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INVESTIGATION ON THE EFFECT OF SLAG AND LIMESTONE POWDER ADDITION IN ALKALI ACTIVATED METAKAOLIN

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Abstract

Alkali activated material has recently attracted great attention as an environmentally friendly building material. Generally, it exhibits higher mechanical properties and better durability when compared with Portland cement based materials. In this article, mixed alkali activated systems consisting of N-A-S-H and C-A-S-H hydrated gels were produced, using metakaolin as a dominated starting material and granulated blast furnace slag as an additional calcium and silicate source. Limestone powder was then added into the blended metakaolin-slag system as a filler. The influences of slag and limestone powder on setting times, strength development and microstructure were analyzed. The results indicate that, with the increasing slag content, the setting time decreases; the compressive strength firstly increases and then decreases after the slag replacement exceeds a critical value. Additionally, an optimum content for limestone powder in the metakaolin-slag blends is around 15%. The FTIR spectra and XRD pattern present the changes in reaction products when slag was added, and the scanning electron microscope results showed a denser matrix of the geopolymeric system when compared to reference sample. However, no significant changes of reaction products were detected when limestone powder was incorporated, which indicates that the limestone powder only works as a non-reactive phase in the alkali activated system.

Keywords: Alkali activation, Metakaolin, Slag, Limestone powder

1. INTRODUCTION

The production of Portland cement is now responsible for nearly 7% of the total CO₂ emission around the world [1], other accompanied negative impacts of cement industry are the consumption of natural resources and high energy cost. Thus it is urgent to find approaches to reduce the environmental impacts of cement industry. Among those studies of finding alternatives to ordinary Portland cement, alkali activated materials have attracted great attention in recent years mainly due to their high mechanical properties, low environmental and energy consumptions [2,3]. Also other excellent properties are reported such as low heat resolution during hydration, fast early strength development, good durability related properties and thermal stability [4-7]. Raw materials used in producing alkali activated binders are usually industrial wastes, other than ordinary Portland cement, thus a promising future of sustainable development can be seen from this new clinker free building material.

Alkali activated materials are usually produced by mixing alkaline activator solutions with solid raw materials. The hydration process is called geopolymerization, which contains a series of chemical reactions including the dissolution of starting materials, polycondensation, condensation and finally the crystallization of new reaction products [8]. Final products can be classified into two types according to different raw material chemistry, for silicate and aluminum enriched precursors like metakaolin or fly ash, final products are amorphous alkaline aluminosilicate hydrate gels with a three-dimensional network of aluminate and silicate units that are charge balanced by alkali cations, or can be simply described as N-A-S-H type gel [9]; for calcium and silicate enriched ones like granulated blast furnace slag, final products are aluminium substituted C-S-H (I) type gel with lower Ca/Si ratio [10]. Sodium hydroxide modified sodium silicate has been widely accepted as an ideal activator in terms of strength and other beneficial properties, the alkaline activator type and dosage have a significant influence on the reaction process and comprehensive properties of the final products [11]. Other important factors are curing conditions such as temperature, time, relative humidity and physicochemical properties of the starting materials such as mineralogical composition and fineness [12].

Commonly alkali activated materials based on Si and Al type have excellent mechanical properties, enhanced reinforcement steel bonding and fire resistance, but at the same time, drawbacks such as long setting time and slow strength development, high requirement of curing are obstacles to their practical application. Several researchers have found that adding slag into Si and Al based geopolymer can improve the compressive strength and decrease the setting time to some degree, also acceptable strength can be achieved [13,14].
Calcium component in slag is responsible for the decrease of setting, the stable coexistence of aluminum substituted C-S-H gel and the N-A-S-H gel contributes to the improvements in terms of strength, permeability and durability [15]. Besides, only a few work focuses on the utilization of limestone powder as additives in alkali activated systems. Limestone has already been used in Portland cement system successfully, the already known influences are improving workability and strength by the filler effect, accelerating the hydration process by the nuclear effect, also lowering the price and energy cost of the overall system. Thus there could be some potential benefits of using limestone in alkali activated systems such as modifying workability, strength, cost and environmental impact. The aims of this paper are to investigate the influences of slag on setting, strength and phase changes on alkali activated metakaolin, also to study the effects of limestone powder on metakaolin-slag blends.

2. EXPERIMENTAL PROGRAM

2.1 Materials

The metakaolin was produced in the laboratory by calcination of kaolinite under 750 °C for 4 hours. The kaolinite containing small amount of quartz and illite as impurities, mineral composition are shown in Table 1, with a specific surface area of 490 m²/kg and more than 90% of particles are finer than 2 µm. The granulated blast furnace slag used in Portland cement system successfully, the already known influences are improving workability and strength by the filler effect, accelerating the hydration process by the nuclear effect, also lowering the price and energy cost of the overall system. Thus there could be some potential benefits of using limestone in alkali activated systems such as modifying workability, strength, cost and environmental impact. The aims of this paper are to investigate the influences of slag on setting, strength and phase changes on alkali activated metakaolin, also to study the effects of limestone powder on metakaolin-slag blends.

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Kaolinite</th>
<th>Quartz</th>
<th>Illite</th>
</tr>
</thead>
<tbody>
<tr>
<td>%</td>
<td>91.91</td>
<td>1.6</td>
<td>6.49</td>
</tr>
</tbody>
</table>

Table 1: Mineral compositions of kaolinite

<table>
<thead>
<tr>
<th>Oxides</th>
<th>Metakaolin</th>
<th>Slag</th>
<th>Limestone</th>
</tr>
</thead>
<tbody>
<tr>
<td>CaO</td>
<td>54.46</td>
<td>38.06</td>
<td>53.42</td>
</tr>
<tr>
<td>SiO₂</td>
<td>10.34</td>
<td>34.14</td>
<td>0.61</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>39.86</td>
<td>10.34</td>
<td>0.49</td>
</tr>
<tr>
<td>MgO</td>
<td>0</td>
<td>7.95</td>
<td>1.62</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>1.98</td>
<td>0.38</td>
<td>0.21</td>
</tr>
<tr>
<td>LOI</td>
<td>2.01</td>
<td>0.03</td>
<td>43.15</td>
</tr>
</tbody>
</table>

Table 2: Major compositions of raw materials (%)

2.2 Mix design and sample preparation

Samples were prepared by firstly mixing NaOH with sodium silicate solutions and then cooling it down to room temperature, then blending the liquid activator with the raw material in the mixer. Slag was added from 0% to 40% with a range of 10% by mass to study the influences of slag addition on setting, strength and reaction products of alkali activated metakaolin. For each sample, sodium hydroxide and sodium silicate were used with the fixed total SiO₂/Al₂O₃ ratio of 4.0 and Na₂O/Al₂O₃ ratio of 1.0. Based on the alkali activated metakaolin-slag mixes with ideal property, limestone powder was added from 0% to 20% by mass of the total binder to analyze its effects on strength and reaction products of the blended system. A overall water/binder ratio of 0.37 was used for all mixes, while the total water includes the free water and water in activator solution, and the total binder was defined as raw materials and the solid phase in activator. All mixes were cured at 25 ± 2 °C with a relative humidity of 90% until the testing age.

2.3 Testing methods

The setting times of the alkali activated pastes were tested using the Vicat needle method in accordance with the Chinese national standard GB/T 1346-2011. The compressive strength testing were carried out according to the Chinese national standard GB/T 17671-2005. The samples were tested after 7 and 28 days curing, the strength values for each sample was obtained from the average of three mixes. The X-ray diffraction was conducted using a CuKα radiation over the 20 range from 5° to 70°, a fourier transform infrared spectroscopy measurement with the wavenumbers ranging from 4000 to 400 cm⁻¹ is performed to investigate the reaction products. All powder samples for XRD and FTIR analysis were prepared by mechanical grinding and tested after 28d of curing. The microstructural characters of different mixes were investigated by a scanning electron microscope.

3. RESULTS AND DISCUSSIONS

3.1 Setting times

The setting times of alkali activated metakaolin pastes with the slag replacement up to 40% are shown in Table 3. It can be seen that as the slag content increases, both the initial and final setting times decrease significantly. The initial setting time of the neat metakaolin sample was 155 mins, while as the slag replacement reached 40%, the initial setting decreased nearly three times to 45 mins. Similar results happened to the final setting time. It seems that the remarkable reduction of setting times was caused by the calcium in the slag. The structure of metakaolin is dominated by silicate and aluminum oxide compounds, the Al-O and Si-O bonds in metakaolin are relatively difficult to be activated due to the nature of the original clay-type structure, thus the initial and final setting times were relatively long. On the other hand, because of the presence of calcium in the molecular structure, the glassy phase of slag is more disordered, thus it is more vulnerable to
breakdown under alkali activation. Finally, the fast release of calcium in solution lead to the reduced setting time.

<table>
<thead>
<tr>
<th>Slag replacement (%)</th>
<th>Initial setting (min)</th>
<th>Final setting (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>155</td>
<td>240</td>
</tr>
<tr>
<td>90</td>
<td>105</td>
<td>215</td>
</tr>
<tr>
<td>80</td>
<td>70</td>
<td>165</td>
</tr>
<tr>
<td>70</td>
<td>55</td>
<td>130</td>
</tr>
<tr>
<td>60</td>
<td>45</td>
<td>115</td>
</tr>
</tbody>
</table>

Table 3: Setting times of metakaolin-slag blends

3.2 Strength development

Figure 1 presents the 7d and 28d compressive strength results of alkali activated metakaolin with slag replacement from 0% to 40%, which shows that as the slag content increases, the compressive strength firstly increases and reaches the highest value at the slag content of 20%, then leads to a reduction of the compressive strength. All samples have a 7d strength of more than 58 MPa and 28d strength of more than 60 MPa, and the highest strength reaches 72.5 MPa with 20% slag replacement by weight. Also, the strength development from 7d to 28d becomes more significant as the increase of slag content.

The increase in compressive strength when slag was added is attributed to the presence of the calcium in the slag and the accompanied formation of calcium enriched hydrated C-A-S-H type gel is widely accepted as an gel that characterized of high strength [16]. The formation of small amount of C-A-S-H gel helped to refine the microstructure of N-A-S-H gel (main hydration products of metakaolin) and generated more crosslinking within the reaction product [17], then resulted in an increase in compressive strength. The decreased strength when higher slag was added indicates that there may exist an ideal balance between the activator character and the raw materials.

Metakaolin and slag mixes that with the highest compressive strength results (80% metakaolin and 20% slag in this case) were chosen and incorporated with limestone powder to study its effect on compressive strength of alkali activated metakaolin and slag blends, and the results are shown in Fig. 2. All samples show similar compressive strength values after 7d curing except the one with 20% limestone content. The 28d compressive strength stably increases as the limestone content increases, the highest strength is obtained with a limestone content of 15%, the further higher content leads to a considerable reduction of compressive strength. This reveals that the addition of limestone powder has limited influence on the early strength but could slightly increase the strength at longer ages within the range of optimum content. Higher content of limestone powder may have detrimental effects on the structure of hydrated gels and then lead to the decrease of compressive strength.

Fig. 1: Compressive strength of metakaolin-slag mixes

3.3 FTIR spectra analysis

Figure 3 shows the fourier transform infrared spectra of alkali activated metakaolin paste samples after 28 days curing with slag contents of 0%, 20% and 40%. The main absorption bands are listed in Table 4. The major absorption band is around 1000 cm⁻¹, which is assigned to the asymmetric stretching vibration of the Si-O-T (T=Si or Al) bonds [18], a typical vibration band of N-A-S-H type gel from metakaolin. The presence of this band also indicates the polymerization process took place. With the incorporation of slag, the wavenumber of this band decreased. Generally, the reduction of wavenumber indicates the decrease of polymerization degree of the reaction products. Thus here it means the addition of slag into metakaolin reduced
the polymerization degree of the hydrated gels to some extent, therefore the structure of the hydrated N-A-S-H gel was changed into a lower polymerization one by the presence of calcium from slag. It proves that the addition of slag indeed has some influences on the final hydrated product of metakaolin. Other significant wavenumber changes also reveal the structure of reaction products has been altered by the addition of slag, such as the bending vibration of T(IV)-O at around 570 cm\(^{-1}\) disappeared as the slag was added and the T-O asymmetric stretching vibration bonds around 1400 cm\(^{-1}\) increased at the beginning and then disappeared as the increase of slag content.

![FTIR spectra of metakaolin-slag blends](image)

**Fig. 3: FTIR spectra of metakaolin-slag blends**

<table>
<thead>
<tr>
<th>Absorption bands</th>
<th>100 MK 0 GGBS</th>
<th>80 MK 20 GGBS</th>
<th>60 MK 40 GGBS</th>
</tr>
</thead>
<tbody>
<tr>
<td>-OH, H(_2)O</td>
<td>3445</td>
<td>3445</td>
<td>3445</td>
</tr>
<tr>
<td>H(_2)O</td>
<td>1645</td>
<td>1647</td>
<td>1647</td>
</tr>
<tr>
<td>O-C-O</td>
<td>--</td>
<td>--</td>
<td>1471</td>
</tr>
<tr>
<td>T-O</td>
<td>1375</td>
<td>1417</td>
<td>--</td>
</tr>
<tr>
<td>Si-O</td>
<td>1015</td>
<td>1001</td>
<td>1000</td>
</tr>
<tr>
<td>Al(IV)-O</td>
<td>702</td>
<td>701</td>
<td>700</td>
</tr>
<tr>
<td>T(IV)-O</td>
<td>566</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Si-O</td>
<td>444</td>
<td>448</td>
<td>448</td>
</tr>
</tbody>
</table>

**Table 4: Corresponding bands of metakaolin-slag blends**

The absorption bands at around 3450 cm\(^{-1}\) and 1650 cm\(^{-1}\) are attributed to the stretching and deformation vibrations of O–H and H–O–H groups, which present the water bonded in the reaction products. No significant changes in the bands at around 700 cm\(^{-1}\), which represent the vibration of four coordinated aluminum mainly from metakaolin, showing that the addition of slag has limited influence on the structure of AlO\(_4\) in the final reaction products. The absorption bands around 450 cm\(^{-1}\) represent the internal deformation vibrations of Si-O bonds [19], which is mainly influenced by the nature of raw materials and thus no remarkable changes take place. The absorption band around 1470 cm\(^{-1}\) appeared only in the samples with large slag content, it represents the asymmetric stretching mode of the O–C–O groups [20], which means the carbonation has taken place to some extent, it may also indicate that samples with higher slag content are more vulnerable to carbonation, but still more study are needed to confirm this conclusion. The infrared spectra of samples containing limestone powder are not shown because there are no significant changes in the major absorption bands when limestone powder was added, which indicates that the incorporation of limestone powder has no influence on the reaction products and it only works as a non-reactive filler in alkali activated system.

### 3.4 XRD analysis

The XRD patterns of alkali activated metakaolin pastes with 0%, 20% and 40% slag replacement are shown in Figure 4. The main reaction products of all mixes are amorphous phases except small amount of crystalline phases that obtained from the raw materials such as quartz and illite. The major hump area for neat metakaolin samples are from 20° to 30°, which is within the typical range of alkali activated metakaolin and reveals the formation of hydrated N-A-S-H type gels from metakaolin. The amorphous phases with humps ranging from 20° to 30° can be regarded as the result of the geopolymerization progress of metakaolin.

![XRD spectra of metakaolin-slag blends](image)

**Fig. 4: XRD spectra of metakaolin-slag blends**

As the slag content increases, the main hump area gradually moves to higher degree, which is in coincidence with the FTIR results and reveals the phase changes in reaction products caused by slag addition. As the slag content continues to increase, the intensity peaks at around 30° appears, which represents the formation of an C-S-H type gel from alkali activated slag. The intensity peaks that represent C-S-H gel
becomes more significant in samples with higher slag content. Combined with the results from FTIR, the available calcium in the solution caused by the addition of slag seems to have significant influences on the formation of reaction products other than N-A-S-H and changing the original N-A-S-H gel structure to some extent such as a lower polymerization degree and changing the major hump areas. The XRD patterns with limestone powder incorporation are not shown because as the limestone content increases in the metakaolin and slag mixes, no changes take place in the original hump areas and intensity peaks, the intensity peaks of limestone still remain unchanged, which combined with the FTIR result reveals again that the limestone powder seems to work as only a filler in the alkali activated system.

3.5 SEM analysis
The SEM images of the 28d alkali activated metakaolin paste samples containing 0% and 20% slag are shown in Figure 5. The left and right column represents the neat metakaolin sample and samples containing 20% slag, respectively.

![SEM images of metakaolin with/without slag addition](image.png)

It can be seen that even though neat alkali activated metakaolin samples have acceptable 28d compressive strength of around 60 MPa, some remarkable pores are still remaining under high magnifications. The reaction products exhibit layered structure in short ranges and disordered in longer ranges. The failure of samples in compressive strength test also showed layered structure, it seems that the nature of clay-type structure of metakaolin should be responsible for the failure. When a suitable content of slag was added, pores between the layers of alkali activated metakaolin were condensed by the hydrated slags. Also, the addition of slag provides calcium and leads to the formation of C-S-H type gel, which is characterized by a high strength, linked together with the original N-A-S-H gel, then contributes to the refinement of microstructure and improves the compressive strength.

4. CONCLUSIONS
The incorporation of slag significantly reduced the initial and final setting times of alkali activated metakaolin, and the setting times continually decreased as the increase of the slag content. The compressive strength firstly increased and then decreased as the slag content increased, and the optimum slag content in this case is 20%, similar results were obtained in limestone powder addition into metakaolin-slag blends and the optimum limestone powder content is around 15%. Both the XRD and FTIR results show significant changes in reaction products as a result of slag incorporation. Both N-A-S-H from metakaolin and C-S-H type gel from slag were detected, the addition of slag induced some changes of the original N-A-S-H gel structure such as the polymerization degree and the disappearance of some typical vibration bonds. Limestone powder seemed to work as an non-reactive filler in the alkali activated system since no significant changes were detected in both XRD and FTIR analysis. The microstructure analysis showed that the original alkali activated metakaolin paste was modified by the addition of slag, resulting in a denser matrix and resulted in an improved compressive strength.

REFERENCES


**BIOGRAPHIES:**

Xu Gao, has received his Master’s degree from Wuhan University of Technology, China. Currently working as a Ph.D. student in the Department of the Built Environment, Eindhoven University of Technology, the Netherlands. His research interests include the modification, modeling and application of alkali activated materials.

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