Assessing the chemical involvement of limestone powder in sodium carbonate activated slag

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Abstract This study aims to investigate the effect of limestone powder (LP) on the reaction of sodium carbonate activated slag. The results show that the incorporated LP up to 30% improves the strength development, especially at advanced curing ages. A slightly accelerated reaction is observed for samples containing low amount of LP (≤5%), while mixture with 10% LP shows the optimized results with respect to the heat release and strength development. Chemical effect of incorporating LP is observed at high replacement levels (≥15%), indicated by the formation of a new phase, natron (Na₂CO₃·10H₂O). Besides, relatively high contents of hydrotalcite-like phases are generated when increasing the dosage of limestone powder. The chemical changes, including the volume changes of generating natron and the transformation of natron to calcite, is potentially responsible for the enhanced mechanical properties.

Keywords Limestone powder · Sodium carbonate activated slag · Chemical effect · Reaction kinetics · Reaction products · Mechanical properties

1 Introduction

Alkali activated materials (AAM) have been attracting worldwide attention during the last few decades because of their good material properties and environmental benefits [1]. Nevertheless, depending on the used raw materials and alkaline solutions, the performance of the resulted materials can be varied [2, 3]. At present, most attentions are paid to the development of mixtures with higher strength, or improved durability, upgrading the-state-of-the-art of AAM [4–7]. However, the effects of supplementary materials on the reaction of AAM are rarely studied, especially their potential chemical involvement.

As a supplementary material, limestone powder (LP) has been widely applied in Portland cement based building materials because of its low price and good performance [8–12]. The effect of LP on the hydration of cement has been intensively investigated, ranging from the filler effect to the chemical involvement in cement hydration: (1) inert filler acting as nuclei sites; (2) accelerating the hydration of C₃S [13, 14] and (3) reacting with C₃A forming calcium carboaluminates [14–16]. However, the function of LP on the reaction of alkali activated materials has not been

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systematically studied yet. Up till now, only limited efforts [17–19] have been devoted to this topic partly because the system of alkali activated material (AAM) is more complex than Portland cement system due to the different activators and raw materials applied [1, 20, 21].

It is reported that in the low-calcium containing alkali activated metakaolin system, LP is dissolved in the sodium hydroxide solution and its presence enhances the release of Al and Si ions from metakaolin, leading to the formation of layered calcium carboaluminates [17]. While in high-calcium containing alkali activated slag system, the function of limestone is not well understood. LP is often regarded as an inert filler in the slag based alkali activated system [13–15]. Besides, intensive Ca\(^{2+}\) ions will be released from slag particles and thus the dissolution of LP releasing Ca\(^{2+}\) ions is actually detained compared to its hydrolysis in the low-calcium containing alkali activated system. As a result, LP is more likely to be restricted to an inert filler, acting as nucleation sites. Gao et al. [18] characterized waterglass activated slag-fly ash-limestone blends, and reported that the incorporation of LP shows good filler effect by giving a slightly higher strength than that of fly ash. However, no trace of chemical involvement can be identified as there are no monocarboaluminate or new phases found on the XRD pattern. Most recently, Rakhimova et al. [22] studied the influence of different LP on the properties of slag activated by sodium carbonate based waste and found that the 28 days strength was not weakened up to 50% addition of LP (Blain fineness ≥400 m\(^2\)/kg). However, they stated that the benefits on the strength development is mainly attributed to the “physical activity” of LP.

Compared to waterglass, sodium carbonate as an activator shows enhanced contribution concerning setting time and shrinkage [2, 23, 24]. Besides, the production of Na\(_2\)CO\(_3\) is more cost-effective and environmentally friendly compared to other activators [25]. It is noteworthy that LP could act differently in sodium carbonate activated slag (SCAS) system due to its particular reaction mechanism compared to slags activated by waterglass or sodium hydroxide [4, 26, 27]. As reported by Bernal et al. [4] and our previous study [26], CO\(_3^{2-}\) anions concentration in the pore solution and the initially precipitated calcium carbonate significantly control the sodium carbonate activation process. According to the previous researches [4, 23, 28–30], the main reaction products of sodium carbonate activated slag are C–(A)–S–H gel, calcium carbonate (CaCO\(_3\)), gaylussite (Na\(_2\), Ca(CO\(_3\))\(_2\)-10H\(_2\)O), hydrotalcite (M\(_{g6}A\(_{l2}CO\(_3\)(OH)\(_{16}-\)4H\(_2\)O), etc. With respect to the carbonates groups in the minerals, the presence of limestone could be potentially involved in the reaction, e.g. Ca\(^{2+}\) can be potentially involved in the formation of C–(A)–S–H gel. Furthermore, the incorporation of LP contributes to the strength improvement and lowered cost. Moseson [19] developed concretes with strength up to around 41 MPa at 3 days and 65 MPa at 28 days while the CO\(_2\) emission and energy consumption is reduced by 97% compared to Portland cement (PC) based materials. However, that research was mainly focused on the economic benefits and CO\(_2\) emission, while the chemical function of LP on the reaction of SCAS was not discussed.

The present research aims to study the effect of LP on the reaction kinetics, reaction products and strength development of sodium carbonate activated slag. The heat evolution of mixtures with different LP incorporation levels was measured, up to 7 days. Furthermore, the specimens was ground for microstructural analyses, including X-ray diffraction (XRD), Thermogravimetry and Derivative Thermogravimetry (TG-DTG) and Fourier transform infrared spectroscopy (FTIR), etc. The strength developments were characterized at different curing ages. The fresh behaviour was evaluated. The chemical involvement of LP especially at a high content in the sodium carbonate activation is observed and the related mechanism is discussed.

2 Materials and experiments

2.1 Materials

Ground granulated blast furnace slags (GGBS) (provided by ENCI B.V, the Netherlands) and limestone powder were applied as raw materials. The oxide compositions of these two materials were determined by using X-ray Fluorescence (XRF), as shown in Table 1. The basicity coefficient of GGBS was calculated based on the chemical composition, following the formula (\(K_b = (CaO + MgO)/(SiO_2 + Al_2O_3)\)), yielding 1.3. The particle size distributions (PSDs) of the GGBS and LP, determined by Mastersizer 2000, are presented in Fig. 1, with the d(0.5) of
Table 1 Chemical composition of the GGBS determined by XRF

| Chemical composition (%) | GGBS | Limestone |
|--------------------------|------|-----------|
| SiO₂                     | 35.50| 0.84      |
| CaO                      | 38.60| 53.96     |
| Al₂O₃                    | 13.60| 0.24      |
| MgO                      | 10.20| 1.01      |
| Fe₂O₃                    | 0.48 | 0.32      |
| SO₃                      | 1.27 | –         |
| K₂O                      | 0.48 | 0.34      |
| TiO₂                     | 1.01 | –         |
| Cl                       | 0.01 | –         |
| L.O.I                    | –    | 43.01     |
| Density (kg/m³)          | 2930 | 2710      |
| Blaine fineness (m²/kg)  | 373  | 546       |

19.37 µm and 10.30 µm, respectively. Sodium carbonate (powder form, analytical grade) was firstly dissolved in water followed by cooling down to room temperature (20 ± 1 °C), prior to further actions.

2.2 Experiments

2.2.1 Flowability and strength test

The starting materials (Table 2) were firstly mixed and the flowability of samples was measured by a Haëgermann cone with the diameters of: top = 70 mm, base = 100 mm and height = 50 mm. Two perpendicular diameters of the resulted paste spread were measured and the flowability of the pastes is interpreted by the average diameter. Afterwards, the fresh pastes were cast into plastic moulds (40 × 40 × 160 mm³) following the preparation procedure suggested by EN 196-1. The specimens were manually vibrated and then covered with plastic foils to prevent the potential moisture loss. After hardening, the samples were demoulded and stored in a climate chamber with the relative humidity of >95% at room temperature (20 ± 1 °C) till the ages of testing.

2.2.2 Sodium carbonate activation kinetics assessment

The heat evolution of the designed mixtures (Table 2) was characterized by an isothermal calorimetry instrument (TAM AIR Calorimeter), set at 20 °C. Fresh pastes were manually prepared and transferred to an ampoule before loading to the calorimeter. It should be noted that the measurement in the initial ~45 min might not be reliable because of the instability of the instrument and the sampling time about 4–6 min after mixing was not measured.

Fig. 1 Particle size distributions of the GGBS and LP
2.2.3 Microstructural analysis

The remaining samples after strength tests were ground into powders (≈63 μm) for the reaction products and microstructural analyses. The reaction products of the samples were analyzed by X-ray diffractometry (XRD) analysis using a Cu tube (20 kV, 10 mA) with the settings of: scanning range 3°–50°, step of 0.02° and measuring time 3 s/step. About 40–70 mg powdered samples were measured by thermogravimetric analysis (TGA), and further evaluated with derivative thermogravimetric analysis (DTG). The specimen was firstly isothermally held at 20 °C for 3 h to stabilize the internal balance of instrument before heated up to 1000 °C with the rate of 1 °C/min, using nitrogen (40 ml/min) as the carrier gas. The FT-IR spectra scanned 12 times from 4000 to 400 cm\(^{-1}\) were collected by applying a PerkinElmer Frontier\textsuperscript{TM} IR/FIR Spectrometer with the attenuated total reflection (ATR) method (GladiATR, crystal diamond) at a resolution of 4 cm\(^{-1}\) using the ground solid powders.

3 Results and discussion

3.1 Flowability

As shown in Fig. 2, the incorporation of LP increases the flowability of the mixtures up to a replacement level of about 15%. When further increasing the dosage of LP to 30%, the workability of the mixtures is slightly decreased. Improvement on the fresh behavior of pastes were also observed in PC based materials after incorporating LP [31, 32]. This can be contributed by two mechanisms. Firstly, the packing of the mixture is improved by the applied LP as it is finer than slag particles (see Blaine fineness), leading to a better flowability. A similar phenomenon was found by Brouwers and Radix [33] and Quercia et al. [34] that an improved packing of mixtures leads to a better spread flow. Secondly, due to the presence of sodium carbonate in the solution, the dissolution of LP is prohibited because of the saturation limitation. The LP consumes less water than slag particles as the dissolution of those particles increases the water demand. However, when increasing the dosage of LP from 15 to 30%, the flowability of the mixture decreases, which can be explained by the reduced volume percentage of water.

It should also be noted that the fresh behavior is also affected by the initial reaction of sodium carbonate activated slag, i.e. precipitation of CaCO\(_3\) and gaylussite [35]. Due to the high concentration of CO\(_3^{2-}\) anion provided by the activators and low saturation limit of CaCO\(_3\), the precipitation and formation of gaylussite will quickly occur when slag starts to release Ca\(^{2+}\) ions. The early age reaction will negatively affect the flowability of the samples. By replacing slag particles, the initial reaction is delayed to a certain content (when the LP dosage ≥10%), as shown in the calorimetric results (Fig. 4). Consequently, the workability of the pastes is improved.

| Table 2 | Designed mixtures with different contents of LP |
|---------|---------------------------------|
| Mixture | Slag content (wt%) | LP (wt%) | Na\(_2\)CO\(_3\) dosage (Na\(_2\)O wt%) | Water/solid ratio (–) |
| Ref.    | 100 | 0 | 4.0% | 0.4 |
| LP 5%   | 95 | 5 | |
| LP 10%  | 90 | 10 | |
| LP 15%  | 85 | 15 | |
| LP 30%  | 70 | 30 | |

![Fig. 2 Spread flow of the mixtures versus the LP content](image)
3.2 Sodium carbonate activation kinetics

Figure 3 presents the heat evolution of sodium carbonate activated slag with different LP replacement levels. It is shown that the reaction process of SCAS incorporating LP is similar to that of PC hydration or other alkali activated materials [36–39], which can generally be divided into five stages. Similar to the previous research results [4, 26], the dormant periods of the mixtures are relatively long due to the initial precipitation of calcium carbonate. As depicted in Fig. 3, with 5% LP addition, the heat release intensities of the samples are slightly improved, indicating a slightly faster reaction of slag particles after incorporating LP. When increasing the LP dosage to 10%, the reaction is delayed but the intensity is increased. However, further increasing the LP content leads to a slower and less intensive reaction as shown in Table 3.

It should be noted that LP with 10% replacement level gives the strongest intensity on the heat evolution, indicating the optimal effects of LP addition (replacement levels between 5 and 30%). The same trend is also found in the compressive strength development of the samples containing different levels of LP (Fig. 4).

Ke et al. [30] investigated the reaction kinetics of SCAS by using calcined layered double hydroxides (CLDH), which removed the CO$_3^{2-}$ anions, yielding a significant rise in the pH and consequently fast reaction. Bernal et al. [4] also reported that the concentration of CO$_3^{2-}$ anions in the pore solution plays a significant role on the early age reaction of SCAS. In the current study, the incorporated LP provides extra surface areas for the precipitation of reaction products, but, in the meantime sufficient Ca$^{2+}$ ions need to be dissolved to consume the same amount of CO$_3^{2-}$ anions which require a further dissolution of slag particles due to the dilution effect, i.e. a deeper dissolution. In this case, the incorporation of LP can accelerate (mixture LP 5%) or intensify (mixture LP 10%) the reaction. However, dilution of binder can also lead to the reduction on the intensity and the time to reach the reaction peak, especially when the replacement level is high ($\geq 15\%$), as shown in Fig. 3. In PC based system, the incorporation of LP has been reported to show acceleration effect on the cement hydration [13], and both intensity and time to reach reaction peak (TTRP) were improved when PC/C$_3$S were blended with LP. The similar results were obtained by Zajac et al. [40] who studied the influence of limestone on the hydration of PC, and found that a mixture with 4% of LP addition showed a slightly faster and intensive reaction than pure PC mixture. As
shown in Fig. 3, the acceleration effect of adding LP on the reaction was found in the sodium carbonate activated slag-LP blends when the LP content is 5%. When increasing the LP content to 10% or higher, different behaviours were observed between PC and sodium carbonate activated system. This will potentially affect the reaction products as well as other hardened behaviour such as strength, and this will be discussed in the following sections.

3.3 Compressive strength

The compressive strength of the mixtures at different curing ages was measured and the results are shown in Fig. 4. The strength in overall ranges between 32.9 and 38.1, 51.5 and 54.5 and 62.0 and 67.6 MPa at the curing ages of 7, 28 and 180 days, respectively. At the early stage, the mechanical properties of the mixtures increase with the increase of LP content up to 10% and then slightly decline. From the long-term strength development point of view, samples containing LP up to 30% are slightly improved compared to mixtures without LP. It is remarkable that the incorporation of LP does not weaken the mechanical strength of the samples, e.g. the compressive strength of mixture containing 30% of LP is only 2.4% lower than the reference sample (28 days).

LP has been widely applied in Portland cement based materials and its effect on the mechanical properties of PC has been intensively studied [12–15]. Kumar et al. [12] studied the replacement levels (up to 50%) of LP on the strength development PC type I/II and found that the compressive strength of samples at different ages is roughly linearly proportional to the PC content. For example, when the PC was replaced by 30% LP, the compressive strength of the mixture at
28 days is decreased from about 44 MPa (reference without LP) to about 30 MPa, namely a relative decrease of about 32%. Kumar et al. [41] investigated the influence of the LP fineness on the PC hydration and reported that after 28 days of curing all samples containing LP (15% replacement) show similar compressive strength and the strength is about 13% lower than the reference samples. Ramezanianpour [10] reported the mechanical properties of PC based samples decrease with the increase of LP replacement levels. Nevertheless, samples with up to 10% LP replacement show comparable properties to PC concrete.

For the alkali activated slags (AAS), since the slags do not contain the ingredients C₃ₐ or C₃ₛ, as a result, theoretically the benefits of incorporating limestone in the AAS are mainly based on the inert filler effect, i.e. providing extra specific surface area acting as nuclei sites. Abdalqader et al. [42] characterized the mixtures of sodium carbonate activated slag-fly ash blenders, and found that the effect of fly ash on the strength development at the early ages is not prominent. Gao et al. [18] studied the properties of alkali activated slag-fly ash blends with limestone addition using waterglass as activators, and found that increasing the LP content with a constant slag content leads to a slightly better strength development. However, in both studies, the main reaction products remain the same and no trace of chemical involvement was observed.

As for sodium carbonate activated slag, due to the initial precipitation of calcium carbonate at the very early stage [2, 4, 26], the incorporated limestone powder (calcium carbonate) can potentially influence the reaction process as they share the same chemical composition, affecting the strength development. It is certain that LP can provide extra surface area and act as nuclei sites, accelerating the reaction of the specimens. However, the dilution effect normally plays a more important role leading to a slower strength development for instance in Portland cement based system. The possible reasons for the strength enhancement of sodium carbonate activated slag blended with LP (optimal LP replacement 10%) will be discussed in the following sections.

3.4 XRD

Figure 5 shows the XRD patterns of the mixtures at different curing ages. Three main phases, calcite (PDF #47-1743), gaylussite (PDF #74-1235) and hydrotalcite (PDF # 14-0191), can be identified for all the samples at the curing age of 7 and 28 days, respectively. The reflections of the calcite in the mixtures containing LP are intensive while the intensities of other crystalline phases are relatively weak. The intensities of the calcite increase with the increasing addition of LP in the mixtures, and no other polymorphous of calcium carbonate, i.e. aragonite and vaterite, are detected, possibly due to their low contents or poor crystallization.

Gaylussite is identified as a secondary reaction product for all samples prepared with different contents of LP at the early ages. No clear difference can be observed on the intensity of the produced gaylussite. When approaching the curing age of 28 days, the intensities of gaylussite for all the mixtures decrease, especially for the samples with high levels of LP replacement (over 15%). After 180 days of curing, the reflections of gaylussite are vanished, indicating it has dissolved and converted to other reaction products [4, 43]:

$$\text{NaCa(CO}_3\text{)}_2 \cdot 5\text{H}_2\text{O} = \text{CaCO}_3 + 2\text{Na}^+ + \text{CO}_3^{2-} + 5\text{H}_2\text{O}$$ (1)

Bernal et al. [4] reported that the entire gaylussite seems to convert to more stable products after 180 days of curing. Kovtun et al. [29] found that the intensity of gaylussite decreased in all accelerated pastes using accelerators such as sodium hydroxide and cement. The formation of hydrotalcite is also observed at the early ages but with only weak reflections (Fig. 5a), and with the ongoing reaction the intensity is increasing. Furthermore, the intensity of the hydrotalcite is improved after 28 days of curing, indicating the continuing alkali activation.

It should be noted that after 28 days of curing, the formation of natron is clearly identified in the mixtures containing over 15% LP, while at the curing age of 7 days only very weak reflections at around 30.8° are shown, indicating that the incorporation of LP chemically affects the reaction process. The potential mechanisms will be discussed in Sect. 4.

3.5 TG-DTG

Figure 6 presents the TG-DTG results of mixtures at the curing ages of 7, 28 and 180 days, and all samples
Fig. 5 XRD patterns of the mixtures at different curing ages as versus the LP content (strongest calcite peak is truncated for clarity)
Fig. 6 TG-DTG results of the mixtures versus the LP content at the curing ages of 

(a) 7 days, (b) 28 days and (c) 180 days, respectively (the mass losses of mixture LP 30% for \( \Delta m_2 \) (400–560 °C) and \( \Delta m_3 \) (560–700 °C) are calculated slightly differently depending on the DTG results)
possess similar patterns by showing 3 main peaks. The first mass loss before 200 °C is caused mainly by the removal of free water and partly by the dehydration of C–(A)–S–H gel [23]. It should be noted that there are several other small separate peaks shown in the low temperature range of 20–200 °C, while in the previous researches only one broad peak centred at around 105 °C is shown [23, 42]. Considering the XRD results, these peaks can potentially be attributed to the dehydration of sodium carbonate salts. For example, natron (Na2CO3·10H2O) is stable at room temperature but dehydrates to sodium carbonate heptahydrate, Na2CO3·7H2O, at 32 °C and then to sodium carbonate monohydrate, Na2CO3·H2O, at above 38 °C. Furthermore, most mixtures show similar trends and intensities at the low temperature range, while the mixture with 30% LP replacement shows an intensive peak at around 50 °C after 28 days of curing, indicating the dehydration of natron. The existence of natron in the reaction products is also confirmed by the XRD results as shown in Fig. 4 for the samples with a high content of LP (≥15%).

The decomposition of hydrotalcite can be generally classified into three stages [44–46]: (1) removal of interlayer water before 200 °C, (2) decomposition of structural hydroxyl groups (∼350 °C) and finally (3) the decomposition of interlayer carbonate anions (∼440 °C). According to the DTG pattern, in this case, the second decomposition peak appearing at around 310 °C can be assigned to the dehydroxilation of brucite-like layers of hydrotalcite and the small shoulder at 400–560 °C could be the decarbonation of hydrotalcite. The decarbonation of the other phases happens at around 600–700 °C, including: calcite, LP and gaylussite after its dehydration happening at approximate 130 °C [47]. In addition to these main peaks, the tiny peak centred at about 950 °C can be regarded as the thermal decomposition of sodium carbonate [48].

Table 4 summarizes the mass loss of mixtures at different temperature ranges, representing different reaction products as discussed above. As can be seen, at the early age (7 days), Δm1 shows that higher contents of hydrotalcite-like phases are generated in the mixtures with lesser LP addition as more Mg/Al are available from the slag particles. However, at advanced curing ages (≥28 days), mixture with 5% LP gives the highest content of hydrotalcite-like phases while the differences among the mixtures are small. It is noteworthy that the mixture with LP 30% only contains 70% slag of the reference mixture. Therefore, the amount of hydrotalcite-like phases should be approximately 70% of the reference assuming a similar reaction degree, while the determined value is 99%, indicating a much higher reaction degree of slag particles in this condition. Besides, a clearly increased mass loss between 400–560 °C (Δm2) and 560–700 °C (Δm3) is observed for the decomposition of calcium carbonate minerals when increasing the LP content in the mixtures. It should be noted that a reduction on the calcite content from 7 to 28 days is observed and it then increases from 28 to 180 days in all mixtures.

3.6 FT-IR

Figure 7 presents the FT-IR spectra of samples with different LP contents at the curing age of 7 days. All mixtures are showing similar patterns by giving similar curves and peak positions while the intensities of the peaks vary, indicating that they all contain similar reaction products but with different contents. In general, corresponding with the peak at approximate 1640 cm⁻¹, the broad peak centred at around 3333 cm⁻¹ due to the scissor bending of molecular water can be considered as the H–O–H stretching vibrations, i.e. the existence of free water in the reaction products [49, 50]. The peaks centred at about 1410, 873 and 711 cm⁻¹ are assigned to be the vibrations of ν3[CO3²⁻], ν2[CO3²⁻] and v4[CO3²⁻],

| Mixture (%) | 7 days | 28 days | 180 days |
|------------|--------|---------|----------|
|            | Δm1    | Δm2    | Δm3      | Δm1    | Δm2    | Δm3      | Δm1    | Δm2    | Δm3      |
| Ref.       | 3.07   | 1.75   | 1.17     | 2.94   | 1.74   | 1.02     | 4.22   | 2.34   | 1.03     |
| LP 5%      | 2.98   | 1.84   | 2.79     | 3.29   | 1.87   | 2.68     | 4.33   | 2.33   | 2.92     |
| LP 10%     | 2.72   | 1.97   | 4.31     | 2.87   | 1.91   | 4.15     | 4.27   | 2.68   | 4.30     |
| LP 15%     | 2.66   | 2.60   | 5.74     | 2.87   | 2.14   | 5.11     | 4.24   | 2.94   | 5.66     |
| LP 30%     | 3.29   | 3.67   | 9.49     | 2.33   | 3.89   | 8.84     | 4.16   | 3.62   | 10.16    |
respectively, which can be attributed to the vibrations of activators used, initially precipitated calcium carbonate or generated carbonate salts such as gaylussite and hydrotalcite [28, 51, 52]. In general, the intensities of the peaks at around 873 cm$^{-1}$, the vibration of carbonate anions, are proportional to the content of LP in the designed mixtures.

The sharp peak at around 946 cm$^{-1}$ is generally observed for sodium carbonate activated slags of the generated C–(A)–S–H [25, 42], while for the water-glass activated slags the position of this peak shifts to a higher wavenumber between 950 and 1000 cm$^{-1}$ [52–54], pointing to a different chemical proportion (Si/Ca ratio) [51, 55, 56]. Nevertheless, the LP incorporation does not affect the peak position, indicating that the gel structure of C–(A)–S–H remains the same regardless of the LP dosages. However, it should be noted that the accuracy of using FTIR to check the gel structure and Ca/Si ratio at micro-level is not fully understood yet and still open for discussion. To reach a more convincing conclusion, further research is still needed.

4 Discussion

The delayed reaction process is caused not only by the dilution effect, but also by the chemical effects. When having a lower slag content, the mixtures will take more time to release sufficient Ca$^{2+}$ to be precipitated with CO$_3^{2-}$ anions forming CaCO$_3$, lowering the CO$_3^{2-}$ concentration as the reaction is controlled by the concentration of CO$_3^{2-}$ anions in the pore solution [4]. Gaylussite (Na$_2$Ca(CO$_3$)$_2$·5H$_2$O) is a secondary reaction product of sodium carbonate activation and will be entirely transferred to more stable products (Eq. 1) [4, 28]. As shown in Table 4, the amounts of calcite of all samples after a long period of curing are remaining at the same levels, indicating that the released CO$_3^{2-}$ anions from gaylussite are transferred to other phases such as hydrotalcite-like phases. However, different from Rakhimova et al. [22], the formation of a new phase, natron (Na$_2$CO$_3$·10H$_2$O), is clearly observed for the samples containing ≥15% LP. In this case, it is most likely that insufficient ions are released from slag particles to react with CO$_3^{2-}$ anions when the dosage of LP is high, leading to the formation of natron. On the other hand, it is also reasonable to assume that the ions released by slag particles are largely consumed due to the presence of LP, indicating a higher reaction degree.

LP in general shows a negative effect on the strength development of PC based materials [9]. However, an improved strength development with up to 30% of LP addition is clearly observed for sodium carbonate-slag activation (Fig. 4). The results are in agreement with Rakhimova et al. [22], however,
they stated that this is mainly due to the “physical activity” of LP. It is clear that the incorporation of LP densifies the microstructure and provides extra surface area for the precipitation of reaction products, which benefits the strength development.

However, it should be noted that the molar volume of natron together with re-precipitated calcite is higher than that of gaylussite which can consequently refine the pore structure (especially for mixtures containing ≥15% LP). Besides, the transformation of natron to calcite during the long period curing (28–180 days) also releases free water to the hydration of slag particles and the gradual precipitated calcite is similar to the carbonation process [57] which can also further modify/alter the pore structure. On the other hand, another potential benefit of incorporating LP is that at advanced curing time the concentrations of both Ca²⁺ and CO₃²⁻ are low, then the hydrolysis of CaCO₃ can provide Ca²⁺:

$$\text{CaCO}_3 \Rightarrow \text{Ca}^{2+} + \text{CO}_3^{2-}$$

which will generate C–(A)–S–H gel while the CO₃²⁻ is consumed by the formation of carbonate-containing phases such as hydrotalcite. All of these contribute to the strength development.

In general, the chemical effects lead to an enhanced strength development, including the formation of natron originated from gaylussite and transition of natron to calcite. It is reasonable to assume that the formation of natron is mainly caused by the lack of sufficient ions from slag particles such as Mg²⁺ and Al³⁺ to react with the released CO₃²⁻ anions from the conversion of gaylussite. Abdelqader et al. [28] characterized the reaction products of sodium carbonate activated slag-fly ash blenders and reported that no new phase was formed with up to 25% addition of fly ash or difficult to be identified. While, in the present study, the presence of LP in the mixture is observed to be involved in the chemical reaction.

5 Conclusions

The effects of limestone powder on the workability, reaction kinetics, reaction products and strength development of sodium carbonate activated slag are investigated. Different analytical approaches are applied to study the microstructure and chemical phases. Based on the obtained results, the following conclusions can be drawn:

1. The flowability of the pastes increases with the increase of LP content till the mass replacement level of 15%.
2. The reaction is accelerated when the LP dosage is lower than 5% and then is delayed due to the dilution effect.
3. Considering the reaction kinetics and strength development, an optimal dosage 10% of LP replacement is concluded.
4. The presence of LP does not affect the main reaction products but chemically affects the decrystallization of gaylussite, resulting in the formation of a new phase especially when the dosage of LP is higher than 15%.
5. A relatively high content of hydrotalcite-like phases is generated for the mixtures containing LP.
6. The formation and transformation process of natron result in the enhanced mechanicals properties.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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