Multi-scale Laboratory Evaluation of the Physical, Mechanical and Microstructural Properties of Soft Highway Subgrade Soil Stabilized with Calcium Carbide Residue

| Journal: | Canadian Geotechnical Journal |
| Manuscript ID | cgj-2015-0245.R1 |
| Manuscript Type: | Article |
| Date Submitted by the Author: | 13-Aug-2015 |
| Complete List of Authors: | Jiang, Ning-Jun; University of Cambridge, Department of Engineering Du, Yan-Jun; Southeast University, Institute of Geotechnical Engineering Liu, Songyu; Southeast University, Institute of Geotechnical Engineering Wei, Ming-Li; Southeast University, Institute of Geotechnical Engineering Horpibulsuk, Suksun; Suranaree University of Technology, Civil Engineering Arulrajah, Arul; Swinburne University of Technology |
| Keyword: | calcium carbide residue, multi-scale, pozzolanic reaction, soil stabilization |
Multi-scale Laboratory Evaluation of the Physical, Mechanical and Microstructural Properties of Soft Highway Subgrade Soil Stabilized with Calcium Carbide Residue

Ning-Jun Jiang
PhD candidate, Department of Engineering, University of Cambridge, UK. Formerly, PhD student in Institute of Geotechnical Engineering, Southeast University, China.

Yan-Jun Du *
Professor, Institute of Geotechnical Engineering, Southeast University, Nanjing 210096, China
(*Corresponding author. PH/FAX: 86-25-83795086, E-mail: duyanjun@seu.edu.cn)

Song-Yu Liu
Professor, Institute of Geotechnical Engineering, Southeast University, Nanjing 210096, China.

Ming-Li Wei
PhD candidate, Institute of Geotechnical Engineering, Southeast University, Nanjing 210096, China.

Suksun Horpibulsuk
Professor, School of Civil Engineering, Suranaree University of Technology, Nakhon Ratchasima 30000, Thailand.

Arul Arulrajah
Professor, Centre for Sustainable Infrastructure, Swinburne University of Technology, Victoria 3122, Australia.

A Research Article Submitted for Possible Publication in

Canadian Geotechnical Journal
Abstract: Calcium carbide residue (CCR) is an industrial by-product, stockpiles of which are rapidly accumulating worldwide. Highway embankment construction has been identified as an avenue to consume huge quantities of CCR as an economical, less energy intensive and environmental-friendly chemical additive for soil stabilization. Previous studies have investigated the mechanical behavior of soils stabilized by CCR or blends of CCR with other additives; however, interpretation of the macro-scale geomechanical behavior of CCR stabilized soft soils from a systematically microstructural observation and analysis is relatively unknown. This paper presents a multi-scale laboratory investigation on the physical, mechanical and microstructural properties of CCR stabilized clayey soils with comparison to quicklime stabilized soils. Several series of tests were conducted to examine the Atterberg limits, particle size distribution, compaction characteristics, unconfined compressive strength, California-Bearing-Ratio and resilient modulus of the CCR stabilized clayey soils. The influences of binder content, curing time, and initial compaction state on the physical and mechanical properties of treated soils are interpreted with the aids of physicochemical and microstructural observations including soil pH, soil mineralogy obtained from X-ray diffraction and thermogravimetric analysis, and pore size distribution obtained from mercury intrusion porosimetry. Soil particle flocculation and agglomeration at the early stage and pozzolanic reactions during the entire curing time, which originate from the finer particle size, greater specific surface area and higher pH value of calcium carbide residue, are the controlling mechanisms for the superior mechanical performance of CCR stabilized soils. The outcomes of this research will contribute to the usage of CCR as a sustainable and alternative stabilizer to quicklime in highway embankment applications.

Key Words: calcium carbide residue; multi-scale; pozzolanic reaction; soil stabilization
Calcium carbide residue (CCR) is a by-product of polyvinyl chloride (PVC), polyvinyl alcohol and acetylene production. CCR is formed through the hydrolysis of calcium carbide, as shown by the following equation (Jaturapitakkul and Roongreung 2003):

\[
\text{CaC}_2 + 2\text{H}_2\text{O} = \text{C}_2\text{H}_2 + \text{Ca(OH)}_2
\]

The dominant component of CCR is Ca(OH)$_2$, with limited amounts of calcium carbonate (CaCO$_3$), SiO$_2$, and trace components of sulfide, metal oxide and organic matters (Kampala and Horpibulsuk 2013). CCR is presently widespread in developing and developed countries alike (Sharma and Reddy 2004; Du et al. 2011, 2015a; Horpibulsuk et al. 2013b; Phetchuay et al. 2014). CCR usually appears as high-alkaline and high-moisture-content slurry. If not handled properly, CCR becomes a source of pollution to surface and underground water (Krammart and Tangtermsirikul 2004; Sharma and Reddy 2004). In recent years, the increasingly large production quantity of CCR due to growing demand has often resulted in serious environmental pollution with stockpile areas (Du et al. 2011). Reuse applications for CCR, particularly in large civil engineering infrastructure applications that can rapidly deplete these growing stockpiles, are urgently being sought. The usage of CCR as a sustainable cementitious binder for soil stabilization has been identified as a low-carbon and less energy intensive means to reuse this by-product and furthermore eliminate negative environmental connotations associated with stockpiling this by-product. Similar approaches have been successfully attempted in recent years for other geomaterials including geopolymer, phosphate-rich materials and demolition aggregates (Du et al. 2011, 2014b; Horpibulsuk et al. 2013b, 2014; Sukmak et al. 2013a, b, 2015; Arulrajah et al. 2014; Cai et al. 2015).

Soft clay deposits, which are widely distributed in East China regions, impose great challenges in the construction of infrastructure projects such as highway embankments (Liu et al. 2011;
Horpibulsuk et al. 2013b). These soft clay deposits are typically of low strength, low stiffness and low permeability, making them difficult to improve and compact. These difficulties often result in the highway embankments having low bearing capacity and furthermore susceptible to excessive settlements (Kodikara and Chakrabarti 2005; Chakrabarti and Kodikara 2007; Han et al. 2007; Gnanendran and Piratheepan 2010). Chemical stabilization is an effective method to improve the engineering properties of soft clayey soils (Shen et al. 2013). Huge amount of chemical additives are needed to stabilize the soft soils in-situ, given that the geometries of typical embankments are of large lengths and widths. The large-scale quantities of CCR by-products generated in Eastern China, creates the opportunity for the recycling of this industrial by-product in the construction of highway embankment (Du et al. 2011), particularly as soft soil deposits are largely prevalent in this region. The reuse of CCR in highway embankment construction is cost-economic and has no extra associated embedded energy consumption compared to conventional cement-based binders, making it an attracting alternative for project contractors and constructors (Horpibulsuk et al. 2012, 2013a; Du et al. 2015a). Similar to other chemical additives (e.g., Portland cement, fly ash and slag) for soft clay stabilization (Jin and Al-Tabbaa 2014; Du et al. 2015b), CCR is mixed with the parent soft clayey soil and reacts with clay minerals and water to improve the strength, stiffness and durability of the stabilized soil (Kampala and Horpibulsuk 2013; Du et al. 2011).

Previous studies focused on the use of CCR alone or the blend of CCR and other chemical additives in soft soils stabilization. For example, Kampala and Horpibulsuk (2013) examined the physical and engineering properties of a problematic silty clay stabilized with CCR and stated that CCR was more effective than lime in soft soil stabilization in terms of engineering, economic, and environmental perspectives. Horpibulsuk et al. (2013a) investigated the strength characteristics of a silty soil treated by the blend of CCR and fly ash and proposed a controlling strength development
mechanism based on different strength improvement zones. Vichan and Rachan (2013) systematically investigated the strength development patterns of the soft Bangkok clay stabilized by blends of CCR and biomass ash, and reported that the properties of both materials significantly affected the strength gains. Phetchuay et al. (2014) used CCR-alkali activated stabilized clay geopolymer as a sustainable pavement subgrade material and examined the influential factors for strength development. Most of these existing studies focused on the examination of engineering properties and associated influential factors affecting the strength of soils stabilized by CCR alone or with blends of CCR and other additives. The interpretation of macro-scale geomechanical behaviors of CCR stabilized soft soils from a systematically microstructural observation and analysis is however relatively unknown. Microstructural analytical methods in geotechnical engineering typically include X-ray diffraction (XRD), scanning electron microscopy (SEM), mercury intrusion porosimetry (MIP) and thermogravimetric analysis (TGA) (Mitchell and Soga 2005). These methods have been extensively employed to produce explicit microstructural supporting evidences for the hypothesis explaining geomechanical behaviors of soft soils or aggregates stabilized by conventional chemical additives (e.g., Portland cement and lime) (Locat et al. 1996; Al-Mukhtar et al. 2010; Stoltz et al. 2012; Du et al. 2014a; Mohammadinia et al. 2014;). For example, Wild et al. (1993) employed the XRD, SEM-EDAX and thermal analysis to investigate the chemical, morphological and microstructural changes occurring during moist curing and soaking of lime stabilized kaolinite. Lemaire et al. (2013) carried out a multi-scale studies of the cement-lime stabilized plastic silty soil and interpreted the mechanical properties from the physicochemical and microstructural perspective using SEM, XRD, and MIP. Similarly, multi-scale observations and analysis would be helpful to understanding the controlling mechanisms of soft clay stabilization by CCR.
This paper presents a multi-scale laboratory evaluation of the physical, mechanical and microstructural properties of CCR stabilized soft clayey soil. Several macro-scale series of tests were conducted to examine the Atterberg limits, particle size distribution (PSD), compaction characteristics, unconfined compressive strength ($q_u$), California-Bearing-Ratio (CBR) and resilient modulus ($M_r$) of the CCR stabilized clayey soils. Quicklime, which is used extensively in stabilizing highway subgrade materials, is selected as a control chemical additive for comparison purposes. The influences of binder content, curing time, and initial compaction state on the physical and mechanical properties of stabilized soils are interpreted with the aids of physicochemical and microstructural observations including the soil pH, soil mineralogy obtained from XRD and thermogravimetric analysis (TGA), and pore size distribution obtained from mercury intrusion porosimetry (MIP) analysis.

### Materials and Methods

#### Soils and binders

The soil used in this study was excavated from the site of West Changzhou Ring Expressway located in Changzhou City, Jiangsu Province, China. The basic physical and engineering properties of the soil are listed in Table 1. The soil is classified as a low plasticity clay (CL) based on ASTM D2487 (ASTM 2011a). The result of X-ray fluorescence (XRF) analysis of the parent soil indicates that it contains 67.9% of silicon dioxide ($\text{SiO}_2$), 14.1% of aluminum oxide ($\text{Al}_2\text{O}_3$), 5% of ferric oxide ($\text{Fe}_2\text{O}_3$), 2.5% of magnesium oxide ($\text{MgO}$), and 1.3% of calcium oxide ($\text{CaO}$) (see Table 2). The particle size distribution curve of the parent soil is shown in Fig. 1.

CCR used in this study was collected from Jiangsu Changzhou Changfei Acetylene Manufacturing Co. Ltd. Its basic physical and chemical characteristics are listed in Table 3 and its major chemical constituents are shown in Table 2. The CCR was air-dried prior to the specimen
preparation for various laboratory-scale tests due to its natural moisture content up to 60.9%.

Quicklime used in this laboratory test was produced by Liyang Shanghuang Yangzhu Tianfu Lime Manufacturing Station. Basic physical and chemical characteristics of the quicklime are listed in Table 3 and its major chemical constituents are presented in Table 2. This quicklime is classified as High-Calcium Lime based on ASTM C51-11 (ASTM 2011b). The particle size distribution curves of the CCR and quicklime are shown in Fig. 1.

**Sample preparation**

Prior to the series of test in this study, the standard Proctor compaction test were conducted to obtain the maximum dry density ($\rho_{\text{dmax}}$) and optimum water content ($w_{\text{opt}}$) of binder-amended soils right after mixture. For Atterberg limits and particle size distribution (PSD) tests, collected soils were air-dried before they were passed through the sieve with 0.5 mm opening size. Then, the air-dried soils were thoroughly mixed with predetermined amount of binders (i.e. CCR or quicklime) and water (approximately $w_{\text{opt}}$). The binder-amended soils were then cured in sealed vinyl bags at 20°C and relative humidity of 95% for 28 d before subjected to the Atterberg limits and PSD tests.

For other tests in this study, the air-dried soils were firstly prepared with predetermined amount of binders (i.e. CCR or quicklime) and water (approximately $w_{\text{opt}}$) in cylindrical iron molds ($\Phi 152 \times H 170$ mm for CBR and $M_r$ tests, and $\Phi 50 \times H 50$ mm for unconfined compression test; soils for pH, TGA and MIP tests were sampled from the $\Phi 50 \times H 50$ mm sample) via the static compaction method to achieve degree of compaction of 93%, 94% and 96%. All inner walls of molds were lubricated with Vaseline to reduce friction. All samples were subsequently cured at 20°C and relative humidity of 95% for 7 and 28 d, respectively, before testing. Table 4 presents the binder dosage, curing time, degree of compaction, and the number of identical samples for different tests in
this study. For unconfined compression, CBR, resilient modulus, and TGA tests, three identical samples were tested. The coefficient of variation (COV) for the results of unconfined compression test, CBR and resilient modulus tests are less than 8% and COV for TGA test is less than 4%, indicating excellent repeatability of the test results.

**Testing methods**

Atterberg limits were conducted according to ASTM D 4318 ([ASTM 2010](#)). PSD tests for the parent soil, CCR, quicklime, and CCR and quicklime stabilized soils were conducted using a Mastersizer 2000 laser particle size analyzer (Malvern Inc., U.K.). Prior to PSD analysis, both stabilized and unstabilized soil specimens were air-dried and grinded through a 0.3 mm sieve. Then, 15 g grinded specimen was mixed with sufficient distilled water and subjected to the PSD analysis. Standard Proctor compaction test was conducted with a standard compaction effect of 600 kN-m/m³ as per ASTM D 698 ([ASTM 2012](#)). The unconfined compression test was performed based on ASTM D4219 ([ASTM 2008](#)). The rate of vertical load remained 1mm/min until the failure of the specimen. It should be noted that the specimen size in this study (Φ50 × H50 mm) is slightly difference from the length-to-diameter ratio (2.5) recommended by ASTM D 4219. The CBR test was conducted according to ASTM D 1883 ([ASTM 2014](#)). A circular piston was used to intrude stabilized soils in a mold at a constant rate of penetration. The CBR was determined as the ratio of the unit load on the piston required to penetrate 2.5 mm or 5 mm of the test soil to the unit load required to penetrate a standard material of well-graded crushed stone. Resilient modulus test was conducted as per AASHTO T307 ([AASHTO 2007](#)), which was also previously adopted by Tastan et al. (2011). The specimen size in this study (Φ152 × H170 mm) is modified from that in AASHTO T307 (Φ102 × H203 mm). $M_r$ was calculated based on the ratio of deviator stress and the
recoverable strain. Different confining and deviator stresses were applied on the specimens to cover the range of expected stresses in the field.

Physicochemical and microstructural observational tests conducted in this study include soil pH, XRD, TGA and MIP. Soil pH measurement was carried out using a HORIBA pH/COND METER D-54 as per ASTM D4972 (ASTM 2013). Both stabilized and unstabilized soils were grinded through a sieve with 2 mm opening size. The liquid to solid ratio of 1.0 was used to mix the soil and distilled water. The pH of the slurry was then measured after 1 hr of retention. In order to examine the chemical and mineralogical compositions of CCR and quicklime stabilized soils, XRD test was conducted for stabilized soils with 6% binder content at 180 d. As a reference, unstabilized soil was also subjected to the XRD test. The XRD tests were performed using a Rigaku D/Max-2500 X-ray diffractometer. Cu-Kα (λ = 1.540538Å) X-ray tube with an input voltage of 40 kV and a current of 200 mA was utilized. Prior to the test, both stabilized and unstabilized soil specimens were freeze-dried and then grinded into a sieve with 0.038 mm opening size. The tests were carried out between two-theta values of 5 to 60° with a step length of 0.02° and a scanning rate of 2°/min.

Thermogravimetric analyses (TGA) was conducted by heating a test specimen continuously from room temperature to 750°C at a heating rate of 20°C/min in a nitrogen environment. In this study, TGA was performed using a differential scanning calorimeter (Perkin-Elmer Pyris 1). After designated curing periods of 28 and 120 d, 3 identical cubic samples (1 cm × 1 cm) were extracted and soaked in absolute ethyl alcohol for 96 hr to terminate the hydration. The specimens were then dried at 30 °C and grinded through a 200-mesh sieve. Approximate 30 ± 0.5 mg sieved specimens were used for the TGA test. The results of TGA are presented as a curve of the mass loss versus temperature. The first derivative of the mass loss curve is recorded as a function of time, which is
known as derivative thermogravimetric analysis (DTG).

The MIP test is based on the fact that mercury is a non-wetting fluid that has to be pressurized in order to penetrate a porous medium (Diamond 1970). In this method, all pores are considered to be of cylindrical shape and therefore the Jurin’s equation which calculates capillary pressure can be applied in MIP method:

\[
d = -\frac{4\tau \cos \theta}{p}
\]

where \(d\) is the diameter of the pore intruded, \(\tau\) is the surface tension of intruded liquid (i.e. mercury), \(\theta\) is the contact angle, and \(p\) is the applied pressure. In this study, MIP test was carried out using an AutoPore IV 9510 mercury intrusion porosimeter (Micromeritics Co. Ltd. USA). The maximum applied pressure is \(6 \times 10^4\) psi (i.e. 413 MPa) and the surface tension of mercury is \(4.84 \times 10^{-4}\) N/mm at 25°C (Mitchell and Soga 2005). The contact angle is taken as 135°. After curing periods, stabilized soil specimens were broken up to about 1 cm \(\times\) 1 cm cubes in a careful manner to eliminate disturbance. Liquid nitrogen was used to freeze the soil specimens, after which the specimens were placed in a freezing unit with a vacuum chamber, and were dried by sublimation of the frozen water at a temperature of –80 °C. (Penumadu and Dean 2000; Li and Zhang 2009).

Results and Analysis

Atterberg limits

The results of Atterberg limits tests are shown in Table 5. It is found that both liquid limit \((w_L)\) and plastic limit \((w_p)\) of CCR and quicklime stabilized soils are higher than those of unstabilized soils, regardless of binder dosage. In contrast, plastic index \((I_p = w_L - w_p)\) values of both CCR and quicklime stabilized soils decrease by approximate 15% as compared to those of unstabilized soils. Kinuthia (1999) and Du et al. (2014a) indicated that formations of flocculate and agglomerate
(short-term), and pozzolanic reactions (long-term) in lime stabilized soils were able to remarkably
modify $w_L$ and $w_p$. Due to the similarity in chemical components between CCR and quicklime, the
mechanism proposed by Kinuthia (1999) can explain the variations of Atterberg limits of both CCR
and quicklime stabilized soil with binder content and curing time in this study. The decrease in $I_p$
values with amendment of CCR or quicklime observed in this study is consistent with those
examined by previous studies (Locat et al. 1996; Du et al. 1999) which found that the addition of
chemical additives (such as lime) could result in an increase in both $w_L$ and $w_p$ but a reduction in $I_p$.

**Particle size distribution (PSD)**

The results of PSD test is shown in Table 6. After 28 d curing time, the addition of 4% and 6%
CCR leads to reduction of clay-sized particle percentage from 13.6% to 7.0% and to 3.8%,
respectively, which are higher than the cases of quicklime addition (from 13.6% to 10.0% and to
8.2%, respectively). The sand-sized particle percentage of CCR and quicklime stabilized soils
increase substantially from 2.5% to 37.8% and to 60.7 for 4% and 6% CCR, as well as from 2.5%
to 26.4% and to 26.7% for 4% and 6% quicklime, respectively. The changes in clay- and sand-sized
particle percentages in the stabilized soils are dominantly attributed to both short-term flocculation
formation and long-term pozzolanic reactions (Nalbantoglu and Tuncer 2001; Tran et al. 2014). The
flocculation contributes to agglomeration of fine particles (Nalbantoglu and Tuncer 2001; Kampala
et al. 2013) and the synthesized pozzolanic products would coat the surface of soil particles (Tran et
al. 2014), both making relatively high fraction of coarse-grained particles.

In comparison with quicklime, CCR stabilized soils exhibit 143% to 228% higher sand-sized
particle percentage after curing for 28 d. The clay-sized particle percentages of CCR stabilized soils
are 29% to 54% less than quicklime stabilized soils at 28 d, respectively. The observations suggest
that CCR yields a superior capability for modifying particle size distribution of the soils as compared to the quicklime.

Compaction characteristics

The results of standard compaction test show that when the binder content increases from 4 to 6%, $w_{opt}$ slightly changes from 13.4 to 14.1% for CCR and from 13.2 to 13.4% for quicklime, respectively. In contrast, $\rho_{d_{\text{max}}}$ reduces from 1.78 to 1.74 g/cm$^3$ and from 1.73 to 1.70 g/cm$^3$ for CCR and quicklime stabilized soils, respectively, as a consequence of increase in the binder content from 4 to 6%. The phenomena are consistent with those reported by Fahoum et al. (1996), who also observed the reduction of $\rho_{d_{\text{max}}}$ with increasing lime content for cohesive soils, and Kampala and Horpibulsuk (2013), who reported the reduction of $\rho_{d_{\text{max}}}$ with increasing CCR content for silty clay. The reason of the phenomena is predominantly attributed to the flocculation and agglomeration, as a consequence of cation exchange, resulting in material bulking (Kinuthia et al. 1999). The $\rho_{d_{\text{max}}}$ values of the CCR stabilized soil are always higher than those of quicklime stabilized soils at the binder contents of 4 and 6%, indicating that CCR stabilized soil can achieve a better compaction performance relative to the quicklime with the same dosage.

Unconfined compressive strength

Figure 2 shows the effects of curing time, degree of compaction and binder content on the $q_u$ of CCR and quicklime stabilized soils. The $q_u$ of unstabilized soils is also marked in Fig. 2 for comparison. It can be noticed that $q_u$ keeps increasing steadily with curing time, irrespective of the degree of compaction and binder content for both stabilized soils. More specifically, the CCR stabilized soil exhibits notable strength development at the initial 60 d than the subsequent 60 d,
regardless of the degree of compaction. However, the quicklime stabilized soil gains predominant strength at the initial 30 d with 94 and 96% degree of compaction. With 93% degree of compaction, the predominant strength development occurs at the initial 60 d. Moreover, it is evident that higher content of CCR or quicklime results in higher $q_u$ values for all the curing times tested. At the same binder content, the $q_u$ values of CCR stabilized soil are higher than those of the quicklime stabilized soil regardless of the curing time and degree of compaction. The strength development for CCR and quicklime stabilized soft clayey soils in this study is consistent with that reported by Kampala and Horpibulsuk (2013). The strength growth at early stage is attributable to the flocculation and agglomeration of the soil particles (Kinuthia et al. 1999) while the long-term strength development is determined by the pozzolanic reactions (Wild et al., 1993).

CCR and quicklime stabilized soils have a discrepancy in the effect of degree of compaction on the strength development. The $q_u$ values at 120 d for CCR stabilized soils are around 2250 kPa regardless of the degree of compaction. In contrast, dependence of $q_u$ on the initial compaction state is noticeable for quicklime stabilized soils cured for 120 d. For example, in the case of 6% amendment, the $q_u$ values at 120 d are 1600 and 2200 kPa for 93% and 96% degree of compaction, respectively. Le Runigo et al. (2009) stated that the impact of compaction energy on the pore size distribution of quicklime stabilized silty soil is margin, indicating a similar soil fabric even under different degree of compaction conditions. However, Osinubi (1998) showed that higher compaction energy was related to higher $q_u$ values for lime stabilized soils, which is consistent with the results of quicklime stabilized soil tested in this study. Since the binder contents adopted in this study are not higher than 6%, the CCR stabilized soils are in the active zone, as suggested by Horpibulsuk et al. (2013). In contrast, the quicklime stabilized soils are in the inert or deterioration zone as suggested by Bell (1996). In the active zone, the long-term $q_u$ of the CCR stabilized soil
increases with increasing binder content, which is due to the fact that all input portlandite is
consumed through pozzolanic reactions (Horpibulsuk et al. 2013). However, in the inert or
deterioration zone, the \( q_u \) of the quicklime stabilized soil ceases to increase or decrease with
increasing binder content, which is caused by internal-structure damage due to presence of
excessive free lime (Horpibulsuk et al. 2013). Therefore, when the stabilized soils are in different
zones (i.e., active zone and inert or deterioration zone), the effect of degree of compaction on their
\( q_u \) would be different. This may explain the discrepancy between CCR and stabilized soils in terms
of the effect of degree of compaction on the \( q_u \).

**California Bearing Ratio (CBR)**

Figure 3 presents the effects of curing time, binder content on CBR values with various degree
of compaction. The CBR values of both CCR and quicklime stabilized soils are dramatically higher
than that of the unstabilized soil, and they increase steadily with increasing curing time, due to the
flocculation and agglomeration of soil particles at early stage and pozzolanic reactions at long term
(Kinuthia et al. 1999). The CBR values of both CCR and quicklime stabilized soils increase with
increasing degree of compaction, which is consistent with that reported by Osinubi (1998). In
contrast, the binder content affects CBR values in a different manner for CCR and quicklime
stabilized soils. The CBR values of CCR stabilized soils at two binder contents (4 and 6%) are
practically the same, while the quicklime stabilized soil exhibits approximate 30% increase in the
CBR when the binder content increases from 4% to 6%.

Similar to the unconfined compression test results, the CCR stabilized soils possess much
higher CBR values relative to the quicklime stabilized soils, regardless of the binder content, curing
time and degree of compaction within the considered range. This result is consistent with that
obtained from the full-scale field trial tests (Du et al. 2015a), confirming that the CCR stabilization brings higher bearing capacity for the highway soft clayey soils relative to the quicklime. In addition, the measured CBR values also satisfy the requirement specified by the China highway construction standard (CBR ≥ 8%) (China MOT 2004).

**Resilient modulus (M<sub>r</sub>)**

**Figure 4** illustrates the development of M<sub>r</sub> for CCR and quicklime stabilized soils with curing time and degree of compaction. It is evident that CCR and quicklime stabilization leads to a significant increase in M<sub>r</sub>, as compared to the unstabilized soil. The M<sub>r</sub> values of both CCR and quicklime stabilized soils at 28 d are approximately two times those at 7 d. Elevation of degree of compaction from 94% to 96% results in approximate 20% increase in M<sub>r</sub> for the CCR stabilized soil, whereas only marginal increase in M<sub>r</sub> for the quicklime stabilized soil. The change of M<sub>r</sub> with curing time and degree of compaction is similar to that of q<sub>a</sub> (see Fig. 2) and CBR (see Fig. 3).

The M<sub>r</sub> values of the CCR stabilized soils are found to be remarkably larger than those of quicklime stabilized soils. At 7 d, the M<sub>r</sub> values of CCR stabilized soils with degree of compaction of 94% and 96% are 15% and 29%, respectively, higher than those of quicklime stabilized soils. This difference increases to 31% and 44% at 28 d for the degrees of compaction of 94% and 96%, respectively.

**Soil pH**

**Figure 5** shows the changes in measured soil pH values of CCR and quicklime stabilized soils with the curing time. It is found that pH values of both CCR and quicklime stabilized soils decrease gradually with increasing curing time, which is attributed to the pozzolanic reactions between
Portlandite (Ca(OH)$_2$) and reactive SiO$_2$/Al$_2$O$_3$ in the soil matrix expressed by the following equation (Kinuthia et al. 1999):

\[
\text{(3)} \quad \text{Ca(OH)}_2 + \text{SiO}_2 + \text{H}_2\text{O} \rightarrow \text{C-S-H}
\]

\[
\text{(4)} \quad \text{Ca(OH)}_2 + \text{Al}_2\text{O}_3 + \text{H}_2\text{O} \rightarrow \text{C-A-S-H}
\]

During the pozzolanic reactions, alkaline portlandite is gradually consumed and transformed to less alkaline secondary cementitious products (e.g., C-S-H and C-A-S-H), leading to reduction in soil pH (Kinuthia et al. 1999; Al-Mukhtar et al. 2010; Du et al. 2014a). It is noted that when pH of stabilized soils is higher than 10 to 11, the reactive SiO$_2$/Al$_2$O$_3$ in soil matrix would keep dissolving (Saride et al. 2010) while the formed secondary cementitious products are still thermodynamically stable (Stronach and Glasser 1997). Moreover, the pH of the CCR stabilized soil is higher than that of the quicklime stabilized one during the entire curing period. This is mainly due to the higher pH of CCR (12.84) than that of quicklime (12.74), as shown in Table 3.

**X-Ray diffraction (XRD)**

Figure 6 presents the XRD diffractograms of the stabilized and unstabilized soils. The minerals in the unstabilized soil are predominantly quartz, kaolinite and illite with trace montmorillonite. For the CCR stabilized soil, formation of C-S-H is detected at 2\(\theta\) of 27.5°, 28.3°, 52.9° and 54.5°, respectively. In contrast, no distinct peaks for portlandite are identified in the CCR stabilized soil. As C-S-H is the secondary cementitious products (see Eqs. (3) and (4)) formed in the pozzolanic reactions, the identified peaks of C-S-H and absence of portlandite in the CCR stabilized soil at 180 d confirms the completed progress of pozzolanic reactions. The XRD diffractogram of the quicklime stabilized soil reveals the formation of both C-S-H (27.5° and 52.9°) and calcite (13.1°, 29.4° and 36.0°). The formation of calcite is attributed to the air exposure of soil during the curing.
period and subsequent carbonation of quicklime and hydrated products (Verbrugge et al. 2011). The vulnerability of the quicklime stabilized soil to the carbonation, as compared to the CCR stabilized soil, primarily occurs at the early stage (Hunter 1988; Al-Mukhtar et al. 2012), during which quicklime stabilized soils develop relatively high porosity (which is demonstrated from the MIP test presented in the later section) and hence relatively large exposure area to carbon dioxide (CO$_2$) in the air.

**Thermogravimetric analysis**

Figure 7 shows the TGA and DTG results for the CCR and quicklime stabilized soils with 6% content at 28 and 180 d. The results of TGA are presented as a curve of the mass loss/first derivative of the mass loss versus temperature. Significant mass losses can be observed at a temperature of 50 to 200°C and 200 to 300°C from peaks in the DTG curves. The peaks in DTG curves (or mass losses in TGA curves) correspond to the presence of pozzolanic reaction products (C-S-H, C-A-H and C-A-S-H) during their thermal decompositions (HaHa et al. 2011). In addition, a sharp peak can be identified at the temperature of 425°C, which is confirmed to be portlandite (HaHa et al. 2011). Since the pozzolanic reaction products (viz. C-S-H, C-A-H and C-A-S-H) have relatively low degree of crystallinity, their contents could not be calculated by the stoichiometric relation. In this study, the loss of hydroscopic water corresponding to the temperature of 50 to 200°C and 200 to 300°C was arbitrarily used to represent the content of C-S-H, and summed content of C-A-H and C-A-S-H, respectively. Table 7 shows the content of C-S-H and summed content of C-A-H and C-A-S-H in the soils tested. As seen in Table 7, the content of C-S-H and summed content of C-A-H and C-A-S-H increase with the increase in curing time. At both 28 and 120 d, the content of C-S-H (1.861% and 2.108% for 28 and 120 d, respectively) and summed content of C-A-H and
C-A-S-H in the CCR stabilized soil (0.95% and 1.01% for 28 and 120 d, respectively) are higher than those in the quicklime stabilized soil (1.804% and 1.98% (C-S-H), 0.86% and 0.864% (sum of C-A-H and C-A-S-H) for 28 and 120 d, respectively). Figure 8 illustrates the correlations between \( q_u \) and summed content of C-S-H, C-A-H and C-A-S-H of the soils with 6% binder content. It is evident that higher summed content of C-S-H, C-A-H and C-A-S-H corresponds to higher \( q_u \) of the soils.

**Mercury intrusion porosimetry (MIP)**

Figure 9 presents the MIP results for the CCR and quicklime stabilized soils with 6% binder content, which are illustrated by the relations between cumulative pore volume and pore size. It is found that the total pore volumes decrease steadily with elapsed curing time for both CCR and quicklime stabilized soils. When the curing time increases from 28 d to 60 d, the reductions of total pore volumes are 0.03 mL/g and 0.08 mL/g for CCR and quicklime stabilized samples, respectively. The total pore volume for both CCR and quicklime stabilized soils, however, changes marginally from 60 d to 120 d. Therefore, pore-filling by the pozzolanic products occurs primarily at the initial 60 d curing.

The CCR stabilized soil has a significantly smaller cumulative pore volume in all ranges of measured pore sizes (Fig. 9). Particularly, the total pore volume of the CCR stabilized soil is approximately 0.054 mL/g lower than that of quicklime (0.23 mL/g) at 28 d. Nevertheless, the total pore volume is almost identical for both soils at 60 d. At 120 d curing, the pore size distribution curves of both stabilized soils are well overlapped, indicating they have similar long-term porosimetry characteristics.

Figure 10 shows the pore volumes of different types of pores in the CCR and quicklime
stabilized soils at different curing time. Horpibulsuk et al. (2009) reported that for the cement and
fly-ash stabilized silty clays, pore diameters of 0.01 and 10 µm are thresholds between
intra-aggregate and inter-aggregate pores, and inter-aggregate and large air pores, respectively. It is
found that the volumetric reduction in the large air pores with curing time, primarily, contributes to
the reduction in the total pore volume for CCR and quicklime stabilized soils. The quicklime
stabilized soil possesses greater volume of large air pores relative to the CCR stabilized soil
regardless of the curing time. This coincides with the higher $q_u$ value of the CCR stabilized soil over
the quicklime stabilized soil. Previous studies also reported the similar relation between strength
and volume of large pores in the soils. For example, Munkholm et al. (2002) found that tensile
strength was conversely linearly correlated to the volume of pores with diameter $> 30$ µm for sandy
loam. Locat et al. (1996) found that the mechanical properties of lime stabilized inorganic clay were
mainly controlled by the volume of pores with diameter $> 0.01$ µm.

Discussion

The results of this laboratory evaluation study demonstrate that the CCR stabilized clayey soil
has superior mechanical performances than the quicklime stabilized soil. This advantage in
mechanical performance can be interpreted from the physicochemical and microstructural points of
view. At the early stage of curing, the binder-soil interaction is dominated by cation exchange
(Hunter 1988). The cation exchange between the calcium ions from the hydration of CCR or
quicklime and the readily exchangeable cations initially adsorbed on the clay particle results in a
reduced thickness of diffusion double layer of clay particles, and formation of flocculation and
agglomeration of soil particles as a consequence (Kinuthia et al. 1999; Saride et al. 2010). Finer
particle size and larger specific surface area of the binder particles are preferable for the formation
of flocculation and agglomeration (Kinuthia et al. 1999). Since the CCR has higher amount of fine
particles and greater specific surface area relative to the quicklime (Table 3), flocculation and
agglomeration of soil particles would be more notable in the CCR stabilized soil. This mechanism is
substantiated by the analysis of particle size distribution of the stabilized soils, in which the particle
size of the CCR stabilized soil is coarser than that of the quicklime stabilized soil at 28 d (Table 6).

The long-term interaction between the CCR or quicklime and soil is dominated by pozzolanic
reactions (Hunter 1988). Portlandite reacts with the reactive SiO$_2$/Al$_2$O$_3$ in soil matrix and produces
2010). Finer particle size and larger specific surface area of the binder particle are more preferable
for the implementation of reactions between portlandite and reactive SiO$_2$/Al$_2$O$_3$ in the soil matrix.
Therefore, higher amount of pozzolanic products is found to form in the CCR stabilize soil relative
to the quicklime stabilized soil at the same curing time (Table 7).

The pH of the raw binder material also contributes to the different mechanical performances of
the stabilized soils, since the pozzolanic reactions are controlled by the pH of binder–soil system
and higher alkaline environment facilitates the dissolution of reactive SiO$_2$/Al$_2$O$_3$ within clay
minerals (Stronach and Glasser 1997; Saride et al. 2010). As the pH values of the CCR stabilized
soil are higher than those of the quicklime stabilized soil (Fig. 5), the amendment of CCR to the
parent soil would generate a faster dissolution rate of reactive SiO$_2$/Al$_2$O$_3$ in the soil matrix and rate
of pozzolanic reactions as a consequence. Because of the essential role of pozzolanic products in
soil pore filling, soil particle bonding and soil strength development, a superior mechanical
performance of the CCR stabilized soil over the quicklime stabilized soil is expected (Figs. 2 to 4).

In summary, it can be postulated that the variation in the physical and mechanical properties
(Atterberg limits, particle size distribution, compaction characteristics, $q_u$, CBR, and $M_r$) of the
CCR stabilized soil could be identified by resorting to systematic investigations to detect changes in the physicochemical and microstructural characteristics via soil pH, XRD, TGA and MIP analyses. The superior mechanical performances for the CCR stabilized soil can be explained fundamentally from the basic properties of the binders and soil-binder interactions (flocculation/agglomeration and pozzolanic reactions). Furthermore, environmental impacts of the use of CCR in soil stabilization including leachability of heavy metals have been evaluated using batch-type leaching tests; the results demonstrated that CCR is an environmental-friendly binder (Du et al. 2015a). Further study is recommended for CCR stabilization of soils with entirely different properties (e.g., silty and sandy soils).

Conclusions

This study presents a multi-scale laboratory investigation of physical, mechanical and microstructural properties of CCR stabilized highway soft clayey soils, with a comparison with quicklime stabilized soils. The following conclusions can be drawn from this research:

(1) The stabilization by CCR or quicklime results in increase in the liquid limit and plastic limit while decrease in the plasticity index. The addition of CCR or quicklime also leads to an increase in sand-sized particle percentage. Under the same curing period and binder content, the CCR stabilized soil attains higher sandy-sized particle percentage relative to the quicklime stabilized soil. The increase in the curing time and binder content facilitates larger $q_u$, CBR and $M_r$. Under the same curing period, binder content and degree of compaction, the CCR stabilized soil exhibits superior mechanical performances relative to the quicklime stabilized soil.

(2) Both CCR and quicklime stabilized soils display a gradual reduction in soil pH with
increased curing time, and pH of the CCR stabilized soil is always higher relative to the quicklime stabilized soil. Pozzolanic products like C-S-H, C-A-H and C-A-S-H are identified in both CCR and quicklime stabilized soils. The summed content of pozzolanic products in the CCR stabilized soil is higher than that in the quicklime stabilized soil. The CCR stabilized soil has a much smaller total pore volume than the quicklime stabilized soil within the initial 28 d, though the difference is almost eliminated at 120 d. The strength of the stabilized soil is found conversely correlated with the large pore volume in the soil.

(3) The fundamental mechanisms for the superior mechanical performances of the CCR stabilized soil over quicklime stabilized soil are the faster and more complete formation of flocculation and agglomeration of soil particles at the early stage as well as pozzolanic reactions within soils for the entire curing time. Finer particle size, greater specific area and higher pH value of CCR than quicklime are the essential contributors to the controlling mechanisms.

Acknowledgements

Financial support for this research was obtained from the National Natural Science Foundation of China (Grant No. 51278100, 41330641 and 41472258) and Natural Science Foundation of Jiangsu Province (Grant No. BK2012022). The fifth author is grateful to the Thailand Research Fund under the TRF Senior Research Scholar program Grant No. RTA5680002.

References

Transportation Officials, Washington D.C., USA.


ASTM 2011b. Standard terminology relating to lime and limestone (as used by the industry). ASTM C51-11, American Society for Testing and Materials, West Conshohocken, USA.

ASTM 2012. Standard test methods for laboratory compaction characteristics of soil using standard effort (12 400 ft-lbf/ft³ (600 kN-m/m³)). ASTM D698-12, American Society for Testing and
Material, West Conshohocken, USA.


https://mc06.manuscriptcentral.com/cgj-pubs


Liu, S.Y., Shao, G.H., Du, Y.J., and Cai, G.J. 2011. Depositional and geotechnical properties of


Penumadu, D., and Dean, J. 2000. Compressibility effect in evaluating the pore-size distribution of


Table 1. Properties of soils tested

<table>
<thead>
<tr>
<th>Index</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Natural moisture content, $w_n$ (%)</td>
<td>29.4</td>
</tr>
<tr>
<td>Specific gravity, $G_s$</td>
<td>2.73</td>
</tr>
<tr>
<td>Liquid limit, $w_L$ (%)</td>
<td>37.8</td>
</tr>
<tr>
<td>Plastic limit, $w_P$ (%)</td>
<td>19.9</td>
</tr>
<tr>
<td>Maximum dry density, $\rho_{d,\text{max}}$ (g/cm$^3$)</td>
<td>1.92</td>
</tr>
<tr>
<td>Optimum moisture content, $w_{\text{opt}}$ (%)</td>
<td>13.5</td>
</tr>
<tr>
<td>Particle size distribution (%) $^c$</td>
<td></td>
</tr>
<tr>
<td>Clay (&lt; 0.002 mm)</td>
<td>13.6</td>
</tr>
<tr>
<td>Silt (0.002 to 0.074 mm)</td>
<td>83.9</td>
</tr>
<tr>
<td>Sand (&gt; 0.074 mm)</td>
<td>2.5</td>
</tr>
</tbody>
</table>

$^a$ Based on ASTM D4318 (ASTM, 2010).

$^b$ Based on ASTM D698 (ASTM, 2012).

$^c$ Measured using a laser particle size analyzer Mastersizer 2000.
<table>
<thead>
<tr>
<th>Chemical composition</th>
<th>Soil (%)</th>
<th>CCR (%)</th>
<th>Quicklime (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CaO</td>
<td>1.3</td>
<td>68.99</td>
<td>68.54</td>
</tr>
<tr>
<td>SiO₂</td>
<td>67.9</td>
<td>2.84</td>
<td>2.54</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>14.1</td>
<td>2.16</td>
<td>1.0</td>
</tr>
<tr>
<td>MgO</td>
<td>2.5</td>
<td>0.12</td>
<td>0.34</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>5.0</td>
<td>0.15</td>
<td>0.62</td>
</tr>
<tr>
<td>SO₃</td>
<td>0.01</td>
<td>0.76</td>
<td>0.11</td>
</tr>
<tr>
<td>Loss of ignition</td>
<td>5.19</td>
<td>24.85</td>
<td>26.51</td>
</tr>
</tbody>
</table>

*a Measured using a X-ray Fluorescence Spectrometer.
Table 3. Basic physical and chemical properties of CCR and quicklime

<table>
<thead>
<tr>
<th>Index</th>
<th>CCR</th>
<th>Quicklime</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific gravity, $G_s$</td>
<td>2.32</td>
<td>3.21</td>
</tr>
<tr>
<td>Specific surface area (m$^2$/g)</td>
<td>24.664</td>
<td>5.020</td>
</tr>
<tr>
<td>pH $^b$</td>
<td>12.84</td>
<td>12.74</td>
</tr>
<tr>
<td>Particle size distribution (%) $^c$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clay (&lt;0.002 mm)</td>
<td>4.2</td>
<td>3.9</td>
</tr>
<tr>
<td>Silt (0.002 to 0.074 mm)</td>
<td>67.6</td>
<td>37.8</td>
</tr>
<tr>
<td>Sand (&gt;0.074 mm)</td>
<td>28.2</td>
<td>58.3</td>
</tr>
</tbody>
</table>

$^a$ Measured using a Quantachrome Autosorb-iQ-AG automated gas sorption analyzer.

$^b$ Based on ASTM 4972 ([ASTM, 2013](#)).

$^c$ Measured using a laser particle size analyzer Mastersizer 2000.
Table 4. Summary of binder dosage, curing time, degree of compaction and number of identical samples for different tests in this study

<table>
<thead>
<tr>
<th>Testing program</th>
<th>Binder dosage (%)</th>
<th>Curing time (day)</th>
<th>Degree of compaction (%)</th>
<th>Number of identical samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atterberg limits</td>
<td>0, 4, 6</td>
<td>28</td>
<td>NA&lt;sup&gt;h&lt;/sup&gt;</td>
<td>1</td>
</tr>
<tr>
<td>PSD&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0, 4, 6</td>
<td>28</td>
<td>NA&lt;sup&gt;h&lt;/sup&gt;</td>
<td>1</td>
</tr>
<tr>
<td>Compaction</td>
<td>0, 4, 6</td>
<td>0</td>
<td>NA&lt;sup&gt;h&lt;/sup&gt;</td>
<td>1</td>
</tr>
<tr>
<td>UCS&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0, 4, 6</td>
<td>7, 28, 60, 120</td>
<td>93, 94, 96</td>
<td>3</td>
</tr>
<tr>
<td>CBR&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0, 4, 6</td>
<td>7, 28</td>
<td>93, 94, 96</td>
<td>3</td>
</tr>
<tr>
<td>$M_r$&lt;sup&gt;d&lt;/sup&gt;</td>
<td>0, 4, 6</td>
<td>7, 28</td>
<td>94, 96</td>
<td>3</td>
</tr>
<tr>
<td>Soil pH</td>
<td>6</td>
<td>7, 28, 60, 120</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>XRD&lt;sup&gt;e&lt;/sup&gt;</td>
<td>0, 6</td>
<td>180</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>TGA&lt;sup&gt;f&lt;/sup&gt;</td>
<td>6</td>
<td>28, 120</td>
<td>100</td>
<td>3</td>
</tr>
<tr>
<td>MIP&lt;sup&gt;g&lt;/sup&gt;</td>
<td>6</td>
<td>28, 60, 120</td>
<td>100</td>
<td>1</td>
</tr>
</tbody>
</table>

<sup>a</sup> Particle size distribution
<sup>b</sup> Unconfined compressive strength (UCS)
<sup>c</sup> California Bearing ratio (CBR)
<sup>d</sup> Resilient modulus
<sup>e</sup> X-ray diffraction (XRD)
<sup>f</sup> Thermogravimetric analyses (TGA)
<sup>g</sup> Mercury intrusion porosimetry (MIP)
<sup>h</sup> Not available
Table 5. Results of Atterberg limits tests for CCR and quicklime stabilized and unstabilized soils

<table>
<thead>
<tr>
<th>Soil</th>
<th>Binder content (%)</th>
<th>Curing time (d)</th>
<th>Liquid limit, ( w_L ) (%)</th>
<th>Plastic limit, ( w_p ) (%)</th>
<th>Plastic index, ( I_p )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Untreated soil</td>
<td>0</td>
<td>NA (^a)</td>
<td>37.8</td>
<td>19.9</td>
<td>17.9</td>
</tr>
<tr>
<td>CCR stabilized soils</td>
<td>4</td>
<td>28</td>
<td>42.7</td>
<td>29.6</td>
<td>13.1</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>28</td>
<td>39.8</td>
<td>26.4</td>
<td>13.4</td>
</tr>
<tr>
<td>Quicklime stabilized soils</td>
<td>4</td>
<td>28</td>
<td>41.6</td>
<td>26.8</td>
<td>14.8</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>28</td>
<td>44.3</td>
<td>28.8</td>
<td>15.5</td>
</tr>
</tbody>
</table>

\(^a\) Not available
Table 6. Results of particle size distribution test

<table>
<thead>
<tr>
<th>Soil</th>
<th>Binder content (%)</th>
<th>Curing time (d)</th>
<th>Clay-sized particle (&lt; 2 µm) (%)</th>
<th>Silt-sized particle (2 to 74 µm) (%)</th>
<th>Sand-sized particle (74 to 2000 µm) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unstabilized soil</td>
<td>0</td>
<td>Not available</td>
<td>13.6</td>
<td>83.9</td>
<td>2.5</td>
</tr>
<tr>
<td>CCR stabilized soil</td>
<td>4</td>
<td>28</td>
<td>7.1</td>
<td>55.1</td>
<td>37.8</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>28</td>
<td>3.8</td>
<td>35.5</td>
<td>60.7</td>
</tr>
<tr>
<td>Quicklime stabilized soil</td>
<td>4</td>
<td>28</td>
<td>10.0</td>
<td>63.6</td>
<td>26.4</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>28</td>
<td>8.2</td>
<td>65.2</td>
<td>26.6</td>
</tr>
</tbody>
</table>
Table 7. The relative content of the hydration products

<table>
<thead>
<tr>
<th>Hydration product</th>
<th>Range of water loss (°C)</th>
<th>28 d (%)</th>
<th>120 d (%)</th>
<th>28 d (%)</th>
<th>120 d (%)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>CSH</td>
<td>50 to 200</td>
<td>1.861</td>
<td>2.108</td>
<td>1.804</td>
<td>1.98</td>
<td>HaHa et al. (2011)</td>
</tr>
<tr>
<td>CAH+CASH</td>
<td>200 to 300</td>
<td>0.95</td>
<td>1.01</td>
<td>0.86</td>
<td>0.864</td>
<td>HaHa et al. (2011)</td>
</tr>
</tbody>
</table>
List of Figure captions

**Fig. 1** Particle size distributions of unstabilized soil, CCR and quicklime.

**Fig. 2** Variations of $q_u$ with curing time for soils with various degree of compaction: (a) 93%; (b) 94%; and (c) 96%.

**Fig. 3** Variations of CBR with curing time and binder content for soils with various degree of compaction: (a) 93%; (b) 94%; and (c) 96%.

**Fig. 4** Variations of resilient modulus ($M_r$) with degree of compaction and curing time for unstabilized soils and soils of 6% binder.

**Fig. 5** Variations of soil pH with curing time.

**Fig. 6** XRD diffractograms of unstabilized soil and CCR and quicklime stabilized soils.

**Fig. 7** Thermogravimetric analysis (TGA) and differential thermogravimetric (DTG) analysis of soils stabilized with: (a) CCR at 28 d; (b) quicklime at 28 d; (c) CCR at 120 d; and (d) quicklime at 120 d.

**Fig. 8** Summed content of C-S-H, C-A-H and C-A-S-H versus $q_u$ of soils stabilized with CCR and quicklime (6% binder content).

**Fig. 9** MIP test results for CCR and quicklime stabilized soils.

**Fig. 10** Distributions of different types of pores in CCR and quicklime stabilized soils.
Fig. 1 Particle size distributions of unstabilized soil, CCR and quicklime.

172x122mm (300 x 300 DPI)
Fig. 2 Variations of $q_u$ with curing time for soils with various degree of compaction: (a) 93%; (b) 94%; and (c) 96%.

203x146mm (300 x 300 DPI)
Fig. 3 Variations of CBR with curing time and binder content for soils with various degree of compaction: (a) 93%; (b) 94%; and (c) 96%.
Fig. 4 Variations of resilient modulus (Mr) with degree of compaction and curing time for unstabilized soils and soils of 6% binder.
131x92mm (300 x 300 DPI)
Fig. 5 Variations of soil pH with curing time.

135x95mm (300 x 300 DPI)
Fig. 6 XRD diffractograms of unstabilized soil and CCR and quicklime stabilized soils.
251x356mm (300 x 300 DPI)
Fig. 7 Thermogravimetric analysis (TGA) and differential thermogravimetric (DTG) analysis of soils stabilized with: (a) CCR at 28 d; (b) quicklime at 28 d; (c) CCR at 120 d; and (d) quicklime at 120 d.
Fig. 8 Summed content of C-S-H, C-A-H and C-A-S-H versus qu of soils stabilized with CCR and quicklime (6% binder content).

Unconfined compressive strength, $q_u$ (kPa)

Fig. 9 MIP test results for CCR and quicklime stabilized soils.

$\rho_d = 1.74 \text{ g/cm}^3$ (CCR 6%)

$\rho_s = 1.70 \text{ g/cm}^3$ (Quicklime 6%)

137x95mm (300 x 300 DPI)
Fig. 10 Distributions of different types of pores in CCR and quicklime stabilized soils.
157x111mm (300 x 300 DPI)