Use of alkali activated high-calcium fly ash binder for kaolin clay soil stabilisation:

**Physicochemical evolution**

Elodie Coudert\(^a,b,c\), Michael Paris\(^a\), Dimitri Deneele\(^a,d*\), Giacomo Russo\(^b\), Alessandro Tarantino\(^c\)

\(^a\) Institut des Matériaux Jean Rouxel (IMN), Université de Nantes, CNRS, 2 rue de la Houssinière, BP 32229, 44322 Nantes Cedex 3, France

\(^b\) Department of Civil and Mechanical Engineering, University of Cassino and Southern Lazio, Via Gaetano di Biasio, 43, 03043 Cassino, FR, Italy

\(^c\) Department of Civil and Environmental Engineering, University of Strathclyde, 75 Montrose Street, Glasgow, Scotland, G1 1XJ, United Kingdom

\(^d\) IFSTTAR, GERS, EE, F-44344 Bouguenais, France.

*Corresponding author

Phone : + 33 2 40 84 58 02

Fax: + 33 2 40 58 57 77

E-mail: Dimitri.Deneele@ifsttar.fr

**Abstract**

This study addresses the use of alkali activated high-calcium fly ash-based binder to improve engineering characteristics of soft clay-rich soils as an alternative to common stabilisers. The physico-chemical reaction sequence has been investigated by considering the binder alone and the binder mixed with kaolin. An insight into the reactivity evidenced that calcium-containing phases derived from high-calcium fly ash represent the reactive phases and, hence, pozzolanic activity is the dominant process. New compounds are
formed, thenardite Na$_2$SO$_4$ and an amorphous silicate consisting of chains combined with calcium probably incorporating three-dimensional four-fold aluminium environments.

Keywords

Soil stabilisation; Alkali activated material; Kaolin; High-calcium fly ash
1. Introduction

Soft clay-rich soils are frequently encountered in construction sites. Their poor mechanical performance represents a critical issue in engineering projects. These soils cannot be directly used as earthfill materials and may cause excessive settlements of foundation structures. To improve their engineering characteristics chemical stabilisation involving the addition of a binder to the soil has been widely practiced. The commonly used stabilisers are Ordinary Portland Cement and lime whose stabilisation mechanisms have been widely reported [1-7]. Nevertheless, a major issue with those conventional stabilisers is a very significant environmental penalty due to high carbon dioxide emissions and energy intensive processes.

In the low carbon agenda, the development of novel technologies that are both cost- and carbon-efficient is of prime importance, particularly in the construction sector for which cement production contributes to at least 5–8% of global carbon dioxide emissions [8]. As an alternative, industrial by-products such as high-calcium fly ash, rice husk ash, and silica fume have been successfully mixed as cementing additives to soft soils resulting in environmental and economic benefits [9-15].

Another alternative gaining attention is the use of Alkali Activated Materials as a viable sustainable binder whose often-claimed advantage is a much lower CO$_2$ emission process compared to traditional Portland cement. Works on alkali activated soils are recent and aim to stabilise different types of soil from clayey soil [16-17], sandy clay [18], Lateritic soils [19] marl, marlstone [20], silty sand [21], road aggregates [22] to mixed soil synthesised in laboratory [23-24]. The overall work shows the potential of alkaline activation for soil improvement, and this for different designed applications i.e. in deep soft soil [18], at shallow depth [24] or in rammed earth construction [25].
Alkali Activated Materials are defined as any binder system derived by the reaction of an alkali metal source (usually alkali hydroxide and alkali silicate solutions) with a solid aluminosilicate powder (commonly metakaolin, fly ash, blast furnace slag or natural pozzolan) [26-27]. It gives a hardened material at room temperature with mechanical properties potentially suitable for Portland cement replacement.

The type of aluminosilicate material needed in the alkali activation process varies as well. In fact, most of the studies were conducted on the use of fly ash [16-18; 20-21;23-25; 28-29]. Nevertheless, Zhang et al. [24] also examined the feasibility of metakaolin based alkali activated soil, and some other studies established on slag based alkali activated soil are as well existing [16, 17, 23].

As stabilisation using alkaline activation is a recent research area, studies about the understanding of the physicochemical reactivity of such systems have received little attention so far [17]. Yet, the molecular structure and the chemical composition of the alkali activated binders is essential to properly assess the resulting strength and durability of the final material.

This work focuses on the use of calcium-rich high-calcium fly ash from coal combustion activated by sodium-based alkaline solution as a binder for clay kaolin stabilisation. High-calcium fly ash was selected in the context of resource-saving being an industrial waste. Kaolin was selected as a model soil to represent a wide class of clays encountered in engineering projects.

The study was designed in three stages. An initial stage consisted in the investigation of the reactivity of the alkali activated high-calcium fly ash binder by itself, including (i) which phases are present and which phases are accessible during alkaline activation, (ii) which compounds are subsequently formed, and (iii) reactivity timescale. A second stage
focused on the interaction of the high-calcium fly ash-based binder with the kaolin clay to understand how the presence of kaolin modifies the reactivity of the system. Kaolinite is generally unreactive to alkali attack at ambient temperature. However, the addition of clay may affect chemical reactions as occurs in clay-cement mix [30]. Finally, the physicochemical evolution occurring in the alkali activated high-calcium fly ash is compared with one occurring in the same kaolin stabilised by i) lime or ii) a mix of lime and the same high-calcium fly ash used in this experimental programme. This is aimed at assessing the potential benefit of high-calcium fly ash-based binder compared to the more traditional lime.
2. Material and methods

2.1 Materials

A Polish high-calcium fly ash derived from hard coal and coal slime combustion in fluidised bed boiler was used. Its chemical analysis is given in Table 1, and consists primarily of SiO₂, Al₂O₃ and CaO. The high-calcium fly ash contains, approximately, 52% of particles sized lower than 45 µm and 41% lower than 10 µm.

Speswhite kaolin provided by Imerys Minerals UK, and whose chemical composition is given in Table 1 was used. It is mainly constituted of kaolinite (95%) and secondarily of muscovite (4%) [3]. The kaolin contains, approximately, 100 % of particles sized lower than 10 µm and 80 % lower than 2 µm.

A unique alkaline solution was used: a sodium silicate with a mass ratio SiO₂/Na₂O of 1.7 and a dry mass percentage of 44%; supplied by Woellner group and named GEOSIL 34417.

Table 1

<table>
<thead>
<tr>
<th></th>
<th>SiO₂</th>
<th>Al₂O₃</th>
<th>Fe₂O₃</th>
<th>CaO</th>
<th>CaO_free</th>
<th>MgO</th>
<th>SO₃</th>
<th>Na₂O</th>
<th>K₂O</th>
<th>H₂O</th>
<th>L.o.I.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fly ash</td>
<td>39.4</td>
<td>19.8</td>
<td>7.4</td>
<td>18.6</td>
<td>5.2</td>
<td>1.8</td>
<td>4.1</td>
<td>2.0</td>
<td>1.8</td>
<td>0.0</td>
<td>1.7b</td>
</tr>
<tr>
<td>Kaolin</td>
<td>49.2</td>
<td>34.5</td>
<td>1.2</td>
<td>0.0</td>
<td>0.0</td>
<td>0.2</td>
<td>0.0</td>
<td>0.1</td>
<td>1.7</td>
<td>13.1</td>
<td>12.0c</td>
</tr>
</tbody>
</table>

*a Free calcium oxide content, *b* from [3], *c* from [33]

2.2 Sample preparation

Sample preparation consisted in (i) mix of liquid sources ie. silicate and water (ii) mix of aluminosilicate powders ie. high-calcium fly ash and kaolin in the case of soil-source sample (iii) mix of (i) and (ii) previously prepared.

Three types of mixes were studied and named F100, KF50 and KF20. F100 is the high-calcium fly ash based alkali activated binder. It corresponds to a solid phase made of
high-calcium fly ash only, whereas KF50 and KF20 are the alkali activated soils. KF50 corresponding to a solid phase made of 50% of high-calcium fly ash and 50% of kaolin in mass, and KF20 corresponding to a solid phase made of 20% of high-calcium fly ash and 80% of kaolin. List of samples is summarized up in Table 2.

To ensure a good workability, the amount of added water with respect to the solid mass (eg. mass of kaolin and high-calcium fly ash) was fixed to 50% for all the samples. Additionally, the mass ratio of alkaline solution to high-calcium fly ash was fixed to 50% for all the samples, giving the initial molar ratios (considering that kaolin is unreactive):

\[
\text{Si/Al} = 2.0, \text{Si/Na} = 3.5 \text{ and } \text{Al/Na} = 1.8. \text{ The Al/Na ratio was not fixed to one because of the presence of calcium ions in high quantity in our system playing a role of charge compensation as well as sodium.}
\]

The paste obtained was poured in closed plastic molds and cured at room temperature (20°C). Samples were finally demoulded and freeze dried at curing times of 1, 3, 7 or 28 days.

Table 2

<table>
<thead>
<tr>
<th>Sample</th>
<th>Fly ash</th>
<th>Kaolin</th>
<th>Water</th>
<th>Alkaline solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>F100</td>
<td>50</td>
<td>0</td>
<td>25</td>
<td>25</td>
</tr>
<tr>
<td>KF50</td>
<td>28.6</td>
<td>28.6</td>
<td>28.6</td>
<td>14.3</td>
</tr>
<tr>
<td>KF20</td>
<td>12.5</td>
<td>50</td>
<td>31.3</td>
<td>6.3</td>
</tr>
</tbody>
</table>

2.3 Methods

A variety of characterisation techniques were complementary used to probe the mineralogical, structural and microstructural sample characteristics.
X-ray diffractograms of powdered samples were obtained with a Bruker D8 Advance diffractometer, using CuKα radiation generated at 40 mA and 40kV. Specimens were step-scanned from 2 to 60° 2θ at 0.017° 2θ steps integrated at the rate of 1s/step. Derivative thermogravimetric curves were obtained on a Netzsch STA 449F3 Jupiter thermal analyser. The samples were heated from 20–1000 °C at a rate of 10 °C/min under argon atmosphere. FTIR spectra were obtained on an FTIR Bruker Vertex 70 spectrometer. Specimens were prepared by mixing 30 mg of sample in 270 mg of KBr. Spectral analysis was performed over the range 4000–400 cm⁻¹ at a resolution of 4 cm⁻¹. Solid-state $^{29}$Si NMR spectroscopy was performed using a Bruker Avance III 300 MHz (7 T) spectrometer and 7 mm MAS probe. $^{29}$Si MAS spectra were acquired with a single $\pi/2$ pulse excitation of 5.5 μs and $^1$H decoupling. The repetition times were 2 s, 120 s and 30 s for the raw high-calcium fly ash, the raw kaolin and all the activated samples, respectively. For all $^{29}$Si spectra, MAS spinning rate was set to 5 kHz. Solid-state $^{27}$Al NMR spectroscopy was performed using a Bruker Avance III 500 MHz (11.7 T) spectrometer and 2.5 mm MAS probe. $^{27}$Al MAS spectra were acquired with a single pulse excitation of $\pi/12$ pulse of 3.3 μs and $^1$H decoupling. Repetition time was set to 1 s and MAS spinning rate to 30 kHz. Spectra were referenced against TMS (tetramethylsilane) for $^{29}$Si and an Al(NO₃)₃ aqueous solution for $^{27}$Al. Lastly, attention should be drawn to the fact that iron initially present in the high-calcium fly ash renders the interpretation more complex, notably spectra from different mixes cannot be quantitatively compared. Finally, samples were studied by SEM from polished section. Freeze-dried samples were impregnated under a vacuum with an acrylic resin (LR White). The polymerisation of the
resin was performed in an oven at 60°C over 48 h. The samples were then polished with diamond powder and coated with carbon before the observation. The observations were done with a HITACHI SU5000 scanning electron microscope equipped with an energy-dispersive X-ray analyser (Quantax microanalyser system composed of X-Flash® SDD detector and the Esprit software). The microscope was operated at an accelerating voltage of 20 kV and working distances of 10 mm.

3. Results and discussion

The first section of results presents the physico-chemical evolution of the binder alone without the addition of kaolin. The second part focuses on the description of the soil source material mixed with binder. To end with, the system investigated in this study is compared with lime-based systems already described in the literature.

3.1 Alkali activated high-calcium fly ash binder

3.1.1 X-ray diffraction (XRD)

Results obtained by XRD show that the original high-calcium fly ash is constituted of a vitreous phase (hump between 17°2θ and 38°2θ), and crystalline phases which include calcium-containing minerals: anhydrite CaSO₄, calcite CaCO₃ and portlandite Ca(OH)₂, and other minerals: quartz SiO₂, feldspar (K,Na,Ca)(Si,Al)₄O₈, hematite Fe₂O₃ and muscovite (Si₃Al)O₁₀(Al₂)(OH)₂K (Fig. 1).
Fig. 1. XRD of raw fly ash and fly ash based alkali activated material F100 as a function of curing time; A=anhydrite; C=calcite; F=feldspar; H=hematite; M=muscovite; P=portlandite; Q=quartz; T=thenardite.

Fig. 1 also shows XRD patterns of the alkali activated high-calcium fly ash as a function of curing time. Regarding the crystalline phases, anhydrite CaSO$_4$ and portlandite Ca(OH)$_2$ are consumed as a function of time. In addition, thenardite Na$_2$SO$_4$ starts forming at 3 days. Its formation can be explained by the release of sulphate anions from anhydrite dissolution, and its subsequent combination with sodium issued from the alkaline solution.

3.1.2 Derivative thermogravimetric analysis (DTG)

Fig. 2 shows derivative thermogravimetric analyses of the original high-calcium fly ash, and the high-calcium fly ash based alkali activated binder as a function of time. The DTG peaks of the original high-calcium fly ash detected at 385, 605 and 965 °C indicate the decomposition of portlandite Ca(OH)$_2$, calcium carbonate CaCO$_3$ and anhydrite CaSO$_4$ respectively.
Fig. 2. DTG curves of raw fly ash and fly ash based alkali activated material F100 as a function of curing time.

The relatively low decomposition temperature of anhydrite compare to its theoretical decomposition at 1214°C [31] is ascribed to the fact that anhydrite is found interlinked with other calcium-rich phases by SEM (as illustrated later in section 3.1.4.1).

Regarding the alkali activated high-calcium fly ash, and complementary to XRD, thermogravimetric analyses confirm the consumption of portlandite Ca(OH)$_2$ over time. DTG curves also show the dissolution of calcium carbonate CaCO$_3$ with time. Finally, above 780 °C, the observed mass losses of the activated samples reveal the decomposition of sulphate minerals: anhydrite CaSO$_4$, thenardite Na$_2$SO$_4$ being formed from 3 days as detected earlier by XRD (see 3.1.1), for which theoretical polymorphic transformation occurs at around 900°C [32-33].

3.1.3 Fourier Transform Infrared spectroscopy (FTIR)
Fig. 3A shows FTIR spectra of the original high-calcium fly ash, and the high-calcium fly ash based alkali activated binder as a function of time in the range of $\text{CO}_3^{2-}$ stretching vibrations.

![FTIR spectra](image)

Fig. 3. FTIR of raw fly ash and fly ash based alkali activated material F100 as a function of curing time in the $\text{CO}_3^{2-}$ stretching vibrations range (A) and raw kaolin and alkali activated kaolin KF50 as a function of curing time in the OH stretching vibrations range (B).

Different features of the $\text{CO}_3^{2-}$ band is observed over time: at 1 day a single band around 1453 cm$^{-1}$ is seen, while at 28 days a doublet positioned at 1426 and 1483 cm$^{-1}$ is observed. Those modifications validate the formation of calcium carbonate during the curing time as observed by DTG.

Finally, Fig. A.1 (see appendices) shows FTIR spectra in the area of sulphate minerals. It confirms the dissolution of calcium sulphate (anhydrite, $\text{CaSO}_4$) along with the subsequent formation of sodium sulphate (thenardite, $\text{Na}_2\text{SO}_4$) as previously seen by XRD (see 3.1.1).
3.1.4 Scanning Electron Microscopy (SEM)

3.1.4.1 Calcium-rich phases

Fig. 4 shows SEM observations of the alkali activated high-calcium fly ash at 1 day. More specifically, it focuses on calcium-rich phases previously detected as reactive phases being dissolved following the alkali attack by XRD, TGA and FTIR.

Calcium-rich phases are initially present in high-calcium fly ash as nodules of large size from 100 to 250 μm. Besides, it is seen that chemical elements such as calcium, sulphur and silicon are not homogeneously spread within nodules suggesting a varying mineralogy. For instance, in area 2, despite a high content of sulphur indicating a prevalence of anhydrite (CaSO₄), the detected percentage of SO₃ with respect to CaO remains too low to be owed to the presence of anhydrite phases only. It is therefore concluded that the various calcium-containing phases such as anhydrite CaSO₄, calcite CaCO₃ and portlandite Ca(OH)₂ (previously detected by XRD and DTG in section 3.1.1 and 3.1.2) are interlaced within those nodule structures which represent reactive structures under alkaline attack.
Fig. 4. SEM observations of a calcium-rich nodule from the alkali activated fly ash at 1 day: (A) SEM micrograph, (B) chemical composition (wt. %) of areas 1 and 2, (C) and (D) chemical mappings of calcium and sulphur respectively.

Overall sample

Fig. 5 additionally shows microstructural observations of the alkali activated high-calcium fly ash binder. At 1 day (Fig. 5A), a porous material along with distinct unreacted high-calcium fly ash particles is seen. In contrast, at 28 days (Fig. 5B), a more compact material with less pores is observed evidencing the formation of new compounds. Moreover, at 28 days, several unreacted high-calcium fly ash particles are still observed especially from the vitreous phase i.e. spherical and vesicular particles (see Fig. 5B, C and D).
Fig. 5. SEM micrographs of (i) the alkali activated fly ash binder F100 (Column 1) (A) at 1 day and (B), (C) and (D) at 28 days, (ii) the alkali activated kaolin KF50 (Column 2) (E) at 1 day and (F), (G) and (H) at 28 days; C=calcium nodules; G=glass; Q=quartz.
Additional chemical analyses revealed that changes in microstructure are more significant around calcium-rich particles which were previously detected as the main reactive part of the raw high-calcium fly ash.

As a matter of fact, SEM micrographs show zones of higher density appearing brighter around calcium-rich nodules (see encircled area in Fig. 5B).

Table 3 gives an average chemical composition of the denser reactive areas. Notably, it indicates a ratio of sodium to sulphur around 2 matching with that of precipitated thenardite whose formula is Na₂SO₄, and implying that most of the sodium from the alkaline solution is taken up to form thenardite.

Table 3

<table>
<thead>
<tr>
<th></th>
<th>Si</th>
<th>Ca</th>
<th>Al</th>
<th>Na</th>
<th>S</th>
<th>K</th>
<th>Fe</th>
<th>Mg</th>
<th>O</th>
</tr>
</thead>
<tbody>
<tr>
<td>F100-28d</td>
<td>22.1</td>
<td>19.2</td>
<td>7.0</td>
<td>3.7</td>
<td>1.7</td>
<td>1.2</td>
<td>0.8</td>
<td>0.5</td>
<td>43.8</td>
</tr>
<tr>
<td>KF50-28d</td>
<td>24.1</td>
<td>14.5</td>
<td>8.8</td>
<td>3.1</td>
<td>1.6</td>
<td>1.6</td>
<td>0.7</td>
<td>0.3</td>
<td>45.3</td>
</tr>
</tbody>
</table>

Furthermore, Table 3 indicates that the massive reactive area is primarily composed of silicon and calcium. Supposing that the new compounds are mainly located in those denser areas, these results suggest that apart from thenardite the new compounds are enriched in silicon and calcium.

3.1.5 Nuclear Magnetic Resonance (NMR)

Nuclear Magnetic Resonance was finally used to follow amorphous phases and precise the structure of the new compounds formed in our investigated system.

Fig. 6 shows $^{27}$Al MAS-NMR spectrum of the original high-calcium fly ash as well as the spectrum belonging to the alkali activated high-calcium fly ash at 28 days. $^{27}$Al NMR spectrum of the original high-calcium fly ash displays two resonances whose
dissymmetrical shapes are due to electric field gradient distribution caused by the distribution of geometries of the AlO₄ and AlO₆ polyhedra. More specifically, the main resonance whose maximum is detected at 55 ppm corresponds to Al(IV) of the vitreous phase. While the resonance located at 3 ppm corresponds to Al(VI) of the vitreous phase.

Fig. 6. $^{27}\text{Al}$ MAS-NMR spectra of the raw fly ash and alkali activated fly ash at 28 days.

Comparatively, the high-calcium fly ash based alkali activated material spectrum at 28 days exhibits a main resonance sharper and a maximum shifted to 59 ppm corresponding to four-fold coordination of aluminium namely $q^4(4\text{Si})$. Those modifications in the spectrum compared to the original high-calcium fly ash indicate that part of the aluminium released from high-calcium fly ash leads to the formation a new aluminium bearing phase whose aluminium is tetrahedrally coordinated. Lastly, the resonance owed to octahedral aluminium at 3 ppm is still present which means that high-calcium fly ash is not totally dissolved after 28 days.
Fig. 7 shows $^{29}\text{Si}$ MAS-NMR spectra of the original high-calcium fly ash as well as the alkali activated high-calcium fly ash at 28 days. $^{29}\text{Si}$ NMR spectrum of the original high-calcium fly ash shows a broad resonance between -85 and -105 ppm attributed to the presence of a wide range of $Q^3$ and $Q^4$ silicon local environments from the vitreous phase.

In contrast, the alkali activated high-calcium fly ash spectrum at 28 days displays a clear additional resonance centred at -85 ppm. This new position can be attributed to both the formation of $Q^4(4\text{Al})$ or else $Q^2$-type silicon environments. Nevertheless, regarding the high value of the Si/Al ratio around 3.2 in the denser area comprising the new products, and measured by SEM (see Table 3) it is unlikely that $Q^4(4\text{Al})$ environments are present. The new resonance at -85 ppm consequently indicates the formation of $Q^2$-type silicon environments in chain structure. Considering that the spectrum does not show any resonance corresponding to $Q^1$ silicon environments the length of those formed chains is high. It is also worth noting that $Q^2$ Si environments in chain possess a charge deficit of
2 - which must be compensated. However, sodium cations are not available as they are associated with sulphur to form thenardite Na$_2$SO$_4$ (see sections 3.1.1 and 3.1.3). Consequently, only calcium cations released from calcium reactive phases of high-calcium fly ash can compensate this charge deficit. Those silicon chain structures are therefore combined with dissolved calcium which matches with the chemical composition of the denser reactive area primarily made of silicon and calcium (see Table 2). The significant broadening of the resonance at -85 ppm also indicates that those chains are not well organised as C–S–H structures generally observed in Portland cement [34]. Considering the low Ca content of the raw fly ash, the line broadening fully agrees with amorphous C-S-H with a low Ca/Si ratio.

Finally, the broadest part of the activated high-calcium fly ash spectrum at 28 days from -90 ppm to -100 ppm indicates the presence of Q$^3$ and Q$^4$ environments mainly issued from the remaining vitreous phase of high-calcium fly ash, and in accordance with the previous observations of unreacted high-calcium fly ash particles at 28 days by SEM (see 3.1.4.2). In summary, both $^{29}$Si and $^{27}$Al MAS-NMR spectroscopy indicates the formation of new signals following alkaline activation. It is of interest to understand whether the new aluminium-containing phase seen in $^{27}$Al NMR correlates with the silicate chain structure observed in $^{29}$Si NMR. As a comparison, aluminium in linear structure such as C–S–H is either found (i) as Q$^2$ environments corresponding to aluminium substituting for silicon atoms, and located around 68-74 ppm in $^{27}$Al NMR or else (ii) as Q$^3$ environments corresponding to crosslinking through alumina bridging tetrahedra positioned around 63-68 ppm in $^{27}$Al NMR [35-37]. However, as mentioned above, the new aluminium resonance observed in our investigation at 59 ppm would rather correspond to q$^4$(4Si) environment. Consequently, if aluminium is incorporated into the silicate chain structure,
it would be in a three-dimensional environment which has not been described in literature yet. To conclude, results concerning the alkali activated high-calcium fly ash binder showed that calcium-rich phases constitute the reactive part of the raw high-calcium fly ash, while its vitreous phase remains mainly unreactive. The new compounds formed are mainly located around calcium-rich reactive particles and present a complex chemistry and structure which differ from cementitious compounds generally encountered in cement or lime based system.

3.2 Interaction between the alkali activated high-calcium fly ash binder and kaolin

The following section aims at understanding the interaction between kaolin and the alkali activated high-calcium fly ash binder previously described. More specifically, it aims at answering the following question: does the presence of kaolin modify the reaction sequence?

Two stabilised soils were studied i.e. KF50 for which the solid phase is made in mass of 50% of high-calcium fly ash and 50% of kaolin, and KF20 made of 20% of high-calcium fly ash and 80% of kaolin. Observations made for these two mixes turned out to be similar for all the techniques used. Therefore, only the results of KF50 will be shown while the results of KF20 can be found in Supporting Information.

Firstly, Fig. 3B shows the infrared spectrum of the original kaolin as well as the spectra belonging to the alkali activated soil KF50 as a function of time. The four bands observed in the 3695–3620 cm\(^{-1}\) range are typical of the presence of kaolinite, and arise from the vibration of its internal OH groups. Notably, disorder in kaolinite is mainly detectable in this OH-stretching region by FTIR [38]. Those bands being still observed over time in...
the alkali activated soils suggest that kaolinite does not react under alkaline conditions.

The smaller heights seen at 7 and 28 days are only due to higher sample densities at higher curing times after the formation of new compounds, leading to smaller probed distance and therefore lesser absorbance.

Fig. A.3 (see appendices) shows XRD patterns of the activated soil KF50 as a function of curing time. Similarly to the alkali activated high-calcium fly ash binder (see 3.1.1), it indicates the dissolution of anhydrite CaSO$_4$ along with the formation of thenardite Na$_2$SO$_4$.

Fig. 5E, F, G and H shows microstructural observations of the alkali activated kaolin KF50. At 1 day, and in contrast with the alkali activated high-calcium fly ash binder (Fig. 5A), KF50 presents a relatively low porosity due to the presence of small-sized kaolinite platelets filling the pores (Fig. 5E). At 28 days, and in a similar way to the activated high-calcium fly ash (Fig. 5B), KF50 presents a more compact microstructure around calcium-rich phases (see encircled area in Fig. 5F).

Furthermore, by comparing the alkali activated high-calcium fly ash binder (Fig. 5C and D) with the activated soil KF50 (Fig. 5G and H), kaolinite platelets are distinctly observed (see red arrows as an example of platelet observation), and homogeneously spread across the whole sample. In fact, kaolinite platelets were observed not merely in the matrix but also in the most reactive massive areas supporting that kaolinite does not react even in the reactive areas but rather acts as a filler.

Table 2 gives an average chemical composition of KF50 massive areas. It shows similar tendencies than for the alkali activated high-calcium fly ash binder F100. Only slightly
higher contents of silicon and aluminium are measured for KF50 due to the presence of kaolinite.

Fig. 8 shows $^{27}$Al MAS-NMR spectra of the alkali activated high-calcium fly ash binder F100 studied in the first section, and the alkali activated soil KF50 at 1 and 28 days. By comparison with the binder F100, $^{27}$Al NMR spectroscopy of KF50 displays two additional resonances due to the presence of kaolin i.e. a main resonance at 4 ppm due to Al(VI) of the octahedral layer of kaolinite, and a resonance at 70 ppm owed to Al(IV) and corresponding to substitution of Al for Si in the tetrahedral layer of kaolinite and muscovite.

At 1 day, KF50 presents a resonance whose maximum is located at 55 ppm and owed to the vitreous phase of high-calcium fly ash, while at 28 days a shift of this resonance to 58 ppm indicates the formation of tetrahedral aluminium in $q^4(4Si)$ environments (as previously described for the alkali activated high-calcium fly ash in section 3.1.5).

![Graph showing $^{27}$Al MAS-NMR spectra of F100 and KF50 at 1 and 28 days.](image)
Fig. 8. $^{27}\text{Al}$ MAS-NMR spectra of the alkali activated fly ash F100 at 28 days and alkali activated kaolin KF50 at 1 and 28 days.

Fig. 9 shows $^{29}\text{Si}$ MAS-NMR spectra of the alkali activated high-calcium fly ash binder F100 studied in the first section, and the alkali activated soil KF50 at 1 and 28 days. $^{29}\text{Si}$ NMR spectroscopy of KF50 samples show an additional thin resonance at -91 ppm corresponding to the silicon of the tetrahedral layer of kaolinite. This resonance does not undergo any modification over time confirming the non-reactivity of kaolinite.

Furthermore, from 1 to 28 days KF50 shows the clear appearance of a resonance at -85 ppm due to the formation of silicon chains combined with dissolved calcium, and as previously described for the alkali activated high-calcium fly ash binder (see 3.1.5).

To conclude, results of this second section showed that kaolin is unreactive during alkaline attack. Besides, a similar reaction sequence than for the alkali activated high-calcium fly ash binder occurs.
Fig. 9. $^{29}\text{Si}$ MAS-NMR spectra of the alkali activated fly ash F100 at 28 days and alkali activated kaolin KF50 at 1 and 28 days.

### 3.3 Comparison with lime treated kaolin

This last section focuses on a comparison with previous studies which reported the physicochemical evolution of the same kaolin treated either by a common stabiliser: lime [6], but also treated by a mix of lime and the high-calcium fly ash used here [39].

Table 4 gives a summary of the reaction sequences for each system highlighting strong differences. This section consequently aims at providing a better understanding of the consequences of those variable reaction sequences on the final material in terms of performance and durability.

#### Table 4

Compared reactivity sequences of kaolin treated either by lime, or a mix of lime and fly ash or an alkali activated fly ash
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Phases dissolved</td>
<td>Kaolin, lime</td>
<td>Kaolin, lime, fly ash</td>
<td>Fly ash</td>
</tr>
<tr>
<td>2. Reactivity timescale</td>
<td>Slow</td>
<td>Fast</td>
<td>Fast</td>
</tr>
<tr>
<td>3. Phases formed</td>
<td>Aluminate Calcium Aluminate Hydrate C₆A₆H Calcium Aluminium Oxide Hydrate C₆A₆O₆C₆H Εττριγίτη Calcium Aluminate Hydrate C₆A₆O₆H Ca₆Al₆(SO₄)₆(OH)₁₂·2₆H₂O</td>
<td>Amorphous silicate consisting of chains combined with calcium and probably incorporating the observed q₄(4Si) aluminium environments</td>
<td></td>
</tr>
</tbody>
</table>

3.3.1 Reactivity of initial phases

Firstly and regarding the reactivity of the phases initially present, it was found that kaolin is inert in our alkali activated samples which contrasts with the two other lime based systems for which kaolinite is dissolved. The limited reactivity of kaolinite in the alkali activated soils herein studied is due to the presence of other more reactive phases. Besides, its limited reactivity is beneficial for the system as no side effects will occur.

For the three systems, calcium-containing phases constitute the main reactive part of the mixes playing a pivotal role in the reaction development scheme. In fact, their dissolution leads to the release of dissolved calcium into the medium and hence pozzolanic activity i.e. formation of new calcium cementitious compounds responsible of the strength improvement. In the case of our alkali activated soils however, it is not lime CaO that constitutes a supply of calcium but calcium-containing minerals from high-calcium fly ash CaSO₄, CaCO₃ or Ca(OH)₂. For all calcium-source types the supply of Ca²⁺ cations remains identical. What changes is the anion simultaneously released from their dissolution: when high-calcium fly ash is present the dissolution of its calcium-containing minerals is accompanied by the release of various anions such as OH⁻, SO₄²⁻ and CO₃²⁻.
influencing the reaction sequence, and with potential negative effects for the durability as seen later.

Finally, concerning the vitreous phase of high-calcium fly ash, although amorphous and hence metastable it showed few reactivity leading to a preferential pozzolanic activity as seen in Portland cement, rather than polymerisation reactions associated with the formation of an aluminosilicate three-dimensional network characteristic of low-calcium alkali activated materials and geopolymers.

3.3.2 Reactivity timescale

Reactions in presence of high-calcium fly ash are fast with new compounds already observed at 28 days for both systems made of kaolin, lime and high-calcium fly ash [39] but also kaolin and alkali activated high-calcium fly ash. Whereas, longer reactivity timescale occurs for lime treated kaolin: new cementitious compounds being previewed from 60 days and clearly detected only after 270 days [6].

For a system made of lime and kaolin its pozzolanic activity depends on the dissolution of kaolinite which constitutes the only source of aluminium and silicon. Considering that kaolinite possesses a stable crystalline mineral structure hard to dissolve it explains the slow reactivity of that system. By contrast, high-calcium fly ash contains reactive phases i.e. calcium-rich phases primarily and to a small extent its vitreous phase (for the high-calcium fly ash used here) thermodynamically less stable than kaolinite and therefore easier to dissolve. That is why reaction sequences are faster for both high-calcium fly ash systems. It is even faster for our alkali activated soils as the alkaline solution brings a mixture of ions ready available. Those fastest reaction times would constitute an advantage in the case where the quickly formed products are as well stable binding phases which will be discussed now.
3.3.3 Stability and structure of the compounds formed

In the case of formerly studied systems made of kaolin and lime as well as kaolin, lime and high-calcium fly ash, a preferential release of aluminium over silicon from kaolinite and/or high-calcium fly ash dissolution occurs conducting to the formation of aluminium compounds primarily. In fact, for a lime treated kaolin Calcium Aluminate Hydrate $\text{Ca}_2\text{Al}_2\text{O}_4\cdot\text{H}_2\text{O}$ and monocarboaluminate hydrate $\text{Ca}_4\text{C}_2\text{Al}_6\text{O}_{16}\cdot\text{H}_2\text{O}$ are formed [6]. In the case of a kaolin treated by a mix of lime and high-calcium fly ash Calcium Aluminate Hydrate $\text{Ca}_2\text{Al}_2\text{O}_4\cdot\text{H}_2\text{O}$ but also Calcium Aluminium Oxide Carbonate Hydrate $\text{Ca}_6\text{Al}_2\text{O}_7\cdot\text{CaCO}_3\cdot\text{H}_2\text{O}$, Calcium Aluminium Oxide Hydrate $\text{Ca}_6\text{Al}_2\text{O}_7\cdot\text{H}_2\text{O}$ and Ettringite $\text{Ca}_6\text{Al}_2(\text{SO}_4)_3(\text{OH})_{12}\cdot26\text{H}_2\text{O}$ are formed [39]. Whereas, in our alkali activated soils investigated here, aluminium dissolved from the vitreous phase of high-calcium fly ash was found in a three-dimensional four-fold environment ($q^4(4\text{Si})$) which strongly contrasts with the six-fold environments found in the Calcium Aluminate Hydrates.

Regarding the formation of silicate compounds, none are formed in a system made of kaolin and lime [6] because the dissolution of kaolinite is slow and starts by the release of its aluminium. Hence, no silicon is made available. Kaolinite being the only source of silicon in that system the limited dissolution of silicon implies the non-formation of silicon compounds. By contrast in a kaolin, lime and high-calcium fly ash system, the dissolution of silicon from the vitreous phase of high-calcium fly ash leads to the formation of Calcium Silicate Hydrate $\text{Ca}_2\text{SiO}_4\cdot\text{H}_2\text{O}$ [39]. Finally, in the alkali activated soils herein studied, the supply of silicon from the alkaline solution primarily and also from the vitreous phase of high-calcium fly ash leads to the formation of silicon chains combined with calcium, but whose NMR signature greatly differs from $\text{Ca}_2\text{SiO}_4\cdot\text{H}_2\text{O}$ commonly observed as described above (see section 3.1.5). It is also likely that aluminium
found in three-dimensional four-fold environment \((q^4(4Si))\) is incorporated into those silicon chains. Calcium Silicate Hydrate C–S–H is the principal binding phase of Portland cement and concrete primarily responsible for its strength [40]. In addition, its structure is more stable than Calcium Aluminate Hydrates [41]. Their presence is consequently beneficial for the performances. The fact that in our alkali activated samples, a diverse structure compared to usual C–S–H is observed cannot easily be assessed in term of stability at the present moment. Indeed, regarding the lack of crystallinity of our silicon chains formed, experience proved that the crystallinity of the binding agent alone constitutes a poor measure of stability over the timescales relevant to the majority of concrete structures [41]. A further investigation of the performances would help apprehending a potential beneficial effect of this uncommon silicon chains structure over time.

Finally, for both high-calcium fly ash systems, sulphate minerals are formed: either Ettringite \(\text{Ca}_6\text{Al}_2(\text{SO}_4)_3(\text{OH})_{12}.26\text{H}_2\text{O}\) for a system made of kaolin, lime and high-calcium fly ash [39], or Thenardite \(\text{Na}_2\text{SO}_4\) in our alkali activated soils. Their formation is due to the dissolution of anhydrite \(\text{CaSO}_4\) initially present in the high-calcium fly ash, and releasing sulphate anions \(\text{SO}_4^{2-}\) that are subsequently recombining with available cations. For the case of a kaolin, lime and high-calcium fly ash system the formation of Ettringite is taking up aluminium and calcium hence slowing down both the simultaneous formation of aluminate and silicate hydrates. Whereas, in our alkali activated system, sodium cations ready available from the alkaline solution combine with sulphate anions. Therefore, the formation of sulphate minerals does not affect the parallel development of the pozzolanic activity i.e. the formation of silicate chains. Finally, a key
point to consider for durability aspects is the high solubility of sulphate minerals which are few stable salts in water. In fact, a previous study in which leaching tests were performed on an alkali activated sulphate-bearing kaolin showed that the uptake of sulphate anions by the gel is low, namely less than 40% [42]. The study of the effect of wetting-drying cycles is consequently warranted to verify a further impact of the presence of thenardite on the long-term performances.

3 Conclusions

Here, the development of a novel soil binder that is an alkali activated calcium-rich high-calcium fly ash for clay soil stabilisation was explored. The study of its reactivity showed that (i) the overall calcium-bearing minerals from high-calcium fly ash constitute the reactive phases while its vitreous phase remains mainly unreactive, (ii) new compounds are formed, thenardite Na₂SO₄ and an amorphous silicate consisting of chains combined with calcium - (iii) reactions happen within 1 to 28 days. The interaction between the binder developed and the model soil chosen i.e. kaolin, showed that kaolin is unreactive. Its presence whatever the proportion does not modify the physicochemical evolution of the system that is neither the dissolved phases, formed compounds, nor reactivity timescale. The inert kaolinite platelets were in addition found homogenously embedded in the matrix acting as a filler. When compared to lime treated kaolin, although pozzolanic activity remains the dominant process reaction sequences are strongly different. In the case of alkali activated soils the formation of calcium-silicon chains phases more stable than calcium aluminium hydrates encountered in lime based systems is beneficial for long-term stability purpose. Those observed silicon chains however show an uncommon structure whose effect on the performances will be checked in a future investigation. Finally, the formation of
thenardite a highly soluble salt in water raises interest about the durability of the material
which will be also further investigated.

Acknowledgements

The authors wish to acknowledge the support of the European Commission via the Marie
Skłodowska-Curie Innovative Training Networks (ITN-ETN) project TERRE 'Training
Engineers and Researchers to Rethink geotechnical Engineering for a low carbon future’
(H2020-MSCA-ITN-2015-675762).

Appendices

Fig. A.1. FTIR of raw high-calcium fly ash, and high-calcium fly ash based alkali
activated material F100 as a function of curing time in the $\text{SO}_4^{2-}$ stretching vibrations
range; 681, 616 and 596 cm$^{-1}$ = anhydrite CaSO$_4$; 639 and 617 cm$^{-1}$ = thenardite
$\text{Na}_2\text{SO}_4$. 
Fig. A.2. FTIR of the raw kaolin and alkali activated kaolin KF20 as a function of curing time in the OH stretching vibrations range.

<table>
<thead>
<tr>
<th>Wavenumber (cm⁻¹)</th>
<th>Absorbance</th>
</tr>
</thead>
<tbody>
<tr>
<td>3600</td>
<td></td>
</tr>
<tr>
<td>3605</td>
<td></td>
</tr>
<tr>
<td>3609</td>
<td></td>
</tr>
<tr>
<td>3613</td>
<td></td>
</tr>
<tr>
<td>3620</td>
<td></td>
</tr>
</tbody>
</table>

Fig. A.3. XRD of alkali activated kaolin KF50 as a function of curing time;

A=anhydrite; C=calcite; F=feldspar; H=hematite; K=kaolinite; M=muscovite;

P=portlandite; Q=quartz; T=thenardite.
Fig. A.4. XRD of the alkali activated kaolin KF20 as a function of curing time; A=anhydrite; C=calcite; F=feldspar; H=hematite; K=kaolinite; M=muscovite; P=portlandite; Q=quartz; T=thenardite.

Fig. A.5. $^{27}$Al MAS-NMR of the raw high-calcium fly ash, raw kaolin and alkali activated kaolin KF20 at 0 and 28 days.
Fig. A.6. $^{29}\text{Si}$ MAS-NMR of the raw high-calcium fly ash, raw kaolin and alkali activated kaolin KF20 at 0 and 28 days.

References


