Experimental Investigation on Carbonation Behavior in Lime-Stabilized Expansive Soil

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The carbonation behavior of lime-stabilized expansive soil is important for assessing the stabilization efficiency from the perspective of durability. In this study, the accelerated carbonation tests, measurement of pH value distribution, and the free swell ratio tests were conducted to investigate the evolutions of carbonation depth, carbonation extent, and expansive potential of lime-stabilized expansive soil. XRD, MIP, and SEM techniques were adopted as supplements to reveal the carbonation mechanism. Results demonstrated that the carbonation depth of lime-stabilized expansive soil increased significantly as time elapsed; however, the rate of increase reduced when the carbonation time increased. Higher carbonation depth was obtained at higher temperature and CO2 concentration and lower relative humidity, which was described by an empirical model. Fully, partly, and noncarbonated zones were subsequently presented with an increase in the depth of the soil. The expansive potential of lime-stabilized expansive soil was partially recovered during carbonation. The obtained linear relationships between the free swell ratio and pH value were adopted to describe the evolution of expansive behavior with carbonation time and depth. In microstructural analysis, the conversion of portlandite into calcium carbonate was significant, which resulted in changes in microstructure and controlled the carbonation behavior.

1. Introduction

Expansive soil is found in tropical semiarid and moderate climatic regions and can trigger serious engineering hazards, such as foundation failure and slope instability, due to its swelling-shrinkage behavior and well-developed fissures during the water uptake/loss process [1–4]. Hence, the expansive potential of the soil is stabilized with the use of cement-based binding materials before constructions in these regions. For assessing the stabilization efficiency, the durability of the stabilized soil after being subjected to factors such as acid rain, sulfate attack, wetting-drying cycles, freezing-thawing cycles, and carbonation process is worthy of a detailed investigation [5–14].

Carbonation is one of the key durability properties that has attracted extensive attention in recent years. When the concrete or cement- and calcium-based materials are exposed to the atmosphere, CO2 will diffuse into the water or air phases in the pore space and reacts with portlandite (Ca(OH)2) and calcium silicate hydrate (C-S-H), to form calcium carbonate (CaCO3) [15–17]. Carbonation starts from the surface and propagates to the center of the material; therefore, the deep understanding of carbonation front and propagation rate, namely, the carbonation depth and rate, is critical in assessing the extension in service life of stabilized soil or a concrete structure [18, 19]. Based on literature review, Ta et al. [20, 21] summarized that the factors affecting carbonation depth and rate were chemical compositions and mix proportions in porous materials, along with temperature, relative humidity, concentration, and partial pressure of CO2 in the external environment [19, 22–24]. Cui et al. [17] performed accelerated carbonation tests to study the effect of CO2 concentration on concrete carbonation, where four stages were identified based on the CO2 concentration ranging from 2% to 100%. The increase in carbonation depth was significant when concrete was exposed to CO2.
concentrations below 20%; moreover, the effect reduced with any further increase in CO$_2$ concentration. Similar carbonation tests by Hyvert et al. [23] confirmed the effect of CO$_2$ pressure on the evolution of carbonation depth. Based on the distributed pH value along the height of the concrete mixture, Lu et al. [19] created fully and partly carbonated zones. Furthermore, although the carbonation depth increased gradually, the carbonation rate decreased with the carbonation time increase. The elevated temperature had minimal effect on the depth of the partly carbonated zones, but it instead resulted in greater depths for the fully carbonated zones. Additionally, the depths of both the fully and partly carbonated zones decreased with an increase in the relative humidity. For prediction of carbonation depth and rate of the porous materials under carbonation, simple empirical and enhanced physical models were proposed [18, 23, 25–27]. Investigations on carbonation mechanism showed that the formation of an insoluble salt of CaCO$_3$ resulted in clogged pores with reduced pore size, which not only changed the carbonation front and propagation rate [17, 23, 28], but also improved the strength and leaching characteristics of the stabilized material [6, 27, 28]. The significant effect of carbonation on the short-term and long-term strengths had been confirmed by Chen et al. [29], Nakarai and Yoshida [30], Du et al. [9], and Ho et al. [31], wherein the strength of cement-treated soil experienced an increase initially followed by a slight decrease over a long period. The microstructural analysis presented that Ca(OH)$_2$ was primarily produced by cement hydration and consumed by carbonation and pozzolanic reactions, whereas CaCO$_3$ was mostly provided by the carbonation of Ca(OH)$_2$ and C-S-H. The rapid increase in the strength initially was attributed to an increase in the amount of CaCO$_3$ provided by Ca(OH)$_2$. In contrast, under long-term conditions, increased Ca(OH)$_2$ provided by C-S-H decreased the strength. Bin-Shafique et al. [32] and Sweeney et al. [33] reported that carbonation could decrease the buffer capacity and change the chemical compositions of the stabilized soil, which can accelerate the release of well-stabilized heavy metals. However, Lange et al. [34], Bertos et al. [35], and Du et al. [9] were of contrasting opinions. They comprehensively performed the acid-neutralizing capacity, toxicity characteristic leaching procedure, and microstructural analysis to confirm the positive effects of carbonation on the leachability of heavy metals in the stabilized soil.

Carbonation behavior investigations into concrete and cement-treated soil have progressed since the 1980s, which include describing the carbonation process, assessing the carbonation effect, and displaying the carbonated mechanism. However, due to the large amounts of Ca(OH)$_2$ and C-S-H existing, forming, or participating during the stabilization process, the carbonation effect on lime-stabilized expansive soil is significant and worthy of investigation. Progressive carbonation, especially, may release the expansive potential of well-stabilized expansive soil, which may conversely influence the carbonation behavior due to changes in its microstructure. Until now, relevant researches on this issue have been rarely reported. Therefore, a series of tests, such as accelerated carbonation tests, pH value distribution measurements, free swell ratio tests, and microstructural analysis, are conducted in this study. Based on the results, the carbonation behavior in lime-stabilized expansive soil is described, the carbonation effect on the expansive potential is investigated, and the carbonation mechanism is revealed.

2. Materials and Methods

2.1. Materials and Specimen Preparation. The soils tested in this study were sampled from a construction site at a depth of 3–3.5 m in Hefei city, China. The physical properties of the soil were tested and are summarized in Table 1. The weak-medium expansibility for the tested soil can be identified. Based on X-ray fluorescence (XRF) analysis, the major chemical components were determined and listed in Table 2. Particle size distribution of the soil was tested using laser particle size analyzer and is presented in Figure 1. As can be seen, nonuniformity coefficient of 18.25 and curvature coefficient of 4.01 can be determined; silt content (2–20 μm) of 59.40% and clay content (<2 μm) of 40.62% were characterized for the tested soil. A maximum dry density of 1.63 g/cm$^3$ and an optimal water content of 24.05% were determined by the Proctor compaction test, in accordance with ASTM D698-12e2 (Figure 2). In this study, the lime purchased from a local building market in Hefei city, China, was used to stabilize the expansive soil. The chemical components of the lime were determined by performing XRF analysis, which were presented in Table 3.

When preparing the lime-stabilized expansive soil specimens, the mass ratio of lime to dry soil was set to 8% according to the excellent stabilization efficiency reported by Cheng et al. [36]. The oven-dried soil and lime were initially pulverized and sieved through 2.0 and 0.5 mm sieves, respectively. Based on the optimal water content (Figure 2), the target quantity of deionized water was sprayed on the lime-soil powder. After thorough mixing, the mixture was weighed and put in a ring mould. Static compaction was achieved with a piston at a rate of 0.1 kN/min to obtain target cylindrical specimens with 50 mm diameter, 100 mm height, and 1.55 g/cm$^3$ dry density, which was equivalent to 95% of the maximum dry density shown in Figure 2. The specimens were demoulded and wrapped with plastic bags and, then, cured under standard curing conditions (relative humidity of 95 ± 4%, temperature of 20 ± 2°C) for 28 days.

2.2. Accelerated Carbonation Test. An accelerated carbonation test was performed on the lime-stabilized specimen according to the Chinese standard GBT 50082-2009. A GangYuan TH-B environment-controlled carbonation chamber was used in the test as shown in Figure 3, where the temperature ($T$), relative humidity ($RH$), and carbon dioxide concentration ($C_b$) could be automatically controlled in the ranges of 15–30°C, 50%–90%, and 0%–20% with corresponding accuracies of ±2°C, ±4%, and ±3%, respectively.

The lime-stabilized specimens were extracted from the curing room after 28 days. One side of the specimen was chosen as the carbonated surface and was exposed to the
external environment, while the remaining five surfaces were coated with epoxy resin, to ensure unidirectional carbonation. Next, the prepared specimen was placed in the carbonation chamber with T as 15°C, 20°C, 25°C, and 30°C, RH as 60%, 70%, 80%, and 90%, and C\textsubscript{c} as 5%, 10%, 15%, and 20%, while being controlled subsequently. In each test exploring the influence induced by changes in one of the T, RH, and C\textsubscript{c}, the other two parameters should be kept constant at the standard carbonation condition (T as 20 ± 2°C, RH as 70 ± 4%, and C\textsubscript{c} as 20 ± 3%). During carbonation, the specimens were extracted from the carbonation chamber at regular intervals and split using a scroll saw. The freshly split surface was cleaned and sprayed with phenolphthalein, known as a typical pH indicator. The indicator used was a phenolphthalein solution 1% in ethanol with 1 g phenolphthalein and 90 mL of 95% (v/v) ethanol diluted in 100 mL of water [37, 38]. The phenolphthalein-alcohol solution was a colorless acid-base indicator, which turned purple when the pH value of the pore solution went above 9; hence, the carbonation front was easy to identify. Typical results obtained from this study are shown in Figure 4. The carbonation depth can be easily determined using the average value at five locations.

### 2.4. Free Swell Ratio Test

For investigating the expansive potential of lime-stabilized specimens undergoing carbonation, two additional stabilized specimens were prepared followed by experiencing standard carbonations. At carbonation times around 2d and 5d, the specimens were subsequently extracted and the corresponding carbonation depths were measured using the phenolphthalein indicator, which were 25 and 44 mm, respectively. Then, each of them was cut into five circular disks at every 20 mm height and then oven-dried at 105°C for 24 h. The samples were pulverized and sieved through a 0.5 mm mesh followed by the free swell ratio tests performed according to the Chinese standard JTG E40-2007.

### 2.5. Microstructural Analysis

Specimens were exclusively prepared and cured under standard curing conditions, followed by carbonation under standard conditions. After a carbonation time around 8d, the specimens were extracted and the corresponding carbonation depth measured using the phenolphthalein indicator was found to be 55 mm. Then, the pH value distribution of the specimen was measured using a pH meter, which was adopted to divide the carbonated zone into a fully carbonated zone (pH value < 9), partly carbonated zone (9 ≤ pH value < 11), and non-carbonated zone (pH value ≥ 11) according to previous researches. Finally, a microstructural analysis was performed in the different zones of the specimen. As shown in Figure 5, samples in fully carbonated and noncarbonated zones were extracted, pulverized, and sieved through a 0.5 mm mesh, to analyze the chemical components using X-ray diffraction (XRD). Samples with dimensions of 5 mm × 5 mm × 5 mm in fully carbonated and noncarbonated zones were adopted to analyze the mineralogical compositions and micromorphology using a scanning electron microscopy (SEM). Samples with dimensions of approximately 3 mm × 3 mm × 3 mm and masses between 1.5 and 2.0 g in fully carbonated, partly carbonated, and noncarbonated zones were adopted to analyze the pore size distribution using mercury intrusion porosimetry (MIP). The referred testing procedures related to XRD, MIP, and SEM are described in detail by Du et al. [9] and Shah et al. [40].

### 3. Results and Discussion

#### 3.1. Evolution of Carbonation Depth of Lime-Stabilized Expansive Soil

Evolution of carbonation depth with time of the lime-stabilized expansive soil, with consideration to changes in temperature, relative humidity, and carbon dioxide concentration, are presented in Figure 6(a)–6(c), respectively.
As shown in Figure 6, the carbonation depth of the lime-stabilized expansive soil increases with time elapsing. The increasing rate is significant in the initial stage followed by a relatively stable trend over a long period of time. Similar results were previously observed through tests performed on normal concrete and fly ash concrete [19]. This is because of the fact that, owing to the CO₂ concentration gradient between the external and internal environment, CO₂ diffuses into the specimen and proceeds with time, during which carbonation occurs in the specimens involved in the interaction between the CO₂ and Ca(OH)₂, or C-S-H. The resulting products CaCO₃ will clog the soil pores and conversely impedes the CO₂ diffusion. Thus, the carbonation rate approaches a stable level with time [19, 28, 41].

Curves in Figure 6 also present that the higher temperature or CO₂ concentration in the environment results in a higher carbonation depth at a given carbonation time, while the higher relative humidity in the environment results in a lower carbonation depth at a given carbonation time. This is because a higher temperature can promote the CO₂ diffusion rate in a porous medium and decrease the degree of saturation of the specimen, which will conversely increase the diffusing capacity of CO₂. Similarly, the higher CO₂ concentration can directly improve the diffusing capacity of CO₂ due to the enhanced concentration gradient in the specimen, while the higher relative humidity increases the liquid saturation in the pore space and reduces CO₂ diffusion accordingly [17, 19, 23].

For the carbonation behavior in concrete or cement- and calcium-based materials, prediction of carbonation depth using simple empirical or enhanced models with implied physical mechanism was proposed in the past decades [18, 23, 26, 27]. For simplicity, the evolution of carbonation depth was normally regarded as a function of the square root of time, as shown in the following equation:

\[ x_c = k_c \cdot \sqrt{t} \]  

(1)

where \( x_c \) is the carbonation depth (mm); \( t \) is the carbonation time (d); \( k_c \) is the carbonation coefficient (mm/d⁰.⁵), which depends on the inherent properties of the material, as well as the external environment such as temperature, relative humidity, and carbon dioxide concentration.

Based on Figure 6, the relationships between carbonation depth and square root of carbonation time in terms of the effect of temperature, relative humidity, and carbon dioxide concentration are shown in Figures 7(a)–7(c), respectively. Linear relationships between the carbonation depth and square root of carbonation time can be observed for conducting tests under different conditions. The slope of a line indicating the carbonation coefficients \( (k_c) \), which are dependent on the test conditions, is summarized and listed in Table 4. Based on this, the relationships between carbonation coefficient and temperature, relative humidity, and carbon dioxide concentration are plotted in Figures 8(a)–8(c), respectively.

As shown in Figure 8, with an increase in temperature and carbon dioxide concentration, and a decrease in relative humidity, the carbonation coefficient is linearly increased. Based on the superposition principle, the carbonation coefficient can be expressed as

\[ k_c = M + m \cdot T + n \cdot RH + l \cdot C_c. \]  

(2)

where \( T \) is the temperature (°C); \( RH \) is the relative humidity (%); \( C_c \) is the carbon dioxide concentration (%); \( M \) is the intercept and \( m, n, l \) are the slopes of the linear relationship between the carbonation coefficient and the temperature, relative humidity, and carbon dioxide
concentration, respectively. Based on the results of Figure 8, $m = 0.3358$, $n = -0.5198$, and $l = 0.582$ can be obtained when $M$ is equal to 35.91, 36.15, and 38.07, respectively. Due to the minimal difference, an average value of $M$ equal to 36.71 is adopted in the present study for simplicity. Finally, evolution of carbonation coefficient with temperature, relative humidity, and carbon dioxide concentration can be obtained as shown in the following equation:

$$k_c = 36.71 + 0.3358 \cdot T - 0.5198 \cdot RH + 0.582 \cdot C_c.$$

(3)

Furthermore, by substituting Equation (3) with Equation (1), evolution of carbonation depth with time can be predicted using the following equation:

$$x_c = \left[36.71 + 0.3358 \cdot T - 0.5198 \cdot RH + 0.582 \cdot C_c\right] \cdot \sqrt{t}.$$

(4)

3.2. Evolution of Carbonation Extent of Lime-Stabilized Expansive Soil. The carbonation extent of lime-stabilized expansive soil can be reflected by the pH value distribution measured using a pH meter. The results for specimens with carbonation depths of 25 and 44 mm are presented in Figure 9. The referenced pH value distribution for fully carbonated and noncarbonated specimens is also shown in the figure.

Curves in Figure 9 show nonuniform distributions of pH value in the carbonated specimens. At the top surface, pH values are significantly lower than 9, even approaching 6, indicating the fully carbonated zones of the specimen. With an increase in depth from the top surface, pH value increases and becomes stable with values almost equal to 11.4, indicating the noncarbonated zones of the specimen. Additionally, it is clear that specimens experiencing a longer carbonation time have lower pH values at a given depth. These evolutions of pH values were attributed to the conversion of $\text{Ca(OH)}_2$ and C-S-H to $\text{CaCO}_3$, which imply that

| Chemical component | CaO | MgO | SiO$_2$ | Al$_2$O$_3$ | Fe$_2$O$_3$ | K$_2$O | LOI | Total |
|-------------------|-----|-----|---------|-------------|-------------|--------|-----|-------|
| Content (%)       | 73.12 | 1.89 | 0.62    | 0.28        | 0.23        | 0.05   | 22.48 | 98.67 |

| Test type | Carbonated specimen | Carbonated zone |
|-----------|---------------------|-----------------|
| XRD MIP SEM | Partially carbonated zone (9 ≤ pH value < 11) |
| MIP | Partially carbonated zone (9 ≤ pH value < 11) |
| XRD MIP SEM | Noncarbonated zone (pH value ≥ 11) |

Table 3: Major chemical components of the used lime.
the carbonation extent reduces with an increase in carbonation depth and a decrease in carbonation time [13–17].

Based on carbonation depths of 0, 25, and 44 mm measured using the phenolphthalein indicator and temperature of 20°C, relative humidity of 70%, and carbon dioxide concentration of 20%, the accurate carbonation time for these tests is determined as 0 d, 1.79 d, and 5.55 d using Equation (4). Therefore, the evolution of carbonation extent (reflected by pH value) with time can be quantitatively plotted as shown in Figure 10.

As shown in Figure 10, carbonation on lime-stabilized expansive soil does not show uniform propagation. First, the top zone of the specimen is carbonated and it reaches the fully carbonated zone quite rapidly in a short time. However, with an increase in depth, from the top of the specimen, the carbonation extent becomes limited, especially at the bottom zone; thus, insignificant carbonation is presented. Consequently, CAO3 resulting from the carbonation blocks the soil pores, leading to a denser structure of the specimen. Subsequently, CO2 diffusion can hardly proceed in a further depth of the specimen. As a result, the carbonation extent is limited during the long-term exposure to CO2 [17, 19, 28, 41]. Furthermore, curves shown in Figure 10 indicate that the carbonation extent is a function of depth and time. On this basis, the evolution of the carbonation extent (reflected by pH value) can be quantitatively illustrated, as in Figure 11.

Figure 6: Evolution of carbonation depth with time of the lime-stabilized expansive soil, effect of (a) temperature, (b) relative humidity, and (c) carbon dioxide concentration.
3.3. Evolution of Expansive Potential of Lime-Stabilized Expansive Soil. The evolution of expansive potential for specimens with carbonation depths of 25 and 44 mm is shown in Figure 12.

As shown in Figure 12, the expansive potential of the specimen recovers after carbonation. A high free swell ratio of 30% can be observed at the top zone of the specimen. With an increase in depth from the top of the
and reduces the density of the soil. Additionally, the hydration products including Ca(OH)$_2$ and C-S-H into CaCO$_3$ releases the expansive potential of the soil. However, in carbonation, the conversion of Ca(OH)$_2$ and C-S-H gel, and enhanced the density of the lime-stabilized expansive soil. Conversely, influence the carbonation behavior. To reveal the carbonation mechanism, XRD, MIP, and SEM were performed in different carbonated zones. The results are presented in Figures 15–17.

As shown in Figure 13, the recovery of expansive potential increases with an increase in the carbonation time. This increasing trend is significant at the top of the specimen. With an increase in depth from the top surface, the increase in expansive potential turns limited. This phenomenon indicates that carbonation has a deteriorative effect on the stabilization efficiency of the lime-stabilized expansive soil. However, this deterioration effect is not uniformly distributed in the specimen. The stabilization efficiency can be guaranteed at a depth far away from the carbonated surface, which may be attributed to the reduced carbonation effect due to pore clogging and shrinkage during carbonation.

3.4. Microstructural Analysis of Carbonation Mechanism. Carbonation behavior is normally accompanied by mineralogical and microstructural changes in the specimen, which conversely influence the carbonation behavior. By performing the MIP tests, the measured pore size distributions in noncarbonated, partly carbonated, and fully carbonated zones are shown in Figure 16, confirming a significantly reduced pore volume after carbonation. Although the amount of medium-sized pores, of around 0.5 μm, increases slightly after carbonation, the amount of pore space decreases significantly. On the other hand, the amount of micropores increases due to the shrinkage of the specimen.

### Table 4: Carbonation coefficients obtained during accelerated carbonation test.

| Test condition                     | $k_c$ | $R^2$  |
|------------------------------------|-------|--------|
| Temperature, 15°C                  | 16.22 | 0.9772 |
| Temperature, 20°C                  | 17.98 | 0.9824 |
| Temperature, 25°C                  | 19.29 | 0.9909 |
| Temperature, 30°C                  | 21.38 | 0.9863 |
| Relative humidity, 60%             | 23.50 | 0.9972 |
| Relative humidity, 70%             | 18.34 | 0.9815 |
| Relative humidity, 80%             | 11.96 | 0.9827 |
| Relative humidity, 90%             | 8.30  | 0.9820 |
| Carbon dioxide concentration, 5%   | 10.84 | 0.9618 |
| Carbon dioxide concentration, 10% | 15.07 | 0.9746 |
| Carbon dioxide concentration, 15% | 16.81 | 0.9730 |
| Carbon dioxide concentration, 20% | 19.96 | 0.9765 |

 specimen, the recovered expansive potential becomes limited. For a given depth, the specimen with a carbonation depth of 44 mm owns a higher recovery of the expansive potential than that with a carbonation depth of 25 mm. From the results shown in Figure 12, it can be concluded that an excellent stabilization efficiency is guaranteed only for depths above 70 mm for specimens with a carbonation depth of 25 mm. Possible reasons could be summarized that, in initial stabilization, the complicated physicochemical process, for example, pozzolanic reactions, constrained the expansive potential and enhanced the density of the lime-stabilized expansive soil. However, in carbonation, the conversion of Ca(OH)$_2$ and C-S-H into CaCO$_3$ releases the expansive potential and reduces the density of the soil. Additionally, the formation and growth of CaCO$_3$ in carbonation would have an expansive ratio of approximately 11.7%, which can increase the solid volume in the specimen as well.

Based on Figures 9 and 12, it can be seen that a higher carbonation extent (a lower pH value) indicated a higher expansive potential of the specimen. Therefore, a relationship between the free swell ratio and the previously determined pH value can be plotted as shown in Figure 13.

As shown in Figure 13, linear relationships exist between the free swell ratio and pH value for specimens with different carbonation depths (time), confirming a strong dependency of expansive potential on carbonation extent. Theoretically, a unique fitted line can be obtained for carbonated specimens with different carbonation depths, a scenario not shown in Figure 13. This discrepancy may be attributed to the different microstructures in specimen preparation and the continuous microstructural changes caused by carbonation. The detailed reasons to this discrepancy could be explored in further researches. However, for qualitative and simple investigation of the swelling potential of the specimen during carbonation process, a fitted line of the average values is obtained through the collected data, which is shown in Figure 13 and expressed in the following equation:

\[
\delta_{sf} = 53.61 - 3.95 \times \text{pH.}
\] (5)

Here, $\delta_{sf}$ is the free swell ratio of the carbonated specimen (%); pH is the pH value measured using a pH meter.

Based on Equation (5) and the results shown in Figure 10, evolution of free swell ratio with carbonation time and depth for lime-stabilized expansive soil is presented in Figure 14.

As shown in Figure 14, the recovery of expansive potential increases with an increase in the carbonation time. This increasing trend is significant at the top of the specimen. With an increase in depth from the top surface, the increase in expansive potential turns limited. This phenomenon indicates that carbonation has a deteriorative effect on the stabilization efficiency of the lime-stabilized expansive soil. However, this deterioration effect is not uniformly distributed in the specimen. The stabilization efficiency can be guaranteed at a depth far away from the carbonated surface, which may be attributed to the reduced carbonation effect due to pore clogging and shrinkage during carbonation.

![Table 4: Carbonation coefficients obtained during accelerated carbonation test.](image)

![Figure 12](image)

![Figure 13](image)

![Figure 14](image)
Figure 8: Relationships between carbonation coefficient and (a) temperature, (b) relative humidity, (c) carbon dioxide concentration.

Figure 9: Distribution of pH value in the carbonated specimen.
Figure 10: Evolution of carbonation extent with time of lime-stabilized expansive soil.

Figure 11: Evolution of carbonation extent of lime-stabilized expansive soil (using MATLAB). (a) Carbonation time-0 d. (b) Carbonation time-1 d. (c) Carbonation time-2 d. (d) Carbonation time-3 d. (e) Carbonation time-4 d. (f) Carbonation time-5 d.
small-sized and large-sized pores notably reduces during carbonation. This phenomenon is consistent with the conclusions reached by Cui et al. [17], Du et al. [9], and Ho et al. [31], which observed reduced pore volume for specimens experiencing carbonation, further leading to improved strength and leaching properties of the specimen. Morandeau et al. [42] and Cui et al. [17] indicated that the formation and growth of CaCO_3 during carbonation clogged the pores and reduced their sizes. As a result, the evolution of the carbonation depth and carbonation extent could be delayed during long-term periods, as shown in Figures 6 and 10.

Mineralogical compositions and micromorphology measured in noncarbonated and fully carbonated zones are shown in Figure 17. Evidently, the plain-like Ca(OH)_2 and amorphous C-S-H gel can be observed in lime-stabilized expansive soil (Figure 17(a) and 17(b)), while CaCO_3 with typical cuneiform crystal structure can be observed in the specimen experiencing carbonation. Interparticle pores are clogged by the produced CaCO_3, resulting in enhanced densification of the specimen as shown in Figure 17(c). Such microstructural changes may delay the evolution of carbonation depth and carbonation extent as shown in Figures 6 and 10.
4. Conclusions

For investigating the carbonation behavior of lime-stabilized expansive soil, the accelerated carbonation test, pH value distribution measurement, free swell ratio, and microstructural analysis were conducted. The conclusions drawn from the study are as follows:

(1) Carbonation depth of lime-stabilized expansive soil increased significantly with time; however, the increasing rate became stable over a long period of time. Higher temperature and CO₂ concentration along with lower relative humidity in the environment resulted in deep carbonation depths. Evolution of carbonation depth with time was predicted by an established empirical model, in which the influences of changes in temperature, relative humidity, and carbon dioxide concentration were considered.

(2) Carbonation extent in lime-stabilized expansive soil was reflected by the measured pH value distribution, through which the nonuniform distribution of carbonation extent was observed. The pH values were significantly lower than 9 (even approaching 6) at the top surface, indicating the fully carbonated zone of the specimen. With an increase in depth, the pH value also increased and became stable at a value of 11.4, indicating the noncarbonated zone in the specimen. Additionally, specimens experiencing longer carbonation time owned a higher carbonation extent at a given depth.

(3) The expansive potential of lime-stabilized soil was recovered after carbonation. A high free swell ratio of 30% was observed at the top zone of the specimen. With an increase in depth from the top of the specimen, the recovered expansive potential became
limited. At a given depth, the expansive potential was enhanced with an increase in the carbonation time.

(4) The acquired linear relationships between the free swell ratio and pH value confirmed a strong dependency of expansive potential on the carbonation extent. The evolution of expansive behavior with carbonation time and depth was finally described based on the measured pH value distribution.

(5) Microstructural analysis confirmed that the conversion of portlandite into calcium carbonate controlled the carbonation behavior in lime-stabilized expansive soil. The total pore volume significantly reduced after carbonation. Although the amount of medium-sized pore, of around 0.5 μm, increased slightly after carbonation, the amount of small-sized and large-sized pores notably reduced during carbonation.

Data Availability

The data used to support the findings of this study are available from the corresponding author upon request.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

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