Research Article

Fly Ash Particle Size Effect on Pore Structure and Strength of Fly Ash Foamed Geopolymer

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The aim of this paper is to study the particle size effect of fly ash (FLA) on pore structure and strength of Fly Ash Foamed Geopolymer (FAFG). Information on the macro-pores such as macro-pore size and distribution of FAFG is captured through binarization processing. Porosity and compressive strength of FAFG are respectively tested by Archimedes density test method and uniaxial compressive strength test method. It can be concluded that the FLA particle size has an effect on the pore structure and strength of FAFG. More specifically, the effect of FLA particle size shows itself macroscopically on the quantity of middle and large macro-pores and the uniformity of macro-pores distribution, and microscopically on the quantity of micro-holes and cracks and calcium silicate (C-S-H) quantity at the early stage of FAFG mixture. All of the properties of FAFG follow some kind of changing rule except at the turning point when FLA particle is of 0.125~0.25 mm in size. To explain clearly the root cause of FLA particle size effect on FAFG, SEM, and XRD are employed to explore the microstructure of FAFG and the component of FLA. It turns out to be the amorphous phase SiO2 content in FLA of different particle sizes which could determine the reaction extent of FAFG mixture.

1. Introduction

Coal combustion or the co-firing of biomass and coal has encountered certain operational problems, such as ash deposits, slugging, fouling and corrosion, reducing the thermal efficiency of boilers [1, 2], and resulted in problems in solid waste disposal, which is a long-stand research hotspot. As the largest amount of solid waste produced by rapid oxidation of coal powder during thermal power generation [3], Fly ash (FLA) pollutes the local environment and increases the cost of the treatment for coal mine. On the other hand, because of its advantages like good thermal insulation, light weight, and low cost [4, 5], FLA has been widely used in some architectural fields like nonbearing wall construction, bathroom backfilling and as functional materials such as thermal insulation material, sound-insulation material, and so on [6, 7]. Apart from being used in the traditional building fields, FLA has been applied in different areas as a good raw material for the preparation of geopolymer [8–12]. For example, to prove the possibility of in-situ resource utilization of FLA, there are a good many scholars having done plenty of basic studies on this aspect [13–15], among which there is one proposing that fly ash foamed geopolymer (FAFG) as backfilling material can be an excellent combination of fly ash consumption and coal mine fire prevention [16, 17]. The complex underground environment demands highly for some specific properties of FAFG, mainly focusing on its pore structure and strength.

As for the preparation of FAFG, the most significant point is to stabilize the suitable pore structure and generate appropriate strength, which are affected by different factors such as temperature, components, active materials, and so on [18–21]. However, there are few studies on FLA particle size effect on the properties of FAFG and the root cause for the effect has not been made clear yet. But plenty of FAFG preparation experiments have made it evident that the different FLA particle sizes do have some influence on the property of FAFG. The main objectives of this research are to observe the FLA particle size effect on the properties of FAFG and to investigate the root cause for the effect.

In terms of the way of foam generation, FAFG can be divided into two categories: mechanical foamed-FAFG [22,
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and chemical foamed-FAFG [24, 25], which are respectively produced by special foaming machine with some kind of foaming agent and by finely divided aluminum or hydrogen peroxide [26, 27]. In this research, FAFG was made of FLA of different particle sizes and other raw materials in the form of chemical foaming with hydrogen peroxide. With the high-quality pictures taken by SLR camera, macro-pore structure characteristics were measured through binarization processing analysis. XRD and SEM were employed to determine the components of FLA and to observe the micro-pore structure of FAFG respectively. The uniaxial compressive strength was tested by the universal testing machine. The novelty of my research lies firstly in the investigation of the FLA particle size effect on the properties of FAFG, and secondly in the finding of the root case for this effect. The components of FLA changed with the particle size, which resulted in different performances and different degrees of strength of macro- and micro-scale pore structure accordingly. It turns out that the crystal/amorphous ratio might be the root cause for the effect.

2. Materials and Methods

2.1. Materials. As is shown in Table 1, fly ash of six size ranges were obtained through using vibrating screen, and they are respectively −0.045 mm, 0.045~0.075 mm, 0.075~0.125 mm, 0.125~0.25 mm, 0.25~0.5 mm, and +0.5 mm. Pictures of fly ash through particle size classification from Dananhu coal mine no.1 are shown in Figure 1. Except for the max size (+0.5 mm) and the min size (−0.045 mm), the fly ash graded by size was used as the raw material for FAFG. Detailed information on other raw materials and additives are listed in Table 2.

2.2. Experimental Procedure. Based on the experience from a trial and error approach in the early stage, the mixed proportion of FAFG was obtained, which is shown in Table 3. All of the FAFG mixtures were prepared according to the uniform ratio, of which the only difference was the grading size of the fly ash. To obtain the initial samples, the mixtures were cast in steel cubic molds (100 mm), foaming naturally because of the inside reaction of hydrogen peroxide. The foaming period lasted about a quarter of an hour, and then the samples were left at 100°C in a thermostatic heater for 2 hours; the consolidation was completed in a room with temperature of 22–24°C till the day for testing. The central cubes (70.7 mm) incised from the untreated samples (100 mm) were prepared for the microstructure observation, porosity measurement

<table>
<thead>
<tr>
<th>Size/(mm)</th>
<th>Mass fraction/(%)</th>
<th>Add up/(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>−0.045</td>
<td>47.03</td>
<td>47.03</td>
</tr>
<tr>
<td>0.045~0.075</td>
<td>10.60</td>
<td>57.63</td>
</tr>
<tr>
<td>0.075~0.125</td>
<td>20.02</td>
<td>77.65</td>
</tr>
<tr>
<td>0.125~0.25</td>
<td>15.00</td>
<td>92.65</td>
</tr>
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<td>0.25~0.5</td>
<td>6.60</td>
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</tr>
<tr>
<td>+0.5</td>
<td>0.75</td>
<td>100</td>
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</tbody>
</table>

Figure 1: Pictures of fly ash through particle size classification from Dananhu coal mine no. 1.
and strength testing. The central cubes incised are shown in Figure 2.

2.3. Test Method. The X-ray diffraction diffractometer (XRD Bruker D8 Advance) was employed to determine the crystalline phase of FLA. The pore morphology pictures of FAFG samples were obtained through the digital single lens reflex (Nikon D5500). With the help of open source software (Image-Pro Plus), the pore size distribution of FAFG samples was analyzed through employing image binarization processing for the pore morphology pictures. The strength testing of FAFG samples was carried out using the universal testing machine (WDW-300). The microstructure of FAFG samples was observed through scanning electron microscope (SEM, ZEISS SIGMA-300). The bulk density of FAFG samples was measured using the standard Archimedes method with distilled water. The total porosity was calculated according to the following Equation (1):

\[ P = \left( \frac{\rho_b - \rho}{\rho_b} \right) \times 100\% \tag{1} \]

where \( \rho \) and \( \rho_b \) represents the bulk density and powder density, respectively.

3. Result and Discussion

3.1. Pore Structure Analysis. To some extent, the cross section of the incised central cubes could reflect the real inner pore structure of FAFG, and therefore, the Nikon D5500 was used to obtain a high resolution image of the cube side. Pore structure data was analyzed after binarization processing \[28\] using the open source software "Image Pro Plus". Raw picture and picture after binarization processing are respectively shown in Figure 3 where pore structure and its distribution can be clearly seen.

Diameter is one of the main characteristic parameters of pore structure, but there are few completely suitable classification standards for such porous materials as aerated or foamed geopolymer (concrete) of largescale (μm-mm). Most of classification standards or methods existing \[29\] are based on the standard of IUPAC in which there is only

![Figure 2: Incised central cubes of FAFG.](image-url)
The porosity ranges from 66.37% to 73.15%, leading to FAFG density range from 300 to 500 kg/m³. As can be seen in Figure 5, with the enlargement of FLA particle size, the compressive strength (7d) decreases and the porosity increases. FAFGs3 is still a turning point where the compressive strength arrives at the bottom and the porosity reaches the peak, and the porosity stops to rise and begins to decrease while the compressive strength shows the converse change. There are many studies on the relationship between porosity and strength, and almost all of their conclusions point out that large porosity means bad strength performance [21, 32]. Two main factors could be used to explain the connection between strength and pore structure; one is the free volume of pores (porosity) and the other is the single pore size (heterogeneously distributed large pores). Both of them are related to FLA particle size. With the enlargement of FLA particle size, the quantity of heterogeneously distributed large macro-pores and the porosity of FAFG increases first and then decreases, as can be respectively seen in Figures 2 and 5. Because of the negative impact of these two factors on strength of FAFG, the compressive strength of FAFG shows a variation trend opposite to that of the two factors.

3.2. Strength Development. Figure 5 shows the result of compressive strength test and porosity determination for FAFG. The early age strength ranges from 0.097 MPa to 0.3241 MPa (percentage difference: 70%), which means the strength difference of FAFG of different FLA particle sizes is large. The porosity ranges from 66.37% to 73.15%, leading to FAFG density range from 300 to 500 kg/m³. As can be seen in Figure 5, with the enlargement of FLA particle size, the compressive strength (7d) decreases and the porosity increases. FAFGs3 is still a turning point where the compressive strength arrives at the bottom and the porosity reaches the peak, and the porosity stops to rise and begins to decrease while the compressive strength shows the converse change. There are many studies on the relationship between porosity and strength, and almost all of their conclusions point out that large porosity means bad strength performance [21, 32]. Two main factors could be used to explain the connection between strength and pore structure; one is the free volume of pores (porosity) and the other is the single pore size (heterogeneously distributed large pores). Both of them are related to FLA particle size. With the enlargement of FLA particle size, the quantity of heterogeneously distributed large macro-pores and the porosity of FAFG increases first and then decreases, as can be respectively seen in Figures 2 and 5. Because of the negative impact of these two factors on strength of FAFG, the compressive strength of FAFG shows a variation trend opposite to that of the two factors.

3.3. XRD Analysis. "Fly ash effect" refers to a series of effects including morphological effect, micro-aggregate effect and active effect. With the enlargement of FLA particle size, the morphological effect and micro-aggregate effect weaken obviously, which might result in some bad impact on FAFG such as more heterogeneous pore distribution, larger porosity and worse strength performance. FAFGs1, FAFGs2, and

![Figure 3: Cross section picture before (a) and after (b) binarization processing.](image)

<table>
<thead>
<tr>
<th>Classification</th>
<th>Macro-</th>
<th>Meso-</th>
<th>Micro-</th>
<th>Supermicro-</th>
<th>Ultramicro-</th>
<th>Submicro-</th>
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<td>0.7−2</td>
<td>&lt;0.7</td>
<td>&lt;0.4</td>
</tr>
<tr>
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<td>200÷400÷3÷3.2</td>
<td>&lt;1.2÷1.4</td>
<td>3÷3.2÷d÷1.2÷1.4</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Cheremskoj</td>
<td>≥2000</td>
<td>—</td>
<td>2000÷d÷200</td>
<td>—</td>
<td>&lt;2÷4</td>
<td>&lt;200</td>
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<tr>
<td>Kodikara</td>
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<td>—</td>
<td>10³÷3×10⁴</td>
<td>25×10³</td>
<td>&lt;3÷4</td>
<td>—</td>
</tr>
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</table>
**Figure 4:** Pore size distribution of FAFG.

**Figure 5:** Curve of porosity and compressive strength to size grading.
SiO₂ content. As can be seen in Table 5, the peak diffraction intensity of SiO₂ first increases and then decreases with the enlargement of FLA particle size. Figure 7 represents the planar model of molecular structure of crystal phase SiO₂ and amorphous phase SiO₂. As can be seen in the figure, the molecular structure of crystal phase SiO₂ was much more compact and orderly than that of amorphous phase SiO₂, which means Si-O bonds of the former were more firmly structured than those of the latter, and that the latter was easier to break by alkaline activator in FAFG mixture. Tennakoon reasoned that amorphous phase SiO₂ was the key to produce reactive SiO₂ in geopolymer formation [35].

To figure out why there is at FAFGs3 always a turning point which leads to the appearance of all these strange phenomena. XRD analysis was conducted on fly ash of different particle sizes. Figure 6 shows the XRD analysis result. As can be seen in the Figure 6, Quartz and Mullite are the main components of FLA. A big “steamed bread peak” appears at the degree of 20~30°; meanwhile, as FLA particle size enlarges, the “steamed bread peak” becomes bigger firstly and smaller then. Size of the “steamed bread peak” reflects the amorphous phase mineral content. The bigger the size of “steamed bread peak” is, the more amorphous phase SiO₂ there is; Taylor and Matulis pointed out that peak heights can proportionally characterize the peak areas when it comes to well-defined peaks without significant peak width [33]. Liu compared the silica content of aluminosilicate with different sources by comparing their peak intensity [34]. Diffraction intensity can reflect the crystal phase mineral content, and therefore, peak diffraction intensity of SiO₂ is conducive to acquisition of crystal phase FAFGs3 accord well with this changing rule as expected, yet FAFGs4 does not follow this changing rule.
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Crystal phase SiO$_2$

Amorphous phase SiO$_2$

**Figure 7:** Planar model of molecular structure of SiO$_2$.

hydrated reaction of FAFG, there is a negative impact on the pore formation and distribution. Molecular structure of amorphous phase SiO$_2$ can be easily broken in the alkaline environment, so that the activated SiO$_2$ could restructure with some other materials in the FAFG mixture; for example, it can react with calcium hydroxide produced by hydration reaction of calcium oxide. This contributes to the uniform spatial and size distribution of pores.

3.4. **SEM Analysis.** Figure 8 shows the SEM (test condition: ETH = 5 kV, Signal A = SE2, Mag = 5.00 KX) figure of FAFG. The original appearance of raw fly ash is a smooth sphere, but many flocs appear on its surface after raw fly ash reacts in the FAFG mixture. After being activated by the alkaline activator, amorphous phase SiO$_2$ reacts with the Ca$^{2+}$ provided by CC and CO$_2$, leading to the formation of a large amount of C-S-H (Calcium silicate hydrate) on the surface of FLA particle. There is often a considerable quantity of Cubilose-shaped or vermicular-shaped C-S-H existing in the fly ash-lime-water hardenite. From Figure 8, it can be clearly seen that the quantity of flocs first decreases and then increases with the enlargement of FLA particle size, whose turning point is still FAFG$. The C-S-H is the main product of FAFG, and the quantity of C-S-H could represents the extent of reaction. Therefore, the reaction extent of FAFG mixture changes with the enlargement of FLA particle size. The change of amorphous phase SiO$_2$ with the FLA particle size is the root cause of the change of reaction extent, and strength of FAFG becomes stronger due to the formation of C-S-H in the early stage.

Microscale cracks and holes can be observed with the help of SEM. Figure 9 shows SEM (test condition: ETH = 5 kV, Signal A = InLens, Mag = 5.00 KX) figure of FAFG. As can be seen in
Results in this study show that the FLA particle size has an effect on the pore structure and strength of FAFG, and the root cause for the effect is the content difference of active substance (amorphous phase SiO$_2$), and the difference is triggered by changes in particle size after sieving. XRD results show that amorphous phase SiO$_2$ content of FLA decreases first and then increases with the enlargement of FLA particle size with a turning point as 0.125~0.25 mm, which means FLA of particle size of 0.125~0.25 mm has the least content of active substance. The content of active substance could determine the reaction extent of FAFG mixture. In short, a fuller reaction of the mixture would produce more C-S-H and less micro-holes and cracks, bringing a better strength performance and more uniform pores of FAFG.

4. Conclusions

Generally speaking, with the enlargement of FLA particle size, the UCS of FAFGs decreases first and then increases, and the porosity and the quantity of middle and large macro-pores increases first and then decreases. Besides, 0.125~0.25 mm is the turning point for both of them. Uniformity of pore distribution gets worse as the FLA particle size enlarges, but it turns better when it comes to 0.25~0.5 mm. According to the micro analysis, with the enlargement of FLA particle size, the content of C-S-H (reaction product of FAFG mixture) decreases first and then increases, and the quantity of micro-pores and cracks increases first and then decrease and their turning points are also 0.125~0.25 mm.

Results in this study show that the FLA particle size has an effect on the pore structure and strength of FAFG, and the root cause for the effect is the content difference of active substance (amorphous phase SiO$_2$), and the difference is triggered by changes in particle size after sieving. XRD results show that amorphous phase SiO$_2$ content of FLA decreases first and then increases with the enlargement of FLA particle size with a turning point as 0.125~0.25 mm, which means FLA of particle size of 0.125~0.25 mm has the least content of active substance. The content of active substance could determine the reaction extent of FAFG mixture. In short, a fuller reaction of the mixture would produce more C-S-H and less micro-holes and cracks, bringing a better strength performance and more uniform pores of FAFG.

Data Availability
No data were used to support this study.

Conflicts of Interest
The authors declare that they have no conflicts of interest.

Acknowledgments
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Supplementary Materials

The graphical abstract introduces preparation process of (fly ash foamed geopolymer) FAFG and some characteristcs testing for FAFG. The whole preparation process includes dry raw materials mixing, alkali-activator adding, foaming and curing. Then the FAFG samples will be incised into cubes (70.7 mm). Finally characteristics of FAFG will be tested with relevant instruments and methods. (Supplementary Materials)

References


