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**Aggregate size effect on the development of cementitious compounds in a
lime-treated soil during curing**

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Abstract

This work aims to investigate the aggregate size effect on changes in mineralogical composition and microstructure of lime-treated compacted soils. Three soil powders with different maximum aggregate sizes ($D_{max} = 5, 1$ and 0.4 mm) were prepared prior to the treatment with 2% of lime. X-ray diffraction (XRD), environmental scanning electron microscope (Env. SEM) coupled with chemical analysis using energy dispersive X-ray spectrometry (EDX) and mercury intrusion porosimetry (MIP) were used to analyse untreated and treated samples at various curing times. Crystallized C-S-H on tobermorite form was identified in the lime-treated soil prepared with large aggregates after one year curing, and an evident increase in nanopores less than $0.1\text{ }\mu\text{m}$ C-S-H was also observed due to C-S-H creation. By contrast, in the case of smaller aggregates, no obvious C-S-H peaks were observed by XRD technique after the same curing time, even though some evidence of such phases are provided by Env. SEM coupled to EDX analysis. But a large amount of undetectable nanopores less than 6 nm (considering the MIP technical limitation) was supposed to be formed and could be attributed to the creation of nanocrystallized C-S-H or poorly-crystallized C-S-H (that may fill the pores larger than $2\text{ }\mu\text{m}$). Such type of C-S-H phases occurred when lime was coated in thin layer on the large surface associated to lime-treated soil prepared with small aggregates.

Keywords: lime-treated soil; aggregate size; curing time; mineral composition; microstructure

1. Introduction

Lime treatment is an effective soil improvement technique widely applied in the field of construction. It largely modifies the soil geotechnical properties through the physico-chemical reactions within the lime-soil-water system (Boardman et al., 2001; Russo, 2005; Al-Mukhtar et al., 2010; Tang et al., 2011; Tran et al., 2014). When quicklime (CaO), soil and water are mixed together, hydration and ionization of quicklime immediately take place. Then, the Ca^{2+} ions in the pore water released by calcium hydroxide ($\text{Ca}(\text{OH})_2$) is adsorbed by ion-exchange at clay minerals surface. The diffuse hydrous double layer surrounding the clay particles can be modified by the Ca^{2+} ion-exchange process, resulting in the flocculation-agglomeration of clay particles (Bell, 1996). These modifications of clay particles induced by lime addition will largely improve the workability of soil by reducing the plasticity, the swelling and shrinkage (Bell, 1989; Russo, 2005). In the long-term, the main reactions between lime and clay minerals are of pozzolanic nature which contributes significantly to the improvement of soil mechanical behaviour in terms of shear strength, shear modulus, compression strength and compressibility (Rajasekaran and Narasimha Rao, 2002; Khattab et al., 2007; Consoli et al., 2009; Tang et al., 2011; Dong, 2013). The mechanical improvement is to be attributed to the creation of cementitious compounds generated by the pozzolanic reaction, coating the soil particles and bonding them together (Bell, 1996; Onitsuka et al., 2001; Nalbantoglu, 2006).

Mineralogical studies of cementitious compounds have been undertaken in recent years. The cementitious compounds can be of various forms due to the different mineralogical composition of soils containing mainly clay minerals such as kaolinite, montmorillonite or illite, and other minerals like quartz and feldspars. Generally, the main cementitious

compounds are calcium silicate hydrate (C-S-H), calcium aluminate hydrate (C-A-H) and calcium alumino-silicate hydrate (C-A-S-H) (Khattab, 2002; Rios et al., 2009; Maubec, 2010; Al-Mukhtar et al., 2010). For lime-kaolinite mixture, the production of C-S-H, C-A-H and C-A-S-H was reported by many researchers (Goldberg and Klein, 1952; Eades and Grim, 1960; Glenn and Handy, 1963; Willoughby et al., 1968; Bell 1996). C-S-H and C-A-H were also detected in the lime-treated montmorillonite (Bell 1996; Hilt and Davidson, 1960). Arabi and Wild (1989) noted that C-S-H hydrates were present in the lime-treated marls containing illite, quartz and feldspar. Eades et al. (1962) also identified the production of C-S-H in the lime-treated quartz.

Even though many studies were performed on lime-treated soils, most of them focused on the lime-treated soil samples prepared in the laboratory. However, often lower performance of lime-treated soils and poor durability of lime treatment are observed in the field conditions. Puppala et al. (2006) reported about 40% lower for stiffness and 20 to 30% lower for strength in the case of treatment in field. Similar results were reported by other researchers (Horpibulsuk et al., 2006; Kavak and Akyarh, 2007). Additionally, higher hydraulic conductivity and swelling potential of lime-treated soils in the field conditions were observed (Bozbey and Guler, 2006; Cuisinier and Deneele, 2008). There are several factors that can contribute to this difference between field and laboratory conditions. In addition to the climatic factors, especially the wetting/drying cycles and freezing/thawing cycles (Pardini et al., 1996; Guney et al., 2007; Tang et al., 2011; Stoltz et al., 2012), the aggregate size may play an essential role in the hydro-mechanical behaviour of lime-treated soils (Tang et al., 2011, Wang et al. 2015). The bender elements testing performed on the lime-treated samples

prepared with different maximum aggregates sizes ($D_{max} = 5, 2, 1$ and 0.4 mm) revealed that lime-treated soils prepared with larger maximum aggregate size presented a relatively lower stiffness (Tang et al., 2011). Dong (2013) also indicated that the lime-treated soil prepared with larger aggregates was more sensitive to wetting/drying cycles. Note that aggregates are assemblages of adjacent soil particles in which the cohesive forces are larger than the disrupting force (Kemper and Chepil, 1965). In the field construction, the scarifying/pulverizing process is performed before the lime treatment to control the soil aggregate size. However, the size of soil aggregates in the field can still reach several centimetres, which is much larger than that of soil aggregates prepared in the laboratory before sample reconstruction. In the laboratory, natural tested soils are usually air-dried, ground and sieved into few millimetres. For example, Du et al. (2014) who studied the engineering properties and microstructure of the cement-stabilized contaminated soil, prepared the samples with kaolin clay which had a maximum aggregate size lower than 2 mm; while Cai et al. (2015) used reactive magnesia to treat in the laboratory a silty soil with the same maximum aggregate size. Jiang et al. (2016) also reported that the used soil was first passed through the sieve with 0.5 mm size prior to treatment.

As the improvement in the mechanical behaviour of soils by lime treatment is proven to be primarily controlled by the cementitious compounds from the pozzolanic reactions, it is expected that the different behaviours of treated soils with various aggregate sizes can be also interpreted from mineralogical analysis. However, no studies have been conducted on this aspect. This constitutes the main objective of the present work. In this study, three different maximum aggregates sizes ($D_{max} = 5, 1$ and 0.4 mm) of soil powders were prepared before

lime treatment. The creation of cementitious compounds was investigated by X-ray diffraction (XRD). In addition, environmental scanning electron microscope coupled with energy dispersive X-ray spectrometry (Env. SEM-EDX) and mercury intrusion porosimetry (MIP) were applied to investigate the microstructure and chemical composition of the treated soils.

2. Materials and methods

The tested soil was taken in Héricourt (France). It is a plastic silt with a clay-size fraction of 27 %. The basic geotechnical properties of this silt given by Wang et al. (2016) are listed in Table 1. This soil corresponds to a silt of high plasticity (MH) following the Unified Soil Classification System (USCS). To prepare soil powders with different maximum aggregate sizes, natural soil was first air-dried, gently ground to crush the block of soil and passed through three target sieves (their maximum diameters, D_{max} , are 5, 1 and 0.4 mm, respectively). The large aggregates which could not pass through the sieve were ground manually until all particles passed through, ensuring no changes in mineralogical compositions during sieving (Tang et al., 2011). Afterwards, soil powders S5, S1 and S0.4 are obtained, with D_{max} equal to 5 mm for S5, 1 mm for S1, and 0.4 mm for S0.4. Figure 1 shows the aggregate size distributions of the three soil powders, determined by dry sieving. Quicklime was used in this study and it has a high purity with a CaO content as high as 97.3 %. Particle size analysis shows that 82.7 %, 95.2 % and 100 % of this lime could pass through 80 μ m, 200 μ m and 2 mm sieves, respectively (Dong, 2013). A lime content of 2 % by mass was selected as binder dosage.

The dry soils were firstly mixed thoroughly with 2 % quicklime. Then, distilled water was added by spray into the dry soil-lime mixture to obtain the target water content ($w = 17$ %, dry

side of the optimum according to the proctor curve). Static compaction was performed after a mellowing period of 1 hour, to prepare soil samples at the target density ($\rho_d = 1.65 \text{ Mg/m}^3$). The samples were carefully wrapped by plastic membrane and scotch tape immediately after compaction. The well-covered sample was stocked in a hermetic box for curing in a chamber at a temperature of $20 \pm 2 \text{ }^\circ\text{C}$. Prior to mineralogical and microstructural analyses, the samples were freeze-dried following the procedure proposed by Delage and Pellerin (1984).

X-ray diffraction (XRD) analysis was performed on both untreated and treated samples. To prepare soil powder for this analysis, freeze-dried and oven-dried samples were crushed and ground to pass through a $32 \text{ }\mu\text{m}$ sieve. After sieving, soil powders were mixed well manually in agate mortar and sprinkled gently in XRD sample holder using a $65 \text{ }\mu\text{m}$ sieve. The top layer was removed carefully by cutting the surface with a thin razor blade leading to a smooth surface without compaction (such preparation allows to decrease the preferential orientation of clay particles). XRD patterns were obtained using a D8 Advance diffractometer from Bruker (θ - θ configuration, Cobalt anode, $E = 35 \text{ kV}$, $I = 40 \text{ mA}$, no monochromator, LynxEye detector). A continuous scan mode, between 3 and $80^\circ 2\theta$, at a rate of $1 \text{ s per } 0.01^\circ 2\theta$ was selected. Diffractograms were exploited with EVA program coupled with the ICPdf2 mineralogical database.

Environmental scanning electron microscope (Env. SEM, Quanta 400 from FEI company) coupled with energy dispersive X-ray spectrometry (EDX from EDAX company) was employed to observe the microstructure and to study the chemical composition of treated samples. The EDX probe which provides an order of magnitude of the chemical content of elements such as Ca, Si and Al was calibrated before the observations. Images were collected

on fresh fractured surface after freeze-drying. Secondary or back scattered electron mode was selected in low vacuum mode (no metal coating was applied before observation).

The mercury intrusion porosimetry (MIP) test was carried out on freeze-dried samples. The applied pressure ranged from 3.4 kPa to 230 MPa. The corresponding mercury intrusion diameter varied from 0.006 μm to 355 μm . In the analysis, the surface tension of mercury was taken equal to 0.485 N/m and the contact angle of mercury was taken equal to 130°.

Table 2 presents the test program. All the 2 % lime-treated samples were freeze-dried before test except a piece of treated sample S1 after 1-year curing that was oven-dried at 105 °C prior to testing. Untreated soil sample was also tested by XRD to obtain data as reference.

3. Experimental results

3.1. Mineralogical analysis

Figure 2 shows the XRD patterns of untreated soil and lime-treated soil S1 after 1-year curing. The results illustrate that the main minerals in untreated soil are quartz and clays with the presence of calcite and feldspars. The identification of clay minerals confirms the presence of kaolinite ($d \sim 7 \text{ \AA}$), illite/muscovite ($d \sim 10 \text{ \AA}$), and montmorillonite ($d \sim 12\text{-}14 \text{ \AA}$). This clay composition is consistent with previous analyses made by Deneele and Lemaire (2012), who studied the same soil as that used in this study. The reflections associated with these clay minerals (with quite similar intensity or position) are also present in the lime-treated soil. For the lime-treated soil S1 ($t = 1 \text{ year}$) after freeze-drying, one new reflection at 2θ equal to 34.2° ($d \sim 3.04 \text{ \AA}$) is identified, suggesting creation of the C-S-H phase. This C-S-H phase would correspond to the main peak of synthetic tobermorite 9 $\text{\AA}$ with the formula $5\text{Ca}_{0.65}\text{SiO}_2$,

2.5H₂O (Pdf file 010-0374) or to the calcium silicate hydrate Ca_{1.55}SiO_{3.5} xH₂O (Pdf file 033-0306). In order to remove the doubt of interpretation related to possible confusion among the phases of unreacted CaO, C-S-H (tobermorite and other C-S-H forms) and magnesian calcite, the XRD pattern of treated soil S1 after oven-drying at 105°C was also collected.

The new peak previously observed disappeared which tends to validate the identified C-S-H phase. Indeed, oven-drying at 105 °C may dehydrate the C-S-H as suggested by Taylor (1997). Gallé (2001) mentioned also that C-S-H decomposition starts at low temperatures and C-S-H can be partially dehydrated at 105 °C, and then its crystallized structure disappears. Rio et al. (2009) also observed that different stages of C-S-H dehydration took place at a temperature in the range from 100 to 250 °C. If oven-drying at 105 °C can cause C-S-H degradation, it is not the case for the magnesium calcite. Indeed, this latter remains stable under temperature effect because decarbonation of calcite occurs at high temperatures (over 700 °C) according to Collier et al. (2008), Rios et al. (2009) and Al-Mukhtar et al. (2014). If the new peak observed on XRD pattern was calcite, this peak would remain present whatever the sample preparation. It was obviously not the case. Moreover, the absence of new peak at $d \sim 3.04 \text{ \AA}$ at short curing time (after just 1-day curing) allows excluding the presence of unreacted CaO after 1-year curing.

Except the C-S-H phase, other minerals in the lime-treated soil seem similar to those of the untreated sample. Note that the crystallized silicate source used to form C-S-H phase could not be clearly identified by XRD because no clear peak decrease was detected, which would signify its dissolution.

The effect of curing time and aggregates size can be specifically observed in a smaller range

from 31° to 39° 2θ where the C-S-H phase involved on the XRD patterns. The results of treated samples S5 and S0.4 are illustrated in Figure 3a and 3b. Similarly to S1, S5 (Figure 3a) also showed an XRD peak associated with the creation of C-S-H in the case of 1-year curing. However, no sign of C-S-H was detected on the sample after a curing time of $t = 60$ days. Surprisingly, no C-S-H reflection was detected on the XRD pattern of treated soil S0.4 after 1 year curing (Figure 3b). This point will be further discussed.

3.2. Microstructure observation and chemical composition analysis

Figures 4a-c present the distribution maps of silicon, aluminium and calcium derived from EDX coupled with Env. SEM observations on treated soil S1 at a curing time of $t = 1$ day. The two first elements belong to aluminosilicates phases such as feldspar and clays.

Areas with high concentration of Si in Figure 4a indicate the presence of quartz grains. The calcium distribution map (Figure 4c) is established to localize hydrated lime after short curing time (in this case, Ca^{2+} has not enough time to diffuse homogeneously into soil aggregates) and the product of lime reaction such as C-S-H (at long time of curing). Even though the protocol to mix the soil with lime was optimized to cover all the soil aggregates, calcium seems to be distributed heterogeneously or rather, some aggregates of hydrated lime appears clearly after 1 day of curing in isolated clusters of several micrometers (Figure 4d). It is however worth noting that isolated small hydrated lime particles below the micron scale cannot be distinguished by Env. SEM. The observations do not allow the detection of the initial soil aggregate border formed by lime coating, whereas slight and continuous borders composed by lime are expected on EDX mapping.

On lime-treated soil S1 after 1-year curing (Figure 5), the calcium distribution map (Figure 5c)

helps to localize the new pozzolanic products as shown in Figure 5d. New phases generally present a typical water sensitive morphology of C-S-H particles characterised by a soft, porous and rounded texture. Such C-S-H phases can be visually detected by operator on Env. SEM images, only when their sizes are large enough to be distinguished, due to the limited resolution of the Env. SEM apparatus applied in this study. If C-S-H phases are present in particle form (but in a size smaller than 1 μm) or in form of thin gel layer, these C-S-H phases are invisible for the operator. However, the C-S-H phases may be present by means of either coating the soil aggregates, or filling the inter-aggregate pores, or binding the adjacent aggregates together.

Finally, even though the XRD did not detect crystallized C-S-H in the lime-treated soil S0.4 after 1-year curing time, the Env. SEM observation revealed local presence of C-S-H particles as shown in Figure 6. The low quantities of such large C-S-H particles may explain the result.

Quantitative EDX analyses complete the Env. SEM observations. Spot analysis at $E = 20 \text{ kV}$ were performed on areas situated on the EDX mapping in Figures 4 and 5 considering their high calcium content. Table 3 gives the atomic percentages of the main elements present in such areas compared to the atomic percentage measured on natural soil before lime addition and in the areas far from the selected calcium spot. The calculated atomic ratio Ca/Si is equal to 2.49 in areas rich in Ca^{2+} in soil S1 at $t = 1$ day while far from such area the Ca/Si ratio is equal to 0.06 (with Si/Al = 2.16). Far from the calcium rich areas, the Ca/Si ratio remains stable (or increases slightly to 0.09) even after 1 year of curing. In soil S1 at $t = 1$ day, the area rich in calcium is associated with local hydrated lime particles $\text{Ca}(\text{OH})_2$. These particles surround soil aggregates and calcium can be expected to diffuse into these aggregates. Note

that usual stoichiometries allow the identification of Si/Al ratios close to 1, 2 and 2.35 for kaolinite, illite and montmorillonite, respectively (and the Ca/Si ratio is normally close to 0.07 in montmorillonite).

After a long time curing (1 year) and for any aggregate size, examination of the chemical composition given by EDX on the new formed phase corresponding to C-S-H shows a decrease and an increase of the calcium and silicon contents, respectively, as compared to the initial spot analysis on hydrated lime particles. The Ca/Si ratio measured on C-S-H particles is about 0.65-0.63 (while Si/Al is close to 2.16 or 2.32) and this ratio is in accordance with the composition of synthetic tobermorite $9 \text{ \AA}$ ($\text{Ca}_{0.65}\text{SiO}_2, 2.5\text{H}_2\text{O}$) found by XRD. This is also in agreement with the observations by Brunauer (1962) and El-Hemaly et al. (1977) suggesting a minimum value of 0.8-0.9 for C-S-H gel produced in the mixture of lime-silica-water. Studies of crystal structures of "tobermorites" (Merlino et al. 1999, 2000, 2001) showed that tobermorite is a series between two endmembers, $\text{Ca}_4\text{Si}_6\text{O}_{15}(\text{OH})_2 \cdot 5\text{H}_2\text{O}$ and $\text{Ca}_5\text{Si}_6\text{O}_{17} \cdot 5\text{H}_2\text{O}$, the Ca/Si ratio varying between 0.66 and 0.83. Wild et al. (1986) also stated that hydration of cement at room temperature created a poorly crystallized C-S-H gel with a high Ca/Si ratio of about 1.5 but the similar C-S-H gel from the reaction in the lime-silica-water mixture had a lower Ca/Si ratio.

3.3. Pore size distribution

Results obtained from MIP tests on the lime-treated soils (S5, S1 and S0.4) during a curing time from 7 to 90 days are presented in Figure 7. Data obtained on soils S5 and S0.4 have been presented previously in Wang et al. (2015). The pore size distributions of all lime-treated samples compacted on the dry side of optimum present a bi-modal behaviour with a

population of macro-pores (inter-aggregate pores) and a population of micro-pores (intra-aggregate pores). Note that the delimitation of the pore entrance diameter between the macro-pores and micro-pores is defined as $d = 2 \mu\text{m}$ in this study (Romero et al., 2011). Generally, macro-pores (larger than $2 \mu\text{m}$) are greatly impacted during compaction by the aggregate size: the larger the aggregate size is, the larger the modal entrance size of the macro-pores appears while aggregate size has negligible effect on the micro-pores (smaller than $2 \mu\text{m}$) (Wang et al., 2015). As an illustration, the modal size of macro-pores for the untreated sample S5 is around $73 \mu\text{m}$ and in comparison, lower value is obtained for the untreated S0.4 soil with a main pore family around $14 \mu\text{m}$. Nevertheless, these untreated samples with different D_{max} values share a similar modal size of micro-pores. The mean value of the pore entrance diameter of the micro-pore family remains around $0.3\text{-}0.4 \mu\text{m}$, as seen in Figure 7.

Lime treatment progressively impacts the microstructure of soil. The quantity of each pore population was given as the ratio of intruded void ratio to the total void ratio equal to $e = 0.68$ (Wang et al., 2016), as illustrated in Table 4. For treated soil S5, an increase in the frequency of the entrance pore diameters between 0.006 and $0.1 \mu\text{m}$ (defined as “nano-pores” in the following text for clarity) can be observed in Figure 7a at a curing time of $t = 90$ days, compared with that after 7-day curing. Particularly, the nano-porosity of sample S5 after 90-day curing is 27 % larger than that of the sample after 7-day curing. Less change took place for macro-pores, except a slight decrease in the modal size of macro-pores. The total intrusion value measured on S5 in Figure 7b seems quite similar for the two curing periods. As illustrated in Figure 7c, similar results are observed on treated soil S1: after 90-day curing,

a small increase (around 14 %) in the frequency of nano-pores ranging from 0.006 to 0.1 μm is detected and the cumulative intrusion curve of S1 at 90 days is similar to that at 7 days. In the case of sample S0.4, a slight increase of 16 % in the quantity of the nano-pores is also detected for the sample after 90-day curing, in comparison with the sample at 7-day curing. In addition, both modal sizes of macro-pores and micro-pores shift slightly to lower values, by keeping similar shape during curing and decreasing both the quantities of macro-pores and micro-pores, as seen in Figure 7e. It is worth noting that the total intrusion value of S0.4 at 90 days decreases significantly compared with that at 7 days, induced by a reduction of both the modal pore entrance size and the quantity of the large macro-pores (in Figure 7f).

4. Discussion

Pozzolanic reactions occur with 2 % lime added to the soil from Héricourt that contains quartz, feldspar and clays, and these reactions are strongly time-dependent. New crystallized phase appeared after a long time of curing (at least 1 year) and the main reflection characterizing this new phase is positioned at 34.2° (2 theta $\text{CoK}\alpha$) (corresponding to $d = 3.04 \text{ \AA}$), close to the main peak of calcite that is present in untreated soil. Comparison of XRD patterns of treated soil S1 after 1-year curing and after oven-drying (at 105°C) or after freeze-drying demonstrated that the new peak at $d = 3.04 \text{ \AA}$ corresponds to the crystallized C-S-H phase (probably at tobermorite $9 \text{ \AA}$) (see Figure 2). The chemical analysis on the C-S-H phase by means of Env. SEM-EDX, gave a Ca/Si ratio equal to 0.63-0.65 for both S1 and S0.4, in agreement with the tobermorite $9 \text{ \AA}$ composition. No significant reflection corresponding to C-S-H was detected at a curing time of $t = 60$ days. This can be explained by the absence of well crystallized C-S-H or by the low detection limit of XRD technique.

309 Indeed, it is difficult to detect any crystalline phase if it is poorly crystallized, amorphous, or
310 when its quantity is too small, generally below 0.5-1 % (Carter et al., 1987; Mitchell, 1993;
311 Moon et al., 2004). In other words, very low quantity of well-crystallized C-S-H phase or
312 amorphous C-S-H gel may be formed at 60 days, whatever the aggregate size.

313 However, at a long curing time ($t = 1$ year), the aggregate size seems to impact significantly
314 on the pozzolanic reactions among the lime-soil-water system. C-S-H phase can be detected
315 on the treated soil prepared with large aggregates, such as S5 and S1 while crystallised C-S-H
316 are difficult to be observed by XRD technique on soil with small aggregates (as seen in S0.4)
317 even though Env. SEM pictures revealed the presence of isolated C-S-H particles in big
318 clusters as for S0.4. Note that the low frequency of such clusters, or the low degree
319 crystallisation of C-S-H phase in S0.4, explains why they were not detected by XRD.

320 Furthermore, when lime powder was mixed with small-sized soil aggregates such as S0.4, a
321 thin layer of lime-aggregate (named transition zone by Ping et al. (1991)) was expected due to
322 the large contact surface of small aggregates. On the contrary, lime powders can be locally
323 concentrated in thick layer on the surface of large aggregates (such as those in S1 and S5)
324 which induce small contact surface area. This is consistent with the experimental results of
325 Ping et al. (1991), who studied the aggregate size effect on the transition zone between
326 granular aggregate and cement paste. The authors reported that the thickness of transition
327 zone decreases with the decrease of the aggregate size. As curing time increased, this area
328 where lime is concentrated is in favour of the formation of larger well-crystallized C-S-H
329 particles. Conversely, a more homogeneous lime distribution in S0.4 should induce either the
330 creation of a poorly-crystallized or amorphous C-S-H phase or gel, or a nanometric

well-crystallized C-S-H particle which is hardly detected by XRD. Even though in S0.4, these C-S-H phases cannot be directly detected by XRD and only localised C-S-H phases were observed by Env. SEM. The appearance of these C-S-H phases was indirectly confirmed by Tang et al. (2011) who conducted bender element tests on lime-treated soils with different aggregate sizes. They reported a higher stiffness for the lime-treated soil with $D_{max} = 0.4$ mm at a long curing, compared to that with $D_{max} = 5$ mm. Comparison with the untreated sample showed that the treated samples had a higher stiffness, whatever the aggregate size. Cementitious compounds bond the soil particles together, inducing the increase of the soil stiffness.

Furthermore, the aggregate size effect on the microstructure of treated soil is also consistent with previous mineralogical analysis. A relatively larger increase in frequency was observed for the nano-pores ranging from 0.006 to 0.1 μm on treated soil with large aggregates, especially for S5 after long curing time. These nano-pores can be attributed to the formation of well-developed C-S-H phase (a hydrated lamellar structure). This formation was favoured by the thick coating of lime around soil aggregates or by the local lime concentration in clusters as observed by Env. SEM. Similar observations were made by Russo and Modoni (2013) who reported that a pore population of 0.007-0.2 μm was developed in a lime-stabilised soil at long term. Alvarez et al. (2013) found a small amount of pores below 0.02 μm , ascribed to C-S-H gel pores. In this study, the formation of C-S-H lead to the creation of nano-pores ranging from 0.006 to 0.1 μm , as observed in the mixture of lime and soils with large aggregates (S5). Such new nano-porous compounds filled the micro-pores (0.1 to 2 μm), decreasing the frequency of these micro-pores. However, less quantity of

nano-pores was found in treated soil S0.4 at the same curing period. While a large reduction in the total intrusion value of S0.4 at 90 days was identified, indicating that more undetectable nano-pores were created (their sizes are lower than 0.006 μm that corresponds to the detection limit of MIP measurement in this study), since the initial void ratio of the sample remains identical (Wang et al., 2016).. These undetectable nano-pores are mainly associated with C-S-H phases, which were generated from the pozzolanic reactions in S0.4 sample. This also agrees with the hypothesis of the formation of nano-crystalline C-S-H or small-sized amorphous C-S-H gels, which is invisible by XRD technique and whose nano-pores are undetectable by MIP measurement. Furthermore, these C-S-H phases in S0.4 soil gradually cover the surface of small soil particles to a large extent, bond the adjacent soil particles together, and also gradually block some entrances of both macro-pores and micro-pores. This may explain the small shift of the modal size of both macro-pore and micro-pore populations. As the new phase filled the pores and blocked the pore entrances, initially un-constricted pores can become constricted pores, leading to a decrease in the total intrusion value (Wang et al., 2015). This is in agreement with the observation by Russo and Modoni (2013).

5. Conclusion

The changes in mineralogical composition and microstructure of a lime-treated compacted soil during curing were investigated, with emphasis on the effect of aggregate size. On the basis of the experimental results, some conclusions can be drawn, as follows:

- 1) XRD peaks of calcium silicate hydrates (C-S-H) on crystallized tobermorite form were

observed on patterns of freeze-dried lime-treated soil sample with large aggregates (S5 and S1) after 1-year curing, while no such C-S-H was observed at shorter curing times. Oven-drying at 105°C allowed confirming the mineralogical nature of the new formed phase as C-S-H.

2) When a given quantity of lime is mixed with large aggregates (S1 and S5),, lime particles during mixing can be concentrated locally within the mixture in order to form lime aggregates or thick coating on soil aggregates. Such calcium-rich areas contribute to the formation of large and detectable C-S-H crystallites as observed by XRD. By contrast, a diffused distribution of the lime powder was expected in the small soil aggregate-lime mixture (S0.4). The low rate of lime aggregation benefits to the development of small (nano) crystallized C-S-H particles, or poor crystallized C-S-H phases, or amorphous C-S-H gels..

3) The decrease of population of micro-pores and the increase of nano-pores in frequency observed in large aggregates lime mixtures (S5 and S1) can be attributed to the formation of visible crystallized C-S-H pores: the new nano-porous compound filled the micro-pores. By contrast, the soil with small-sized aggregates S0.4 mixed with lime behaved differently: very small or nano-sized C-S-H phase homogeneously spread on large soil aggregate surface was expected around soil aggregates (because of the initial thin coating with lime). As a result, the nano-sized pores expected to be less than 0.006 μm in the nano-sized C-S-H particles were too small to be detected by MIP.

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537

538 **Table 1 ESEM-EDAX element analysis of S1 at a curing time of $t = 1$ day, both on the spots where lime**
 539 **and clay particle located**

Element	Wt %		At %		K-Ratio	
	Lime	Clay	Lime	Clay	Lime	Clay
C	6.37	4.28	10.85	7.23	0.0171	0.0068
O	50.54	49.47	64.67	62.74	0.0959	0.1652
Na	0.14	0.21	0.12	0.19	0.0004	0.0007
Mg	0.77	0.73	0.65	0.61	0.0035	0.0036
Al	3.65	10.18	2.77	7.65	0.021	0.0624
Si	7.71	22.88	5.62	16.53	0.0529	0.1496
K	0.39	1.76	0.2	0.91	0.0037	0.015
Ca	27.38	2.06	13.99	1.04	0.2574	0.0185
Ti	0.28	0.59	0.12	0.25	0.0022	0.0051
Fe	2.78	7.83	1.02	2.85	0.0236	0.0683

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543 **Table 2 ESEM-EDAX element analysis of S1 at a curing time of $t = 1$ year, on the spot where CSH located**

Element	Wt %	At %	K-Ratio
C	3.33	5.61	0.0064
O	52.49	66.42	0.1295
Na	0	0	0
Mg	2.32	1.93	0.0113
Al	6.83	5.12	0.0407
Si	16.54	11.92	0.1112
K	0.41	0.21	0.0037
Ca	15.55	7.85	0.1421
Ti	0.17	0.07	0.0014
Fe	2.37	0.86	0.0203

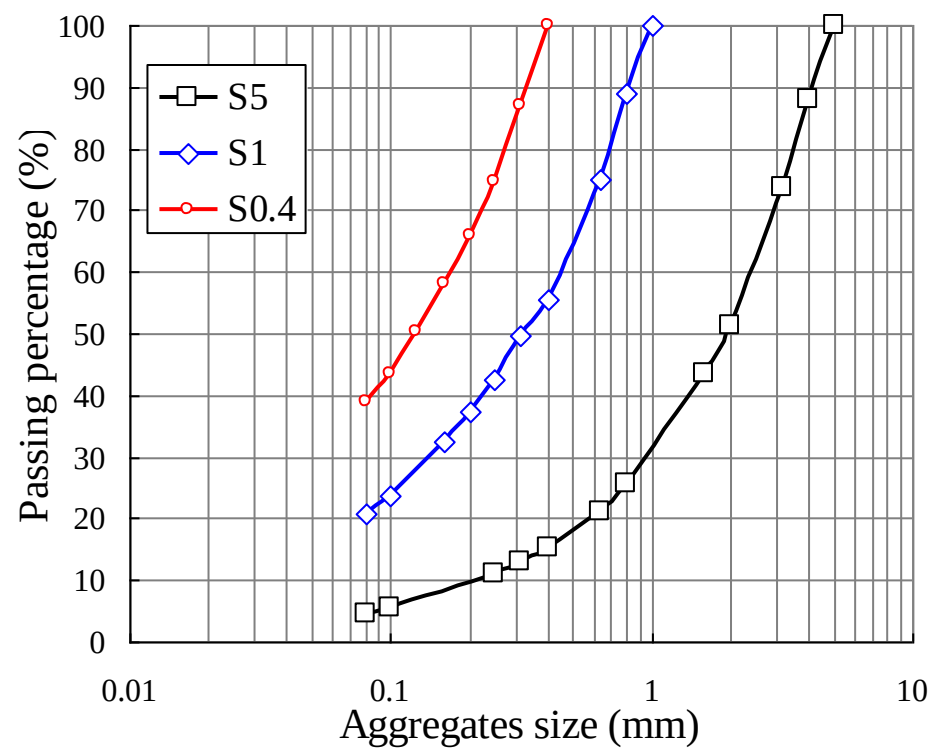
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Table 3 ESEM-EDAX element analysis of S0.4 at a curing time of $t = 1$ year, on the spot where CSH located

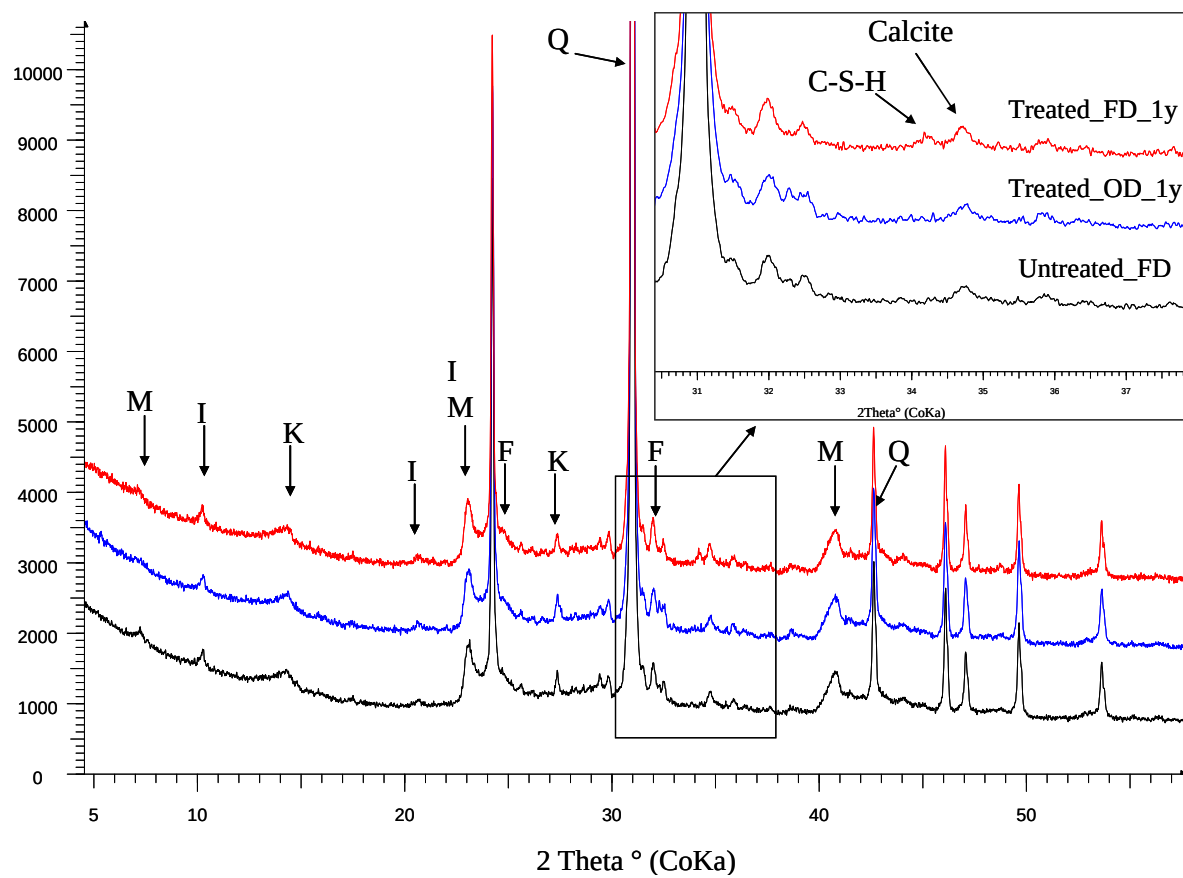
Element	Wt %	At %	K-Ratio
C	7.1	11.32	0.0147
O	55.15	66.01	0.1435
Mg	1.89	1.49	0.009
Al	6.3	4.47	0.0372
Si	14.2	9.68	0.0953
K	0.25	0.12	0.0022
Ca	12.78	6.11	0.1171
Fe	2.34	0.8	0.0201

Figure 1 Aggregate size distribution of soil S5, S1 and S0.4



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555 **Figure 2 X-ray diffraction patterns of the untreated soil S1**



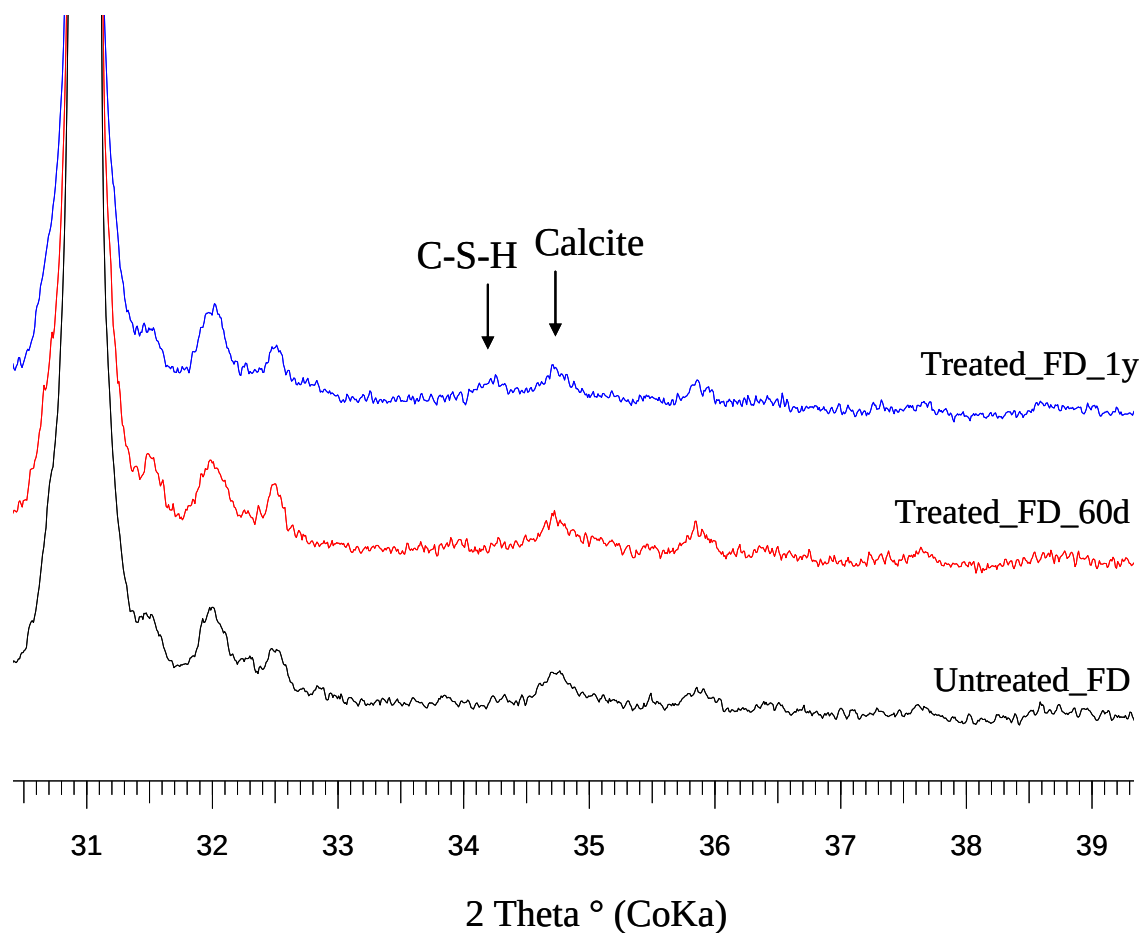
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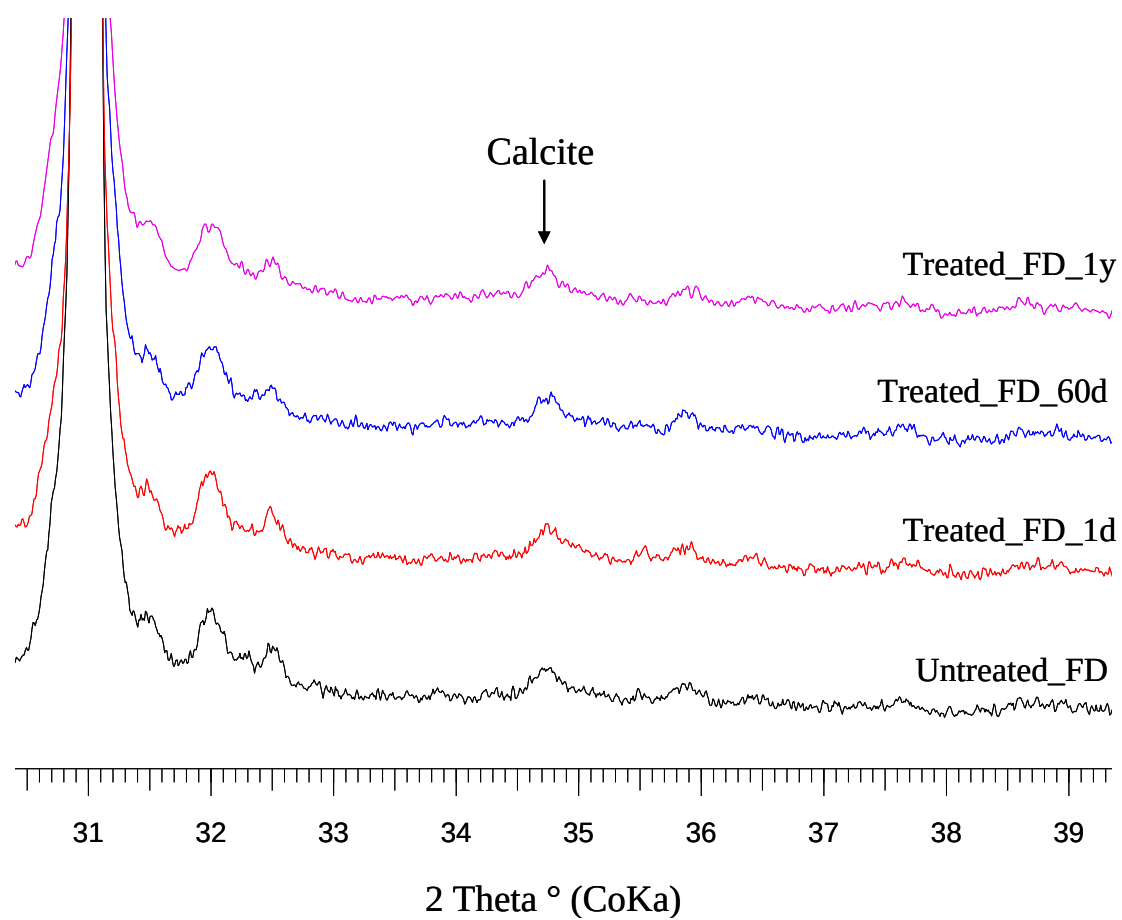
559 **Figure 3 X-ray diffraction patterns of lime-treated soils during curing: (a) S5; (b) S0.4**

(a)



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(b)



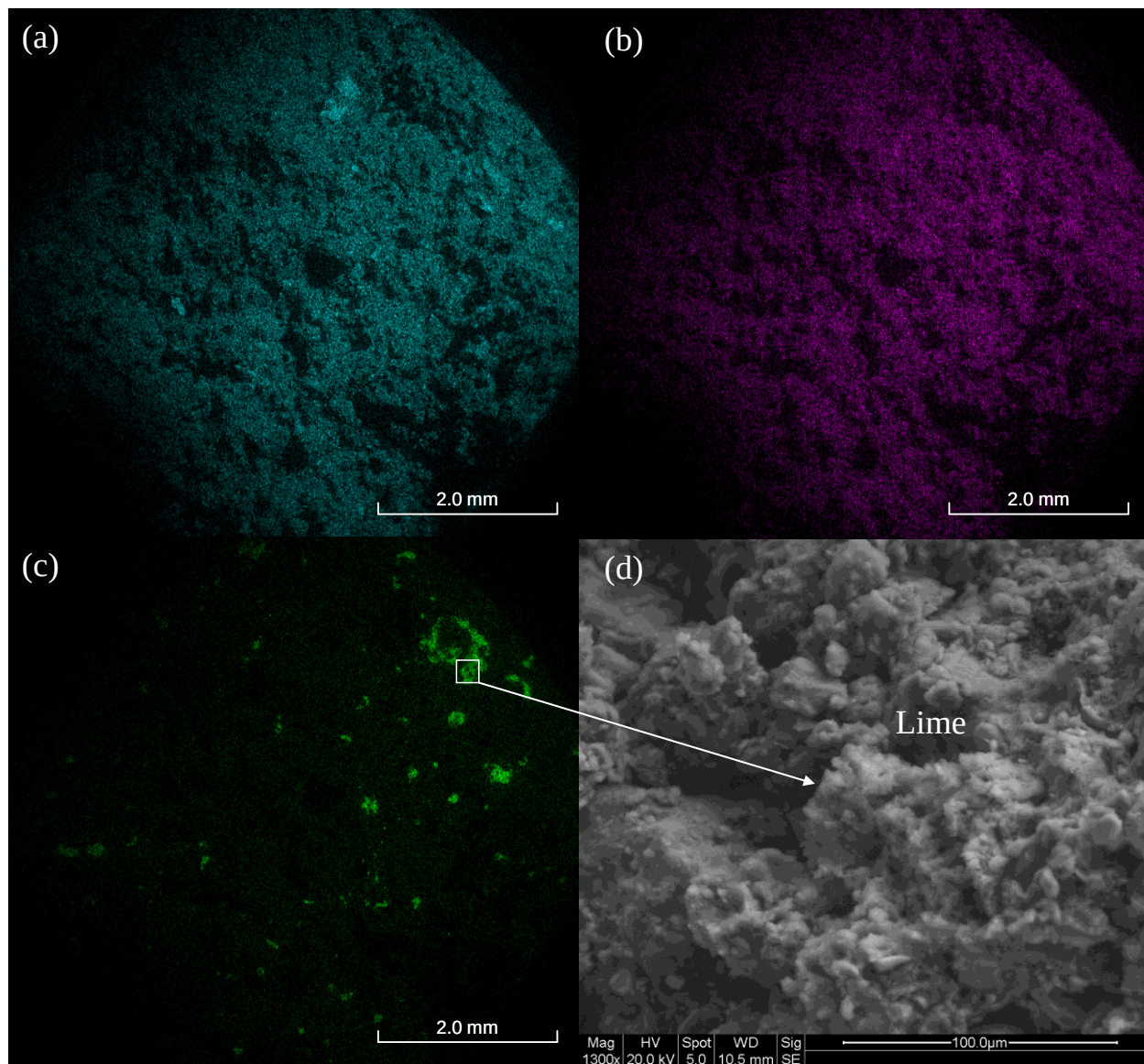
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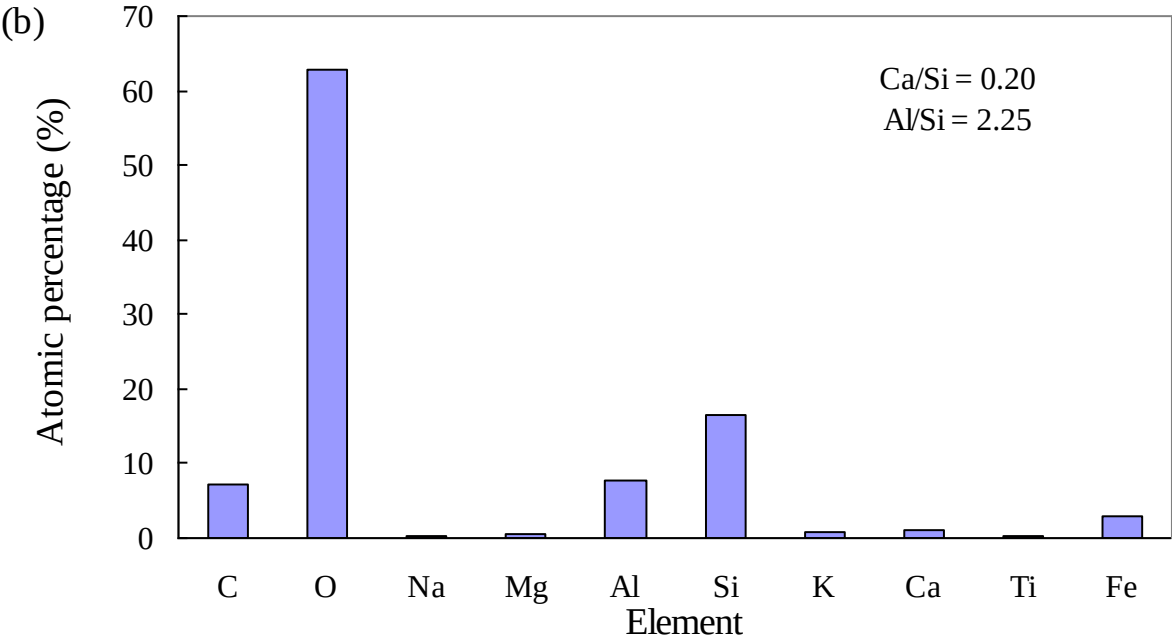
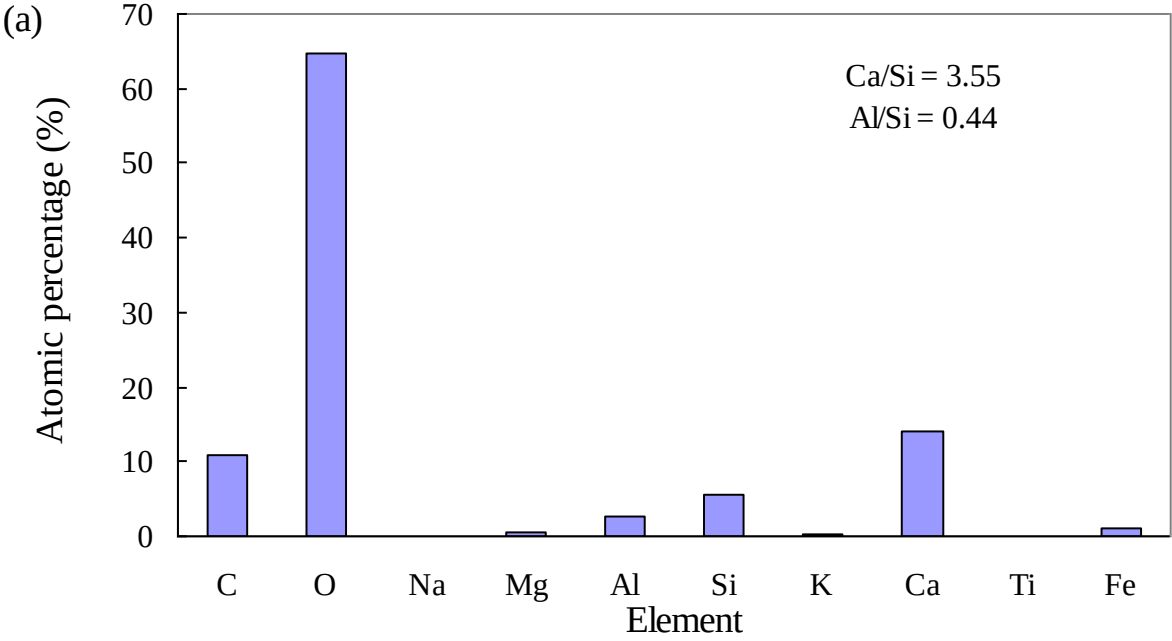
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565 **Figure 4 Elementary distribution maps of a lime-treated soil, S1, at a curing time of $t = 1$ day: (a) Silicium;**
566 **(b) Aluminium; (c) Calcium, and SEM picture of the spot in the calcium concentration area**



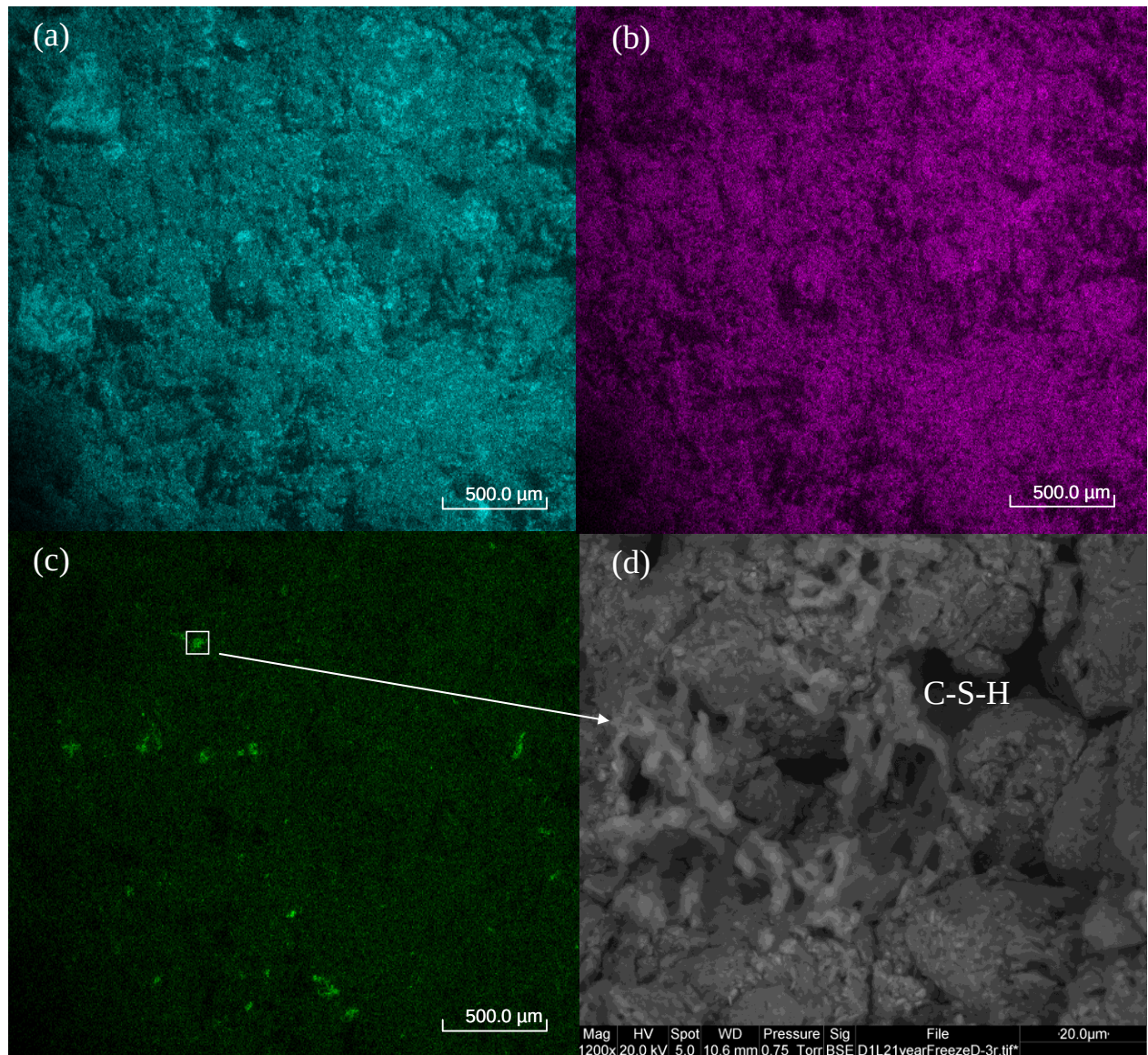
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Figure 5 Element examination by EDAX on lime-treated soil S1 at a curing time of $t = 1$ day: (a) on the spot in the calcium concentration area; (b) on the spot in the clay particle area



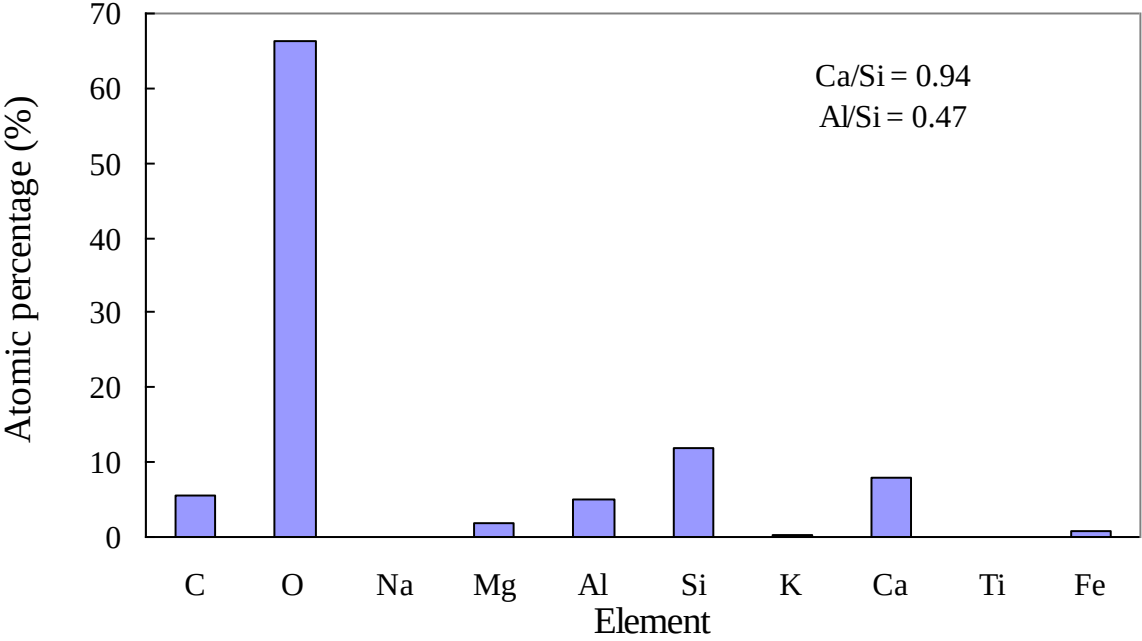
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575 **Figure 6 Elementary distribution maps of a lime-treated soil, S1, at a curing time of $t = 1$ year: (a) Silicium;**
576 **(b) Aluminium; (c) Calcium, and SEM picture of the spot in the calcium concentration area**



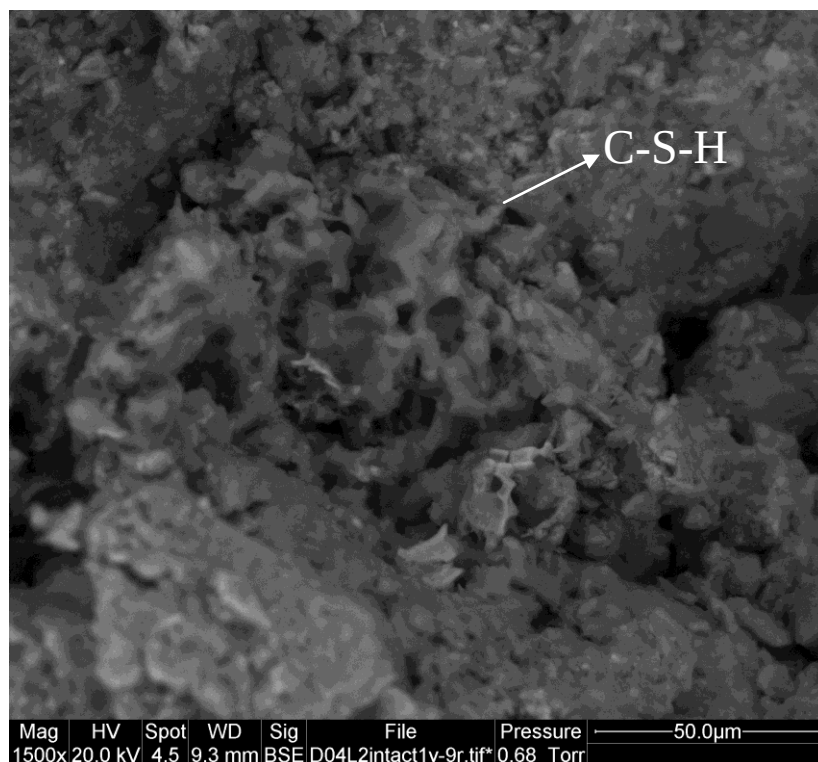
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Figure 7 Element examination by EDAX on lime-treated soil S1 at a curing time of $t = 1$ year, on the spot in the calcium concentration area



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584 **Figure 8 SEM picture of the lime-treated soil, S04, at a curing time of $t = 1$ year**

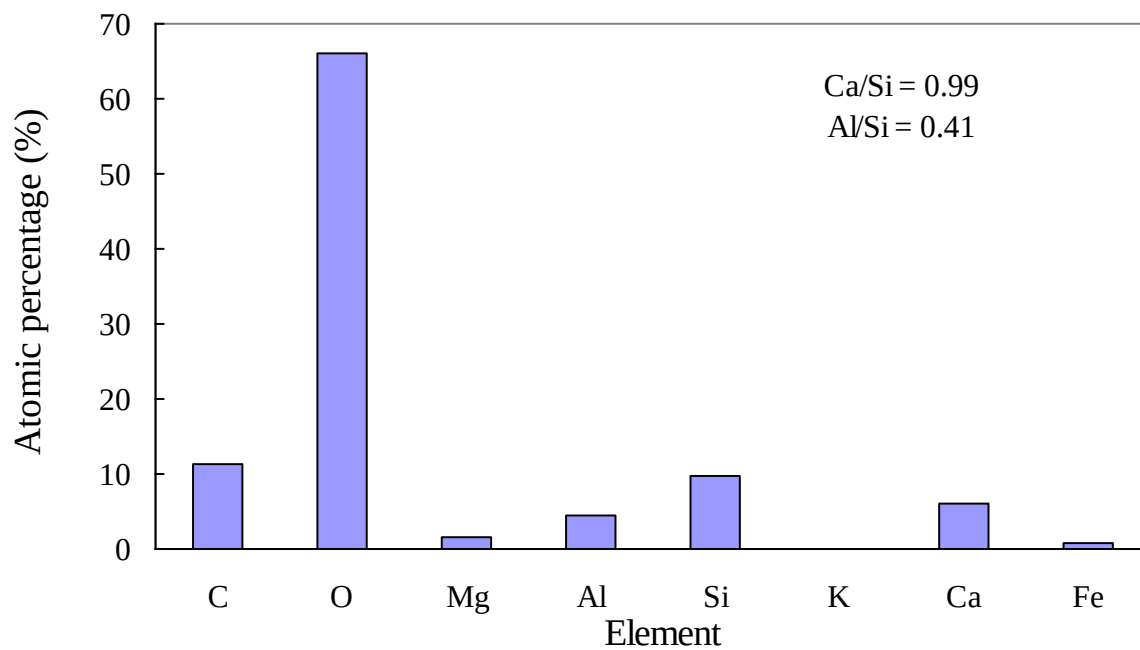


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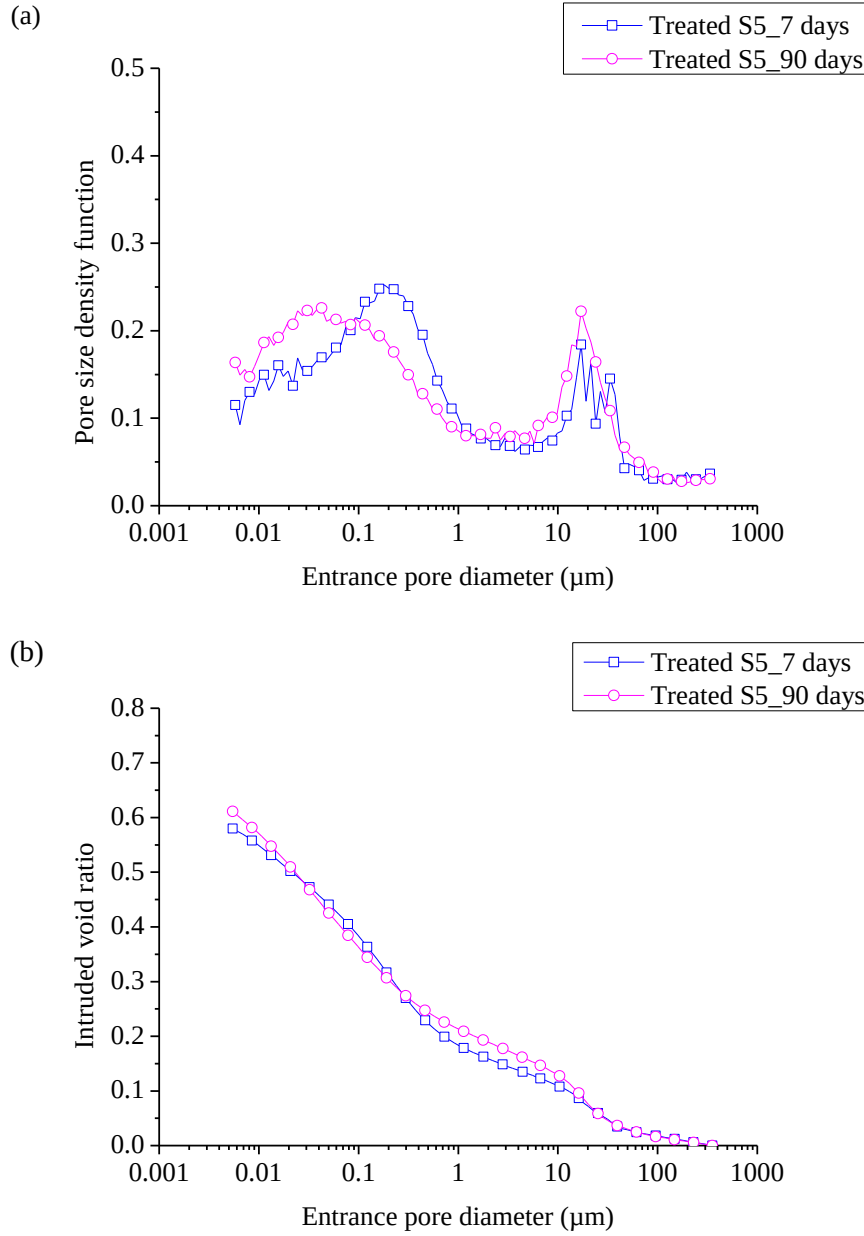
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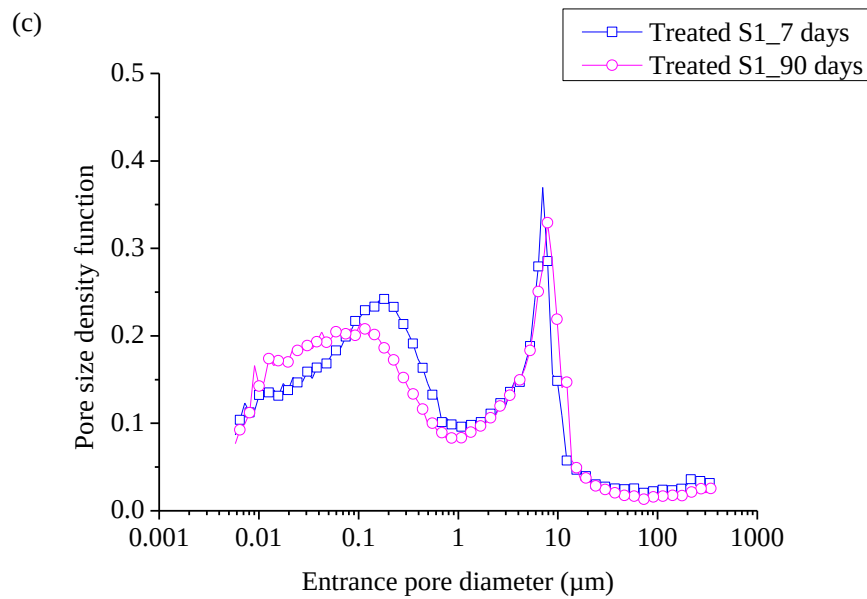
588 **Figure 9** Element examination by EDAX on lime-treated soil S0.4 at a curing time of $t = 1$ year, on the spot
589 **in the calcium concentration area**



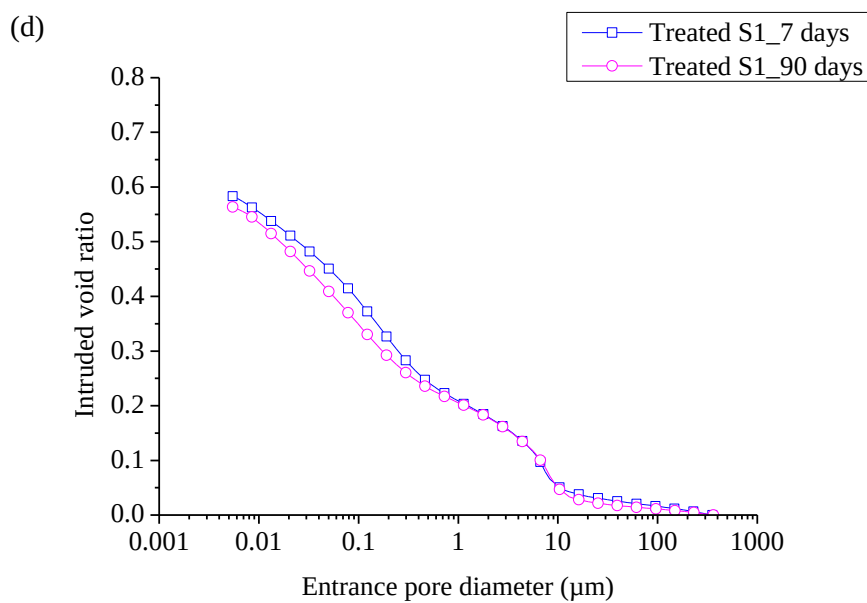
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Figure 10 Pore size distributions and cumulative intrusion curves of lime-treated soils with different aggregate sizes (S5, S1 and S0.4), obtained by MIP





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