Conference Paper

The Effect of the Activator/Precursor Ratio on the Rheological Properties of the Alkali-activated Mining Waste Mud Paste

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Abstract

To determine the properties of paste, mortar or concrete, it is necessary to understand its rheological behaviour first. This study discusses the effect of the activator/precursor ratio on the rheological properties of the alkali-activated paste. The pastes consisted of a mix of 70 % of tungsten mining waste mud, 15% waste glass and 15% of metakaolin. This mix was activated by combining sodium hydroxide and sodium silicate. Five activator/precursor (a/p) ratios were studied: 0.37, 0.38, 0.39, 0.40 and 0.41. The result obtained shows that the rheology of the pastes is affected by the activator/precursor ratio. The rheological behaviour of the paste fits the Bingham model. The yield stress ($\tau_0$) and plastic viscosity ($\mu$) increase inversely with the activator/precursor ratio (e.g. a/p=0.37 gives $\tau_0=84.19$ and $\mu=0.4185$; a/p=0.41 gives $\tau_0=30.389$ and $\mu=0.2937$). The workability increases proportionally with the activator/precursor ratio (e.g. a/p=0.37 gives a slump=133 mm; a/p=0.41 gives a slump=158 mm). The compressive strength decreases when the activator/precursor ratio increases (e.g. at 28 days for a/p=0.37, the compressive strength was 19.6 MPa; for a/p=0.41, the compressive strength was 13 MPa). Finally, the ideal ratios were 0.38 and 0.39.

1. Introduction

Currently, the building industry is facing a great challenge: the protection of the environment and the usage of waste and recycled materials [1]. The production of Portland cement (PC) is responsible for the emission of between 6% to 8% of CO$_2$ greenhouse gases [2]. Besides, in Europe, there is a major concern about the quantity and diversity of hazardous solid waste production.

The increasing concern about the environmental consequences of waste disposal has led to new investigations on new utilization (reuse) possibilities. For example, large quantities of sludge and by-products, such as fly ash, are produced during the combustion of coal used for electricity generation [3]. Also, in Europe, mining and quarrying industry waste are very relevant. These activities produce about 55% of the total industrial wastes according to recent Eurostat data [4].
For a better understanding of the behaviour and performance of the alkali-activated (AA) materials, it is important to study their rheological properties for the manufacturing of AA-based products.

Rheology is the study of the fluidity and deformation of matter in the fresh or hardened state, and the emphasis on flow means that it is concerned with the relationship between stress, strain, rate of strain, and time [5].

In (PC), the rheological testing, using a viscometer to break down the interparticle forces formed by the hydration of cement grains, consists in exposing the pastes to shear stress with known shear rates.

Several different models describe the rheological behaviour of cement pastes. Generally, the cement paste fits the Bingham model[5] Eq. (1), as presented as follows. The Bingham model:

\[ \tau = \tau_0 + \mu \gamma \]  

(1)

Where
- \( \tau \): shear stress,
- \( \tau_0 \): yield stress,
- \( \mu \): plastic viscosity and
- \( \gamma \): shear rate.

Other models can be used to analyse the rheological behaviour of cement pastes, such as the Ostwald de Waele and the Herschel–Bulkley models equations. (2) and (3), respectively, as presented next (Tattersall GH, 1983).

The Ostwald de Waele model:

\[ \tau = k \gamma^n \]  

(2)

The Herschel-Bulkley model:

\[ \tau = \tau_0 + k \gamma^n \]  

(3)

Where
- \( k \): the consistency coefficient (Pa.s\(^n\)), and
- \( n \): the dimension less fluidity index.

Besides the studies on the rheology of cement pastes, there are a few studies on the rheology of alkali-activated materials. Such main research studies are presented in Table.1. Since the purpose of this paper is to contribute to the knowledge of this scientific field, a short review of the existing studies on the rheology of AA materials is presented next.
Table 1: Some research activities regarding the rheology of alkali-activated materials (2010 to 2016).

| Studies                                                                 | References |
|------------------------------------------------------------------------|------------|
| Rheological behavior of fly ash-based geopolymer concrete              | [7]        |
| Rheological behavior of alkali-activated Metakaolin during geopolymerization | [8]        |
| Rheology of geopolymer by DOE approach                                | [9]        |
| Rheological properties of alkaline-activated fly ash used in jet grouting applications | [10]       |
| The interrelationship between surface chemistry and rheology in alkali activated slag paste | [11]       |
| Rheological behavior of a fresh geopolymer based on Metakaolin: Effect of the introduction of calcium carbonate | [12]       |
| Rheology of alkali-activated slag pastes. Effect of the nature and concentration of the activating solution | [13]       |
| Observations on the rheological response of alkali activated fly ash suspensions: the role of activator type and concentration | [14]       |
| Alkali-activated slag cements using waste glass as alternative activators. Rheological behavior | [6]        |

Puertas et al [12] compared the rheological parameters of PC and AA materials, namely the concentration of alkali solutions. Torres-Carrasco et al. concluded that the rheological behaviour of alkali-activated slags pastes fitted the Herschel-Bulkley model when the activator was a commercial water glass solution or NaOH/Na$_2$CO$_3$ combined with waste glass [14].

This study aims to discuss the role of the a/p on the rheological parameters in fresh and hardened state and choose the ideal ratio, which is the one that has the best behaviour in a fresh state with good mechanical performance.

2. Materials and Methods

2.1. Materials

The materials used in this study to prepare the AA paste were Metakaolin (Mk), tungsten mining waste mud (TMWM) and waste glass (WG) as a precursor. The metakaolin was supplied by BASF with the reference name "MASTERLIFE Mk", mud waste was collected from the Panasqueira tungsten mine located in Covilhã, Portugal, and the waste glass was obtained by crushing glass bottles. The chemical composition (determined by SEM-EDX) is given in Table 2 and the physical properties (bulk density and Blaine finesses) are presented in Table 3.

The sodium hydroxide (SH), NaOH, and sodium silicate (SS), Na$_2$SiO$_3$, are the activators. The sodium hydroxide solution was prepared by dissolving sodium hydroxide
The AA paste was a mix of 70% (TMWM), 15% (WG) and 15% (Mk) mixed in the dry state for 5 min to assure the homogeneity of our precursor. The (NaOH) with a molarity =10M and (Na<sub>2</sub>SiO<sub>3</sub>) were mixed for 3 to 5 minutes to form our activator. The alkali-activator solution with a ratio (Na<sub>2</sub>SiO<sub>3</sub>)/ (NaOH) =4 was added to the precursor. The paste was stirred for 2.5 min at 200 rpm, followed by 2.5 min at 400 rpm.

### Table 2: Chemical composition of (Mk), (TMWM) and (WG) in %.

| Chemical composition | Mud waste | Glass | Metakaolin |
|----------------------|-----------|-------|------------|
| Na<sub>2</sub>O      | 0.85      | 12.52 | 0.32       |
| SiO<sub>2</sub>      | 46.67     | 68.13 | 52.28      |
| Al<sub>2</sub>O<sub>3</sub> | 17.01   | 2.80  | 42.99      |
| K<sub>2</sub>O      | 4.90      | 0.86  | 0.94       |
| CaO                  | 0.69      | 10.52 | /          |
| SO<sub>3</sub>      | 7.90      | 0.23  | /          |
| Fe<sub>2</sub>O<sub>3</sub> | 15.47   | 2.90  | 1.49       |
| MgO                  | 4.83      | 2.04  | 0.47       |

### Table 3: Physical properties of (Mk), (TMWM) and (WG).

| Physical properties | Mud waste | Glass | Metakaolin |
|---------------------|-----------|-------|------------|
| Bulk density (g/cm<sup>3</sup>) | 3.10  | 2.49  | 0.2683     |
| Blaine fineness (cm<sup>2</sup>/g) | 742   | 2665  | 4467       |

pellets (98% purity obtained from Fisher Scientific, Schwerte, Germany) in deionized water and allowed to cool before use. Sodium silicate was obtained from Solvay SA, Póvoa de Santa Iria, Portugal. Table 4 presents the chemical composition and density of activators by weight.

### Table 4: Chemical composition and density of (SH) and (SS) in %.

| Chemical composition | Sodium hydroxide | Sodium silicate |
|----------------------|------------------|----------------|
| Na<sub>2</sub>O      | 13.02            | 19.37          |
| SiO<sub>2</sub>      | 0.00             | 62.60          |
| Al<sub>2</sub>O<sub>3</sub> | 0.00   | 0.90           |
| H<sub>2</sub>O       | 43.27            | 142.32         |
| Density (g/cm<sup>3</sup>) | 1.0192  | 1.5725         |

### 2.2. Methods

#### 2.2.1. Synthesis of samples

The AA paste was a mix of 70% (TMWM), 15% (WG) and 15% (Mk) mixed in the dry state for 5 min to assure the homogeneity of our precursor. The (NaOH) with a molarity =10M and (Na<sub>2</sub>SiO<sub>3</sub>) were mixed for 3 to 5 minutes to form our activator.
rpm at a temperature of 20°C for different activator/precursor (a/p) ratios, namely 0.37, 0.38, 0.39, 0.40 and 0.41.

### 2.2.2. Determination of rheological parameters

For the rheological study, a Viskomat NT rheometer (Rheometer for mortar and paste, produced by Schleibinger Testing Systems) was used. This apparatus automatically measures a series of data points of torque (T) and speed (N).

Immediately after mixing the paste, it was transferred into the rheometer cup to start measuring the rheological parameters.

In the Viskomat NT rheometer, as the cylindrical sample container rotates (Fig.1), the paste flows through the blades of the impeller and exerts a torque which is measured by a transducer. For the testing conditions, the rotation speed of the container was set to vary with time as a ramp speed profile. In the ramp profile (Fig. 2), the rotation speed was adjusted to vary with time, increasing from an initial value of zero to 120 rpm for 5 minutes and then decreasing from 120 rpm to zero in other 5 minutes. This speed profile was used to determine plastic viscosity and yield stress-related coefficients (μ and τ₀, respectively).

### 2.2.3. Paste fluidity (mini slump)

The paste fluidity was determined with the mini-slump test [15] on the samples, 7, 16, 25, 34 and 52 min after mixing. The slump cone has an upper diameter of 19 mm, a lower diameter of 38.1 mm, and a height of 57.2 mm. The cone is positioned in the center of the plate. After pouring the paste into the cone without causing it to overflow, the upper part of the cone is tamped lightly to bleed off any entrapped air pockets, and the cone is then gently lifted. After the pastes were dropped 10 times on a flow table, the diameter was measured in three directions: the values used were the arithmetic means of these measurements. Fig. 3 presents a spread of the paste with a/p= 0.41.

### 2.2.4. Compressive strength

The compressive strength tests were performed on a 3000 KN electro-hydraulic mechanical testing machine, the “ADR Touch 3000 BS EN Compression Machine with Digital Readout and Self Centring Platens”, in accordance with EN 196-1. Compressive strength data was obtained from testing five specimens with 25 mm cubic in size.
3. Results and Discussion

The influence of the a/p ratio on the rheological behaviour of the AA paste is discussed in two parts in this section: fresh and hardened states.
3.1. Fresh state

3.1.1. Evaluation of torque and rotational speed

Fig. 4 presents the changes of torque versus rotational speed of the AA pastes for different a/p ratios. From the graph, it can be observed that all the pastes tested fit the Bingham model (Eq (1)). Also, it can be seen that the increase of torque along with the increase of rotational speed, for example in the mix 0.41, needs a smaller torque (around 97.43327 N.mm). In opposition, mix (a/p) =0.37 needs the highest torque (170.53838 N.mm). These results are explained by the following: when the paste is very liquid, the stress to break the interparticle forces is smaller when compared to a higher viscous paste.

The results obtained in our study are comparable to the results obtained by other authors using slag-based AA pastes [12]. However, their study was based on changing the activator concentration while in this study this parameter was kept constant and the a/p ratios varied from 0.37 to 0.41.

3.1.2. Evaluation of yield stress

The relative yield stress was graphically determined from the curve of torque versus rotational speed. Fig. 5 shows such results. It is verified that the relative yield stresses of the mixes increases with time which is explained by the formation of C-S-H gel (the paste is more hardened; for example, in the first 7 min, the value of yield stress in the paste with the 0.37 a/p ratio was 84.19 N.mm and became 98.368 N.mm in 52 min). In opposition, it was observed that the increase of the a/p ratios lets the relative yield stress decrease which is explained with the decrease of the interparticle forces between
grains of paste (for the ratio a/p= 0.37, the yield stress was 98.368 N.mm in 52 min, but in the 0.41 a/p was 40.084 N.mm in the same time).

3.1.3. Evaluation of relative plastic viscosity

In the presented models, many factors affect the viscosity. The solid content and the interparticle forces between particles are one of these factors. The relative viscosity can be described by the empirical expression, proposed by [16], as follows:

\[ \mu = \mu_0 (1 - (\Phi/\Phi_{\text{max}}))^n \]  

(4)

Where \( \mu_0 \) is the solution viscosity, \( \Phi \) is the actual solid volume fraction \( \Phi_{\text{max}} \) is the maximum solid volume fraction.

The variation of plastic viscosity versus the time in different a/p ratios is represented in Fig. 6. The graph shows that the relative plastic viscosity increases with the decrease
of the a/p ratio and increases proportionally with time. This can be explained due to
the solid content and the stress to break the interparticle forces: the plastic viscosity
increases for pastes with a lower liquid phase which requires higher stress to break the
interparticle forces and higher value of solid content (Eq (4)). For example, the paste with
a 0.38 a/p ratio after 52 min has a higher value of plastic viscosity (0.4696 N.mm.min)
than the paste with a 0.40 a/p ratio (0.3989 N.mm.min).

![Relative plastic viscosity versus time.](image)

**Figure 6**: Relative plastic viscosity versus time.

### 3.1.4. Slump flow

Fig. 7 shows the variation of the slump (fluidity) of different pastes with a different ratio
of a/p versus time. From this graph, the reduction of the slump diameter with time and
the decrease of a/p ratios can be observed. This reduction of the diameter of the slump
occurs due to the formation of C-S-H gel with time (the paste is more hardened, 145
mm for the paste with a 0.39 a/p ratio in 7 that becomes 126 after 52 min).

![Slump versus time.](image)

**Figure 7**: Slump versus time.
3.2. Hardened state

3.2.1. Compressive strength

Fig. 8 presents the compressive strength results measured at 7, 14, and 28 days. When comparing the results obtained for the different mixtures, it was observed that the compressive strength decreased with the increase of a/p ratios. This can be explained by the increase of paste porosity which represents a compressive strength reduction of about 35% at 28 days for 0.37 to 0.41 a/p ratios. Moreover, it also decreases because of the increase of the a/p influence on the cohesion between the grains of the paste.

![Compressive strength results](image)

**Figure 8:** Compressive strength as a function of the a/p ratio.

4. Conclusions

This paper experimentally investigated the rheological properties of AA pastes of different mixes of tungsten mining waste mud, grounded waste glass and metakaolin. All mixes were activated using sodium hydroxide and sodium silicate. The main conclusions that can be drawn from this study are listed below:

1. The rheological behaviour in AAM pastes fits the Bingham model. The torque increases proportionally with the increase of rotational speed. Besides, the torque x rotational speed curves were found to be dependent on the activator/precursor ratios (i.e. the increase in the activator/precursor offset increased torque and relative yield stress).

2. The relative yield stress and relative plastic viscosity increased inversely with the increase of the a/p ratio and proportionally with time. This can be explained by the solid content of the mixes and the quantity of the liquid activator used (i.e. more...
liquid decreases the interparticle forces that are needed to break the interstitial cohesion).

3. The workability (fluidity) decreases with the decrease of the a/p. This can be explained by the decrease of the cohesion between the grains of the paste.

4. The compressive strength was affected inversely by the a/p ratio (the increase of compressive strength with the decrease in the a/p ratio). This can be explained by the increase in paste porosity.

5. Finally, the ideal a/p ratios in this study were a/p=0.38 and a/p=0.39 because these ratios have a higher mechanical performance with good rheological behaviour in the fresh state.

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