Effects of Light-Burnt Dolomite Incorporation on the Setting, Strength, and Drying Shrinkage of One-Part Alkali-Activated Slag Cement

In Kyu Jeon, Jae Suk Ryou, Sadam Hussain Jakhrani and Hong Gi Kim *

Department of Civil and Environmental Engineering, Hanyang University, Seoul 04763, Korea
* Correspondence: dmkg1404@hanyang.ac.kr; Tel.: +82-2-2220-4323

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Abstract: This study investigates the potential of light-burnt dolomite (LBD) as a supplementary cementitious material with ground granulated blast furnace slag (GGBFS) and Ordinary Portland cement (OPC). In this work, LBD was substituted for up to 20% of GGBFS in sodium sulfate-activated slag systems. The effects of LBD incorporation on the flow, setting time, compressive and flexural strength development, and drying shrinkage were explored with, X-ray diffraction and thermogravimetric analyses. LBD incorporation resulted in greater strength development of an alkali-activated slag system. The optimum LBD content for strength development was 10%, regardless of ordinary Portland cement content. In addition, LBD decreased the drying shrinkage, accelerated the hydration process, and induced hydrotalcite formation, which can be attributed to the reactive MgO inside LBD.

Keywords: light-burnt dolomite; GGBFS; setting; strength; drying shrinkage; mortar

1. Introduction

Ordinary Portland cement (OPC) is a common raw material for concrete structures. However, the cement manufacturing process is considered to be a major factor in global warming and air pollution because it emits a large amount of carbon dioxide [1]. In alkali-activated cement (AAC), all or part of the cement binder is replaced with supplementary cementitious materials (SCM); thus, AAC is a potential eco-friendly alternative that can reduce carbon dioxide emissions [2,3]. AAC is generally obtained from two-part reactions with solid aluminosilicate precursors and concentrated aqueous alkali activators [4]. However, two-part reactive AAC with corrosive and viscous alkali activators, such as sodium silicate solution has high costs, due to handling and application problems [5,6]. Recent studies have focused on the development of user-friendly one-part AAC, in which only water is added [7–9].

Studies have recently been conducted on alkali-activated slag (AAS), which is a type of AAC that is an alternative to OPC [10–12]. Slag has a similar chemical composition to OPC, and several studies indicate that AAS has good strength development with lower calcium content [13,14]. In AAS procedures, the alkali activation mechanism consists of the combined reaction of destruction-condensation, where the primary material is destroyed in low stable structural units and interacted with condensed structure. As a result of this reaction, low crystalline calcium silicate hydrate (C-S-H) with a low CaO/SiO$_2$ ratio is produced in the activation process [15,16]. AAS has different chain structures than OPC: While, OPC consists of linear finite chains with SiQ$^1$ and SiQ$^2$, AAS has a long chain organized with SiQ$^2$ or SiQ$^2$ (1Al) [17].

Previous studies have attempted to enhance the mechanical properties of AAS by incorporating different admixtures. Chang et al. showed that incorporating gypsum in AAS pastes increased...
the compressive strength and decreased the setting time. Park et al. demonstrated that AAS activated by clinker-free CaO, substituted with gypsum, had positive effects on strength development, due to ettringite formation.

However, AAS has larger shrinkage values than OPC, which has led to a decrease in its use [18,19]. Previous research [20,21] found that the drying shrinkage of AAS was 2 to 3 times higher than OPC under the same curing environment. Thomas et al. [22] indicated that capillary stress, surface tension, and disassembling pressure were the major factors in the drying shrinkage of cementitious materials. Ye et al. [23,24] found that the structural combinations of alkali cations, in C-S-H and C-A-S-H, which led to a diminished stacking uniformity of C-S-H and C-A-S-H, also made it easier to collapse and repartition against drying conditions. To decrease the negative effects of shrinkage in an AAS system, Neto et al. [25] showed that the presence of sodium silicate and silica in AAS decreased shrinkage, due to additional ettringite formation.

Dolomite [CaMg(CO$_3$)$_2$] is obtained from mineral dolomite rock, consisting of calcium, magnesium, and carbonate. In high-alkaline conditions, dolomite undergoes a de-dolomitization reaction, reacting with portlandite to produce calcite and brucite [26,27]. The light-burnt dolomite (LBD) is produced by raw dolomite through a calcination process at 900–1000 °C and consists mainly of reactive CaO and MgO. LBD is an economical powder-phased lime material that is used as an activator. A previous study [28] used LBD, up to 2%, as an activator with sodium silicate solution, in order to analyze the hydration and strength development of AAS. However, other studies have not used LBD as an SCM. If LBD could be activated like GGBFS and shows higher, not only mechanical properties, but shrinkage compensate performance, than the GGBFS mixture, the feasibility of using LBD as SCM will be demonstrated. Additionally, large amounts of reactive MgO can play a role in mechanical and durability performance of cement composite [29]. Therefore, this study aimed to investigate the influence of LBD used as a SCM on the hydration process, strength development, and drying shrinkage of AAS. For this purpose, workability, compressive and flexural strength, isothermal conduction calorimetry (ICC), and drying shrinkage were explored with X-ray diffraction analysis (XRD) and thermogravimetric analysis (TGA).

2. Experimental Process

2.1. Materials

OPC complying with ASTM C150 [30], ground granulated blast furnace slag (GGBFS), and light-burnt dolomite (LBD), obtained from a local R company in Gwangyang, Republic of Korea were used as binders in this study. Sodium sulfate anhydrous with a purity of 99% was used in this study as an activator. The chemical and physical properties of the OPC, GGBFS, and LBD were determined with X-ray fluorescence (XRF) (S8 Tiger, Bruker, Billerica, MA, USA), as presented in Table 1. Figure 1 shows the particle size distributions of OPC, GGBFS, and LBD, which were measured with a particle size analyzer (Malvern Mastersizer 2000, Malvern, Malvern, England). The specific surface area of the raw materials was obtained from BET method using a surface area analyzer (3 Flex, micromeritics instruments, Micromeritics, GA, USA). Figure 2 shows the mineralogical composition of LBD, obtained from XRD analysis. Natural sand with a maximum size of 5 mm, absorption of 1.05%, and fineness modulus (F.M) of 2.7 was used as a fine aggregate in this study.
Table 1. Chemical and physical properties of Ordinary Portland cement (OPC), ground granulated blast furnace slag (GGBFS), and light-burnt dolomite (LBD).

| Composition | Weight (%) | OPC | GGBFS | LBD |
|-------------|------------|-----|-------|-----|
| SiO₂        |            | 20.8| 34.1  | 3.0 |
| Al₂O₃       |            | 6.3 | 16.1  | 1.3 |
| Fe₂O₃       |            | 3.2 | 0.4   | 0.6 |
| CaO         |            | 62  | 42.3  | 52.2|
| MgO         |            | 3.3 | 4.1   | 25.3|
| SO₃         |            | 2.2 | 2.5   | 0.4 |
| Na₂O        |            | -   | -     | 0.06|
| K₂O         |            | -   | -     | 0.2 |
| P₂O₅        |            | -   | -     | 0.03|
| TiO₂        |            | -   | -     | -   |
| Loss on ignition |    | 1.3 | 0.05  | 1.85|
| Specific surface area [cm²/g] | | 3200 | 4893  | 3038|

Figure 1. Particle size distributions of Ordinary Portland cement (OPC), ground granulated blast furnace slag (GGBFS), and light-burnt dolomite (LBD).

2.2. Mix Proportions and Fabrication of Mortar Specimens

The mortar mix proportions used in this study are presented in Table 2. The mortar mixture had a fine aggregate-to-binder weight ratio of 2 and a water-to-binder weight ratio of 0.4. Sodium sulfate is an alkali activator, with 4% of the binder weight used. The amount of cement in the mixtures was fixed at 10% or 20%. LBD was replaced (by weight) with 0%, 10%, and 20% of GGBFS. The values are denoted as C10S90, C10S80L10, C10S70L20, C20S80, C20S70L10, and C20S60L20. For example, C10S80L10 indicates a mortar mix made with 10% cement, 80% GGBFS, and 10% LBD.

Mortar cubes (50 × 50 × 50 mm³) and prisms (40 × 40 × 160 mm³) were prepared in accordance with ASTM C109 [31], and ASTM C348 [32], respectively. After mixing, the cube and prisms molds
were vibrated for 30 s using a vibration table to remove air voids. Mortar samples were then sealed in plastic wrap to prevent moisture evaporation and stored in ambient conditions (20 ± 5 °C and 60 ± 5% relative humidity) for 24 h. After 24 h, the mortar samples were demolded and kept in water at 23 ± 2 °C for curing until the desired duration was reached.

![Figure 2. XRD patterns of LBD.](image)

**Table 2. Mix composition of the specimens.**

| Specimen Code | OPC (wt%) | GGBFS (wt%) | LBD (wt%) | Sodium Sulfate | Sand | Water-to-Binder Ratio |
|---------------|-----------|-------------|-----------|----------------|------|----------------------|
| C10S90        | 0.1       | 0.9         | 0         | 0.04           | 2    | 0.4                  |
| C10S80L10     | 0.1       | 0.8         | 0.1       | 0.04           | 2    | 0.4                  |
| C10S70L20     | 0.1       | 0.7         | 0.2       | 0.04           | 2    | 0.4                  |
| C20S80        | 0.2       | 0.8         | 0         | 0.04           | 2    | 0.4                  |
| C20S70L10     | 0.2       | 0.7         | 0.1       | 0.04           | 2    | 0.4                  |
| C20S60L20     | 0.2       | 0.6         | 0.2       | 0.04           | 2    | 0.4                  |

**2.3. Testing Methods**

In this study, the properties of fresh mortar mixtures were determined by testing their workability. The workability of mixtures was tested using the flow table according to ASTM C230 (ASTM, 2004b) [33], and the procedure followed ASTM C1437 (ASTM, 2001) [34]. To observe the setting time of fresh mortar, the initial and final setting times were investigated using Vicat needles in accordance with ASTM C191 (ASTM, 2004a) [35].

To evaluate the mechanical properties of AAS, the compressive strengths of cube samples were measured using a universal testing machine (Shimadzu, CCM-200A; Shimadzu Corporation, Kyoto, Japan), in accordance with ASTM C109 [31]. A flexural strength test was conducted on the prism samples, according to the ASTM C348 [36] after 3, 7, and 28 days of water curing. For each test, three replicates were measured; the average value is presented. The heat evolution was investigated using a semi-adiabatic calorimeter according to a previous study [37]. For XRD and TGA analysis, powder specimens were obtained from the AAS mixtures after 7 days of hydration using an X-ray diffractometer (RINT D/max 2500, 40 kV, 30 Ma, scanning speed 2°/min, wavelength 1.54 Å) and a thermogravimetric analyzer (TGA7 PERKINELMER, TA instruments, PerkinElmer, Waltham, MA, USA). The drying shrinkage of AAS mixtures was tested using a dial gauge in accordance with ASTM 490 [38].
3. Results and Discussion

3.1. Flow Test

Figure 3 shows the flow diameters of alkali-activated slag mortar mixtures. The flow value of AAS mortar was influenced by the replacement levels of LBD and OPC in the binder. For the OPC replacement levels, a reduction of OPC content in the mixture enhanced the GGBFS content and decreased the fluidity value, which is a similar trend to that seen in a previous paper [39,40]. The fluidity of C10S90, C10S80L10, and C10S70L20 decreased by 5.4%, 6.2%, and 3.1%, respectively, compared with C20S80, C20S70L10, and C20S60L20. Regardless of OPC content, an increasing amount of LBD led to a reduction of fluidity for all mortar mixes. In a mixture containing 10% OPC, the fluidity of C10S80L10 and C10S70L20 declined by approximately 10.7%, and 17%, respectively, compared with C10S90. With increasing replacement levels of LBD, a higher fluidity reduction effect appeared. The decrease in flow value is similar to the results reported in previous studies [41]. H. Ye et al. indicated that the dolomite incorporation decreases the workability of AAS, thus, leading to increased water demands to achieve a similar flow value.

![Figure 3. Fluidity of alkali-activated slag mixture.](image)

3.2. Setting Time

Figure 4 illustrates the initial and final setting times of an AAS mixture, incorporated with LBD. Setting time is an important factor in the strength development of cementitious materials, which normally occurs in a short time [19]. Generally, an AAS system has shorter initial and final setting times than OPC-based mixtures. In comparing 10% OPC mixture with 20% OPC mixture, the 20% OPC-based AAS showed slightly longer initial and final setting times, due to the amount of GGBFS content. Less OPC content led to increased GGBFS content in the same amount of binder; greater GGBFS content also decreased the setting time [42]. In addition, both the initial and final setting times decreased with enhanced LBD content. When the LBD content was increased from 10% to 20% in a 10% OPC-based AAS system, the initial setting values decreased by 37.8% and 96.15%, and the final setting values decreased by 32.3% and 67.9%, compared with only OPC, and a GGBFS-based AAS system, respectively. A 20% OPC-based AAS system has similar tendencies. When LBD increased from 10% to 20%, the initial setting values decreased by 33.3% and 86.6%, and the final setting values decreased by 28.8%, and 54%, respectively. The setting time can be affected by the generation heat release of
materials. LBD has more reactivity than GGBFS and produces more heat release. The cumulative heat release of different LBD content in an AAS system is discussed in Section 3.3.

Figure 4. Initial and final setting time of alkali-activated slag (AAS) mixtures.

3.3. Isothermal Heat Release

Generally, the effects of temperature on reaction kinetics were investigated using isothermal calorimetry. Figure 5 shows the calorimetric curves of the heat release of AAS mixtures. Figure 5a shows the heat release curve of 10% OPC-containing mixtures, and Figure 5b represents the heat release curve of 20% OPC-containing mixtures, which both have two distinct peaks. The first sharp peak occurred immediately after mixing all the mixtures. Bernal et al. [43] believed that this initial peak was produced from the dissolution of the dry mixture. With increasing LBD content, the samples showed higher maximum heat evolution rates for the dissolution peaks compared with 0% LBD samples. Thus, a higher LBD content promotes dissolution. A similar heat evolution enhancement was observed in a previous study of an AAS system with dolomite [28,44]. Yang et al. [28] found that the increase in heat evolution with LBD was due to the hydration procedure for LBD. Reactive CaO and MgO inside LBD, produced calcium hydroxide, C-A-S-H gel, and magnesium hydroxide, which affected the heat evolution of the AAS system. Ye et al. [41] also reported that the replacement of slag with dolomite, increased the Na₂O content and caused the slag to react at a more advanced level.

The second peak occurred between 20 and 30 h. Zhang et al. [45] found that the acceleration peaks are affected by the reorganization of the C-A-S-H gel phase, which is related to the polymerization degree of the binder. Figure 5a,b show a higher second peak value with increasing LBD content. In both mixtures, increasing the amount of LBD caused the peak time to be slightly delayed, due to the high MgO content of LBD. LBD reacted with water and produced magnesium hydroxide, which is an insoluble product. During this formation process, the dissolved Mg²⁺ led to a decrease of hydroxyl ions in the solution, then caused a reduction in the alkalinity of the mixture [46].

Figure 6 shows the cumulative heat release of AAS mixtures. The cumulative heat release sequence by mixture was as follows: C10S70L20, C20S60L20, C10S80L10, C20S70L10, C10S90, and C20S80. LBD incorporation enhanced the cumulative heat release compared to specimens without LBD. These increases in the cumulative heat release can affect the setting time, as was discussed in Section 3.2.
3.4. Compressive Strength

Figure 7 represents the compressive strength results of AAS specimens with different LBD and OPC replacement levels. The compressive strength development showed similar trends but different magnitudes for each mixture. The compressive strength of AAS mortar was enhanced significantly at 7 days, but then the rate of compressive strength increased less rapidly. Regardless of LBD addition, mortar mixes with 20% OPC had higher compressive strength values, compared with 10% OPC mortar mixes. Similar results were observed by Yang et al. [28].

The incorporation of LBD in AAS at a GGBFS replacement ratio of 10% had a significant effect on the compressive strength for all different ages, regardless of OPC content. For example, C20S70L10 showed the highest compressive strength value of 52.4 Mpa (range 52.1 Mpa to 52.7 Mpa) at 28 days of curing, which is 31.7% greater than C20S80 without LBD. In addition, C10S80L10 had a compressive strength value of 49.05 Mpa (range 48.7 Mpa to 49.4 Mpa) at 28 days of curing, which is approximately 60% greater than C10S90. Although mortar specimens containing 20% LBD showed lower compressive strength values than did those with 10% LBD, the C10S70L20 and C20S60L20...
specimens had strength values that were 44% and 13.3% greater than those of the C10S90, and C20S80 specimens, respectively.

Hailong Ye et al. [41] found that the incorporation of dolomite induced porosity reduction and strength development, by enhancing the reaction degree of slag because of a filler effect. Machner et al. [47] showed similar enhancement of compressive strength results and attributed the compressive strength development to the use of dolomite through carbonate addition. The addition of dolomite induced an ettringite stabilization effect and the formation of carbonate AFm phases.

Figure 7. Compressive strength of alkali-activated slag mortar with different mixtures.

3.5. Flexural Strength

Figure 8 shows the influence of LBD incorporation on the flexural strength development of AAS mortar mixtures. Flexural strength showed a different tendency than did compressive strength, which increased steadily over time. In all mortar mixtures, flexural strength tended to increase until 7 days of curing, but decreased somewhat by 28 days of curing. The mixtures with 10% LBD, in AAS, showed higher flexural strength values, compared with 0% LBD mixtures for all curing ages. This is similar to compressive strength development. C10S80L10 specimens showed the highest flexural strengths at all curing ages: 10.88 Mpa (range 10.33 Mpa to 11.43 Mpa), 12.19 Mpa (range 11.54 Mpa to 12.84 Mpa), and 10.5 Mpa (range 10.25 Mpa to 10.75 Mpa) at 3 days, 7 days, and 28 days of curing. The tendency for reduced flexural strength after 7 days in an AAS system was also demonstrated in a previous study [42]. In addition, Nedeljković et al. [42] found that large deviations in flexural strength values were mainly due to microcracks. In AAS systems, Hubler et al. [48], Thomas et al. [49], and Collins and Sanjayan [50] observed microcracks both inside, and on the surface of, the AAS samples. With increasing hydration, the mixtures showed greater compressive strength regardless of the microcracks. However, the microcracks in specimens resulted from several factors; for example, surface drying shrinkage played a major role in reducing flexural strength. The drying shrinkage of AAS mortar is discussed in Section 3.6. Duxson et al. [51] proposed a different concept to demonstrate flexural strength behavior through the defect density of unreacted material. Duxson believed that an enhanced amount of unreacted material would increase the defect density. Thus, potential methods of failure would increase due to the increased defect density.
3.6. Drying Shrinkage of Mortar Specimens

The drying shrinkage values of samples with different OPC and LBD content are shown in Figure 9. Regardless of LBD content, greater drying shrinkage was shown in all mixtures containing 10% OPC, compared with mixtures containing 20% OPC. Increased alkali-activated binder content improved the drying shrinkage. These trends were also reported in previous studies. Collins and Sanjayan [21] found that AAS had drying shrinkage values that were approximately three times greater than those of the OPC concrete. Aydin et al. [52] investigated the drying shrinkage of AAS with different hydration products. The common hydration products of AAS are C-S-H gels, which have low Ca/Si ratios (0.8–1.1), and lack of crystal phases, such as Ca(OH)₂, leading to higher drying shrinkage. All specimens showed a similar trend, with higher LBD content being associated with a lower drying shrinkage value. After 3 weeks, the drying shrinkage values decreased by 26.3%, and 29.8% for C10S80L10, and C10S70L20, respectively, compared with C10S90 (which only consists of OPC and GGBFS). In addition, C20S70L10 and C20S60L20 had drying shrinkage values that were approximately 31% and 42% less than that of C20S80, respectively. These results are consistent with previous studies [53,54]. The decreased drying shrinkage associated with LBD could be attributed to the role of reactive MgO in LBD. The reactive MgO inside LBD can accelerate the hydration of the AAS system and fill the pores, so it helps to reduce the drying shrinkage. Fang et al. [55] and Jin et al. [56] showed that the use of reactive MgO in alkali-activated binder can reduce the drying shrinkage.
3.7. XRD Analysis

Figure 10 shows the XRD patterns of the alkali-activated slag mixtures, with different OPC and LBD content, at 7 days of curing. In sodium sulfate-activated slag mixtures, the main reacted products are ettringite and C-S-H [57]. Regardless of LBD content, ettringite and C-S-H could be identified in all AAS mixtures. Akermanite was identified due to the mineralogical properties of GGBFS. Calcium hydroxide peaks were measured in all mixtures, unlike typical sodium sulfate-activated slag mixtures, which can be attributed to the replacement of OPC content. The peak intensity of hydrotalcite in LBD-containing specimens was stronger than that of specimens without LBD. In addition, increasing LBD content was associated with greater hydrotalcite peak intensity. A high amount of reactive MgO in LBD can help to produce Mg-Al type layered double hydroxide. A distinct peak of C/CSH, which overlaps with C-S-H and calcite, could be seen in all mixtures; the intensity of these peaks was slightly enhanced with LBD content. This may be part of the reason for the increased heat evolution discussed in Section 3.3. LBD may accelerate the polymerization and hydration process of an AAS system and lead to the production of more C-S-H type gel. The increase in calcite peak intensity, due to the increase in LBD content, can be attributed to the large amount of calcite in LBD. The change in peak values from OPC content was not as evident as the apparent change in peak values from LBD content. It can be demonstrated that reactive CaO in LBD can produce the Ca(OH)\(_2\) with water and then the hydration proceeds with SiO\(_2\) in GGBFS to produce C-S-H gel, which leads to accelerate hydration procedure. Additionally, reactive MgO in LBD can form voluminous products, such as hydrotalcite and can produce dense microstructure [58,59].
Figure 10. XRD patterns of AAS mixtures with different OPC and LBD content at 7 days. Key: E–ettringite \( \text{Ca}_6\text{Al}_2(\text{SO}_4)_3(\text{OH})_{12}\cdot 26\text{H}_2\text{O} \), Q–Quartz, Ht–hydrotalcite \( \text{Mg}_6\text{Al}_2\text{CO}_3(\text{OH})_{16}\cdot 4\text{H}_2\text{O} \), C–Calcite, CSH–Calcium silicate hydrate, A–akermanite \( \text{Ca}_2\text{Mg}(\text{Si}_2\text{O}_7) \), CH–Calcium hydroxide.

3.8. TGA Analysis

Figure 11 shows the derivative thermogravimetric (DTG) analysis curves of 20% OPC-based AAS mixtures, with increasing LBD content. Regardless of LBD inclusion, all AAS mixtures showed mass loss peaks at four different temperature regions. The first mass loss peak is below 150 °C is related to the evaporation of free water [60], the dehydration of C-S-H [19], and ettringite [61]. The second mass loss peak is located between 350 and 400 °C, which can be attributed to the thermal dehydration of the hydrotalcite phase [62]. With increasing LBD content, a higher mass loss was indicated in the DTG curve of 20% OPC-based AAS mixtures because the higher amount of reactive MgO inside LBD helps to produce a hydrotalcite phase. The progressive mass loss, above approximately 400 °C, is related to the dehydration of a C-S-H type gel, such as C-A-S-H gel [60]. A higher amount of LBD in AAS mixtures shows a significant mass loss in this temperature region. When LBD was incorporated in an AAS system, LBD accelerated the polymerization process and assisted the hydration product in an AAS system, which is associated with cumulative heat release, as discussed in Section 3.3. The last mass loss peak is between 600 and 750 °C, which is attributed to the thermal decomposition of a carbonate product, such as calcite. In the DTG curve between 600 and 750 °C, the biggest mass loss peak is found in C20S60L20, followed by C20S70L10 and C20L80.

LBD has more carbonate than OPC and GGBFS. Therefore, LBD in an AAS system might play a role in the formation of carbonate salt, due to the large amount of reactive CaO, MgO, and \( \text{CO}_3^{2-} \), compared with GGBFS. Figure 12 shows that the mass loss fractions contribute to the hydrotalcite (350–400 °C), C-A-S-H gel (400–500 °C), and carbonate salt (600–750 °C). Regardless of OPC content, all AAS mixtures with 20% LBD had the greatest hydrotalcite, C-A-S-H gel, and carbonate salt mass loss fraction among the specimens that included LBD. Yang et al. [44] found that, using calcined dolomite consumes \( \text{CO}_3^{2-} \) ions to produce the hydrotalcite phase, and then increases the alkalinity to accelerate C-A-S-H gel formation.
4. Conclusions

In this study, one-part sodium sulfate alkali-activated slag cement mixture was prepared using OPC, GGBFS, and LBD as binder materials, with solid sodium sulfate powder as the alkali activator. These experiments were conducted to investigate the effects of LBD incorporation on setting, strength, and drying shrinkage of an AAS system. From the results, the following conclusions can be made:

1. The replacement of up to 20% of GGBFS with LBD has minimal negative effects on the fluidity of AAS mixtures. As the replacement ratio increased to 20%, the fluidity reduction was approximately 10% to 17% compared to specimens without LBD based on 10% OPC specimens. The groups containing 20% OPC had slightly higher fluidity compared with groups containing 10% OPC.

2. Increasing LBD content tends to enhance the heat evolution of an AAS mixture. It also accelerates the polymerization and hydration process of an AAS system, which is due to the formation of calcium hydroxide from reactive CaO in LBD, resulting in a reduction of the initial, and final, setting times of AAS mixtures.
3. Replacing GGBFS with LBD has positive effects on compressive and flexural strength. Regardless of OPC content, LBD-containing specimens showed greater strength development than mixtures without LBD. Reactive CaO in LBD accelerate the hydration process in producing more C-S-H gel and reactive MgO in LBD produce voluminous hydration products to make dense microstructure. The highest compressive and flexural strength values were obtained from 10% LBD mixtures, followed by 20% and no-LBD content in both 10% and 20% OPC-based mixtures.

4. In terms of drying shrinkage, increasing the OPC content from 10% to 20% causes a reduction of GGBFS content, resulting in decreased shrinkage values. Replacing LBD with GGBFS can effectively reduce the drying shrinkage of sodium sulfate-activated AAS mixtures, regardless of OPC content.

5. The main hydration products of ettringite, C-S-H, and Ca(OH)$_2$ were detected by XRD and TGA in all mixtures. Increasing amounts of C-S-H, hydrotalcite phase, and calcite were observed in LBD-containing specimens, which can be attributed to the reactive MgO, CaO, and calcite inside LBD.

**Further Research**

Future research should focus on the influence of LBD incorporation on microstructure and mechanical durability. Studies should also be conducted to investigate the microstructure, chloride penetration, and carbonation of AAS mixtures.

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