INFLUENCE OF THE CARBONATE-FREE CLAY CALCINATION TEMPERATURE
AND CURING CONDITIONS ON THE PROPERTIES OF ALKALI-ACTIVATED
MORTAR

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Abstract. Alkali-activated materials (AAMs) are one of the most perspective alternatives to the traditional Portland
cement based materials. AAMs could reduce the environmental pollution and greenhouse gas emission due to the lower
heat consumption in production (up to 80% compared to the ordinary Portland cement). In the current research alkali-
activated mortar was developed from the locally available low-carbonate Illite clay. Experimental work consisted of Illite
clay calcination at different temperatures and regimes (from 500-800°C) and incorporation of the obtained material into
the mixture of alkali-activated mortar. Four curing regimes were applied to alkali-activated mortar. The results indicate
that hardened mortar properties are mainly affected by curing temperature and regime and the compressive strength up to
28.8MPa could be achieved for the mortar samples aged 2 months.

Keywords: Alkali activated mortar, curing conditions, Illite clay, calcination temperature

Introduction

Today the cement and concrete industry has become a large scale industry due to the significant consumption of
cement. The total cement production in the world reached 3.7 million tons in 2012 (U.S. Geology Survey, 2013). Concrete is the second most widely used commodity in the world, after water (Alfinc, 2000). By producing such amount of cementitious materials, serious environmental pollution has been caused. Cement production contributes at least 5-8% of the global carbon dioxide emissions (Screvener and Kirkpatrick, 2008).

Ordinary Portland cement (OPC) is usually produced by heating the mixture of raw materials in a rotary kiln to about 1,450°C (Damtoft et al.). (Damtoft et al., 2008) have calculated that 0.53 t of CO₂ per ton of clinker is released along with an average of 0.34 t CO₂ per ton of cement associated with the energy consumption in the production process. Alkali activated materials (AMMs) are believed to be the most promising substitute material for OPC due to the much lower CO₂ emissions compared to OPC. The low CO₂ emission level is reached by using lower temperature for forming AAMs compared to the high sintering temperature used in OPC production. Calculation of CO₂ emissions using various alkali-activated fly ash and metakaolin binders has been done by Duxson et al. (Duxson et al., 2007). Calculation included CO₂ emissions during production of Na₂O in the chloric-alkali process; SiO₂ as aqueous sodium silicate was used as the primary inputs and the calculation showed that CO₂ saving was up to 80% compared to CO₂ emissions in OPC production.

Alkali activation of the materials containing amorphous Al₂O₃ and SiO₂ is a way in which specific cementitious materials are produced. Alkali activation of raw material composition with the concentrated NaOH solutions produced an amorphous material—a zeolitic precursor with excellent cementing properties in terms of mechanical strength (Palomo, 1999). The rate of chemical reactions was influenced by various parameters such as curing temperature, alkali concentration, initial solids content, etc. When the activator concentration increased, a delay in material structure formation was observed, whereas the increasing temperature accelerated its formation.

In the present research low calcium alkali activated binder has been developed. The fundamental binder structure in low-calcium alkali-activated systems is known to be a highly disordered, highly cross-linked aluminosilicate gel. Both Si and Al are present (Duxson et al., 2007). Alkali hydroxides are usually combined with an aluminosilicate source in the form of an aqueous solution (Provis, 2009). Given the complexity of the gel structure, which is formed through silicate activation of fly ash or metakaolin, and in particular its dependence on a large number of compositional and processing parameters, various approaches to the analysis of the gel binder were developed and implemented in the past years (Rahier et al., 1997, Palomo et al., 2005, Yao et al., 2009).

Researchers have shown that kaolinite clay is not the only natural source material to form alkali-activated systems. Illite-smectite clays have also shown strong potential for utilization in alkaline activation applications and the high Si content of the clays favours the use of a low-silicate alkaline activation solution in these systems (Buchwald et al., 2009). Other researches have shown that AAMs could be obtained from different types of grass ashes and waste materials together with calcined Illite clay at elevated curing temperatures and compression strength results could be up to 23 MPa (Bajare et al. 2011).

In this research carbonate-free Illite clay will be investigated in order to obtain alkali-activated mortar. The effect of calcination temperature of clay minerals will be tested and optimal curing conditions of forming AAM will be obtained.
Experimental setup

The basic raw materials used in the current study in order to obtain alkali-activated mortar were carbonate-free Illite clay (IC), lead silicate glass waste (LSG), sand, NaOH flakes and water. Alkali – activation of raw material composition was done by 6M sodium hydroxide solution. Materials were processed before application in the AAM.

Materials

Carbonate-free Illite clay

Carbonate-free Illite clay (IC) was used as a source of aluminium and silica dioxides needed in the activation process. Chemical composition of IC is given in Table 1.

Table 1. Chemical composition of Illite clay, lead-silicate glass and sand (w%).

<table>
<thead>
<tr>
<th></th>
<th>Illite clay</th>
<th>LSG</th>
<th>Sand</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al₂O₃</td>
<td>15.5</td>
<td>1.0</td>
<td>15.1</td>
</tr>
<tr>
<td>SiO₂</td>
<td>62.2</td>
<td>69.1</td>
<td>76.5</td>
</tr>
<tr>
<td>CaO</td>
<td>0.8</td>
<td>1.3</td>
<td>1.2</td>
</tr>
<tr>
<td>SO₃</td>
<td>-</td>
<td>-</td>
<td>0.1</td>
</tr>
<tr>
<td>TiO₂</td>
<td>1.9</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Na₂O</td>
<td>0.1</td>
<td>8.0</td>
<td>-</td>
</tr>
<tr>
<td>K₂O</td>
<td>4.1</td>
<td>1.2</td>
<td>-</td>
</tr>
<tr>
<td>MgO</td>
<td>1.3</td>
<td>-</td>
<td>0.4</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>7.2</td>
<td>0.2</td>
<td>0.6</td>
</tr>
<tr>
<td>PbO</td>
<td>-</td>
<td>19.0</td>
<td>-</td>
</tr>
<tr>
<td>LOI, 1000°C</td>
<td>4.7</td>
<td>-</td>
<td>5.9</td>
</tr>
<tr>
<td>Total</td>
<td>97.8</td>
<td>99.8</td>
<td>99.8</td>
</tr>
</tbody>
</table>

IC was calcined in four temperature regimes – from 500 to 800°C – by using slow and rapid calcination methods. The slow calcination regime was applied by using laboratory rotary furnace at 500 to 800°C. Rapid calcination regime was applied by using laboratory rotary furnace for sample 700° (R). Sample 700° (R) was calcined for 15 minutes at the maximum temperature (700°C). The specific test methodology was created at the Institute of Silicate Materials, Riga Technical University, to detect reactivity of calcined clays used in the AAM. The amounts of amorphous SiO₂ and R₂O₃ in the calcined IC are given in Table 2.

Table 2. Amount of amorphous SiO₂ and R₂O₃ in carbonate-free Illite clay calcined at different temperatures and regimes, (w%).

<table>
<thead>
<tr>
<th></th>
<th>500°</th>
<th>600°</th>
<th>700°</th>
<th>700° (R)</th>
<th>800°</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total SiO₂</td>
<td>72.70</td>
<td>72.65</td>
<td>73.15</td>
<td>72.94</td>
<td>73.20</td>
</tr>
<tr>
<td>Total R₂O₃ (Al₂O₃+Fe₂O₃+TiO₂) Fe₂O₃/6,15 TiO₂/1,21</td>
<td>19.00</td>
<td>19.40</td>
<td>19.08</td>
<td>19.35</td>
<td>19.48</td>
</tr>
<tr>
<td>SiO₂ active</td>
<td>1.10</td>
<td>1.05</td>
<td>0.89</td>
<td>1.05</td>
<td>1.11</td>
</tr>
<tr>
<td>R₂O₃ active</td>
<td>5.42</td>
<td>7.14</td>
<td>7.92</td>
<td>7.79</td>
<td>6.75</td>
</tr>
<tr>
<td>Fe₂O₃/1,16+1,23 TiO₂/0</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The highest amount of active R₂O₃+SiO₂ (8.81%) was detected for IC calcined at temperature 700°C, but the lowest result was for IC calcined at 500°C (6.52%). The rapid calcination of IC at temperature 700°C provided the total R₂O₃+SiO₂ of 8.84%, which was similar to slow calcination at the same temperature. It means that the maximal calcination temperature is in the range from 500 to 800°C and the speed of calcination does not influence the formation of active R₂O₃+SiO₂ significantly. Compared to kaolinite clays calcined at the same temperatures (700-800°C), the active R₂O₃+SiO₂ was 33-34% (according to data from the Institute of Silicate Materials). The calcined IC was ground in a planetary ball mill Retsch PM400 for 30min with 300rpm. The specific surface of the obtained IC was in the range from 20.12 to 21.42 m²/g. The micrograph of 700°C calcined IC is given in Fig. 1. Small particles with size from 0.5-1 µm were observed on larger particles with the size up to 10 µm.

Fig. 1. 700°C calcined IC particle size and shape (ground in the planetary ball mill for 30 min.).

The XRD analysis of calcined IC at 700°C is given in Fig. 2. The main minerals were quartz SiO₂, microcline KAISIO₂ and Illite K(ALFe)₂AlSi₃O₁₀(OH)₂H₂O.

Fig. 2. XRD analysis of IC calcined at 700°C.

Lead-silicate glass waste (LSG)

Lead-silicate glass waste used in this study comes from an outworn fluorescence lamp recycling plant. As it is not possible to recycle utilized glass by traditional methods due to specific chemical composition and high amount of lead, it poses a serious disposal problem, because glass is
not biodegradable and land filling is not an environmentally friendly solution. Usage of this type of glass waste in the production of AAMs is a waste management solution.

Fluorescent lamp recycling process includes lamp classification, glass separation, cleaning from harmful components and glass grinding. The chemical composition of waste glass LSG is given in Table 1.

In order to increase the fineness of waste glass, it was additionally ground in the laboratory planetary ball mill Retsch PM 400. The applied grinding period was 30 minutes. The particle size distribution in the powder material was determined by laser diffraction method. The results showed that the additionally ground material contains a wide range of particle sizes: from 8 µm to 30 µm with the average grain size of 26.3 µm. The specific surface of 30 min ground LSG was 0.63 m²/g.

Alkali activator solution
Commercially available sodium hydroxide flakes with 97% purity were also used. The alkali-activated mortar was prepared using 6M NaOH solution. The solution/solid (IC+LSG) mass ratio for all mixtures was constant – 0.49.

Mixture design and test methods
The main goal of research was to determine the appropriate calcination temperature for IC to reach highest mechanical properties of alkali-activated mortar. Four different IC calcination temperatures (500, 600, 700 and 800°C) and two regimes (rapid and slow) were used for calcination of IC. The mixture composition consisted from calcined IC, LSG, fine sand: from fraction 0.3/2.5 mm and alkali activator solution. The mass ratio of components is presented in the Table 3.

Table 3. Alkali-activated mortar mixture composition.

<table>
<thead>
<tr>
<th>Component</th>
<th>Mass ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcined Illite Clay</td>
<td>1.0</td>
</tr>
<tr>
<td>Ground lead-silicate glass waste</td>
<td>3.0</td>
</tr>
<tr>
<td>Fine sand 0.3/2.5mm</td>
<td>4.3</td>
</tr>
<tr>
<td>Alkali activator solution, 6M NaOH</td>
<td>2.0</td>
</tr>
</tbody>
</table>

All dry components were mixed together for 2 minutes using electrical one shaft hand mixer. Then 6M NaOH alkali solution was added to the mixture and mixing continued until homogenous mixture was prepared. The properties, such as consistency of fresh mortar (by flow table) according to LVS EN 1015-3 and fresh mortar density (LVS EN 12350-6), were determined. Alkali-activated mortar mixture was cast in 40x40x160mm prismatic moulds. Moulds were sealed with plastic sheet to avoid rapid water evaporation before specific curing regime was applied.

Four curing conditions were chosen: temperature regime of 75°C for 24h and 72h and of 100°C for 24h and 72h. After the curing period samples were cooled at room temperature and demoulded. The samples were kept in ambient environment until testing on the 4th and 60th day.

Flexural and compressive strength of the hardened alkali-activated mortar bars were tested according to LVS EN 1015-11 at the age of 4 days and 60 days.

Results

Fresh alkali-activated mortar results indicated that IC calcination temperature does not affect fresh mortar density (it was in the range from 2005-2025 kg/m³) (Table 4). In the fresh mortar the lowest consistency was for mixture composition with IC calcined at 500°C – 157 mm. For other mixtures fresh mortar consistency was in the range from 180-198 mm.

Table 4. Fresh alkali-activated mortar properties.

<table>
<thead>
<tr>
<th>IC calcination temperature and regime, °C</th>
<th>Fresh mortar density, kg/m³</th>
<th>Consistency (flow table) mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>500º</td>
<td>2010</td>
<td>157</td>
</tr>
<tr>
<td>600º</td>
<td>2011</td>
<td>181</td>
</tr>
<tr>
<td>700º</td>
<td>2009</td>
<td>195</td>
</tr>
<tr>
<td>700º (R)</td>
<td>2005</td>
<td>180</td>
</tr>
<tr>
<td>800º</td>
<td>2025</td>
<td>198</td>
</tr>
</tbody>
</table>

The flexural strength of the hardened alkali-activated mortars is given in Figure 3. The lowest flexural strength was for samples cured at temperature regime 75°C for 24h. It was in the range from 1.5-2.0 MPa. By extending the curing period from 24h to 72h at temperature 75°C flexural strength of all specimens increased almost twice and reached 2.6-5.5 MPa. The highest flexural strength was reached in the specimen with IC calcined at temperature 800°C – 5.5 MPa followed by 700°C – 4.5 MPa.

![Fig. 3. Flexural strength of the hardened alkali-activated mortar samples at the age of 4 days cured under different temperature regimes.](image-url)
Results of compressive strength for 4 day old alkali-activated mortar samples are given in Figure 4. The compressive strength of all 4 days old specimens did not depend on the curing temperature (75°C or 100°C, 24h) and were between 5 and 8 MPa. With longer curing period of 72h at the temperatures 75°C and 100°C the compressive strength of all 4 days old specimens increased significantly – two to three times. The average compressive strength of specimens cured at elevated temperatures (75 and 100°C) for 72 h was between 12 and 19 MPa. The highest result was reached for the mixture with IC calcined at temperature 800°C and cured 72h at maximum temperature 75°C (18.7 MPa).

Samples cured at the higher temperature (100°C) showed strength increase by 20-50% compared to the samples cured at 75°C.

According to these results it was possible to conclude that the period of curing at elevated temperature is the main factor influencing increase of early age compressive and flexural strength for alkali activated mortars made from IC.

The long term compressive strength has provided significant strength increase for all mixtures and curing conditions (Fig. 5). At the age of 60 days samples cured at 75°C for 24h demonstrated compressive strength from 8.2 to 11.6 MPa. Highest result was shown by the samples with IC calcined at temperature 700°C (11.6 MPa). The strength increase of 60 days old samples was between 70-90% compared to the samples cured at the same regime but tested on the 4th day.

The compressive strength for samples hardened for 72h at 75°C was from 18.8 to 28.8 MPa. The highest result was for the mixture composition with IC calcined at temperature 800°C (28.8 MPa). Compressive strength increase was from 60 to 80% compared to the 4th day strength results. Summarizing the test results, it can be observed that samples continued to gain strength significantly over time with no relation to the time period of curing (24 or 72h), but the tendency was that the prolonged curing period (72h) led to higher strength even on the 60th day of testing.

Samples cured at 100°C have demonstrated lower strength increase compared to the samples cured at 75°C – 28-49%. Testing on the 60th day shows strength from 8.5 to 10.6MPa for mixtures hardened for 24h. If compared to samples cured at 75°C for the same period of time, the compressive strength results were more stable for all mixtures. Samples cured at 100°C were with lower strength increase compared to samples cured at 75°C. The reaction processes were enhanced during primary curing, therefore further strength increase was with lower increase rate.

Samples hardened at 100°C for 72h at the age of 60 days show insignificant strength increase – from 16 to 20 MPa. Strength increase was from 10 to 20% comparing to 4 day test results. The average results for samples cured at 100°C were lower compared to the samples cured at 75°C for all mixtures, but these results were much more stable.

Conclusions

Alkali-activated mortar could be created using carbonate-free IC calcined at different temperatures between 500-800°C. Amount of amorphous R₂O₃ in the calcined clay does not have direct correlation with the strength results of the obtained alkali activated mortar. The mechanism of alkali activation must be investigated properly in the future. The highest strength results for alkali-activated mortar were obtained with IC calcined at 800°C with amount of amorphous R₂O₃ +SiO₂ 7.86% – 18.7MPa at 4 days and 28.8MPa at 60 days.

Curing conditions are an important issue for creating AAMs. By choosing appropriate curing conditions and regime cementitious material with similar properties to cement based materials could be created. The optimal curing temperature could be in the range from 70 to 80°C. By increasing temperature from 75 to 100°C small compressive strength increase was observed. The initial strength of alkali-activated mortar increases and results are distributed much more evenly between mixture compositions compared to the results obtained from samples cured at 75°C.

The prolonged curing period leads to strength increase at both temperature regimes. The 4 day strength could be
increased by 100-200% just by increasing curing period from 24h to 72h.

Alkali-activated mortars continue to gain strength during time. All samples demonstrated strength increase between 4th and 38th day. The highest strength increase was for samples hardened at 75°C for 24h – 70-90% and for samples cured at 75°C for 72h it was from 60 to 80%. By increasing curing temperature and time hardening acceleration was observed, but further strength increase decreases with time.

The initial temperature increase rate and final cooling rate should be controlled to avoid thermal shock for the material and to avoid thermal cracks in the structure of the material. Best compressive rate should be controlled and tested in order to avoid thermal shocks for the material and to avoid thermal cracks in the structure of the material. The highest strength increase was achieved for IC calcined at temperature 800°C and curing conditions were 75°C for 72h – 28.8MPa at the age of 60 days.

The obtained material could be compared to the traditional cement-lime mortars considering test results for density and mechanical and physical properties.

References


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