Grinding Method for Increasing Specific Surface Area of Fluidized Bed Fly Ash

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ABSTRACT

In this study, fly ash of a fluidized bed boiler produced in a power plant was stabilized by hydration and carbonation reaction. Then, each raw material was pulverized by two kinds of grinding equipment (Planetary mills and pot mills); the degree of grinding and the agglomeration behavior were observed. It was found that there were changes of specific surface area and particle size distribution according to grinding time. The surface of the raw material was observed using an optical microscope. As a result, agglomerates of about 75 \( \mu \text{m} \) or more due to electrostatic phenomenon were formed as the grinding time became longer; it was confirmed that the crushing efficiency slightly increased with use of antistatic agent.

Key words: Grinding, Fluidized bed fly ash, Agglomeration, Hydration, Carbonation

1. Introduction

Construction waste is dominantly comprised of waste concrete, which has contributed to increasing environmental pollution due to illegal dumping as the previous method of simple landfills has begun to reach its limits. Not only that, recently growing interest in the lack of natural aggregates, conservation of national resources, and environmental conservation has brought recycled aggregates, derived from the reprocessing of waste concrete, to the forefront from an environmental and economic perspective. However, the usage of recycled aggregates is limited because of concerns regarding concrete strength and durability degradation, along with the occurrence of alkali-aggregate reactions due to cement paste existing in the recycled aggregate. In order to resolve such problems, the use of fly ash as admixture is actively being researched.\(^1\)

The fly ash produced in coal-fired power plants is used as an admixture to enhance the properties of concrete, including improving the concrete durability as well as concrete fracture reduction and long-term strength enhancement by reducing the hydration heat. As such, fly ash has recently been used widely both domestically and internationally.\(^2\) Most construction sites still use ordinary Portland cement (OPC) and, as a result, the performance required for various structures is not attained and problems such as crack propagation and strength degradation have occurred due to excessive hydration heat from the use of large amounts of cement.\(^3\) In an effort to resolve these problems, active research and development using fine fly ash and blast-furnace slag as admixtures are being carried out.

The combustion process of using coal in a thermal power plant to produce energy includes the pulverized coal combustion method and the circulating fluidized bed combustion method. The circulating fluidized bed combustion method has recently been a center of interest as a power generation technology because the combustion method exhibits excellent diversification of applied fuels, and economic feasibility and environmental friendliness of the facilities. Also, since the conventional desulfurization system is very costly, a fluidized bed boiler with a long bituminous coal combustion time can be used to perform desulfurization within the furnace and a circulating fluidized bed boiler is used for low combustion temperature and NOx generation minimization. However, the circulating fluidized bed combustion method carries out desulfurization in the furnace, so a large amount of CaO compounds is contained in the coal ash and the SiO\(_2\) content does not conform to the KS standard (KS L5405). When it is used as a concrete admixture, the free-CaO component that does not participate in the concrete hydration reaction induces critical problems regarding the concrete durability, expansion, weathering, and cracks, resulting in everything being landfilled and not being recycled.\(^4\)

In domestic and international studies, fly ash is mixed and then ground before usage. Especially, while there are a lot of cenospheres of spherical particle form in conventional pulverized coal ash so that fluidity is high, fluidized bed coal ash has almost no cenospheres and cannot satisfy the fluidity of KS L 5405; thus, the specific surface area (Blaine) has to be increased through pulverization. Fly ash is used by categorizing it by type according to fineness based on KS L 5405.\(^5\) Chindaprasirt et al.\(^6\) studied the overall strength...
improvement effect when increasing the basic Blaine value by using fly ash mixture as a concrete admixture. Additionally, the fineness of OPC was varied to investigate the hydration reaction properties and compressive strength. In order to use fly ash in concrete, quality criteria like SiO$_2$ content, fineness, density, moisture, ignition loss, flow ratio, and activity factor are selected for maintenance of KS L 5405. Among these KS quality criteria, fly ash types including type 1 (4,500 cm$^2$/g and above), type 2 (3,000 cm$^2$/g and above), type 3 (2,500 cm$^2$/g and above), and type 4 (1,500 cm$^2$/g and above) are classified based on the fineness as measured using an air permeability apparatus. Fineness is a factor that determines the quality of fly ash and is considered the most important criterion that determines whether a material can be utilized as a concrete additive.

In this study, literature review showed that factors of fly ash that affect the strength, including the particle size distribution, specific surface area, chemical composition, and mineral composition impact the pozzolanic reaction rate. In order to recycle the fluidized bed coal ash as a concrete admixture, hydration and carbonation reactions were carried out to increase the Blaine value of the coal ash stabilized through hydration/carbonation of free CaO; this was followed by planetary and ball milling to investigate the pulverization effect according to the pulverization equipment measurement. The particle size distribution measurement, as well as the pulverization efficiency according to the particle size distribution variation during pulverization. Moreover, the pulverization efficiency with respect to time was studied by varying the pulverization duration. The agglomeration behavior due to static electricity of the fine powder raw material became a problem; so, in order to resolve the electrostatic agglomeration phenomenon between particles due to pulverization, the antistatic agent DEG was added to increase the pulverization efficiency of the fluidized bed coal ash and increase the Blaine value.

### Table 1. Chemical Composition of Various Types of Fly Ash

|                  | As-received Fly Ash (RFA) | Hydrated Fly Ash (HFA) | Carbonated Fly Ash (CFA) |
|------------------|---------------------------|------------------------|--------------------------|
| SiO$_2$          | 24.73                     | 24.37                  | 21.17                    |
| Al$_2$O$_3$      | 10.33                     | 10.48                  | 8.57                     |
| Fe$_2$O$_3$      | 4.95                      | 4.31                   | 4.36                     |
| CaO              | 38.41                     | 32.36                  | 31.15                    |
| MgO              | 4.54                      | 3.85                   | 3.72                     |
| Na$_2$O          | 4.03                      | 3.28                   | 2.64                     |
| K$_2$O           | 0.84                      | 0.87                   | 0.66                     |
| TiO$_2$          | 0.51                      | 0.48                   | 0.44                     |
| P$_2$O$_5$       | 0.13                      | 0.12                   | 0.11                     |
| C                | 0.32                      | 0.85                   | 4.69                     |
| S                | 10.15                     | 9.05                   | 10.5                     |
| Ig. Loss         | 1.05                      | 9.98                   | 11.99                    |
| Total            | 100                       | 100                    | 100                      |

#### 2. Experimental Procedure

##### 2.1. Raw Material Analysis

RFA produced in the circulating fluidized bed combustion boiler was used as the raw material in this study. The chemical composition of the raw material was analyzed using XRF (ZSR-100e, Rigaku, Japan); the results are shown in Table 1. The CaO content and ignition loss of the fluidized bed coal ash were 38.35% and 1.05%, respectively, where the ignition loss was very small. This result signified that a large amount of Ca exists in CaO form. Thus, there was a need to stabilize the free CaO for the stable use of this raw material as the concrete admixture; as such, stabilization was carried out through hydration and carbonation reactions. The hydrated fly ash was referred to as HFA and the carbonated fly ash was referred to as CFA. The ignition loss of the stabilized fly ash increased significantly at 9.98.
wt% and 11.99 wt% for HFA and CFA, respectively. The increase in the ignition loss due to the dehydration and decarbonation reactions at high temperature during stabilization is a natural phenomenon. Fig. 1 shows the TG/DTA measurement results. For HFA, most of the mass loss occurred at around 400 ~ 500°C; for the CFA, most of the mass loss occurred at around 700 ~ 800°C. This result shows that the temperatures of crystallization water and decarbonation are different. Mass loss was greater for CFA because, stoichiometrically, the removal of CO₂ results in greater mass loss than the removal of H₂O.

2.2. Raw Material Pulverization and Analysis

The pulverization of the raw materials was performed using a pot mill (PM-002, MonotaRO, Japan) and a planetary mill (PM 100, Retsch, Germany). The pot mill was a ball mill that is commonly used in ceramic processing and involves placing the raw material with a ball in a pot and pulverizing the material by uniaxial rotation. The planetary mill is a pulverization apparatus that is more advanced than the pot mill; in the planetary mill, a small pot is installed in a large, rotating cylindrical disk. The large cylindrical disk rotates while the pot also rotates on its own and this rotation method induces greater ball motion than that of the pot mill. Thus, the planetary mill has a shorter pulverization time than the pot mill. Here, porcelain was used as the material for the jar and media of the pot mill, while alumina was used for the planetary mill. By varying the pulverization time for each piece of equipment depending on the efficiency of its pulverization apparatus, particle size distribution and Blaine value were observed for the pot mill and planetary mill. For the pot mill, pulverization was carried out for 20, 40, and 60 minutes, while pulverization was carried out for 5, 10, 15, and 20 minutes for the planetary mill.

Particle size analysis (Mastersizer 3000, Malvern, U.K) was conducted to analyze the particle size distribution and Blaine value. The analysis results were compared with the particle size distribution measurements obtained using a sieve. For the Blaine measurement, an air permeability apparatus (Auto Blaine tester, Hanil Lab. Tech, Korea) was used in accordance with KS L5106¹⁰ to compare the size of the raw material; a Camscope (Dr. Camscope, Mikey, Korea) was used to observe the raw material and determine its agglomeration state.

In order to observe the electrostatic agglomeration phenomenon, the antistatic agent diethylene glycol (DEG) was used, added at a level of 0.01%. Pulverization was carried out using the planetary mill in the same manner as explained earlier.

3. Results and Discussion

3.1. Particle Size Distribution of the Starting Material

Figure 2 shows the particle size distributions of the RFA, HFA, and CFA. The RFA had an average particle size of approximately 15 µm, while the average particle sizes of the HFA and CFA were approximately 9 µm and 5 µm, respectively. These results show that the free CaO of the RFA reacted with Ca(OH)₂ for HFA and RFA divided due to the expansion stress to become finer particles. For CFA, there was further expansion as Ca(OH)₂ reacted with CaCO₃, so that additional division occurred to result in fine particles.

Table 2 shows the Blaine values and sieve particle size distributions for the starting materials. The Blaine value of RFA was found to be 1900 cm²/g, falling short of 3000 cm²/g. For HFA and CFA, although it was expected that they would have high specific surface areas as their average particle sizes were smaller than that of RFA, their Blaine values were not significantly different from that of RFA at 2100 cm²/g and 2200 cm²/g, respectively.

3.2. Particle Size Analysis

Table 2 shows the particle size distributions analyzed through sieving of RFA, HFA, and CFA. Compared to RFA, HFA and CFA had fewer particles below 45 µm, while there were more particles of 45 ~ 106 µm. This result contrasted with the result of the particle size distribution measurement shown in Fig. 2. Generally, as the agglomeration of fine particles is reflected, measurement through sieving results in more larger particles being measured compared to
PSA analysis, which is an instrumental analysis. Thus, this result was determined to be due to the agglomeration of particles, and the agglomeration phenomenon occurred for CFA with the smallest average particle size, so that the increase in particles of 45 ~ 75 µm was largest. For HFA, there were increases in the 45 ~ 75 µm and 75 ~ 106 µm intervals. This result was thought to be due to the agglomeration of fine particles during drying in the hydration and carbonation processes; this phenomenon causes the particle size to increase more from agglomeration the finer the raw material is. So, this result revealed that an increase in the specific surface through pulverization was necessary for usage of this material as a concrete admixture.

3.3. Pot Mill Pulverization

In Fig. 3, it is difficult to observe significant changes in the specific surface area for the pulverization using the pot mill up to about 60 minutes. This result was thought to be due to the pulverization limit of the pot mill. For the pulverization time of 20 minutes, an increase of around 1000 cm²/g was observed for HFA when compared to before pulverization, while an increase of around 2000 cm²/g was observed for CFA. However, the change in specific surface area was not significant even when the pulverization time was extended to 60 minutes.

As can be observed in the particle size distribution shown in Fig. 4, the 50 µm large particles decreased for HFA compared to F/A, while the particles of about 20 µm increased as the pulverization progressed. For CFA, the particle size appeared to increase for the interval of around 10 µm; however, for both cases, there were no changes in the particle size below 1 µm, so it was determined that there was no impact on the specific surface area variation.

For the particle size distribution when using the pot mill for the same pulverization time, CFA exhibited an overall fine result compared to HFA and CFA had a higher efficiency for the same pulverization time. Also, looking at the above graph, the 20-minute pulverization case for CFA resulted in an observation of a higher ratio of fine particles compared to the pulverization time of 60 minutes. This result was thought to be due to the extended pulverization time inducing electrostatic agglomeration phenomenon of the pulverized particles and resulting in an increased size. When materials were pulverized through the pot mill method, increase of the specific surface area with respect to time was not apparent. Thus, additional pulverization experiments were conducted using the more advanced planetary mill method.

3.4. Planetary Mill Pulverization

Figure 5 shows the specific surface area measurement of the specimen pulverized using the planetary mill. Since the pulverization efficiency of the planetary mill is excellent compared to that of the pot mill, changes in the specific surface area of the powder were investigated by making the pulverization time relatively short. The specific surface area was measured and found to be around 4000 ~ 5000 for CFA and 3500 ~ 4000 for HFA. These values tended to increase

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**Fig. 3.** Change in specific area of HFA and CFA pulverized by pot mill with time.

**Fig. 4.** Particle size distribution of various kinds of fluidized bed fly ash.
gradually with time. Also, the specific surface area was found to be relatively higher than that of the specimens pulverized using the pot mill.

Figure 6 shows the results of the sieved pulverized raw materials. The measurement results reveal that, as the pulverization time increased, although there was an interval in which the overall particle size increased due to particle agglomeration, no decreases in the specific surface area were observed in the particle size distribution data.

Pulverization of the raw material at 5-minute intervals using the planetary mill resulted in an increase in the amount fine particles as the time increased and the refinement of the particles from pulverization continued. However, agglomeration of the raw material occurred in a specific interval, and this was thought to be due to the electrostatic agglomeration phenomenon of the raw material. In the case of HFA, the amount of aggregates in the 75 \( \mu \text{m} \) interval increased as the pulverization time increased. In the case of CFA, the 45 \( \mu \text{m} \) particle size distribution decreased as the pulverization time increased, while the particles above 75 \( \mu \text{m} \) showed an increasing trend due to agglomeration. Due to the high specific surface area, it was difficult to accurately observe the particle size distribution through the
sieve, despite the increase in fine particle content as the pulverization time increased. Despite the particle size being sufficient to pass through the sieve, the electrostatic agglomeration between particles caused the amount of specimen that could not pass through the sieve to increase, and so naturally the ratio of large particles increased. The agglomeration mainly occurred at the 75 µm interval, so it was difficult to observe pulverization efficiency enhancement using the sieve method.

Figure 7 provides an image of the particles for each specimen, taken through an optical microscope. The observations show that carbonation particles were larger, which was thought to be due to the HFA particle agglomeration being weaker and leading to a rougher surface than was the case for CFA; as such, it was thought that in actuality CFA particles were smaller, making agglomeration occur more easily. In this way, the electrostatic agglomeration phenomenon of the particles occurred as the pulverization time increased; thus, the particle pulverization efficiency decreased.

Growth due to particle agglomeration did not exceed 75°C because when this size is exceeded, the agglomeration is pulverized again by the external force. Comparison of pulverization results for the pot mill and the planetary mill showed that the pulverization efficiency was higher for the planetary mill; also, for both pulverization methods, the pulverization efficiency decreased due to the electrostatic agglomeration phenomenon of particles when pulverization time increased.

3.5. Antistatic Agent

In the earlier experiment, it was found that the pulverization efficiency decreased because of particle agglomeration due to static electricity. So, an antistatic agent was added to prevent electrostatic agglomeration and the specific surface area was measured (Fig. 8). For both HFA and CFA, the specific surface area was found to significantly increase when the antistatic agent was used compared to when it was not used. The increase was greater for CFA and this was due to CFA being finer than HFA. Under the same conditions, the effect of the antistatic agent was more pronounced for the CFA specimen. Compared to CFA particles, HFA particles were larger and agglomeration occurred less; hence a greater effect was observed. Under the same conditions, specific surface area values increased by about 30% and 40% for HFA and CFA, respectively. For most intervals, the specific surface area value of the CFA raw material was higher than that of the HFA raw material and it was determined that the pulverization efficiency was higher for the CFA raw material than for that of HFA.

Figures 9 and 10 show the sieve experiment results for the specimen with the antistatic agent added. When the antistatic agent was used, the particle size distribution according to the pulverization time showed that when the same pulverization time was implemented, the antistatic effect for the agglomerated particles, compared to the case in which the antistatic agent was not added, was similar in that agglomeration occurred at 75 µm for HFA; however, weight decreased from approximately 60% to 45%, amount of fine particles increased, and the Blaine value accordingly increased. For CFA, large aggregates of 75 µm decreased and small aggregates of 45 µm increased slightly.
4. Conclusions

The pulverization efficiency for various conditions and types of pulverization equipment and processing state of raw materials was observed using circulating fluidized bed coal ash; the following conclusions were obtained.

1) The planetary mill showed a higher pulverization efficiency than that of the pot mill and the particle size distribution of the planetary mill was more uniform.

2) As the pulverization time increased, agglomeration between particles tended to degrade the pulverization efficiency for the particle size of approximately 75 µm.

3) The particle size distribution and specific surface area measurement results showed that when the electrostatic agglomeration phenomenon was excluded, CFA had the highest specific surface area, followed by HFA and RFA. Also, the agglomeration phenomenon between particles appeared to increase as the particles became finer.

4) When the antistatic agent DEG was used to overcome the particle agglomeration phenomenon, an increase in the specific surface area of about 30 ~ 40% was observed and the pulverization efficiency increased, improving the reactivity of the fly ash.

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