The Effect of Heat Treatment on Alkali Activated Materials

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crossref http://dx.doi.org/10.5755/j01.ms.23.3.16280

Received 23 September 2016; accepted 06 December 2016

The primary object of the present research was to investigate the porous low calcium alkali activated material (AAM). Traditionally Na+ ions for alkali activation solution ensure highly alkaline media which enhances the dissolution of amorphous phase in the raw materials forming solid cementitious material with sodium aluminosilicate hydrate (N–A–S–H) structure afterwards. Almost all alkali ions are partially hydrated filling the pores in the gel structure (N–A–S–H gel, type zeolite precursor) and neutralizing the charge on Al(OH)4 groups. These alkali ions are available for leaching in water environment. Due to this property the application of porous AAM in this research is related to the water treatment systems similar to those of natural zeolites which are considered as effective sorbent because of their porous structure, high specific surface and ion exchange. Porous AAM was obtained from metakaolin, sodium silicate glass, modified sodium silicate solution with Ms = 1.68 and diethylene glycol (DEG) aluminium paste as pore forming agent. The density of AAM was 1150 ± 12 kg/m³ and compressive strength fc > 12 MPa. The effect of heat treatment to microstructure and structural properties of AAM was investigated. Heat treatment is an effective method for changing the alkali leaching kinetic form AAM structure.

Keywords: alkali activated material, heat treatment, alkali leaching, water treatment.

1. INTRODUCTION

The quality of water resources is a major issue in the modern society due to the excessive increase in volumetric production of industrial and household wastewater and inadequate treatment of polluted water which could lead to the environmental problems. Traditionally heavy metals from wastewaters are removed using chemical precipitation, ion-exchange and electrochemical deposition which have many disadvantages such as high energy requirements and toxic sludge production during these processes [1]. The precipitation is considered to be a simple and effective method for wastewater treatment if low cost sorbents is used. It is considered that natural zeolites are effective sorbent material for separation and purification processes because of their porous structure and mobility of alkaline metals [2]. Also alkalizing agents, such as lime or caustic soda, have been used to raise the pH levels needed to induce chemical clarification of wastewaters, i.e. magnesium precipitation begins at approximately pH 9.5, becomes significant above pH 10.5, and is essentially complete at pH 11.0 – 11.5. Good clarification is usually not achieved until pH 11.0 – 11.5 is reached [3]. Flocculation at high pH media results in efficient removal of particles, colloids and certain dissolved materials present in the wastewater.

Natural zeolites could be altered with artificial alkali activated materials (AAM). AAM could be obtained from Al2O3 and SiO2 rich raw materials by using alkali activation technology [4]. Most common raw materials are metakaolin or fly ash which is activated with NaOH or sodium silicate solutions at relatively low temperature. The general formula for the reaction product is 2SiO2·Al2O3·Na2O·2H2O [5] which is expressed as N–A–S–H gel ensuring good durability and mechanical properties [6–8]. The principle of AAM structure formation is given in Fig. 1. When sodium silicate solution with low Ms is used as activator, the solid material formed is amorphous and cementitious, but its structure and composition are different from the product, which is formed using only NaOH. The amorphous N–A–S–H gel has similar chemical composition as natural zeolite materials but without the extensive crystalline zeolite structure [10, 11]. The optimal SiO2/Al2O3 ratio in the raw materials varies between 3.0 and 4.0 while Na2O/Al2O3 ratio is around 1.0 [12].

Alkali activation solution ensures Na+ ions and creates a strong alkaline media for the dissolution of amorphous phases from the raw materials. According to some authors [13, 14] the sodium taken up in these materials to balance the charge deficit in the gel structure, sodium can neutralize the negative charge in two ways: sodium associated with aluminium inside the gel structure (compensation of the charge deficit), more difficult to leaching and sodium partially hydrated that fills the pores in the gel structure, where it neutralizes the charge on Al(OH)4 groups and must easy to exchange. More recently Skvara et al. have indicated that Na+ ions are weakly bound in the nanostructure of the N–A–S–H gel and therefore are almost completely leachable. NaOH easily diffuses to the surface, where it reacts with atmospheric CO2, while forming visible salts available for leaching, such as Na2CO3·H2O, NaHCO3,
The leaching rate of Na\(^+\) and OH\(^-\) from the AAM activated with NaOH or with sodium silicate solution is high for the first few weeks, decreasing over time [16]. A positive factor is the fact that alkalis could be leached out of the material structure without compromising its mechanical properties; therefore, this renders Na\(^+\) as an unnecessary load-bearing element in the nanostructure of the N–A–S–H gel and demonstrates A–S–H to be a load-bearing structure [15].

Fig. 1. Microstructure: a – aluminosilicate sources + Na\(^+\) (aq), OH\(^-\) (aq); b – N-A-S-H gel 1 (Si/Al=1); c – N-A-S-H gel 2 (3D) (Si/Al=2) [9]

The porous structure of AAM would enclose alkalis inside the material structure resulting in a gradual alkali leaching over longer period of time. The porous structure ensures wider surface, where Na\(^+\) ions can diffuse. Alkali release over a longer period of time is an advantage for using material in water treatment systems [17]. The porous material structure can be obtained by mixing gas-releasing agents like aluminium paste [18]. However, waste materials, like non-metallic residues from aluminium scrap recycling industry, can be used as well [19].

The DTA/TG analysis of AAM indicates that the heat treatment ensures loss of adsorbed water at 90 °C. The loss of hydroxyl species depending on the alkaline element used for the synthesis proceeds at around 200°C and the loss of structural water from crystalline structure – at around 400–500 °C [20]. The dehydration reaction could affect properties of the material in water media and hydration reaction occur to reattach the lost water therefore affecting leaching properties in water media.

The present paper suggests using low-calcium AAM obtained by alkali activation technology as the passive system for stabilization of the pH (pH 10 to 11) in the water treatment systems. The effect of heat treatment on the structure of AAM and leaching properties were studied. The prepared AAM was amorphous to semi-crystalline material that is formed by transformation of metakaolin and sodium silicate glass raw materials with sodium silicate solution at low temperature in a very short time.

2. EXPERIMENTAL DETAILS

Porous alkali activated material (AAM) was created using metakaolin (MK) and sodium silicate glass (SSG) activated with modified sodium silicate solution (Ms 1.67) (activator). Activator was prepared using sodium silicate solution with Ms 3.22 and sodium hydroxide flakes with the mass ratio 9:1. Suspension of the diethylene glycol aluminium paste (AlP) mixed with H\(_2\)O was used as a pore structure forming agent (mass ratio 1:5). The chemical composition of raw materials is given in Table 1.

**Table 1. Chemical composition of raw materials**

<table>
<thead>
<tr>
<th>Compound</th>
<th>Amount, wt.%</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MK</td>
</tr>
<tr>
<td>SiO(_2)</td>
<td>57.2</td>
</tr>
<tr>
<td>Al(_2)O(_3)</td>
<td>33.9</td>
</tr>
<tr>
<td>TiO(_2)</td>
<td>2.9</td>
</tr>
<tr>
<td>Fe(O(_3))</td>
<td>1.5</td>
</tr>
<tr>
<td>CaO</td>
<td>0.3</td>
</tr>
<tr>
<td>MgO</td>
<td>0.4</td>
</tr>
<tr>
<td>Na(_2)O</td>
<td>0.3</td>
</tr>
<tr>
<td>K(_2)O</td>
<td>2.7</td>
</tr>
<tr>
<td>Al</td>
<td>–</td>
</tr>
</tbody>
</table>

The mixture composition of AAM is given in Table 2. The electrical hand mixer was used for controlled mixing. All dry raw materials were mixed for 15 seconds, then activator was added to the mixture and mixed for 2 minutes until homogenous paste was obtained. Suspension of AlP was added to the paste to create porous structure and mixed for 10 seconds. Right after the mixing, paste was casted into sealed prismatic moulds measuring 40 × 40 × 160 mm. Moulds were placed in a room temperature (20 ± 2 °C) for 20 minutes to provide pore formation. Then the moulds were covered with plastic film and cured at 80 °C for 24 h. The prepared samples will be referred as A10-T-SSG1 further in the text. After demoulding the following physical properties of the prepared porous AAM samples were tested in accordance with EN 1097-6 and EN 1097-7: bulk density, water absorption (wt.%), open and total porosity (vol.%). The heat treated samples were prepared at 200 °C for 3 h with heating rate 10 °C/min and will be referred as A10-T-SSG1-200 further in the text.

Na\(^+\) ion exchange in water media was determined by using titration method. Alkalis hydrated within the pores of the gel, more weakly bound, are exchanged; in this case - with the water protons (Na\(^+\) ion is exchanged for a H\(^+\) ion). The phenomena resulted in an increased alkalinity of the medium. The pH of water solution after 24 h immersion of the prepared AAM (weight 3.0 ± 0.2 g) in the 100 ml of deionized water was determined with portable pH/mV meter HI 991003 with sensor check. Samples were moved to the new batch with deionized water (100 ml) every 24 h. Obtained leachate was titrated to determine OH\(^-\) group in (OH-mol·(l·g)\(^{-3}\)) with hydrochloric acid 0.01M to pH 7.0 for determination alkalinity.
Table 2. Mixture composition of AAM

<table>
<thead>
<tr>
<th>Compound</th>
<th>A10-T-SSG1 composition, mass ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Suspension of AlP</td>
<td>0.02</td>
</tr>
<tr>
<td>MK</td>
<td>1</td>
</tr>
<tr>
<td>SSG</td>
<td>1</td>
</tr>
<tr>
<td>Activator</td>
<td>1.3</td>
</tr>
<tr>
<td>Activator to solid ratio</td>
<td>0.65</td>
</tr>
<tr>
<td>SiO₂/Al₂O₃</td>
<td>4.5</td>
</tr>
<tr>
<td>Na₂O/Al₂O₃</td>
<td>1.0</td>
</tr>
</tbody>
</table>

The thermogravimetric-differential thermal analysis (DTA/TG) was carried out by using Stanton Redcroft STA 781 thermal analyser from 0 °C to 700 °C at a heating rate of 5 °C/min. The mineralogical and microstructural characteristics of the obtained material before and after heat treatment, as well as before and after leaching test were studied with FTIR, XRD and SEM/EDX.. ATIMATTSON FTIR-TM was used to obtain FTIR spectra. Test specimens were prepared by mixing 300 mg of KBr with 1 mg of AAM sample. The spectral analysis was performed in a range from 2000 to 400 cm⁻¹. The crystalline phases of material before and after treatments were identified with XRD BRUKER AXS D8 ADVANCE X-ray diffractometer and collected with CuKa1, α2 radiation in the range 5 – 60° (2θ). JEOL JSM 5400 scanning electron microscope (SEM) with LINK-ISIS energy dispersive analyser (EDX) was used for the microstructural characterization.

3. RESULTS

Physical properties of the prepared AAM A10-T-SSG1 are given in Table 3. The macro- and microstructure of AAM is given in Fig. 2. The porous structure of the obtained samples is similar to those of aerated autoclaved concrete. A10-T-SSG1 and A10-T-SSG1-200 have homogenous macro pore structure with pore size ranging from 200 to 600 µm (Fig. 2). Micro level pores with size up to 50 µm were detected. The crystalline substances were observed on the surface of pore walls and inside the structure (broken cross-sections) (Fig. 2 b). The size of these crystalline substances was 10 µm. EDX results indicate that these substances contain high amount of Na (35 – 40 wt.%), O (37 – 40 wt.%), Si (14 – 20 wt.%) and Al (3 – 9 wt.%). After the heat treatment at 200 °C (A10-T-SSG1-200) the macro scale changes were not observed (Fig. 2 c). The size of previously observed crystalline substances (found on the pore surface Fig. 2 d) decreased from 10 µm to 1 – 5 µm.

Table 3. Physical and mechanical properties of AAM (A10-T-SSG1)

<table>
<thead>
<tr>
<th>Property</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk density, kg/m³</td>
<td>1150 ± 12</td>
</tr>
<tr>
<td>Water absorption, wt.%</td>
<td>9.5 ± 0.5</td>
</tr>
<tr>
<td>Open porosity, vol.%</td>
<td>10.6 ± 0.8</td>
</tr>
<tr>
<td>Total porosity, vol.%</td>
<td>57.3 ± 0.5</td>
</tr>
<tr>
<td>Compressive strength, MPa</td>
<td>12.8 ± 1.2</td>
</tr>
</tbody>
</table>

Fig. 1. Microstructure of prepared samples: a and c – AAM A10-T-SSG1; b and d – heat treated AAM A10-T-SSG1-200
DTA/TG results indicate that the prepared AAM A10-T-SSG1 has endothermic peak at around of 100 to 120°C and small endothermic effect at around 200°C (Fig. 3). TG curve shows even weight loss during heating resulting in 95.3 % at 400 °C. To study this endothermic effect, AAM was heated at 200 °C for 3 h (A10-T-SSG1-200). Afterwards DTA/TG was repeated and previously detected peaks below 200 °C were not detected.

After 10 days leaching test DTA/TG analyses were repeated for both the heat-treated and untreated samples (A10-T-SSG1-AL and A10-T-SSG1-200-AL). The endothermic effect at around 110 °C was still detected while at 200 °C it diminished. It was detected that the weight loss at 120 °C for A10-T-SSG1-AL was 6 % while for A10-T-SSG1-200-AL were 2.5 % (Fig. 4). At 200 °C the weight loss increased to 11 % and 4.5 % respectively.

Fig. 2. DTA/TG results of prepared A10-T-SSG1 and A10-T-SSG1-200

Fig. 3. DTA/TG results of AAM after leaching test (A10-T-SSG1-AL and A10-T-SSG1-200-AL)

Structural changes in macro and micro level after the heat treatment were not detected. Both heat-treated and untreated samples were tested on alkali leaching kinetic form AAM structure (Fig. 5). It was observed that the untreated sample provides two times higher alkali leaching rate comparing to the sample treated at 200 °C (Fig. 5). The alkali leaching in both cases was gradual with slight trend to lower leaching intensity by the increase of leaching time. After 10 day leaching untreated AAM the sum of alkalinity was 0.026 OH-mol·(l·g)^{-1} while the heat treatment reduced leaching more than 2 times and was 0.012 OH-mol·(l·g)^{-1}. The initial pH level depending from the heat treatment was pH 11.9 for A10-T-SSG1 and pH 11.7 for A10-T-SSG1-200. Gradual pH decrease was observed for A10-T-SSG1 and after 10 days leaching period pH decreased to pH 9.9. The heat-treated sample (A10-T-SSG1-200) showed initial pH decrease up to 5th day reaching pH 11.2 and remained constant up to the 10th day.

Fig. 4. Cumulative leaching rate of OH ions and pH level during leaching test of A10-T-SSG1 and A10-T-SSG1-200

EDX analysis indicates that the element changes in the structure of AAM samples (Table 4). 15 to 30 spectral points for each element were analysed. The heat treatment did not change elemental analysis significantly (A10-T-SSG1 and A10-T-SSG1-200): Al was 6.8 wt.%, Na 14.9 – 15.1 wt.%, Si 30.4 – 32.2 wt.% and O from 45.6 to 46.7 wt.% respectively. After 10 days leaching the element relation changed and the amount of Na reduced noticeably to 5.7 wt.% for A10-T-SSG1-AL and to 9.2 wt.% for A10-T-SSG1-200-AL. The amount of Si and O remained approximately in the same level. This means that a large part of sodium leached as indicated by Skvarka et al. [15]. Sodium that still remains in the matrix could be leached over time; however according to Duxson et al and Criado et al a small amount could be retained as a part of the gel N-A-S-H [15, [25]. Studies at a later leaching age would be needed to clarify this issue.

Table 4. EDX element analysis of AAM at different treatment regimes

<table>
<thead>
<tr>
<th>Element</th>
<th>A10-T-SSG1</th>
<th>A10-T-SSG1-200</th>
<th>A10-T-SSG1-AL</th>
<th>A10-T-SSG1-200-AL</th>
</tr>
</thead>
<tbody>
<tr>
<td>O</td>
<td>45.6 ± 3.0</td>
<td>46.7 ± 1.5</td>
<td>48.9 ± 2.0</td>
<td>46.7 ± 3.4</td>
</tr>
<tr>
<td>Na</td>
<td>14.9 ± 2.9</td>
<td>15.1 ± 3.4</td>
<td>5.7 ± 1.3</td>
<td>9.2 ± 2.1</td>
</tr>
<tr>
<td>Al</td>
<td>6.8 ± 2.1</td>
<td>6.8 ± 2.6</td>
<td>11.3 ± 2.1</td>
<td>8.5 ± 2.6</td>
</tr>
<tr>
<td>Si</td>
<td>30.4 ± 4.1</td>
<td>32.2 ± 3.6</td>
<td>31.5 ± 3.0</td>
<td>32.3 ± 4.1</td>
</tr>
</tbody>
</table>

The FTIR spectra are given in Fig. 6. The bands between 1017-1076 cm⁻¹ are characteristic for the internal vibrations in aluminosilicates associated with T-O (T = Al,
Si) asymmetric stretching vibrations. This band moves to lower frequencies in AAM, which is associated to the formation of a N–A–S–H gel [21]. However, in the leaching process it returns to slightly higher frequencies. The intensity of the band at 452–465 cm\(^{-1}\), associated in all cases with T–O bending vibrations, barely changes its position throughout the activation process [22, 23]. This band provides an indication of the degree of “amorphisation” of the material, since its intensity does not depend on the degree of crystallisation. The band appearing at 780–790 cm\(^{-1}\) corresponds to the quartz doublet present in the original MK. The band at 558 cm\(^{-1}\) corresponds to the Si–O–Al\(^{4+}\) vibrations in dehydroxylated muscovite [24], the intensity of this band decrease in AAM, indicating that this phase shows weak reaction, which probably could be explained with the dihydroxylation. The aluminum coordination environment in the (Al\(_6\)) octahedron gradually changed to five-coordination leading to the relaxation of the crystal and increase of its reactivity. Intense C–O band asymmetric stretching vibration band appeared at 1451 cm\(^{-1}\) [21] for sample A10-T-SSG1, while for sample A10-T-SSG1-200 this band absorption intensity decreased and for samples after leaching (A10-T-SSG1-AL and A10-T-SSG1-200-AL) did not appear at all. This band is associated to the present of a little sodium carbonate that has been solubility in the leaching process.

![Fig. 5. FTIR analysis for raw materials and samples (absorption spectra)](image)

Results of the X-ray diffraction analysis for raw materials and obtained samples are given in Fig. 7. The halo present in SSG diffractogram in the 20–30° 20 intervals is indicative of vitreous component, in MK diffractogram a hump is also observed associated to the structurally disordered phase such as metakaolinite. Some minor crystalline phases were detected in MK: quartz (SiO\(_2\)), mica KAl\(_2\)Si\(_3\)O\(_{10}\)(OH)\(_2\), and muscovite (K\(_{0.6},\)Na\(_{0.1}\))(Fe\(_{0.03},\)Al\(_{1.97}\)). In the obtained AAM the halo shifts to higher 2\(\theta\) values, which indicates the formation of N–A–S–H gel [25] and has the halo in the 2\(\theta\) = 15–40° region, which is typical also for the A10-T-SSG1-200 and samples after leaching test (A10-T-SSG1-AL and A10-T-SSG1-200-AL). The crystalline muscovite peaks were not detected in AAM after alkali activation. Other crystalline materials from MK such as quartz remained in AAM after alkali activation.

![Fig. 6. XRD diffractograms of raw materials and prepared AAM samples](image)

4. DISCUSSION

Porous AAM with the bulk density of 1150 kg/m\(^3\) was obtained using AIP as pore forming agent and the bulk density of material ensures material that does not float in water. The low open porosity (9.5 wt.\%) and water absorption (10.6 vol.\%) provides gradual water penetration into the structure of AAM; therefore, ion exchange could be reduced comparing to the materials with high open porosity around 30 % [26]. However, the fractioned AAM with open pore structure and increased surface could be prepared to enhance leaching [27]. The compressive strength of 12.8 MPa allows to use this material in water treatment systems with pressure up to 1 bar without damaging the AAM matrix.
Two distinct endothermic peaks in A10-T-SSG1 were detected in the DTA spectra. The first endothermic peak appeared at 110 °C, while the second peak appeared at 190 °C. The first endothermic peak was primarily caused by the dissipation of free water from the AAM surface. The water evaporated within this temperature range, and similar behavior was observed by Cheng et al [28]. The second endothermic peak was attributed to the presence of zeolitic water in the reaction products, water bonding tightly with AAM, or water needing higher temperature to reach the surface of AAM [29]. The weight loss of A10-T-SSG1 was 4.0 % at 120 °C and increased to 7 % at 200 °C whereas heat treated AAM (A10-T-SSG1-200) had weight loss of only 0.5 % at 120 °C and 0.8 % at 200 °C according to TG analysis. Heat treatment effectively removed free water which could retard ion exchange due to the low hydration process, which takes place after immersing material into the water. This effect was detected by DTA/TG after the 10 day, leaching test in water. After leaching DTA/TG was repeated and it was observed that two endothermic effects remained for both treated and untreated AAM (A10-T-SSG1-AL and A10-T-SSG1-200-AL). The weight loss was 6 % at 120 °C and 11 % at 200 °C for A10-T-SSG1-AL, while for A10-T-SSG1-200-AL the weight loss was 2.5 % and 4.5 % respectively. The 10 day, immersing period increased the amount of water hydrated in the structure of AAM; therefore weight loss increased.

The leaching test reveals the importance of AAM heat treatment on ion exchange kinetics. Untreated AAM (A10-T-SSG1) with attached alkaline crystalline substances on pore surface (Fig. 2) provides high initial pH; however, over the time pH decreases and amount of Na in structure of AAM decreases which was detected with EDX (Table 4). The ion exchange trend line also declines to reduced leaching intensity at the end of the leaching test. Sample A10-T-SSG1-200 has lower leaching rate comparing to A10-T-SSG1; however, the leaching process is more gradual and stabilizes over time and the pH level of leachate becomes predictable at value pH 11.2. According to EDX more Na remains in the structure after leaching test which indicates the long lasting alkali leaching properties. Also EDX indicated that heat treatment did not affect the amount of Si, O, Na, Al in the structure of AAM, while AAM after leaching provided reduced amount of Na and increased amount of Al.

The FTIR spectra of A10-T-SSG1 shows that N-A-S-H gel formation has occurred (Fig. 6) [23, 30]. The absorption shoulder shift to higher frequencies indicates the bond transformation and alkali activation of raw materials. The heat treatment of AAM removes the C-O (CO$_2^-$) bond absorption spectra at 1451 cm$^{-1}$ which could indicate the decomposition of bicarbonates and carbonates such as NaHCO$_3$ (decomposition temperature 100 to 200 °C). The effect of leaching has narrowed the absorption band shoulder for both AAM and this phenomenon should be investigated more closely.

The obtained AAM is „x-ray amorphous” demonstrated by the halo in the 2$\theta$ = 15–40° region that is usually described as geopolymeric gels formation [31]. Substantial impact of the heat treatment and leaching on obtained material was not detected by XRD.

5. CONCLUSIONS

DTA/TG analysis indicate that the heat treatment of AAM at 200 °C removes free water from the structure of AAM; therefore, the hydration time increases, when AAM is subjected to the water and this effectively reduces ion exchange and the exchange rate is even.

The heat treatment has an important role regarding the leaching rate and pH changes of the leachate. The heat-treated materials provide even leaching rate at least 10 days and pH level stabilizes after 5 leaching days reaching pH 11.2 which remained constant over a period of up to 10 days, while the pH level of untreated AAM decreases and ion exchange rate was up to 3.2 times higher compared to heat-treated AAM.

The content of Na decreases in the structure of AAM during leaching. This would allow predicting the effective leaching time of the material during its immersion into the water, where duration of leaching for the heat-treated material would be considerably longer.

FTIR and XRD analysis did not indicate substantial changes of AAM structure during the heat treatment and leaching confirming that Na leaching is bound to the structure with weak bonds.

The use of heat treated AAM would be preferable in water treatment systems due to the prolonged leaching period and more stable pH level during leaching.

Acknowledgments

This work was supported by the Latvian Council of Science Fund Project Z12.0491 “Smart biofiltration technology” and also by the Spanish Ministry of the Economy and Competitiveness under the projects BIA2013-43293-R.

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