and mortars.

Materials and Methods

Materials

The two residues used as precursors, EAFS and PS, are generated in the metallurgical and metal-mechanic industry, respectively, and were provided by Portuguese companies. They were both presented as a white powder, and their particle size distribution (PSD) curves reveal a maximum particle size of 250 μm. The PSD curve of the river sand, used to fabricate the mortars, was rectified to comply with the maximum grain size defined by EN 196-1:2016 [20]. The PSD curves of the residues are shown in Figure 1 (Figure 1).

The chemical composition of the precursors is shown in Table 1. The EAFS revealed a significant calcium content (49%) and a low aluminium content (7.4%), while the PS is mostly constituted by phosphorus (44%) and iron (35%). Even if the chemical composition of the EAFS revealed some potential for the precursor...
role in alkali activation reactions, it was decided to incorporate a secondary additional residue, namely the PS, which is not an aluminosilicate, in order to assess the possible recycling of the high phosphorous content of the PS in alkaline activation-based materials [21-23] (Table 1).

Table 1: Chemical composition (wt. %) of the EAFS and PS obtained through XRF analysis.

<table>
<thead>
<tr>
<th>Element</th>
<th>EAFS</th>
<th>PS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na₂O</td>
<td>0.13</td>
<td>3.6</td>
</tr>
<tr>
<td>SiO₂</td>
<td>18.8</td>
<td>-</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>7.4</td>
<td>-</td>
</tr>
<tr>
<td>MgO</td>
<td>5.4</td>
<td>-</td>
</tr>
<tr>
<td>K₂O</td>
<td>0.03</td>
<td>-</td>
</tr>
<tr>
<td>CaO</td>
<td>49.5</td>
<td>1</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.4</td>
<td>-</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>10.1</td>
<td>35.2</td>
</tr>
<tr>
<td>ZnO</td>
<td>1.25</td>
<td>12.9</td>
</tr>
<tr>
<td>MnO</td>
<td>1.43</td>
<td>-</td>
</tr>
<tr>
<td>BaO</td>
<td>0.1</td>
<td>-</td>
</tr>
<tr>
<td>SO₃</td>
<td>3.42</td>
<td>-</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>0.07</td>
<td>43.7</td>
</tr>
<tr>
<td>Others</td>
<td>2</td>
<td>3.6</td>
</tr>
</tbody>
</table>

Based on UNE 80103:2013 [24], specific gravity values of 3.11 g/cm³ and 2.66 g/cm³ were obtained for the EAFS and PS, respectively, using a Le Chatelier volumenometer. Mineralogy was analyzed with a BRUKER D8 ADVANCE diffractometer, with CuKα radiation, 40kV and 30mA, fitted with a LYNXEYE super speed detector. The resulting diffractograms of the original materials are presented in Figure 2. The scans covered a 2θ range of 5 to 60⁰, with a nominal step size of 0.01973⁰ and 0.5 s/step. The X-Ray diffractogram of the EAFS showed calcium silicate, gehlenite, and calcium-magnesium-iron as the main phases, with some traces of magnetite and larnite. The halo between the 30 and 40 (°2θ) angles reveals the presence of some vitreous mass. Mineralogically, the PS identified is essentially strength. The presence of amorphous material in the PS was also detected, in several ‘bumps’ throughout the baseline (Figure 2).

Two different activators were used: a solution containing commercial sodium hydroxide and sodium silicate (HS), with a hydroxide / silicate ratio of 2; and a solution used to clean the sand moulds from the Portuguese aluminium processing industry (CS). The sodium hydroxide, supplied in pellets with a specific gravity of 2.13 at 20°C (99 wt.%), was dissolved in deionized water to achieve a concentration of 5M. The silicate presented a unit weight of 1.464 g/cm³ (at 20°C), a SiO₂/Na₂O weight ratio of 2.0 (molar oxide ratio of 2.063) and a Na₂O concentration in the solution of 13.0%. The CS was used as received in the laboratory, after homogenization, and a concentration of approximately 7M was determined.

Testing procedures and specimen production

Fabrication of the specimens comprised an initial mixing stage, performed on a counter mixer, followed by a 30-second vibration...
phase, on a vibrating table. The pastes were then casted into prismatic moulds (4x4x16 cm), which were left to cure for 20 hours, at 80°C. Tests were conducted 24 hours after the start of the mixing stage. The composition of the pastes is shown in Table 2. Due to the nature of the residues and activators, it was not possible to set the same activator / solids ratio for all mixtures. These ratios were adjusted considering the workability exhibited by the mixtures. Two main groups were defined, based on the activator (HS or CS). For each group, one paste and three mortars were studied, each with a different aggregate (sand) content of 10%, 20% and 30%, by dry weight. Also included is the activator / solids ratio for each mixture (the solids were defined as the sum of the precursors and the aggregate) (Table 2).

Table 2: Composition of the mixtures (Ac: activator; Sol: precursor + aggregate).

<table>
<thead>
<tr>
<th>Label</th>
<th>EAFS (wt.%)</th>
<th>PS (wt.%)</th>
<th>SA (wt.%)</th>
<th>Ac / Sol</th>
</tr>
</thead>
<tbody>
<tr>
<td>BHS</td>
<td>100</td>
<td>-</td>
<td>-</td>
<td>0.39</td>
</tr>
<tr>
<td>MHS1</td>
<td>90</td>
<td>-</td>
<td>10</td>
<td>0.35</td>
</tr>
<tr>
<td>MHS2</td>
<td>80</td>
<td>-</td>
<td>20</td>
<td>0.35</td>
</tr>
<tr>
<td>MHS3</td>
<td>70</td>
<td>-</td>
<td>30</td>
<td>0.39</td>
</tr>
<tr>
<td>BCS</td>
<td>50</td>
<td>50</td>
<td>-</td>
<td>0.5</td>
</tr>
<tr>
<td>MCS1</td>
<td>45</td>
<td>45</td>
<td>10</td>
<td>0.39</td>
</tr>
<tr>
<td>MCS2</td>
<td>40</td>
<td>40</td>
<td>20</td>
<td>0.38</td>
</tr>
<tr>
<td>MCS3</td>
<td>35</td>
<td>35</td>
<td>30</td>
<td>0.32</td>
</tr>
</tbody>
</table>

Uniaxial compressive (UCS) and flexural (FS) strength tests were conducted based on the recommendations of EN 196-1:2016 [20]. Compressive strength tests were carried out on the two halves resulting from the prismatic specimen which was initially submitted to a flexural test. A servo-hydraulic testing machine, fitted with either a 10 kN or a 50 kN load cell, depending on the flexural or compressive test being performed, respectively. The tests were carried out under monotonic displacement control, at rates of 0.4 mm/min (UCS) and 0.2 mm/min (FS). The relative displacement between the loading plates in the compressive tests was acquired by a linear variable differential transformer (LVDT), enabling the registration of the full stress-strain curve for each test. Prior to testing, all specimens were weighted and measured. The strength values presented in this work represent the arithmetic average of three (flexural tests) and six (compressive tests) specimens. The determination of the bulk density of fresh mixtures was carried out in accordance with EN 1015-6:1998 [25]. A 1-liter capacity metallic bowl was filled, in excess, with the paste, and vibrated (on a vibrating table), until no further setting was detected. The remaining excess paste was removed with a pallet knife, after which both the mass and the volume were registered (Figure 3). The bulk density of the hardened mixtures was assessed during the porosity test, as described further ahead (Figure 3).

Considering the lack of a specific standard to evaluate the porosity of this type of mixtures, the tests were carried out based on the procedures displayed in NP EN 1936:2008 [26]. For each mixture, three cubic specimens (4 cm) were prepared and dried, in an oven, at 70°C, until constant mass was reached. Test procedures included three main steps, carried out in three consecutive days: i) installation of the specimens in a desiccator (Figure 4a), connected to a vacuum pump, which produced a -26 bar pressure for 24 hours; ii) the container, still under vacuum, was filled with water until the specimens were submerged; iii) after 24 hours, the vacuum was eliminated, and the specimens remained submerged for further 24 hours. Throughout the test, the dry, saturated and submerged (hydrostatic weighing, Figure 4b) weights were registered. The porosity, as well as the bulk density, was then estimated based on the equations presented in NP EN 1936:2008 [26] (Figure 4).
Drying shrinkage tests were carried out based on the procedures established in BS EN 1015-18:2002 [28]. For each mixture, three prismatic specimens, with dimensions 16x4x4 cm, were prepared for each mixture, and submitted to a total curing period of 28 days. In the first two days the specimens remained in the moulds, under a temperature of 20°C. After demoulded, the curing conditions were maintained for further 5 days. For the remaining 21 days, specimens were submitted to a relative humidity of 65%, while still at 20°C. After this 28-day curing period the specimens were sealed, with aluminium tape, broken in two halves, and placed in an oven at 60°C, until constant mass was reached. The next step was to install the specimens in perforated steel supports, with the broken surface facing down, which were then placed inside trays with a 10 mm thick water layer (Figure 6). After 10 and 90 minutes, the specimens were removed from the trays and weighed, and restored back to the tray. The weighing was then repeated after 24 hours, and the specimens were, once again, broken in two halves, in order to measure the height of water penetration(Figure 6).

The water absorption coefficient of the hardened mixtures, due to capillary, was assessed using the procedure described in EN 1015-18:2002 [29]. Three prismatic specimens, with dimensions 16x4x4 cm, were prepared for each mixture, and submitted to a total curing period of 28 days. In the first two days the specimens remained in the moulds, under a temperature of 20°C. After demoulded, the curing conditions were maintained for further 5 days. For the remaining 21 days, specimens were submitted to a relative humidity of 65%, while still at 20°C. After this 28-day curing period the specimens were sealed, with aluminium tape, broken in two halves, and placed in an oven at 60°C, until constant mass was attained. After drying, the specimens were crushed in order to obtain 50 g of particles below 125 µm, from which 10 g were collected and thoroughly mixed with 100 ml of distilled water. The mixture was then left for 24 hours, before filtration and subsequent titration, using ammonium thiocyanate as the titrant. Several tests were also performed to analyze the workability and fluidity of the pastes and mortars. The standard consistency and setting time were evaluated in accordance with EN 196-3:2016 [30], using a Vicat apparatus. The consistency test consists of releasing a plunger into a mixture sample casted in a Vicat mould. Different mixtures, with different water contents, should then be tested, until a distance of 6±2 mm between the plunger and the baseplate is obtained, so that the corresponding consistency can be defined as ‘normal’. However, the role of this test in the framework of the present study was the comparison of the relative performance between the four selected mixtures and, thus, no additional water contents were tested. The initial setting time is measured between the ‘zero’ instance (when the needle punctures the full height of the specimen) and the instance when a distance of 6±3 mm is registered between the tip of the needle and the baseplate. The final setting time is then established when the needle, now with a 5 mm diameter ring attached, penetrates only 0.5 mm into the sample.

The fluidity and/or the wetness of the fresh mixtures were assessed using the flow cone and the flow table tests, based on the recommendations of NP EN 445:2008 [31] and EN 1015-3:1999 [32], respectively. The flow cone test consists on measuring the time that 1 litre of mixture requires to flow through a standard cone (Figure 7a). However, it was not possible to perform this test with any of the mixtures since none flowed through the cone in useful time. Nevertheless, it should be noted that the mixture showing the largest quantity of material flowing through the cone was the BCS, while no passing volume was observed for either of the HS-based mixtures. The flow table test, for assessing the consistency of the fresh state materials (Figure 7b), is started by introducing a sample in a truncated conical mould, in two compacted layers. After careful cleaning the visible disk surface, the mould is slowly lifted and the sample is spread by jolting the flow table 15 times (1/s). Finally, the diameter of the spread sample is measured, in two directions(Figure 7).

**Figure 6: Water absorption test: specimens broken in two halves and submerged in 10 mm thick water layers.**

**Figure 7: Fluidity analyses: a) set up of flow cone test; b) flow table test.**

**Result and Discussion**

**Flexural and compressive strength tests**

Figure 8 displays the flexural and compressive strength results. The most remarkable aspect resulting from these tests is that, as expected, the mechanical performance of the mixtures improved...
with the addition of the aggregate, with an optimum content between 10% and 20%. In terms of flexural strength (Rf), an aggregate content of 20% was the most effective for the HS binder (MHS2, Rf = 1.27 MPa), while a 30% aggregate content produced the best performance among the CS binder mixtures (MCS3, Rf = 1.53 MPa). In terms of compressive strength (Rc), a lower optimum aggregate content (10%) was obtained, for both types of activator. For the HS activator, the compressive strength of the MHS1 (10%, Rc = 6.58 MPa) and MHS2 (20%, Rc = 6.80 MPa) mortars was very similar. These values represent a strength increase of 52%, relatively to the strength obtained by the BHS paste. Considering the mixtures manufactured with the CS activator, the optimum aggregate content was 10% (MCS1), with a compressive strength of 10.20 MPa, which represents an increase of 34%, relatively to the value obtained by the corresponding paste (BCS). Mortars MHS1 and MCS1 were selected to the next phase of the experimental campaign, based on the compressive strength results. The former was selected even if it showed a slightly lower strength (0.22 MPa) than the mortar MHS2, because it represents a lower aggregate content and also because it allows a direct comparison with the CS-based mortar, also with a 10% aggregate content (Figure 8).

![Figure 8: Flexural and compressive strength results on HS-based mixtures and CS-based mixtures, after 20 hours curing at 80°C.](image)

**Bulk density of fresh mixtures**

Figure 9 presents the bulk density of the four mixtures selected for this second phase. The HS-based mixtures showed higher bulk densities than the CS-based mixtures, which can be justified by both the higher specific gravity of the EAFS, relatively to the PS, and the lower activator / solids weight ratios that were used to prepare the HS mixtures. The addition of the aggregate did not represent a significant effect in the bulk density, with increase values of 0.5% and 3.5%, for the HS and CS mixtures, respectively (Figure 9).

![Figure 9: Bulk density of the mixtures](image)

**Porosity**

Figure 10 shows the open porosity ($p_0$) and the bulk density ($\rho_b$) of the hardened mixtures, which were determined with the following equations:

$$p_0 = \frac{m_s - m_i}{m_s - m_d} \times 100 \quad (1)$$

$$\rho_b = \frac{m_d - m_s}{m_s - m_i} \times \rho_{rh} \quad (2)$$

where $\rho_{rh}$ is the water density (980 kg/m$^3$), and $m_d$, $m_s$ and $m_i$ represent the dry, saturated and immersed weight.

Mixtures prepared with CS were more porous, which can be related with the higher particle size of the PS, compared with the EAFS particles, and with the higher amount of activator (CS) that was used in the preparation of these mixtures. The addition of sand leads to a decrease in porosity, for mixtures prepared with HS and CS. Therefore, the durability of the mixtures might be improved when sand is added, since the amount of fluids that might propagate through the alkali activated composites will be lower. As expected, the bulk density values of the hardened pastes follow the trends observed for the bulk density of the fresh mixtures, i.e. the HS-based mixtures presented a higher density than the CS-based mixtures. It should also be mentioned that the differences between the bulk density of the hardened mixtures prepared with the same activator are more pronounced than those observed for the bulk density of fresh mixtures (Figure 10).
Shrinkage

Shrinkage values of mixtures MHS1, BCS and MCS1 are shown in Figure 11. Moments before demoulding, some cracks were observed in the BHS specimens, which made them unsuitable for testing. Shrinkage was more significant on mixture MHS1, meaning that the addition of an aggregate only slightly improved the behaviour of the BHS paste. Another important feature regarding the MHS1 mixture is that shrinkage increased with time and, after 56 days, values of more than 10000 microstrains were registered, which is significant, compared with the values of 1750 to 2250 microstrains obtained by Ye et al. [33] and Jia et al. [34], when studying alkali activated ground granulated blast furnace slag mortars. The mixtures containing phosphate sludge showed significant lower shrinkage values after 14 days, of 2600 and 3400 microstrains, for mixtures BCS and MCS1, respectively. These values were then maintained until the 28-day mark, which further reinforces the idea that the use of PS improved the shrinkage behaviour. The CS-based results, after 56 days, should be regarded with some reservation, since, at that stage, both the top and bottom of these specimens displayed significant damages, which influenced the measurements that were carried out on the digital length comparator (Figure 11).

Water absorption
The coefficients of water absorption are presented in Figure 12. The standard used in this work introduces two methods to determine these coefficients, either for mortars other than renovation mortars \(C_m\) and for renovation mortars \(C_r\), which are given by:

\[
C_m = 0.1 \times (M2 - M1) \quad (3)
\]

\[
C_r = 0.625 \times (M3 - M0) \quad (4)
\]

where \(M1\) and \(M2\) are the masses of the specimens after soaking for 10 and 90 min, respectively, whereas \(M0\) and \(M3\) are, respectively, the dry mass and the mass of the specimens after soaking for 24 hours (Figure 12).

Considering the first case, HS-based mixtures exhibited a lower \(C_m\) than the CS-based mixtures. The inclusion of an aggregate did not produce a significant effect on the water absorption of the mixtures. As water can be a mean of transporting for aggressive agents, it is acceptable to state that mixtures BHS and MHS1 presented a better behaviour regarding this property. Regarding the method for renovation mortars, all mixtures showed a similar behaviour in terms of the coefficient of water absorption. However, after 24 hours, all the water was absorbed, thus not allowing for a proper observation of the height of water penetration in the specimens’ surface. It should also be mentioned that at the end of that period, both HS-based specimens released vapors with an acute odor, after being removed from the tray. Additionally, the aluminium tape that was used to seal the specimens was considerably damaged (Figure 13).

### Water-soluble chloride content

The results of the tests carried out to determine the water-soluble content of the mixtures are depicted in Table 3 and Figure 14. The percentage chloride was computed through the following expression:

\[
\% Cl = \frac{(V2 - V1) \times f \times 3.545}{10 \times m} \quad (5)
\]

where \(V1\) and \(V2\) are, respectively, the sample and the blank titres of 0.1 M ammonium thiocyanate solution (which are shown in Table 3), and \(m\) is the mass of the sample (10 g were weighed for each mixture). The factor of molarity, \(f\), which in the present work was 10.7, is given by:

\[
f = \frac{ml of AgNO_3 (0.1mol/l)}{ml of ammonium thiocyanate} \quad (6)
\]

Mixtures BHS and MHS1 showed an appreciable behaviour regarding this particular test: the chloride contents were considerably low and significantly lower than those achieved by the mixtures BCS and MCS1, suggesting that the presence of the phosphate sludge had a significant influence. Nevertheless, the chloride contents exhibited by mixtures BCS and MCS1 are still acceptable. Another important evidence is that the presence of sand leads to higher values of chloride contents (Table 3)(Figure 14).

### Workability and fluidity tests

**Standard and flow table test consistency:** The consistence is an important feature to understand the fluidity of the fresh mixtures. Consistency results, determined with a Vicat apparatus and with the flow table, can be observed in Figure 15. Regarding the standard consistency (Vicat apparatus), no significant differences were found between mixtures BHS and BCS, since, in both cases, the distance between plunger and baseplate was approximately 35 mm, very far from the required distance of 6 (±2 mm). It is interesting that considerably different mix designs, both in terms of the EAFS and PS contents and the Ac/Sol weight ratios, still produced similar and comparable results. Regarding the consistency behavior assessed by

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**Table 3:** Water-soluble chloride content test results.

<table>
<thead>
<tr>
<th>Label</th>
<th>(V_1) (ml)</th>
<th>(V_2) (ml)</th>
<th>% Cl</th>
</tr>
</thead>
<tbody>
<tr>
<td>BN5</td>
<td>4.8</td>
<td>5.1</td>
<td>0.00994</td>
</tr>
<tr>
<td>MN5.1</td>
<td>4.1</td>
<td>5.1</td>
<td>0.03313</td>
</tr>
<tr>
<td>BCS</td>
<td>1.75</td>
<td>5.1</td>
<td>0.11099</td>
</tr>
<tr>
<td>MCS1</td>
<td>0.2</td>
<td>5.1</td>
<td>0.16234</td>
</tr>
</tbody>
</table>

---

**Figure 13:** General view of a specimen soaked for 24 hours.

**Figure 14:** Water-soluble chloride content.
the flow table test mixtures, the CS-based mixtures showed higher flow values than the HS-based mixtures. Even though the hardening process is faster in the pastes prepared with slag and phosphate sludge, these mixtures were prepared with a higher liquid content. Therefore, CS mixtures showed a greatest flow value. Nevertheless, these mixtures attained a higher compressive strength than HS-based mixtures, which is related with the intrinsic properties of the materials and with the type of activator used. Finally, it is also worth noting that flow values decreased when the aggregate was added (Figure 15).

**Figure 15:** Consistency tests results by the Vicat apparatus (a) and the flow table (b).

Setting times: The initial and final setting times of the studied mixtures are presented in Figure 16. Mixtures that were prepared with slag and phosphate sludge have lower setting times, especially the final setting time, which is of significance since the final setting time is a sign that the mixture has started to harden. Mixtures BCS and MCS1 required less than 50 minutes to achieve such moment. Indeed, during the experimental campaign, casting the mixtures with phosphate sludge was a sensitive task, since the hardening process was significantly faster than with the slag-only mixtures. This strongly affects the workability of the mixtures and, hence, the possible applications for this type of materials might be restricted. On the other hand, the hardening process is slower in mixtures composed only by slag. These results allow to identify the phosphate sludge has a setting time retarding for alkali activated composites (Figure 16).

**Figure 16:** Setting times of the mixtures.

### Conclusion

The present work examines the non-structural performance of alkali activated mixtures incorporating electric arc furnace slag (EAFS) as the main precursor, as well as phosphate sludge (PS) as a potential additive, which were activated using either a commercial solution (HS) or a by-product from the aluminium moulding industry (CS). Different tests were carried out to scrutinize the following properties: bulk density, porosity, shrinkage, water absorption, water-soluble chloride content, workability and fluidity. One of the most relevant findings was that the alkali activated mixture composed exclusively by EAFS exhibited a significant shrinkage after a curing period of 56 days, which is a handicap that should not be disregarded at the moment of defining the possible applications that could be developed with this industrial by-product. Other relevant aspects are related with the workability: mixtures activated with CS, and having PS in its composition, were significantly more fluid and attained lower setting times than the remaining cases. The results of this experimental campaign suggest that non-structural applications can be designed with this kind of alkali-activated mixtures.

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Conflict of Interest

No conflict of interest.

References


