

# Comprehensive Methodology for Soil Stabilisation

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ISSN: 2576-8840



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**Submission:**  November 20, 2025

**Published:**  December 19, 2025

Volume 22 - Issue 4

**How to cite this article:** Gabriel Barbetat. Comprehensive Methodology for Soil Stabilisation. Res Dev Material Sci. 22(4). RDMS. 0010401 2025.  
DOI: [10.31031/RDMS.2025.22.001041](https://doi.org/10.31031/RDMS.2025.22.001041)

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## Abstract

A simple, complete and comprehensive method is proposed to build with soil in an effective and sustainable manner, to ensure the architectural benefits and durability sought, consistent with current standards of quality and control. This work stems directly from a doctoral thesis and vast experience in construction and research over thirty years, and from the latest results obtained from chemical stabilisations using enzymes, carob, Chelidonium majus and various limes. The method consists of applying the most efficient and optimal stabilization to each type of clay present, based on its physicochemical characteristics and the moisture content specific to the construction technique used. This involves characterizing the particle size through granulometry and sedimentometry, its crystallographic nature through thermal gravimetric analysis and/or X-ray diffraction, and testing its strength and shrinkage behavior with different stabilizers. Finally, a working table is presented showing the optimal stabilization methods for each type of clay present.

**Keywords:** Soil stabilisation; Clay; Earth construction; Soil cement; Rammed earth

**Abbreviations:** CEB: Compressed Earth Block; T: Tetrahedra Layer; O: Octahedral Layer

## Introduction

When an architect sets out to design and build an earth structure - using local soil or material extracted from excavations and the foundation itself - they often find themselves disoriented and alone in deciding how to proceed. Numerous standards and standardised tests in the fields of geotechnics and construction provide us with data, but we need to know which are essential and how they correlate most effectively. Working methodologies, when consulted, tend to be limited to a specific soil type, technique, or stabilisation system - and they often avoid considering the crystallographic nature of the clays that actually bind the soil particles together.

This article also argues for the necessity of stabilising the soil to achieve the essential architectural performance required of any modern building material. Today, we have sufficient knowledge to construct earthen buildings resistant to flooding and earthquakes. The degree of improvement in water resistance, volumetric shrinkage and/or increased mechanical strength is determined by the chemical nature, particle size distribution and plasticity of the available soil. Reducing shrinkage, porosity and permeability allow for a general increase in mechanical properties and a reduction in sensitivity to water action - swelling and shrinkage, reduced cohesion and stiffness, erosion and frost shattering. Capillary water or rainwater, upon entering the binding structure of clays, breaks the weak van der Waals forces of surface tension in the interlayer water. Subsequently, it breaks the electrovalent bonds and, under certain alkaline or acidic conditions, even the strong covalent bonds. Therefore, it is essential to coat the material or to physically and chemically stabilise its structure to generate covalent and irreversible bonds. The stabiliser causes cation exchange and a transformation of the clay's peripheral bonds. The soil structure changes, and a new structural matrix is physically formed, intermixed with new elements that reinforce it and improve its flexural strength.

## Methodology

### Local popular technology analysis

An analysis of local technologies and resources is necessary to understand existing experience and traditional knowledge regarding earthen materials. This will allow us to develop a type of local action better suited to native resources, the environment and existing sociocultural values - always in line with the global perspective of sustainable development. This focuses primarily on:

- 1) Locating local earthen constructions.
- 2) Visually identifying construction quality.

**Table 1:** Sieved by weight %.

Sieve Size. Ta	Sieved by Weight % *MOPT (Spain) CRATerre-U.Grenoble [10]	Sieved by Weight Bolomey Tendance, Modified by Adding Clays $X = 110 - 100 * (1 - \sqrt{(Ta/5)})$
0.001	14.9	11.41
0.0046	21.1	13.03
0.01	27	14.47
0.08	44	22.65
0.16	52	27.89
0.32	60	35.3
0.63	71	45.5
1.25	82	60
2.5	98	80.71
5	100	100

Analysis Methodology:

1. Particle size analysis - with washing and deflocculation to remove silt and clay adhering to the aggregates.
2. Quantification of crystallographic types of clays present in the soil - via X-ray diffraction and sedimentation (Brockville method).
3. Shrinkage - 5x70cm box method.
4. Stabiliser determination
5. Performance verification - durability (six wetting-drying-frost cracking cycles); compressive and shear strength

### Preliminary field assessment of organoleptic properties

The following aspects will be observed during sample extraction:

- a. Presence of a musty scent, particularly if the odour intensifies when the soil is moistened or heated. Such soil will be discarded.
- b. Type of clay deposit, distinguishing between allochthonous (transported) and autochthonous (formed in place); in

- 3) Detecting the presence of the ceramic industry, which uses clays rich in illite and/or kaolin.

### Analysis

A simple stabilisation method for any type of building soil should be based on knowledge of its clay mineralogical composition and certain laboratory and construction site data. Stabilisation involves modifying the soil's composition physically and chemically to increase its durability and strength. The order of tests presented here is the simplest and most efficient for stabilising soil and obtaining a suitable building material, without overlooking the characteristics of the soil and the available stabilisers (Table 1).

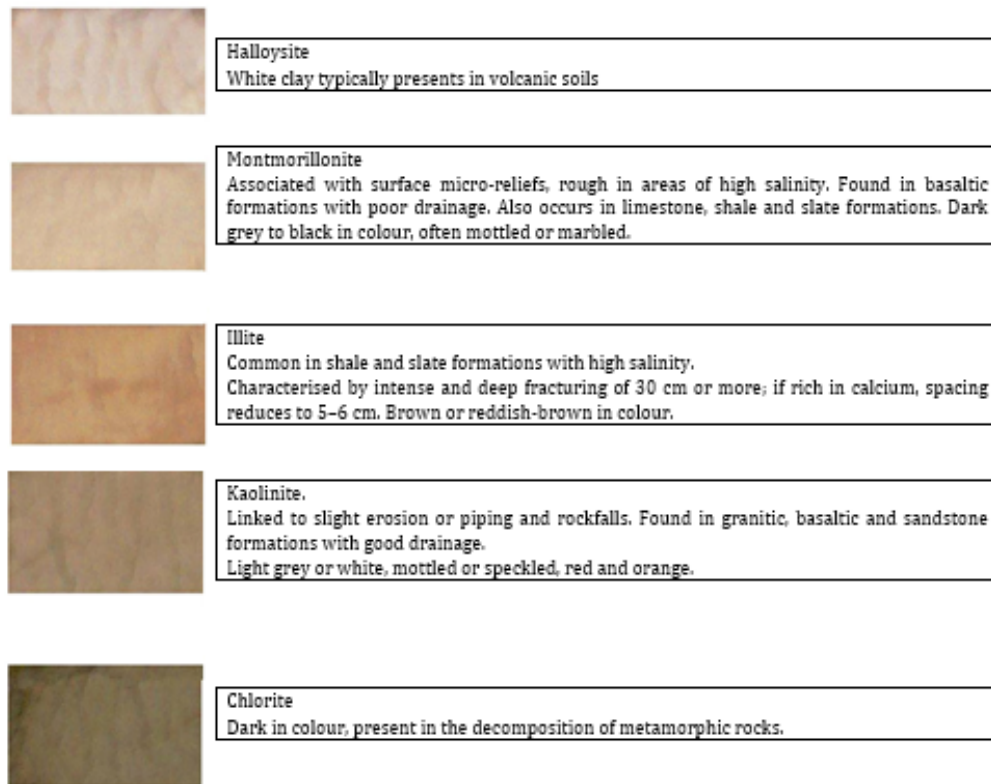
the latter case, and under sufficiently reducing conditions, montmorillonite clays may be present.

c. Geological profile and soil layer depth, including rock types or formations in the region and their erosion, slope stability, potential landslides and microrelief formation. Highly stable, compact and erosion-resistant slopes are favourable for the use of the material in rammed earth and poorly stabilised Compressed Earth Blocks (CEB).

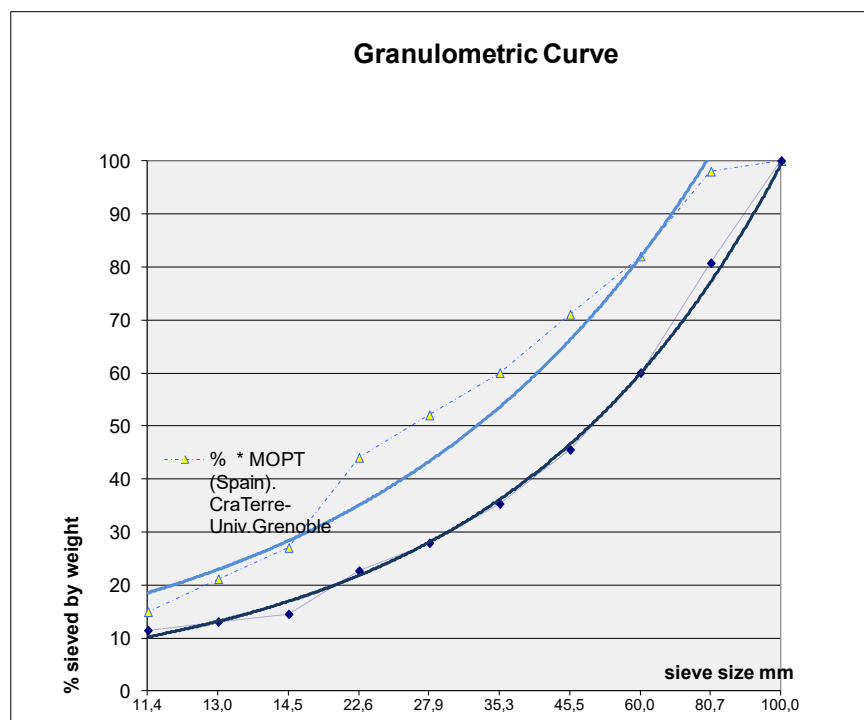
d. Clay behaviour in water, including dispersion, colour, turbidity, waterlogging, and the presence of expansive or fissurable clays in puddles and flood-prone areas (Figure 1 & 2).

e. Texture and consistency, as determined by field tests (5-15cm cylinder test, tongue test and tactile test): sand may or may not be visible but is easily felt by hand. A sample with low moisture content is easily compacted by hand (tap test) and becomes plastic when moisture content increases.

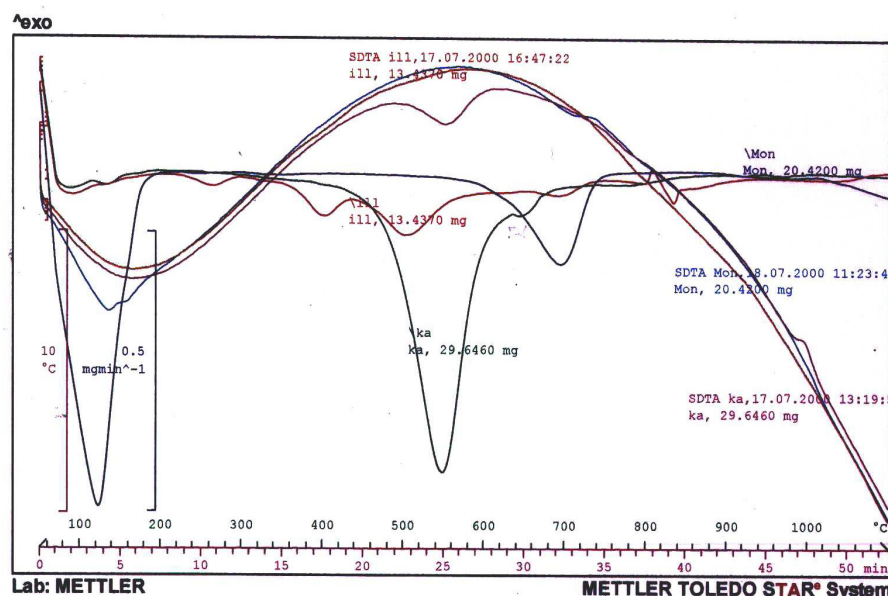
f. Colour of the dried and sieved sample (<88µm), compared with Figure 3, to help identify clay types and detect the presence of iron sulphides, organic matter or oxides. Any specks or inclusions should be identified as carbonates, iron, roots or organic matter.



**Figure 1:** Clay colours.



**Figure 2:** Normalised granulometric curve.



**Figure 3:** ATD-ATG curves and first derivative for the main contrasting clay species: Ka. kaolinites, Mon. montmorillonite, and Ill. Illite.

### Determination and evaluation of the granulometry curve

The determination of particle size distribution curves is carried out through the granulometric [1] and sedimentometric analysis of the various fractions defined by the maximum particle sizes. Thus, for the production of Compressed Earth Blocks (CEB), where the maximum aggregate size is below 10mm, only 500g of sample is required, which greatly facilitates the cleaning and settling of fines. The general granulometric criterion adopted for soil used in construction is that it should contain a clay proportion between 15% and 18%. However, if we increase the granulometric threshold to 5 microns - the fraction where the most strongly flocculated clays are found - we observe values between 21.1% and 24%. Silts appear in almost the same proportion, between 20% and 23%, as shown in the table below. It can also be noted that the limits established by Bolomey's continuous parabola, intended to guarantee a lower pore index, are appropriate for coarse fractions. However, when applied to silts and clays, they diverge significantly from the limits previously adopted.

We compared this with the particle size analysis of several natural soils [2]. The results were highly variable, with the silt-to-clay ratio ranging from 0.0091 and 0.4 up to 20.3. Even in seemingly identical soils, it fluctuated between 0.8 and 20.3. Despite this, 50% of the soils tested showed ratios approaching 1:1. Thus, the ideal silt-to-clay ratio is established as 1:1.

The silty and clayey particle size fraction (<80µm) is separated from the sands by decantation with distilled water, and some of the suspended particles are removed by absorption using a heated sodium-glass capillary tube. The solutions are then placed under an infrared lamp at 60 °C and 60% relative humidity to avoid disturbing the outermost atomic hydration layers, thus allowing for accurate readings by DAT or other physicochemical analyses.

Drying is stopped when a solution with an approximate density of 1.050g/cm<sup>3</sup> is obtained. Sedimentation analysis is then initiated to differentiate the fines using the Brockville method. The 1,000cm<sup>3</sup> solution is stirred manually or mechanically at a rate of 1 to 3 cycles per second, and times, temperatures (20-25 °C is recommended) and readings from a calibrated and corrected sedimentation meter (adjusted for pressure, temperature and deflocculant use) are recorded. The ASTM Soil Hydrometer 152H model is recommended. The best way to take readings with this instrument is from above, noting the value at the top of the meniscus. The initial reading corresponds to the fine sand level; the remainder is silt and clay. Readings are then taken at 1, 2, 3, 4, 8 and 16 minutes (the point at which the clay fraction generally begins), followed by 30 and 60 minutes, 2 hours, 16 hours, 24 hours and finally at 48 hours. The Stokes formula  $D = K\sqrt{L/t}$  is used to calculate the size and amount of sedimented granulometric fraction at each instant [1-3].

### Characterisation of the basic clays present in the sediment fraction

The main clay mineral species present in the soil are:

-Kaolinite (Nacrite, Dickite, Halloysite). Size ≈ 1µm. Consists of 100 layers 720µm thick, held together by hydrogen bonds between OH- groups of the Gibbsite layers and the oxygen atoms of the adjacent silicon layers (T+O), which do not dissociate upon contact with water. Cation exchange capacity: 2.2-15meq. It is non-expansive, has low plasticity and cohesion, and is difficult to disperse - even under acid attack. Thixotropic limit: 70-95g water/100g. Its light brown colour indicates high alumina content, low iron content, and 1% gypsum. Water content is 0.5% in a 5° layer, with a specific surface area of 10cm<sup>2</sup>/g. Kaolinite is a product of the weathering of orthoclase feldspar, derived from granite, and is typically found in sedimentary environments.

-Illite. . Size  $\approx 0.1\mu\text{m}$ . Composition is similar to mica - sextuple layers (T+O+T, T+O+T) - and to xanthophyllite, but with lower potassium content (3-7%), higher combined water (5%) and reduced aluminium substitution (9-32%) by silica in the tetrahedral layer (38-53%). This is attributed to the low potassium content and partial disruption of the  $\text{K}^+$  alkali bond, part of which is replaced by  $\text{Ca}^{2+}$  and  $\text{Mg}^{2+}$ . As a result, illite exhibits a high exchange capacity compared to micas (10-40meq) and a high silica index. It is expansive, with medium plasticity and low permeability. If sodic, it dissolves rapidly in water.

Thixotropic limit: 110-170g water/100g. Effervescence under acid attack indicates the presence of calcium or magnesium illite. Its colour ranges from creamy white, beige to earthy greenish-grey, with white streaking and an unctuous texture. Illite is typically associated with montmorillonite or kaolinite. It reacts slowly with lime and is completely inhibited in the presence of .

-Montmorillonite (nontronite; Fe-rich; Mg saponites with low Al; hectorites with Li; beidellite: low Si, Al-rich). Size  $\approx 0.01\mu\text{m}$ . Triple-layer system O+T+O. Its structure is similar to Brucite and Gibbsite, but in the layers adjacent to these units there are only silica sheets, which lack external hydroxyl bonds. As a result, the units are linked by van der Waals forces. Montmorillonite is highly expansive, with high plasticity, sticky and soft texture, and very low permeability. It has a high cation exchange capacity (75-150meq, though limited with Mg and Al, as these elements are already part of

the structure. According to the Emerson test, it shows the greatest dispersion in water. Thixotropic limit: 700-1350g water/100g. Effervescence under acid attack indicates calcic or magnesian montmorillonite. Its colour is greenish white, becoming much whiter when pulverised. Water content reaches 50% in a 5° layer, with a specific surface area of 1,000  $\text{cm}^2/\text{g}$ .

-Chlorite:  $\text{Si}_3\text{O}_{10}\cdot\text{Mg}_5\text{Al}_2(\text{OH})_8$ . Size  $\approx 0.1\mu\text{m}$ . Laminated structure of six layers, comprising two triple layers (T+O+T) linked by an alkali ionic bridge via the  $\text{K}^+$  cation. This configuration combines the charged basic unit of mica (talc) with the charged unit of Brucite. Possible substitutions of Al for Si in the tetrahedral layer, and  $\text{Fe}^{2+}$  or  $\text{Fe}^{3+}$  for Al in the octahedral layer. Chlorite exhibits low expansiveness, low shear strength and elasticity, and is difficult to disperse in water, even under acid attack. Its colour is bluish grey. Cation exchange capacity: 10-40meq. It commonly associates with Illites.

### Clay content determination by ATD/ATG thermogravimetry or XRD diffraction

Thermogravimetric analysis (TGA) records the mass changes of a given material - whether in solid, amorphous or crystalline phase - as a function of temperature, relative to an inert reference. These results are summarised in Table 2, which highlights the thermal peaks observed, quantifies the corresponding mass losses, and links them to specific clay minerals or binding agents.

**Table 2:** Summary of endothermic peaks associated with elements identified in soil analyses.

Gypsum	Phases SCH	Ettringite	Halloysite	$\text{Ca}(\text{OH})_2$	Quartz	Magnesite	Serpentine
130 °C	150 °C	180 °C	240-580 °C	460 °C	573 °C	600 °C	Strong endo-exo 700-800 °C
				Portlandite		45% MgO	
$\text{CaCO}_3$	$\text{CaCO}_3$	Dolomite	Tridymite quartz		Cristobalite quartz	$\text{FeO} - \text{Fe}_2\text{O}_3$	Talc
750 °C	820-850 °C	720 °C+-800 °C	117 °C, 160 °C, 870 °C		900 °C	800-900 °C	1000 °C
Amorphous Precipitate	Calcite	30 % CaO + 21 % MgO					
Vaterite	Vaterite						
Kaolinite		Montmorillonite			Beidellite	Saponite	
590 °C (<680 °C Dickite-Nacrite), endo-exo 980 °C		700 °C (or 600-610-620-660 °C)			550 °C; endo-exo 700-800 °C	179 °C (double peak), 240 °C, 289 °C, 634 °C, 734 °C, 904 °C	
Exo 950-1050 °C mullite ceramisation		Endo-exo at 900 °C if $\text{Mg}^{2+}$ present (var. 840-870-880 °C); Na/Ca.Mg				Observe $\text{Mg}^{2+}$ content	
Hectorite		Illite			Chlorite	Vermiculite	
180 °C, 680 °C, 780 °C		180-500 °C (520-480 °C or 510 °C)			3 peaks between 750-850 °C	2 double peaks	
strong exo 900 °C		. + endo-exo 900 °C (890-930 °C / 850-910 °C)			+ exo 850 °C	200-250 °C and 850-900 °C	
		Na/Ca.Mg					

For the mineralogical analysis of the clays, a Bruker D8 ADVANCE X-ray diffractometer with Advanced Geometry was also

used, operating in Bragg-Brentano reflection mode with a  $\theta$ -2 $\theta$  configuration and  $\text{CuK}\alpha$  radiation ( $\lambda = 1.5406\text{\AA}$ ) (Figure 4).





**Figure 4:** Bruker D8 ADVANCE X-ray diffractometer. Diffraction cell showing the X-ray monochromator (left) and the detector collecting the sample's reflection (right).

The instrument operates across a range of diffraction angles ( $2\theta$ ) to capture reflections from multiple crystallographic planes, as minerals - particularly clays - exhibit several diffraction peaks of varying intensity. The typical basal spacings (d-values) for the three most common clay minerals are:

- A. Kaolinite 7Å (semiquantitative calculation coefficient 0.14)
- B. Illite 10Å (0.16)
- C. Smectite 10-11Å (0.18)

In the case of kaolinite (always prepared as an oriented aggregate on glass) it is treated with  $Mg^{2+}$  (generally a 1M  $MgCl_2$  solution, where  $Mg^{2+}$ , with a high hydration radius, tends to expand the lamellar layer) and the clay maintains a value of 7Å. The kaolinite is then treated with glycerol or ethylene glycol (which is chemically similar to monoethyl ether, an organic molecule with a large surface area that should force the clay to expand), but the kaolinite still maintains a value of 7Å. Subsequently, the kaolinite is treated with  $K^+$  (a 1M KCl solution, which should force the closure of the clay - in case it had expanded - since potassium has a low ionic potential and can be easily adsorbed by the clay) but the kaolinite again maintains a value of 7Å. The final treatment involves placing the clay sample (oriented aggregate) in an oven at 550 °C. In this case, the clay degrades, and when analysed by X-ray diffraction, it shows no diffraction signal. In this way, we have identified this clay as kaolinite.

The same occurs with illite, which maintains a constant value of 10Å throughout the treatments. With smectite, identification is more complicated because it initially shows a d-value of 10-11Å

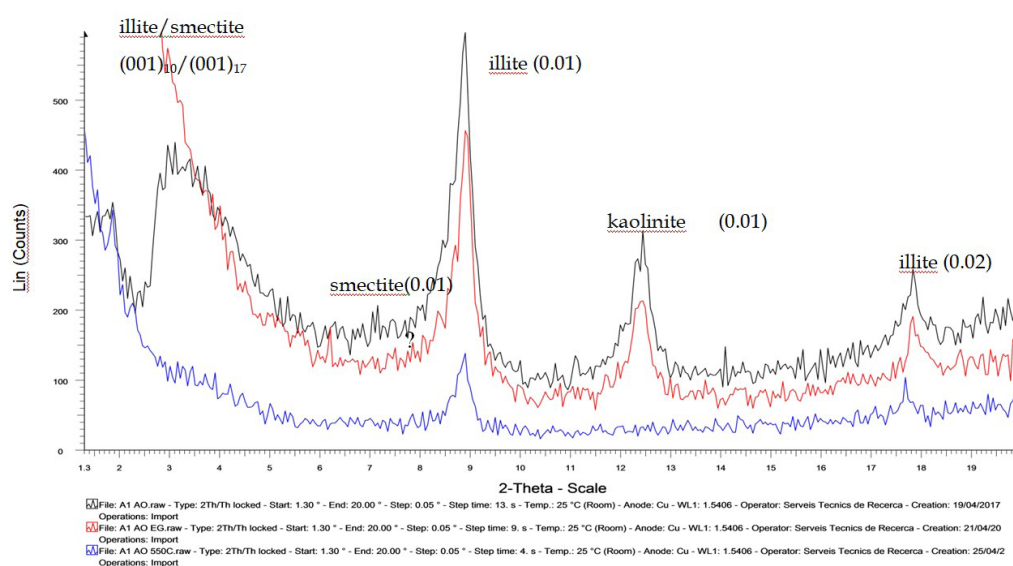
under normal conditions. After treatment with  $Mg^{2+}$ , the d-value increases to 14 - 16Å; with glycerol it increases to 18Å; and when treated with  $K^+$ , it closes to 12Å due to the action of this cation. Finally, after heating at 550 °C, the clay further closes at 10Å, as all hydration water has been lost and diffraction only detects the tetrahedral (2 x 2.4), octahedral (1 x 2.4) and lamellar (1 x 2) layers.

Identifying other types of clay can be even more complicated, as mixed or overlapping clay phases may require density separation treatments. When the clay phases show thin, clean and stylised peaks, this indicates high crystallinity.

#### **Physical analyses. Evaluation of volumetric stability using the shrinkage test**

Using a sample of clay with standardised plasticity, or with the moisture content specific to each technique (drop ball test for CEB-rammed earth) [4], a cubic test mould - preferably plastic or finely milled metal, 50cm long by 5cm wide and high - is filled. The mould is previously greased with oil. The sample is left to dry under laboratory conditions for seven days. After this period, drying is completed in an oven at 60 °C for 24 hours. The dry shrinkage value is determined by measuring the cracks in the prismatic soil sample and the separations from the mould. From this, two comparative values are obtained: the linear shrinkage, expressed as a percentage of shortening relative to length, and the type of cracking, evaluated by the number of cracks in the sample and the distance between them.

The presence of significant cracks will indicate (Figure 5), a priori, the need to repeat the test with already stabilised soil and verify compliance with the following linear shrinkage limits:



**Figure 5:** XRD diagram with the most representative profiles of the clays. Treatments: normal (black line), treated with ethylene glycol monoethyl ether (glycolate, red line), and heated to 550 °C (blue line).

Linear shrinkage of the stabilised rammed earth: <1.0mm/m

Linear shrinkage of the CEB wall: <3% (1)

### Optimal stabiliser

Stabilisation is defined as a physical, physicochemical or chemical process that enables soil to respond satisfactorily and permanently to the demands imposed for its use in construction. Maintaining earth-rendered walls without protective covering is extremely difficult and costly.

This process - aimed at improving water resistance, reducing volumetric shrinkage and increasing mechanical strength - is determined by:

- A. The chemical nature of the base soil, including its particle size distribution and plasticity.
- B. The desired improvements: with respect to the three points above, although the authors of CRA Terre refer more to reducing porosity - that is, the void volumes between solid particles - than to shrinkage [5-13]:
- C. The available products, materials or technological processes.
- D. The economic and environmental impact assessment.
- E. Maintenance and usage conditions. Reducing shrinkage or porosity, permeability and increasing mechanical resistance allows for an overall improvement in mechanical characteristics and a reduction in sensitivity to the action of water: swelling and shrinkage, reduction of cohesion and rigidity qualities, erosion and gelation. The method described here consists essentially of selecting a stabiliser based on local resources, the technique to be used and the characterisation of the soil and

clay types of present. In addition, chemical analysis using Dr Ferran Gomà's method [14], which assesses soluble fractions, cation exchange capacity and pH, provides further identifying data about the material. Activating as much soluble silica as possible - whether from the soil itself or externally through additives - is considered fundamental to the stabilisation process. The increase in soluble silica enables the formation of resistant HCS phases (hydrated calcium silicates), which is associated with increased strength when using cements and pozzolanic admixtures. This also occurs, although to a lesser extent, with the HCA (hydrated calcium aluminates), SA (sulfoaluminates) and FA (ferroaluminates) phases, generated by gypsum, sulphates, aluminous cement, natural cements and lime. Expression of the relationship between soluble silica and strength:

$$\sigma^+ = 31.2 * (\% \text{ soluble SiO}_2) + 37 \pm 5\% \text{ kP} / \text{cm}^2 \quad (2)$$

From there, the stabilisation table is consulted based on the clay's mineralogical type to determine which stabilisers are likely to be most effective. In case of doubt, the best approach is to experimentally test all available stabilisers that appear to yield the best results. The stabilisation table is based on laboratory tests with pure clays conducted in several of our own studies [2,7,9,10,12].

In the final phase of the analytical method, the effectiveness of the stabilised mixture as a construction material must be verified. This involves assessing the strength and durability of various series of five test specimens, using the established stabilisers and proportions (Appendix A and B). Strength measurements - typically compressive and shear in seismic zones - will be performed using 4 x 4 x 4 cm RILEM-type micro-specimens, and ultimately, for final decision-making, with actual CEB specimens (Table 3 & Figure 6).

**Table 3:**

Clays	Stabilisers in Order of Performance to Increase Resistance. Stabilisations Shown in Red Indicate Poor Durability [2]			
Increase in resistance	>100%	100-50%	<50%	Avoid, despite improved water performance
Kaolinite (acidic soils: granites, amphibole granodiorites.)	C. Portland [2]		Cellulose [10]	
	Natural cement and gypsum	CH 10%	Nanofibres [10]	
	<i>Ceranton siliqua</i> L [7]			
	Hydraulic lime 10%+Enzymes 3% [10]	NaOH 0.5%		
			Air lime.	Poor compaction if not stabilised
			Ethyl polysilicate; Sivonal 40 [6]	
Montmorillonite			<i>Ceranton siliqua</i> L [7]	Sodium silicate, prickly pear cactus, <i>Chelidonium majus</i> [12]
	C. Portland (improves with mixed lime)	Caustic soda NaOH 1.5% [9]	Enzymes 0.6% + water 15% [10]	
(basic soils: gabbros, diorites and limestones)	Natural cement and gypsum		Superionised and magnetised water (40%) [9]	Avoid hydraulic lime
Halloysite			Natural cement and gypsum	Avoid soda ash and cement
Chlorite (ultrabasic soils: crystalline schists)	C. Portland	Natural cement and gypsum		Avoid soda ash
			Caustic soda NaOH 1.5 %	Avoid lime alone
Illite	C. Portland		Paval and lime	
	Fly ash and lime, ratio 2 : 1		Gluten	
			<i>Ceranton siliqua</i> L (1)	
			Hydraulic lime	

**Figure 6:** Shrinkage test on three soil samples for rammed earth.



code	Microtest tube Composition Group I	Compression kP/cm2	content % SiO <sub>2</sub> + Al <sub>2</sub> O <sub>3</sub>	Total soluble fraction SiO <sub>2</sub> + Al <sub>2</sub> O <sub>3</sub> , Trivalents, oxides and sulfates content %
Ka	microtest tube Kaolinite 50%.	53.57 kP/cm2	0.83	1.54
Ka14P	microtest tube Kaolinite 50%.14% Portland Cement	180.37 kP/cm2	7.76	9.98
Ka NaOH-	microtest tube Kaolinite 50%.1.5% NaOH	82.65 kP/cm2	2.40	2.71
Ka14CN7Y	microtest tube Kaolinite with Gypsum +Natural Cement	124.83 kP/cm2	6.06	12.85
CL Chlorite	microtest tube 50%Chlorite	39.76 kP/cm2	1.81	1.87
CL NaOH-	microtest tube 50%Chlorite with 1,5% sosa	22.33 kP/cm2	2.35	2.45
CL14P OH-	microtest tube 50% Chlorite with 14% Portland Cement	105.01 kP/cm2	6.72	7.42
CL14CN7Y OH-	microtest tube 50%Chlorite 14% Natural Cement+ 7%Gypsum	67.60 kP/cm2	8.64	15.79
ILL Illite	microtest tube 50% Illite EE.UU	96.57 kP/cm2	2.02	2.12
ILL NaOH-	microtest tube 50% Illite with 1,5% sosa	127.64 kP/cm2	4.38	4.49
ILL14P OH-	microtest tube 50% Illite with 14% Portland Cement	247.10 kP/cm1	7.68	8.38
ILL14CN7Y OH-	microtest tube 50% Illite with 14% Natural Cement + 7% Gypsum	85.73 kP/cm2	5.53	12.17
HALL Halloysite	microtest tube 50% Halloysite pure Indiana. EE.UU	38.13 kP/cm2	17.32	17.33
HALL NaOH-	microtest tube 50% Halloysite pure with 1,5% sosa	32.25 kP/cm2	25.22	25.23
HALL14P OH-	microtest tube 50% Halloysite with 14% Portland Cement	36.44 kP/cm2	28.49	29.26
HALL14CN7Y OH-	microtest tube 50% Halloysite with 14% Natural Cement+ 7% Gypsum	45.18 kP/cm2	28.04	35.99
MON	microtest tube 50% Mon.Montmorillonite EE.UU	79.34 kP/cm2	1.28	1.48
MON NaOH-	microtest tube 50% Montmorillonite with 1,5% sosa	144.89 kP/cm2	1.99	2.20
MON14P	microtest tube 50% Montmorillonite 14% Portland Cement	206.72 kP/cm2	6.70	7.50
MON14CN7Y	microtest tube 50% Montmorillonite 14% Natural Cement +7% Gypsum	164.32 kP/cm2	5.94	12.70

### Appendix A.

Name test tube	Dry density gr/cm3	Compresión Test kP/cm2.	Name test tube	Dry density gr/cm3	Compresión Test kP/cm2.
		Breaking speed 4mm x min			breacking speed 4mm x min
NaOH- 14CN7Y 14P	Contents 1.5% caustic soda Contents 14% Portland Cement Contents 14% Natural Cement				
Kaolinite from Cañada de Verich (Teruel, Spain) and WARD'S EE.UU ref. 49H 0290			Illite from USA Ref. Ward's Catalog. 46E0315,46E4100		
Ka	1.89	12.31	ILL'	1.74	21.22
Ka	1.91	11.80	ILL'	1.74	22.23
Ka	1.82	21.15	ILL	1.74	39.03
Ka	1.89	13.78	ILL	1.89	28.98
Ka NaOH-	1.82	19.59	ILL	1.67	10.21
Ka NaOH-	1.85	17.60	ILL NaOH-	2.02	27.75
Ka14CN7Y	2.10	7.34	ILL NaOH-	2.05	29.68
Ka14CN7Y	1.71	5.98	ILL14P	1.84	54.08
Ka14P1	1.93	60.68	ILL14P	1.91	57.38
Ka14P2	1.89	70.76	ILL14P	1.92	45.90
Ka14P3	1.96	74.40	ILL14P	1.94	66.05
Ka14P4	1.87	62.22	ILL14P	2.02	55.31
Ka14P4	1.82	58.17	ILL14P x	2.02	45.46
Chlorite from Collserola (Barcelona, Spain)			ILL14P xx	1.93	44.80
			ILL14CN7Y	1.83	12.05
CL	1.62	8.50	ILL14CN7Y	2.01	18.93
CL	1.60	8.21	ILL14CN7Y	1.86	27.92
CL '	1.72	9.82	Halloysite from Bedford ( Indiana, USA)		
CL "	1.67	9.27			
CL NaOH-	1.80	4.70	HALL	1.56	8.71
CL NaOH-	1.66	4.47	HALL	1.54	7.94
CL NaOH-	1.82	5.90	HALL	1.59	8.56
CL14P	1.67	20.17	HALL	1.65	9.10
CL14P	1.69	17.24	HALL NaOH-	1.52	8.41
CL14P	1.70	22.82	HALL NaOH-	1.57	8.71
CL14P	1.75	29.67	HALL NaOH-	1.60	7.79
CL14P	1.76	23.71	HALL NaOH-	1.56	6.07
CL14P	1.87	30.22	HALL NaOH-	1.54	5.18
CL14P	1.77	23.03	HALL14P	1.54	9.35
CL14P '	1.79	24.45	HALL14P	1.59	11.57
CL14P '	1.80	22.80	HALL14P	1.65	14.75
CL14CN7Y'	1.46	15.68	HALL14P	1.58	11.59
CL14CN7Y'	1.63	18.55	HALL14CN7Y	1.53	13.88
CL14CN7Y	1.55	16.67	HALL14CN7Y	1.70	8.99
CL14CN7Y	1.52	18.95	HALL14CN7Y	1.70	15.76
CL14CN7Y	1.49	12.76	HALL14CN7Y	1.60	16.72
CL14CN7Y	1.59	11.99	HALL14CN7Y	1.49	12.24
Montmorillonite from USA . Ref. Ward's Catalog 49 H2600					
MON	1.83	23.19	MON14P	1.89	63.79
MON	1.86	30.73	MON14P	1.97	99.99
MON	1.88	30.85	MON14P	1.88	167.41
MON NaOH-	2.24	67.74	MON14CN7Y	1.84	67.80
MON NaOH-	1.96	84.42	MON14CN7Y	1.75	52.58
			MON14CN7Y	1.75	59.62

### Appendix B.

## Stabilisers

The sustainable selection criteria to be applied are summarised in UNE Standard 41410, Annex C - whose drafting we contributed to - and in reference [2]. These include:

- Use of a local product.
- Technological processes appropriate and adapted to the sociocultural environment; it is essential to adopt a technology that is simple to implement and readily assimilated by developers and a workforce accustomed to conventional architecture.
- Economic evaluation.
- Maintenance and use conditions.

Environmental impact must be minimised throughout the stabiliser's life cycle assessment (LCA): minimal consumption of energy, water and non-renewable resources, and minimal emission of toxic substances into the environment during extraction,

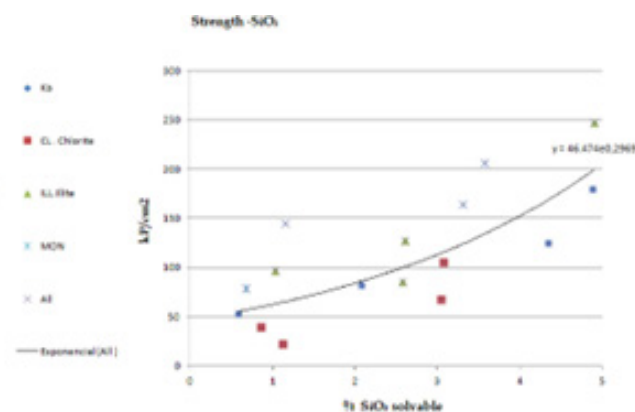
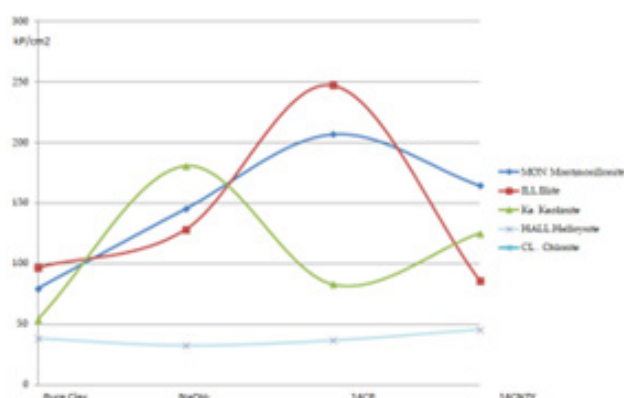
manufacturing, transport, application and throughout its useful life. An initial inventory of inputs and outputs is carried out, along with a comparison with other materials. The calculated values for CEB (stabilised compressed earth block) range from 0.4 to 0.82MJ/kg - a value three times lower than if the block were baked.

e. Curing, deflocculation and hardening factors derived from the local climate.

f. Environmental and structural risks: frost, wind, temperature fluctuations, risk of cracking, and curing conditions.

## Simple and diagonal compressive strength

The defines the condition of the test specimen after immersion for four hours, followed by a flatness check of the faces without sulphur coating. Specimens are typically 10 x 10cm, but we continue to use 4 x 4cm RILEM specimens, tested at a loading rate of 1mm/min (equivalent to 3/7kP/cm<sup>2</sup>s), which yields an average accuracy of approximately 6%. The tests are documented in Appendix A and B (Figure 7).



**Figure 7:** Compressive strength results using mini-specimens of pure clay samples [2].

We have verified that the calculated strength of the masonry, when tested using diagonal compression on CEB specimens, approximates the formula established in the CTE (Spanish Technical Building Code):

$$f_{v_k} = f_{v_{k0}} + 0.36 \times \sigma_k = 0.065 \times f_b \quad (3)$$

## Durability tests. absorption

The first stage of the durability tests consists of saturating the test specimens in distilled water under laboratory conditions. To evaluate the most superficial capillary pores, an initial reading is taken one minute after immersion. Total saturation may also be estimated after 48 hours of immersion, although this step can be omitted due to the similarity of results observed in previous tests. Any method involving boiling has been discarded to avoid the pressure exerted by water currents and the resulting loss of sample integrity. There is an inversely proportional relationship between the percentage of absorption and the soluble silica content. The trend follows an exponential pattern, as expressed by the following equation: [2]

$$y = 1.9x - 0.37 \quad (3)$$

## Cyclic Wetting-Freezing-Drying Test

The proposed durability evaluation method combines, in a single test, six cycles of freezing, wetting, drying and aggressive attack, including sulphate analysis in cases where stabilisation involves non-sulphur-resistant cements. The procedure is derived from ASTM D-559 and D-560, NLT-303-72 and UNE 7033 for freezing tests, and consists of 20 cycles alternating between 10 °C (one hour in water) and -15 °C (four hours).

Following the absorption test, specimens are dried in a muffle furnace at 60 °C for 48 hours. After this period, the specimens are brushed using a tool with a diameter equal to 50% of the surface area of the specimen faces, applying a pressure of 15N. This pressure is verified by placing the sample on the balance pan and applying the brush until a reading of 1.5kg is obtained. After the first wetting and drying cycle, a frost shattering (G) cycle is carried out at -15 °C for eight hours, followed by drying in an oven at 60 °C for the same duration to avoid structural alterations to the clays.

The third cycle consists of saline wetting (Ws) and drying (S). The sodium salt is dissolved to saturation under laboratory conditions. The fourth cycle combines saline wetting, frost shattering and drying in eight-hour periods. The aim is to exacerbate the expansive stresses caused by salt attack with those resulting from ice crystal formation.

## Conclusion

Our 35 years of experience in the earth construction sector have enabled us to develop a more coherent and effective working method, and to share a summary of key findings. We highlight the considerable variation in the behaviour of clay samples when exposed to lime, specialised cements and gypsum. However, improvements in mixing and compaction water, the use of fluidisers and the application of mechanical pressure can be achieved without relying heavily on additives. From the interrelation of the results obtained using the described method, a direct relationship emerges between the crystallographic and chemical nature of the soil and its most effective stabilisation system. There is also a correlation between strength, void ratio  $e$ , degree of stabilisation %, the proportion of fines smaller than 0.4mm (modulus % M), and the liquid limit of the soil (WL) (Figure 8).



**Figure 8:** Earth block shear test conducted at Girona University Laboratory [17].

Water resistance remains a critical factor - its limitations have historically undermined the viability of this construction method and contributed to its decline. This issue must be addressed with even greater urgency given the current scenario of increasingly extreme rainfall. Stabilisation methods that improve water resistance have, in some cases, led to reduced strength. Nonetheless, they are preferable and more durable than relying solely on external hydrophobic treatments. As research into new

soil types and stabilisation methods continues, it is essential to interrelate the data to produce simpler yet more comprehensive working tables that integrate the scientific knowledge relevant to this construction material today.

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