Chemo-physical evolution and microstructure features of lime treated soils

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Abstract. In the paper some results on the effects of chemo-physical evolution of clay-lime-water suspensions on the microstructure of a lime treated kaolin have been presented. A multi-scale investigation on the sedimentation behaviour of clay suspensions under different pore water chemistry has been developed highlighting the chemo-physical mechanisms controlling particle arrangement and the soil fabric formation. The results evidenced the key role of ionic exchange in the short term on the microstructural features of the lime treated soil.

1 Introduction

The addition of lime to a clayey soil has a strong impact on the physical and mechanical properties in the short and long term after treatment [1-7]. Several studies on the basic lime-clay-reaction mechanisms have been widely reported in literature [8-14] allowing a better understanding of chemo-physical evolution of the system and microstructural features induced after the addition of lime [15-17].

When lime is added to the soil water system, either in the form of quicklime (CaO) or as hydrated lime (Ca(OH)_2), a change in the pore water chemistry occurs. For quicklime the first reaction taking place is hydration reaction, highly exothermic, leading to the formation of hydrated lime (portlandite) in the soil water system. The dissociation reaction of the hydrated lime leads to an increase of the pH and to a high concentration of calcium ions in the pore water. The ions dissolved in the clay water system tend to counteract the charge on the basal face and lateral face sites of clay particles surface, controlling both the chemo-physical interactions and the arrangement of clay particles. The high pH environment induced by lime addition promotes the dissolution of the alumino-silicate constituents of clay minerals. Cementitious compounds, such as hydrated calcium silicates (C-S-H) or hydrated calcium aluminates (C-A-H), are formed as products of pozzolanic reactions between silica and alumina and lime.

The effects of the chemo-physical evolution of the clay-lime-water systems on microstructure play a key role in the mechanical improvement of lime treated soils. In the paper some results on microstructure are reported and discussed. The chemo-physical evolution of the suspensions has also been studied at particle scale level. The electro-kinetic properties of clay particles governing the interaction forces have been inferred from Zeta-potential measurements. Soil fabric evolution and its effects on the physical properties have been investigated by means of Dynamic Light Scattering measurements. A discussion section has been finally devoted to the interpretation of results linking different scale level of the experimental analysis.

2 Experimental procedures

2.1 Soil

Speswhite kaolin was provided by Imerys Minerals UK. The specific gravity G_s is 2.6, and its surface area is 14 m^2/g. The pH value is 4.6. Atterberg limits are respectively w_L = 70% and w_p = 32%, with a plasticity index IP equal to 38%. The mineralogical composition is mainly formed by kaolinite clay minerals with a small amount of quartz and muscovite. No swelling clay phases were detected after the ethylene glycol treatment.

2.2 Sedimentation tests

The sedimentation behaviour of kaolin suspensions has been investigated as a function of lime content and cation valence. Suspending were prepared in graduated cylinders with mixture of 50 g of dry kaolin at 100% of water content. Different amounts of CaO and KOH were added to reach the required pH and electrolytes concentrations. Additional water was added up to a initial volume of 1000 ml. The pH value was increased by means of KOH. During the test the suspensions heights were monitored as a function of time.

Two typical curves can be identified (Figure 1) from sedimentation tests, depending on the degree of particles association in the suspension. Figure 1a shows the...
settling of a flocculated system, with the suspension height progressively reduced over time leaving a clear supernatant fluid. Figure 1b shows a dispersed system maintaining its configuration over time. The height of the suspension in the latter case is not relevantly modified and a partition between the suspension height and a clear supernatant fluid is not well recognisable.

In a flocculated system three stages of the sedimentation process can be identified (Figure 2). During the initial stage (induction period), the electrochemical interactions [22] induce the aggregation from small clusters to aggregates depending on the surface charge distribution of particles. Its duration is linked to the degree of particles or aggregates association [25]. A slight translation of the interface between suspended solids and water can be appreciated in this stage. The clear separation between the suspended solids and the supernatant liquid indicates that sedimentation and self-weight consolidation stages are ongoing. The flocculated soil particles settle under gravity forming a deposit of increasing height which undergoes self-weight consolidation. Settling and self-weight consolidation take place simultaneously, as evidenced in Figure 2 with the time \( t^* \). The soil formation line (Figure 2) delimitates the sedimentation and the self-weight consolidation regions and is characterised by a constant value of the void ratio \( e_{\text{in}} \), the soil-formation void ratio, which represents the void ratio characterising the soil skeleton at zero effective stresses [26]. At time \( t_c \), corresponding to the intersection between the soil formation line and the line representing the upper interface of the suspension, an abrupt reduction of settling velocity occurs, indicating the completion of the sedimentation stage [23, 27]. During the subsequent self-weight consolidation stage the sediment reduces its height with a decreasing rate until the final height. During the self-weight consolidation the effective stresses gradually develop due to the mechanical interaction between aggregates [23, 28] until the soil skeleton is in equilibrium under its own weight. The duration \( t_f \) of primary consolidation process generally lasts a time interval of two order of magnitudes larger than \( t_c \), time necessary to complete the free settling stage [23, 27, 29].

2.3 Microstructure analyses

The Zeta potential measurements are based on the electrophoretic mobility of particles under an induced electric field. Mobility is then converted to zeta potential applying the Henry equation with the Smoluchovski approximation [30]. Diluted dispersions with 100 mg/l solid content and different concentration of calcium were prepared. The grain size fraction lesser than 1 \( \mu m \) was extracted and used for zeta potential measurement. The pH of dilute dispersions were adjusted between 2 and 12.4 by adding KOH. The measurement was performed at 25 °C in a capillary cell (DTS1061) with Malvern Nano Zetasizer apparatus. For each sample at least 3 measurements were performed assuring the repeatability of the results.

The hydrodynamic size of soil particles was determined using dynamic light scattering (DLS) technique, based on the interaction between a set of particles and an incident radiation. Details of the technique are referred in [31].

The grain size fraction lesser than 1 \( \mu m \) was extracted by dispersing about 10 g of sample in 1 L ultrapure water and centrifugation method. The suspension (100 mg/l) was prepared at different concentrations of Ca(OH)\(_2\) and

Figure 1. Sedimentation behaviour of suspensions a) flocculated system, b) dispersed system

Figure 2. Stages of sedimentation process (after Vitale, 2016)
KOH. Hydrodynamic size measurements were then performed at 25 °C and laser scattering at 173° in a capillary cell with Malvern Nano Zetasizer apparatus.

3 Results

Sedimentation curves of kaolin suspensions prepared at increasing lime percentages are reported in Figure 3. The evolution of the suspension height is shown as a function of time with reference to two different time scales, namely the whole duration of the test (up to 13000 min, Figure 3a) and the initial stage of the test (up to 200 min, Figure 3b). Not treated kaolin suspension is characterised by a relatively long initial stage (induction period) during which the clayey particles show a negligible settling. A small percentage of lime (1% CaO) promotes a more efficient particles association process due to alkaline environment and divalent cations, as showed by shorter induction period. It is noteworthy that no remarkable changes of the final sediment height were recorded (Figure 3a). As shown in Figure 3b, the higher amount of lime for 3% CaO mixes promotes a higher degree of flocculation with respect to 1%CaO mixes. As a consequence, a shorter induction period has been detected (Figure 3b) with an increase of the final sediment height after the self-weight consolidation stage (Figure 3a). No significant effects on the sedimentation behaviour and final suspension height were observed for samples treated with higher lime percentages (namely 5% CaO and 7% CaO). Therefore, 3% CaO can be regarded as the minimum amount of lime required to promote an effective aggregation of particle in the short term.

The influence of cation valence on the sedimentation characteristics of kaolin suspensions is shown in Figure 4. Sample prepared with KOH at pH = 12.4 showed a long induction period (Figure 4a), with particles dispersed in suspension and a not well recognizable sediment at the bottom of the cylinder (Figure 4b).

A further insight in the effects of pore water chemistry on the aggregation of kaolin particles has been highlighted by microstructural analyses results. In Figure 5 the zeta potential measurements of not treated and lime treated kaolin suspensions prepared at increasing pH have been presented. An increase of the zeta potential magnitude with increasing pH has been detected as consequence of deprotonation reactions respectively on the lateral and octahedral basal faces of kaolinite. The addition of lime (calcium hydroxide at increasing concentration) induces a decrease of the zeta potential decreases with increasing pH. In particular, the reduction is sharp for pH higher than 9. Calcium undergoes hydrolysis at higher pH (pH > 10) leading to the formation of monovalent species [33, 34]. According to [14], the observed behaviour can be linked to the formation and specific adsorption of first hydrolysis product of calcium (i.e. CaOH

The evolution of the size of aggregated particle structure with time was monitored by Dynamic Light Scattering (DLS) measurements. For kaolin suspension in de-ionized water (pH = 4.6) (Figure 6), the measured average size of aggregations increases with time as a consequence of positively charged edge and negatively charged basal face mode of particle aggregation referred to as edge-to-face [35]. As the pH increased to 12.4 by adding KOH, the size of aggregates is smaller and remains unchanged over time. The kaolinite particles are
not flocculated and their mutual interaction is weak (i.e. dispersed state). The high negative surface charge, as indicated by higher zeta potential (Figure 5), increases the electrostatic repulsive forces preventing particles collisions and aggregations. The presence of calcium (Ca(OH)\textsubscript{2}) and the alkaline environment (pH=12.4) favour the larger average size of kaolinite aggregates and a strongly flocculated structure. Calcium ions counteract the effect of higher surface charge and repulsive forces, promoting particles collisions and aggregations.

4 Discussion

A pH dependent surface charge behaviour of kaolin suspension was observed from zeta potential measurements. In acidic environment the interaction between particles is the result of Coulombian attractions between positively charged edges and negatively charged faces (edge-to-face arrangement). In alkaline environment the absolute value of zeta potential increases due to the increase of negative charges on the kaolinite edge surfaces (deprotonation reaction). The results of sedimentation test showed a flocculated behaviour of kaolin suspensions in deionized water at pH=4.6, confirmed by DLS results highlighting the increase of the average aggregate size as a consequence of the edge-to-face flocculation. The increase of pH provided by the addition of KOH induces a suspension stability. The repulsive forces between charged particles prevent flocculation, as confirmed by the constant size of kaolinite particles over time detected by DLS measurements.

The addition of lime induces a change in the pore water chemistry, with an increase of the pH and the divalent cation concentration, affecting the sedimentation behaviour of kaolin suspension. A larger decrease of the absolute value of zeta potential (Figure 5) is the consequence of calcium adsorption at the surface of the negatively charged particles. The repulsive forces are no longer able to counteract the van der Waals attractions [36], and an increase of the flocculated aggregates size is detected, as showed by DLS measurements results.

The aggregation of particles of lime treated suspensions can be considered not affected by the ongoing of pozzolanic reactions with reference to the time intervals analysed during the sedimentation tests (about 9 days). As reported by [31], thermo-gravimetric analyses performed on 3% CaO treated kaolin samples showed a low reactivity in the short term, as showed in Figure 7, where the mass losses of hydrated lime (390°C - 460°C) and cementitious compounds (110-350°C) after the treatment have been represented as function of time. A significant consumption of portlandite and formation of new cementitious phases has been detected only after times longer than 28 days.

The increase of the average dimensions of clay aggregates in saturated lime solution is consistent with the increase of the soil-formation void ratio e\textsubscript{so} determined from the sedimentation curves. The soil formation void ratio e\textsubscript{so} measured at the top of the sediments layer corresponds to the soil skeleton void ratio under zero effective stresses. Its value is representative of the soil skeleton void ratio induced only by the pore water chemistry. In order to estimate its value for each suspension, the end of sedimentation stage time t\textsubscript{s} and the corresponding thickness of the sediment h\textsubscript{s} were evaluated from the sedimentation curves. A lower limit of soil-formation void ratio e\textsubscript{so} can be inferred by considering that the total thickness of the solid matter h\textsubscript{z} (i.e. the material height) for each of the sedimentation tests is
constant throughout the sedimentation/consolidation process [27], that is

\[ h_s = \frac{h_0}{1 + e_0} = \frac{h_f}{1 + e_s} = \text{constant} \quad (1) \]

where \( h_0 \) and \( e_0 \) are respectively the initial height and the initial void ratio of the suspension. The value of \( e_m \) must satisfy the following relationship:

\[ e_m \geq e_s = \frac{h_f}{h_s} - 1 \quad (2) \]

Since for each treated sample the time to achieve full consolidation \( t_f \) was found to be two order of magnitude larger than \( t_1 \) [27] the difference between the calculated \( e_s \) and the soil-formation void ratio \( e_m \) has been considered negligible. Therefore, \( e_m \) was assumed as the representative value of void ratio when the soil skeleton is completely formed without any mechanical effect (i.e. zero effective stresses). The estimated values of \( t_f \) and \( h_s \) and the calculated values of \( e_s \) for 1% CaO and 3% CaO mixes have been reported in Table 1.

**Table 1.** Estimated and calculated values from sedimentation curves

<table>
<thead>
<tr>
<th>Test</th>
<th>( e_0 )</th>
<th>( h_0 )</th>
<th>( t_1 )</th>
<th>( h_s )</th>
<th>( e_s )</th>
<th>( t_f )</th>
</tr>
</thead>
<tbody>
<tr>
<td>K_1%CaO</td>
<td>50</td>
<td>34.5</td>
<td>52.5</td>
<td>19.1</td>
<td>27.3</td>
<td>1530</td>
</tr>
<tr>
<td>K_3%CaO</td>
<td>50</td>
<td>34.5</td>
<td>25</td>
<td>28</td>
<td>40.4</td>
<td>2833</td>
</tr>
<tr>
<td>K_5%CaO</td>
<td>50</td>
<td>34.5</td>
<td>20</td>
<td>27.5</td>
<td>39.7</td>
<td>2886</td>
</tr>
</tbody>
</table>

The final height of the sediments after the soil skeleton formation, at the end of the tests, is affected by the pore water chemistry. The negative charge neutralization induced by the addition of lime causes a reduction of inter-particle repulsions promoting van der Waals attractions between particles. During the consolidation stage, the effective stresses increase inducing the reduction of the soil skeleton volume, due to the self-weight effect and the van der Waals attractions. The contribution of the attractive forces results in lower volume strains during consolidation, and thus higher sediment heights, as showed by lime treated suspensions which experience a consolidation stage characterized by lower settlements due to the higher contribution of van der Waals attractions.

**5 Conclusions**

In the paper some results on the effects of chemo-physical evolution of clay-lime-water suspensions on the microstructure of a lime treated kaolin have been presented. A multi-scale investigation on the sedimentation behaviour of clay suspensions under different pore water chemistry has been developed highlighting the chemo-physical mechanisms controlling particle arrangement and the soil fabric formation.

The results evidenced the key role of ionic exchange in the short term in terms of microstructure features of the treated soil. No effects of pozzolanic reactions were overlapped to the observed behaviour due to the low reactivity of kaolinite. The arrangement of kaolin particles in suspension is strongly pH dependent, with a flocculated face to edge arrangement at pH 4.6 turning to a dispersed one at higher pH. The addition of calcium ions in suspension induces the formation of larger particles aggregates, formed by face to face stacking of particles grouped face to edge. The sedimentation behaviour in presence of calcium is consistent with the effects of particle arrangement, highlighting a shorter induction period, a faster settling stage and a shorter consolidation stage due to the aggregation induced by the pore water chemistry.

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