

Open access • Journal Article • DOI:10.1007/S11440-013-0251-6

Soil treatment with organic non-traditional additives for the improvement of earthworks — Source link [2]

Gaëtan Blanck, Olivier Cuisinier, Farimah Masrouri

Institutions: University of Lorraine

Published on: 01 Dec 2014 - Acta Geotechnica (Springer Berlin Heidelberg)

Topics: Compaction

Related papers:

- Stabilization Mechanisms of Nontraditional Additives
- · Stabilization of Clay Soils with Nontraditional Additives
- · Physicochemical behavior of tropical laterite soil stabilized with non-traditional additive
- Analysis of strength development in non-traditional liquid additive-stabilized laterite soil from macro- and microstructural considerations
- Effects of Xanthan gum biopolymer on soil strengthening





Soil treatment with organic non-traditional additives for the improvement of earthworks

Gaëtan Blanck, Olivier Cuisinier, Farimah Masrouri

▶ To cite this version:

Gaëtan Blanck, Olivier Cuisinier, Farimah Masrouri. Soil treatment with organic non-traditional additives for the improvement of earthworks. Acta Geotechnica, Springer Verlag, 2014, 9 (6), pp.1111–1122. 10.1007/s11440-013-0251-6. hal-01418290

HAL Id: hal-01418290 https://hal.univ-lorraine.fr/hal-01418290

Submitted on 5 Jul 2019

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

- 1 Soil treatment with organic non-traditional additives for the improvement of earthworks
- 2
- 3 Gaëtan Blanck^{1,2}, Olivier Cuisinier¹, Farimah Masrouri¹
- 4 ¹ Laboratoire Environnement, Géomécanique & Ouvrages
- 5 École Nationale Supérieure de Géologie bâtiment E
- 6 Rue du Doyen Marcel Roubault B.P. 40
- 7 F-54501 VANDOEUVRE-LES-NANCY Cedex
- 8
- 9 ² Agence de l'Environnement et de la Maîtrise de l'Énergie
- 10 20, avenue du Grésillé BP 90406
- 11 F-49004 ANGERS Cedex 01
- 12 gaetan.blanck@univ-lorraine.fr
- 13 Tél : + 33 (0)3 83 59 63 01
- 14 Fax : + 33 (0)3 83 59 63 00
- 15
- 16

17 Abstract

18 The aim of this study was to characterise the technical and environmental implications of 19 non-traditional treatments of a low-plasticity compacted silt (PI = 14) and to investigate their 20 actions both at the micro- and macro-structural scales. Three non-traditional additives 21 derived from industrial vegetal by-products were studied. These additives are classified as an 22 acid solution (AS), an enzymatic solution (ES) and calcium lignosulfonate (LS). The first step 23 was to characterise the effects of these treatments on Proctor compaction, bearing capacity, 24 unconfined compressive strength and stiffness. The index properties of the treated samples 25 were also measured to assess changes in the interaction between soil minerals and the 26 additives. These experimental results showed that the 0.002% ES and 2.0% LS treatments 27 were the most effective at improving the soil dry density and allowing the soil to reach 28 optimum density using 3% less water or 25% less compaction energy. In a second step the 29 mechanisms involved by the treatment were investigated. A microscopic study was 30 conducted including scanning electronic microscopy (SEM) and mercury injection porosity 31 (MIP) tests, measurements of the surface tension of non-traditional additives mixed with 32 water completed the study and demonstrated that the ES treatment has the same surfactant 33 properties as sodium dodecyl sulfate, a common surfactant. Compaction tests confirmed that 34 the behaviours of the soil after ES and sodium dodecyl sulfate treatments were similar.

35 Key words: Soil treatment, non-traditional additives, compaction, bearing capacity, surfactant

36

37 **1. INTRODUCTION**

38 In earthworks, sustainable development will play a larger role in future project design and the 39 assessment of technical solutions by contractors. Currently, the reduction of the 40 environmental impact of earthworks is a growing issue. One of the major challenges facing 41 earthworks companies is the need to use a majority of natural materials extracted within the 42 construction site, including materials with very low geotechnical characteristics. The common 43 industry solution is to use lime, cement or fly ash to improve the soil geotechnical properties; 44 however, sustainable development principles also require a reduction in the consumption of 45 water and non-renewable energies as a way to limit greenhouse gas emissions. In this 46 context, soil treatment with various non-traditional additives has been tested. These additives 47 are diverse in nature and some of them derive from the industrial transformation of 48 renewable raw materials. Due to their organic nature, some products are biodegradable. 49 Thus, the treated soils can recover their untreated composition after the end of the 50 biodegradation process. These products are also beneficial due to their efficiency at low 51 dosages, thereby limiting the environmental impact and cost of their use. Various additives 52 have been tested for improving the compaction of dry soils to avoid dust generation and for increasing the durability of uncovered roads [5, 22, 24, 27]. Additionally, additives have been 53 54 tested to stabilise erodible soils [33]. Mechanical properties, such as the unconfined 55 compressive strength [23, 28] or the swelling potential of plastic soils [11, 19], have also 56 been studied.

57 Tingle and Santoni [28] listed seven non-traditional categories based on the major active 58 component and classified them as salts, acids, enzymes, lignosulfonates, petroleum 59 emulsions, polymers and tree resins. Acids are low pH aqueous solutions containing 60 sulfonated molecules such as naphthalene or limonene [13]. The presence of proteins in 61 enzymatic solutions was confirmed by Velasquez et al. [32]. Lignosulfonates are organic

62 polymers derived from lignin extracted by the sulfite processing of cellulose in the wood-pulp 63 and paper industries. According to Tingle et al. [30], petroleum emulsions contain asphalt or 64 synthetic iso-alkane fluids suspended in emulsions by a surfactant, polymer emulsions 65 contain vinyl acetates or acrylic copolymers and tree resins are diverse emulsified byproducts of the timber and paper industries. Nevertheless, many of these products also 66 67 include secondary additives such as surfactants or ultraviolet inhibitors, which may react with 68 the soil minerals or modify the soil water properties and thus influence the mechanical 69 behaviour of the treated soils [30]. The emphasis of this study was put on non-traditional 70 additives that would lower the environmental impact of soil treatment in the earthworks 71 industry. Therefore, this study focused on the following products which are derived from the 72 transformation of renewable organic matters: acids, enzymes and lignosulfonates. For each of these categories, a review of the literature concerning the effects on geotechnical 73 74 properties and suspected mechanisms is presented. Then, geotechnical properties such as 75 compaction, bearing capacity, unconfined compressive strength and stiffness were 76 investigated to define the potential application of these treatments. In the last part, the action of the treatments on the microstructure of the soil was studied by performing scanning 77 78 electronic microscopy (SEM) and mercury injection porosity (MIP) tests to identify the 79 mechanisms involved by the different products.

80 2. LITERATURE REVIEW

81

2a. Acid solutions

Rauch et al. [20] studied the effects of an acid product on the index properties of three pure clay phases (kaolinite, illite and montmorillonite) and two plastic soils. They concluded that the acid treatment had little effect on soil plasticity, swelling potential and shear strength. The authors explained that the observed modifications were induced by initial differences in properties of the compacted samples. Tingle & Santoni [28] and Santoni et al. [23] tested the

87 application of an acid solution to three soils ranging from non-plastic to highly plastic in nature. They found no change in the unconfined compressive strength. The authors 88 89 determined that the acid treatment had no effect on the compaction properties of the soils. 90 However, another study [21] showed that changes in the compaction properties of acid-91 treated soils can occur. Scholen [24] reported in situ studies undertaken by the US Forest 92 Service between 1991 and 1993 which tested the addition of acid solutions. In 1995, no 93 particular failure or degradation of the treated sections was reported. Scholen [24] proposed 94 that acidic non-traditional additives act by altering the electrolyte concentration of the pore 95 fluid, thereby reducing the double layer thickness of clay minerals. Cation exchange was also 96 suspected to occur and to flocculate the clay minerals. Moreover, important lowering of the 97 pH of a soil can also lead to the dissolution of clay minerals [26, 36]. For example, to start the dissolution of kaolinite, the pH values have to be lower than 2 [16, 35]. However, the X-ray 98 99 diffraction, specific surface area and cation exchange capacity were not modified by the 100 treatments, even for dosages as high as 50% [13, 21]. Scanning electron microscope 101 pictures obtained by Katz et al. [13] showed similar structures for the acid-treated and 102 untreated soils. According to the literature, acid solutions modify the compaction and bearing 103 capacity properties of treated soils without modifying the soil mineralogy.

104

2b. Enzymatic solutions

For enzymatic treatments, Rauch et al. [20, 21] concluded that most of the variations in the Atterberg limits they observed were insignificant compared to the typical variations of such measurements. Unconfined Compressive Strength (UCS) tests of the treated soils demonstrated modifications from -30 to +80%, depending on the product and soil tested [18, 28]. Velasquez et al. [32] mentioned an increase in shear strength resistance of 9 to 39% for two enzymatic treatments but detailed analysis of samples properties showed that modifications can be attributed to differences in water content and dry density after 112 treatment. For some enzymatic solutions, limited changes in the compaction references were 113 found [19, 21]. Tingle and Santoni [28] considered that the modifications in the compaction 114 properties after enzymatic treatments were too small to be significant; however, Velasquez et 115 al. [32] mentioned that samples treated with enzymatic solutions achieved a higher density 116 for the same compaction energy. In two of the three field studies reported by Scholen [24], 117 the treated section of uncovered forest roads showed less degradation than untreated 118 sections. Visser [34] performed in situ California Bearing Ratio (CBR) measurements on four 119 soils ranging from sandy to clayey treated with an enzymatic solution. CBR improvement 120 occurred in 4 cases out of 8. For enzymatic treatment, Scholen [24] hypothesised that 121 enzymes bond the organic matter of the soil with clay minerals. These additional bonds 122 should reduce the water affinity of the clay and limit the swelling potential of the soil. 123 However, Harris et al. [11] showed that the 3-D swelling potential of a plastic clay (PI = 51) 124 was not modified by the addition of 0.07% of an ES treatment. Another stabilisation 125 mechanism was suggested by Velasquez et al. [32], who observed that an ES could have 126 surfactant properties and thus stabilise the soil by improving its compaction ability. This 127 property was confirmed by Berney et al. [4] and Park et al. [17], who showed that the dry 128 density of compacted soils is improved by the addition of surfactants. In conclusion, ES seems to induce compaction modifications, but the stabilisation mechanisms of this family of 129 130 products remain in question.

131 **2b. Lignosulfonates**

The addition of lignosulfonates modified the index properties of treated soils in different ways. Lambe [14] showed that small quantities of calcium lignosulfonate (< 0.5%) reduce the liquid limit of a low-plasticity soil. Gow et al. [10] treated a silty sand with 0.25 to 1.0% of lignosulfonate and observed that the plastic limit was reduced when the dosage increased, but no consistent trend was found for the liquid limit. A maximum improvement in the UCS 137 was observed for the addition of 5% of lignosulfonates [23, 28]. Compaction properties were 138 also affected by the treatment. Gow et al. [10] showed that the optimal water content of a 139 compacted soil was reduced and the maximum dry density increased especially for dosages 140 between 0.5 and 1.5%. CBR values measured for each dosage at optimal water content and 141 maximum dry density were improved in comparison to untreated samples but decreased with 142 increased dosages. In situ evaluations of the mechanical performances of the lignosulfonate 143 treatment are limited. Surdahl et al. [27] mentioned a stiffness improvement of a granular soil 144 after treatment. Lignosulfonates are thought to stabilise the soil by coating together its 145 particles [30, 33]. However, lignosulfonates also have surfactant [31] and dispersive 146 properties [14, 25]. Thus, the stabilisation mechanism of lignosulfonates is not yet well 147 understood.

148 In general, soil treatment with non-traditional additives can improve the engineering 149 properties of soils; however, these results vary depending on the tested products, soil nature 150 and at times, the dosages. According to Rauch et al. [20] the variability of the engineering 151 properties may be due to the modification of the optimal density and water content of 152 compacted soils. Thus, as for traditional treatments such as lime or cement, the addition of 153 chemical additives may induce modifications of maximum dry density and optimal water 154 content, which must be determined for accurate field applications. Moreover, the 155 strengthening mechanisms of non-traditional treatments are not yet well understood and 156 where data are available, no correlations between microscopic and macroscopic 157 observations have been established. Therefore, some experiments were conducted to 158 identify the mechanisms of each additive.

159 3. MATERIALS AND METHODS

160 **3a. Non-traditional additives**

The acid solution (AS) tested in this study was a concentrated solution of sulfuric acid mixed with D-limonene, a by-product of citrus processing. For this kind of products, typical dosages tested in the literature are comprised between 0.01 and 0.2%. As shown by different authors, UCS, compaction and clay mineralogical properties were not affected by dosage increase [13, 21, and 28]. Thus, a dosage of 0.01% of concentrated product by dry weight of soil was tested in this study (Tab. 1).

The tested enzymatic solution (ES) is a biodegradable brown aqueous solution with an acidic pH of 4.6. For this type of product, typical dosages are comprised between 0.002% and 0.1%. However, according to Tingle & Santoni [28] and Velasquez et al. [32], the geotechnical properties of soils treated with an enzymatic additive were independent of the dosage. Therefore a dosage of 0.002% was selected.

The tested lignosulfonate (LS) was a biodegradable, organic, water soluble calcium salt used in powder form. The insoluble components were less than 0.5%, the moisture content was approximately 7% and the reducing sugar content was approximately 7% by dry weight of product. Experimental results indicated an optimal additive content of 5% [23, 28] for improving the UCS. On the other hand, Gow et al. [10] had demonstrated that maximum density and bearing capacity values were obtained for a dosage of 0.5%. For that reason, three dosages (0.5, 2.0 and 5.0%) were tested.

The tested surfactant was a powder with a SDS content of more than 95 % by dry weight.
The required quantity of SDS for a 0.01% dosage was diluted in distilled water and then
added to the soil during mixing operations.

182 3b Tested soil

A natural fine-graded silt of low plasticity often encountered in earthworks projects in the Paris Basin was tested in this study (Tab. 2). X-ray diffraction investigations indicated that the main minerals were quartz and calcite (8.0% by weight determined by calcimetry). The <2 μ m fraction contains quartz and clay minerals (chlorite, illite, kaolinite and smectites) and the sulphur content of the soil was less than 0.02%.

188 Before testing, the soil was dried at 60°C and pulverised to pass through a 2 mm sieve. 189 Depending on the liquid or solid nature of the treatment product, a specific protocol was 190 followed to ensure the application was representative of field application procedures. For the 191 AS and ES treatments, the initial quantity of water was mixed with the dry soil material to 192 achieve a water content 3% lower than the target moisture content for compaction. The 193 samples were placed in a container at 20°C for 24 hours to reach the equilibrium of moisture 194 content. Then, the additive was mixed with the quantity of water required to obtain the total 195 desired water content and added during the mixing operation. For LS treatment, the soil was 196 directly brought to the desired water content and set aside for 24 hours to equilibrate. Finally, 197 the powder was added to the soil during mixing. For all treatments, a curing time of one hour 198 in sealed containers was chosen before conducting further tests.

199

200

4. ALTERATION OF GEOTECHNICAL CHARACTERISTICS AFTER TREATMENT

201 4a. Index properties

The liquid limit (LL) and plastic limit (PL) of the treated soil were determined (Tab. 3). For the AS, ES and SDS treatments, the Atterberg limits were not modified; however, a reduction in the LL and PL occurred after the addition of lignosulfonate (LS) at dosages of 2.0 and 5.0%. Despite these changes in the consistency limits, the plasticity index remained constant for all tested treatments and dosages.

207 **4b. pH**

The pH of the treated soil was determined according to ISO 10390 [12]. Measured values showed that the pH of the AS and ES treated soil was not affected despite the acidic features of both products. This phenomenon might be related to the high dilution of the products combined with the increased buffering capacity of the soil due to the presence of calcium carbonate. In contrast, the addition of the LS treatment reduced the pH by approximately one pH unit for the 5.0% dosage. This action is due both to the acidic carboxylic and sulfonate functions of lignosulfonate and to the high dosages.

215 **4c. Compaction**

216 Compaction tests were performed according to ASTM D 698-91 [1] using a standard Proctor 217 test. After compaction, the instant bearing index (IPI) was measured following the same 218 procedure as for the un-soaked Californian bearing ratio (ASTM D 1883-92 [2]) but without 219 placing surcharge weights. A penetration piston of 49.6 mm diameter is used to apply the 220 load on the compacted soil. The penetration rate was 1.27 mm/min. The compaction and 221 instant bearing index curves are plotted in Fig. 1 for AS and ES treated soil. Fig. 2 shows the 222 results for the addition of LS at 0.5, 2.0 and 5.0%. The optimum water content (wopt) and the 223 maximum dry unit weight (γ_{dmax}) for each treatment are summarised in Tab. 4.

Fig. 1 shows that for the AS treatment, no changes in the compaction and IPI curves occurred. After ES treatment, the maximum dry unit weight increased and the optimum water content reduced from 18.2 kN/m³ and 15.5% to 18.6 kN/m³ and 14.5%, respectively. On the dry side of the compaction curve, a maximum increase in the dry unit weight of 0.8 kN/m³ was measured. This means that the compaction ability of the treated soil was improved over a range of water contents from 8 to 15.5%. The IPI curve was also affected by the treatment and shifted towards lower water contents. Thus, the bearing capacity of the soil was reduced 231 for water contents in the vicinity of w_{opt}. However, the measured IPI at the respective 232 optimum water content for both ES treated and untreated soil were roughly equal (Tab. 4). 233 Fig. 2 shows that the LS treatment increased the dry unit weight and reduced the optimum 234 water content of the soil for the three dosages. A maximum densification was obtained with 235 the addition of 2.0% LS, which corresponded to a reduction of 2% in the optimum water 236 content (13.5% for 2.0% LS treatment as compared to 15.5% for untreated soil). For this 237 same treatment, dry unit weights were increased on the dry side of the compaction curve and 238 γ_{dmax} increased from 18.2 to 18.5 kN/m³. Despite an improvement in density, the addition of 239 lignosulfonate reduced the IPI at constant water content, and the IPI reduced further with 240 higher doses of LS. If the soil was compacted at w_{opt} to reach the maximum dry density, the 241 IPI measured for the 2.0% LS treatment was 14, compared to a value of 15 measured at wort 242 for the untreated soil (Tab. 4). These results illustrate the importance of considering the 243 entire compaction curve when assessing the effects of ES and LS treatments.

244 After 0.002% ES and 2.0% LS treatments, an increase in dry unit weight and a reduction in 245 w_{opt} were observed, implying that a minimum dry density may be achieved for lower water 246 contents. For example, the γ_{dmax} of untreated soil may be achieved at 3% lower water 247 content. Moreover, both treatments could also reduce the energy needed to compact the soil 248 to a minimum desired density. An estimation of the potential energy savings was made 249 according to the results and is presented on Figs 3 and 4. These graphs show the 250 compaction and bearing capacity curves for the ES and LS treated soil compacted at 100% 251 and 75% of the standard Proctor energy. For samples compacted on the dry side of the 252 compaction curve, the measured dry unit weights were the same as for untreated soil despite 253 the 25% energy reduction. For the ES and LS treatments, however, the bearing capacity of 254 the soil compacted at 75% of the standard Proctor energy was reduced compared to the 255 untreated soil.

256 4d. Unconfined compressive strength

257 To study the evolution of the mechanical behaviour of treated soil, UCS values were determined at 7 and 28 days for samples compacted at their respective w_{opt} and γ_{dmax} after 258 259 treatment (Tab. 4). Three replicate samples were statically compacted in one layer in a 260 cylindrical mould 100 mm high and 50 mm in diameter. Samples were wrapped in a plastic 261 film, covered by an aluminium sheet and stored in a plastic container at 20°C. Geometrical 262 parameters, sample weight and water content were controlled after the sample compaction 263 and just before testing. The results of the UCS tests at 7 and 28 days are presented in Fig. 5. 264 In comparison with the untreated soil, the resistance of the AS and 2.0% LS treated soil was not significantly modified. The UCS of ES treated soil was slightly improved due to a higher 265 266 maximum dry unit weight obtained after treatment, but the addition of 0.5 and 5.0% of 267 lignosulfonate reduced the unconfined compressive strength by 20 and 50%, respectively. 268 Moreover, UCS values only displayed minor changes as a function of curing time, implying 269 that the treatments mainly have short term effects. For samples compacted at w_{opt} and γ_{dmax} 270 determined for each treatment after 7 days of curing, the mean modulus for untreated, ES 271 and 2,0% LS treated soils were respectively 13, 15 and 22 MPa (Tab. 5). Samples were also 272 compacted at a target water content of 12.5% and a dry unit weight of 18.5 kN/m³. The 273 results showed that only the 2.0% LS treatment significantly reduced the stiffness of the soil.

274 **4e. Swelling potential and hydraulic conductivity**

The determination of one dimensional free swell potential of treated soil was carried out on samples of 70 mm diameter and 15 mm thickness statically compacted in oedometric cells at different initial compaction states. The samples were saturated with distilled water.

Fig. 6 shows the one dimensional swelling potential (Δ H/H₀) for untreated, 2.0% LS and ES treated samples. For each treatment, at optimal conditions of compaction, the swelling potentials were not significantly modified. Moreover, when the untreated soil was compacted at the same dry density and initial water content than treated samples, the swelling potentials were of the same order of magnitude. Therefore, it could be concluded that the studied products did not significantly alter the swelling potential of the tested soil.

After the stabilisation of swelling, the saturated hydraulic conductivities of the samples were determined (Fig. 7). Treated samples compacted at their respective maximum dry densities showed a small decrease of the hydraulic conductivity. For treated and untreated samples on the dry side of the compaction curves, the hydraulic conductivities were very similar.

This section showed that the non-traditional additives led to several modifications of the geotechnical behaviour of the tested material. The modifications did not appear to be timedependant since no significant modification of unconfined compressive strength occurred over the curing period considered. Nevertheless, a key question is the permanence of the improvement brought by these products over a long period of time. In this context, the interaction mechanisms between the soils particles and the additives were investigated.

294

5. THE STUDY OF INTERACTION MECHANISMS

295 **5a. Microstructure of compacted soil**

The investigation of the microstructure of compacted soil was conducted using a scanning electron microscope equipped with an energy dispersive X-ray spectrometer (EDS). Images obtained by collecting the secondary electrons were taken at different magnifications between 100 and 10,000x. All samples were compacted at their respective optimum water content and maximum dry unit weight. After compaction, they were wrapped in a plastic film, covered by an aluminium sheet and stored in a plastic container at 20°C for 28 days. Then, samples 2 to 3 cm³ in volume were carefully extracted and freeze-dried to remove the water and minimise sample deformation. Fragments of the freeze-dried samples were extractedand coated with gold before the SEM analyses.

305 To study the organisation of the internal structure of the compacted soil, SEM investigations 306 were conducted with mercury injection porosity (MIP) tests. Tests were conducted on freeze-307 dried samples compacted at the maximum dry density and optimal water content for each 308 treatment following the same procedure as for SEM preparation. The pore size distribution is 309 obtained by derivation of the injected mercury volume (Vini) function. To compare the four 310 samples, the injected volume was normalised such that the surface under the pore size 311 distribution curve was equal to 1. The void ratio investigated by the mercury (einv) can be 312 defined as the ratio between the injected volume of mercury and the volume of solids. When 313 einv is equal to etot, the entire porosity of the sample is filled by mercury. Thus, the calculated 314 e_{inv} (Tab. 6) reveals that the majority of the existing pores in the samples were filled with 315 mercury during MIP tests.

Fig. 8 shows the surface of the untreated soil. The mineralogical nature of the bigger grains, generally measuring 10 to 50 μ m, was determined by the EDS. These grains were identified to be quartz associated with smaller clay minerals of 1 to 5 μ m diameter. No preferential orientations of the clay minerals were detected. MIP results indicated that the pore size of untreated soil presents a uni-modal distribution with a dominant radius of 0.6 μ m.

Results obtained by SEM showed a slightly more aggregated structure and a face to edge organisation of clay minerals for the soil treated with 0.01% AS (Fig. 9). Individual clay minerals are less visible but inter-aggregates pores size remained similar after treatment as confirmed by MIP data (Figs 12 and 13). Therefore, the structure of the treated soil was not significantly altered in this case.

After 0.002% ES treatment, the same structure as for the untreated soil is observed from the SEM images (Fig. 10). The MIP study (Fig. 12) showed that less mercury was injected in the ES treated sample. This phenomenon is linked with the higher dry unit weight of the ES treated sample, which reduced its total void ratio (e_{tot}) and thus the total injected volume (V_{inj}). This result is associated to a reduction of the amount of pores comprised between 3 and 10 µm (Fig. 13).

332 Fig. 11 shows the soil after the addition of 2.0% of lignosulfonate. In the picture, additional 333 structures can be observed. These structures resemble fibres and are probably 334 lignosulfonate associations forming a network between the soil minerals (arrows). Point 1 on 335 Fig. 11 was identified by the EDS to be a clay mineral, but as the fibres were too small to be 336 analysed by the EDS, their exact nature is unknown. The SEM results also showed larger 337 inter-aggregates pores, large clay minerals were organised in face to face structures 338 orientated in the plane of the picture. Compared to untreated samples, MIP results confirmed 339 the increase of the frequency of pores having radii between 0.3 and 5 µm and also showed a 340 reduction of pores with a radius <0.3 µm. The increase in larger pores radius might be 341 caused by aggregation of clay mineral and dispersion of the created structure due to 342 adsorption of lignosulfonates molecules. The reduction in intra-aggregates pores may be 343 explained by the clogging of the smallest pores by lignosulfonates.

5b. Comparison of the treatment additives with a common surfactant

Some authors, for example, Velasquez et al. [32] demonstrated that enzymatic solutions could have surfactant properties. As enzymatic solutions, lignosulfonates also have surfactant properties, which could alter the mechanical behaviour of soils [10, 31]. To determine if the tested non-traditional additives have surfactant properties, the surface tension was measured using the Du Noüy ring method [9]. The results were compared with 350 the surface tensions obtained for aqueous solutions of sodium dodecyl sulfate (SDS)351 (Fig. 14).

Similar to SDS, low dosages of the enzymatic solution reduced the surface tension of the water. At a concentration of 1.0 g/L the surface tension was reduced by more than 50%. Lignosulfonate also exhibited surfactant properties, but the additive was less efficient in reducing the surface tension of the solution. At higher concentrations, the surface tension of the LS solution stabilised approximately 42 mN/m. Only the acid additive did not exhibit any surfactant properties.

To study the influence of the addition of a surfactant on the compaction behaviour, 0.01% of SDS by dry matter weight was added to the soil (Fig. 15). The required quantity of SDS was first diluted in water, and the same protocol used for liquid non-traditional treatments was followed. The resulting compaction and IPI curves were compared to those obtained after the ES treatment.

With regard to the ES treatment, the SDS addition improved the dry unit weight on the dry side of the compaction curve. The IPI curve for the SDS-treated soil was also slightly shifted towards smaller water contents as was observed after the ES-treatment. Thus, the SDS and ES treatments have a similar effect on the mechanical behaviour of soils. This finding implies that both products may utilise the same stabilisation mechanism. Indeed, the reduction of the surface tension of the soil's interstitial water should lower the matrix suction according to the Jurin law.

The suction of the treated soil was investigated for the three non-traditional treatments. The filter paper methodology was used to measure matrix suction of compacted soil according to ASTM D 5298-94 [3]. After compaction of two 70 mm diameter for 15 mm thickness samples, three filter papers were inserted between each sample. Thereafter, the samples were kept in

sealed container for 7 days to achieve the suction equilibrium. As the calibration curve of the
filter paper was known, the measurement of its water content gave the matrix suction of the
soil.

The results showed that the acid solution (AS) did not change the suction of the soil. This is consistent with the absence of surfactant features of the additive.

As ES treatment has surfactant properties, a reduction of the suction could be expected. In this work, the addition of the enzymatic solution (ES) did not lead to suction modifications (Tab. 7). This can be explained by the biodegradability of the additive. An equilibrium period of seven days may allow the additive to be totally degraded before suction measurements.

383 The LS treatment led to an increase in the matrix suction of the soil. As LS is an organic 384 calcium salt, the increase of the suction might be explained by the hygroscopic nature of 385 lignosulfonates.

386 6. DISCUSSIONS

387 Herein, the effects, mechanisms and applications of the three studied treatments are388 discussed.

AS treatment did not change geotechnical properties of the soil but small modifications in the clay minerals organisation were observed in SEM pictures (Fig 9). The observed modifications could be due to changes in the ionic composition of the pore fluid after addition of the product since this parameter is known to induce fabric changes according to the literature [6-8, 15, 16]. No particular application was evidenced for AS treatment of the studied soil.

395 Treatment with ES demonstrated an increase in the compaction ability of the soil. The 396 increase in maximum dry density induced small modifications of the UCS, stiffness, swelling

397 potential and permeability of the soil. Moreover, the microstructure of the soil was not 398 modified. Therefore, it can be concluded that the observed hydro-mechanical modifications 399 are mostly related to the effect of treatment on compaction behaviour of the soil. This 400 hypothesis was confirmed by the absence of pH and index properties changes. As shown on 401 the Fig. 15, the compaction behaviour of the ES treated soil is similar to the behaviour 402 observed after a treatment with a surfactant. Results shown on Fig. 3 indicated that treated 403 soil had a smaller bearing capacity than the untreated soil at given water content and dry 404 density. This phenomenon may be explained by the surfactant properties of the ES. Strong 405 modifications of the surface tension for low additive dosages may also explain why the 406 compaction ability of the ES treated soil was improved despite a low dosage (0,002%). As 407 the enzymatic solution is biodegradable, the product has disappeared in a few days. During 408 the degradation stage of the ES product, increase in bearing capacity may be expected and 409 after degradation of the additive, no long term modifications are expected beyond those 410 caused by the initial densification. According to the experimental results, the ES treatment 411 can be used to increase the compaction ability on the dry side of the compaction curve and 412 thus compact the soil at higher densities for lower water contents.

413 LS treatment reduced the pH and modified the pore-fluid composition, thus, the organisation 414 and interactions between soil particles were changed as shown by mechanical results and 415 SEM pictures. Experimental results showed dosage dependent effects on geotechnical 416 characteristics. The 2% amount of lignosulfonate added to the soil also reduced the stiffness 417 and the index properties. This observation is consistent with dispersive properties of 418 lignosulfonates which tend to increase repulsive interactions between soil constituents [5, 14, 419 25]. LS treatment also reduced the friction between the grains of the soil, which facilitate its 420 deformation under loading, and thus explain the observed reduction in bearing capacity 421 (Figs. 2 and 4). However, surfactant properties and suction modification after treatment 422 cannot explain alone why the bearing capacity was reduced by an increase in the

423 lignosulfonate dosage (Fig. 2). This observation is probably due to the formation of an 424 organic film increasing lubrication between the particles after the addition of lignosulfonate 425 [9]. An increase in dosage led to thicker films, reducing the bearing capacity of the LS treated 426 soil by facilitating the deformation under loading. As for ES, addition of the product modified 427 the behaviour of the soil during the compaction. Nevertheless, field and laboratory testing 428 confirmed the ability of lignosulfonate to be leached out of the soil [11, 22, 27]. So, it is to be 429 expected that the soil will recover its initial characteristics after a certain period of time. The 430 uses of LS treatment are similar to those of ES but higher dosages are needed and attention 431 must be paid on bearing capacity which is altered by the treatment.

432 **7. CONCLUSION**

433 This paper presented an experimental multi-scale study of a low-plasticity soil treated with 434 three non-traditional organic products classified as an acid solution, an enzymatic solution 435 and lignosulfonate. Hydro-mechanical investigations including compaction, bearing capacity, 436 UCS, free swelling and hydraulic conductivity tests were conducted. The effect of the 437 treatments on the microstructure of the compacted soil was investigated using SEM and MIP 438 tests. The study also proposed a reflexion on the modification of the interactions between soil 439 particles induced by the treatments. The following conclusions can be drawn based on the 440 obtained results.

To evaluate the performances of non-traditional products used for soil treatment, it is
important to consider the entire compaction and bearing capacity curves and not only
the values obtained at the optimal conditions. In fact, the water content of compacted
soil has a strong influence on the dry unit weight obtained after treatment.

The ES and LS treatments reduced the optimal water content and increased the dry
 density of the compacted soil allowing the use of 3% lower water content and a 25%

reduction in the compaction energy, required to achieve the same density when
compared to untreated soil. Therefore, the treatments can improve the environmental
impact of earthwork projects and are particularly well suited for dry soil compaction in
areas where the access to water resources is limited.

Before in situ applications, careful laboratory investigations should be conducted to
determine if the treatment genuinely improves the soil behaviour; some nontraditional products have little or no impact on the mechanical properties of treated
soils. The changes in compaction parameters must be characterised to determine the
optimal application conditions in terms of dosage, desired water content and bearing
capacity.

Treatments with organic products have showed immediate modification of the compaction parameters; however, no evolution of the resistance of the soil over time was observed. Thus, treatments essentially act through a physical modification of the soil behaviour. Moreover, as AS and ES were added at low dosages and were biodegradable, intrinsic characteristics of the soil are not changed and the long term behaviour of the soil should not be modified. For LS treatment, leaching of the additive can be expected in case of water transfer in the compacted structure.

Some non-traditional products act like surfactants. This property appears to be an
important parameter of the effectiveness of non-traditional products when they are
used as compaction aids.

467 ACKNOWLEDGMENTS

The research presented in this paper was funded by the French Environment and Energy
Management Agency (ADEME), Egis Géotechnique and DTP Terrassement.

470

471 **REFERENCES**

- 472 1. ASTM D 698-91 (1991) Test method for laboratory compaction characteristics of soil
 473 using standard effort.
- 474 2. ASTM D 1883-92 (1992) Standard test method for CBR (Californian Bearing Ratio) of
 475 laboratory-compacted soils.
- 476 3. ASTM D 5298-94 (1995) Standard test method for measurement of soil potential
 477 (suction) using filter paper.
- 478
 4. Berney ES IV, Peters JF, Newman JK, Smith DM (2003) Effect of surfactant on the
 479
 dry-side compaction of silty sand. J Transp Res Board 1819:57-62.
- 480 5. Bolander P, (1999) Laboratory testing of nontraditional additives for stabilization of
 481 roads and trail surfaces. Transp Res Record: J Transp Res Board 1652:24–31.
- 482 6. Chen J, Anandarajah A, Inyang H (2000) Pore fluid properties and compressibility of
 483 kaolinite. J Geotech and Geoenvironmental Eng 126(9):798-807
- 484
 7. Di Maio C, Santoli L, Schiavone P (2004) Volume change behaviour of clays: the
 485 influence of mineral composition, pore fluid composition and stress state. Mech of
 486 Mater 36:435-451
- 487 8. Dong X, Wang Y-H (2008) The effects of the pH-influenced structure on the dielectric
 488 properties of kaolinite-water mixtures. Soil Sci Soc Am J 72:1532-1541
- 489 9. Du Noüy PL, (1919) A new apparatus for measuring surface tension. J Gen
 490 Physiology, 1(5):521-524

- 491 10. Gow AJ, Davidson DT, Sheeler JB (1961) Relative effects of chlorides,
 492 lignosulfonates and molasses on properties of a soil-aggregate mix. Highw Res board
 493 Bul, 282:66–83
- 494 11. Harris P, Holdt Jv, Sebesta S, Scullion T (2006) Recommendations for stabilization of
 495 high-sulfate soils in Texas. Transp Res Record: J Transp Res Board 1952:71–79
- 496 12. ISO 10390 (2005) Soil quality Determination of pH
- 497 13. Katz LE, Rauch AF, Liljestrand HM, Harmon JS, Shaw KS, Albers H (2001)
 498 Mechanisms of soil stabilization with liquid ionic stabilizer. Transp Res Record: J
 499 Transp Res Board 1757:50–57
- 500 14. Lambe TW (1956) Improvement of soil properties with dispersants. Boston Soc Civil
 501 Eng J, 41(2):184-207
- 502 15. Mitchell JK (1993) Fundamentals of soil behavior. John Wiley & Sons, New York
- 503 16. Palomino AM, Santamarina C (2005) Fabric map for kaolinite: effects of pH and ionic
 504 concentration on behavior. Clays and Clay Minerals 53(3):211-223
- 505 17. Park J, Vipulanandan C, Kim J, Myoung Hak Oh (2006) Effects of surfactants and 506 electrolyte solutions on the properties of soil. Environ Geology 49(7):213-220
- 507 18. Parsons RL, Milburn JP (2003) Engineering Behavior of Stabilized Soils. Transp Res
 508 Record : J Transp Res Board 1837:20–29
- 19. Rajendran D, Lytton RL (1997) Reduction of sulfate swell in expansive clay subgrade
 in the Dallas district. Res Rep TX-98/3929-1, Texas Transportation Institute, College
 Station

- 512 20. Rauch AF, Harmon JS, Katz LE, HM Liljestrand (2002) Measured effects of liquid soil
 513 stabilizers on engineering properties of clay. Transp Res Record: J Transp Res Board
 514 1787:33–41
- 515 21. Rauch AF, Katz LE, Liljestrand HM (2003) An analysis of the mechanisms and
 516 efficacy of three liquid chemical soil stabilizers: volume 1. Rep FHWA/TX-03/1993-1,
 517 Center for transportation research, University of Texas, Austin
- 518 22. Sanders TG, Addo JQ (1993) Effectiveness and environmental impact of road dust
 519 suppressants. Dep of Civ Eng, Colorado State University.
- 520 23. Santoni RL, Tingle JS, Webster SL (2002) Stabilization of silty sand with 521 nontraditional additives. Transp Res Record: J Transp Res Board, 1787:61–70
- 522 24. Scholen DE (1995) Stabilizer mechanisms in nonstandard stabilizers. Conf Proc 6: 6th
 523 Int Conf on Low-Volume Roads, Vol.2, TRB Natl Res Council, Washington, D.C.,
 524 pp. 252–260
- 525 25. Söjström E., (1993), "Wood chemistry, fundamentals and applications" second ed, 526 293 p.
- 527 26. Sokolovich VE (1973) Acid as chemical stabilizers of clay soils. Scientific Research
 528 Institute of Foundations, 4:22-23
- 529 27. Surdahl RW, Woll JH, Marquez HR (2007) Stabilization and dust control at the
 530 Buenos Aires national wildlife refuge, Arizona. Transp Res Record: J Transp Res
 531 Board 1989:312 321
- 532 28. Tingle JS, Santoni RL (2003) Stabilization of clay soils with nontraditional additives.
 533 Transp Res Record: J Transp Res Board 1819:72–84

- 534 29. Tingle JS, Larson SL, Weiss CA, Newman JK, Peters JF, Tardy B, Berney ES (2004)
 535 Constitutive analyses of nontraditional stabilization additives. Technical Report ERDC
 536 TR-04-5, US Army Engineer Research and Development Center, Vicksburg
- 30. Tingle JS, Newman JK, Larson SL, Weiss CA, Rushing JF (2007) Stabilization
 mechanisms of nontraditional additives. Transp Res Record: J Transp Res Board
 1989:59–67
- 540 31. Trufanova MV, Parfenova LN, Yarygina ON (2010) Surfactant properties of
 541 lignosulfonates. Russ J Appl Chem 83(6):1096–1098
- 542 32. Velasquez RA, Marasteanu MO, Holzalski RM (2006) Investigation of the 543 effectiveness and mechanisms of enzyme products for subgrade stabilization. Int J 544 Pavement Eng 7(3):213–220
- 545 33. Vinod JS, Indraratna B, Mahamud MAA (2010) Stabilisation of an erodible soil using
 546 a chemical admixture. Proc Inst Civil Eng: Ground Improvement 163:43-51
- 547 34. Visser AT (2007) Procedure for evaluating stabilization of road materials with 548 nontraditional stabilizers. J Transp Res Board 1989:21–26
- 549 35. Wieland E, Stumm W (1992) Dissolution kinetics of kaolinite in acidic solutions at
 550 25°C. Geochimica et Cosmochimica Acta 56:3339-3355
- 36. Yeoh NS, Oades JM (1981) Properties of soils and clays after acid treatment. I Clay
 Minerals. Aust J Soil Res 1881(19):147-158

- **Tab. 1** Dosages tested for the three non-traditional additives

| Non-traditional additives | Dosages by dry weight of soil |
|-----------------------------|-------------------------------|
| Acid solution (AS) | 0.01% |
| Enzymatic solution (ES) | 0.002% |
| Calcium lignosulfonate (LS) | 0.5 ; 2.0 and 5.0% |

Tab. 2 Properties of the tested soil

| Property | Low plastic clay |
|-------------------|------------------|
| Passing 2 mm (%) | 100 |
| Passing 80 µm (%) | 87 |
| Passing 2 µm (%) | 19 |
| Specific gravity | 2.67 |
| Liquid limit (%) | 34 |
| Plastic limit (%) | 20 |
| Plasticity index | 14 |
| рН | 9.2 |

Tab. 3 Index properties and pH of the treated soil

| Treatment | рН (+/- 0.1) | PL (%) | LL (%) | PI |
|-----------|-----------------|--------|--------|----|
| Untreated | 9.2 | 20 | 34 | 14 |
| 0.01% AS | 9.2 | 19 | 33 | 14 |
| 0.002% ES | 9.3 | 19 | 33 | 14 |
| 0.5% LS | 8.8 | 19 | 32 | 13 |
| 2.0% LS | 8.4 | 17 | 30 | 13 |
| 5.0% LS | 8.1 | 13 | 27 | 14 |
| 0.01% SDS | 9.1 | 19 | 34 | 15 |

Tab. 4 Geotechnical properties of the treated soil

| Treatment | West (%) | (kN/m^3) | IPI at w | UCS (MPa) | |
|-----------|-----------|------------|-----------|-----------|---------|
| nouthone | wopi (70) | | πιατινορι | 7 days | 28 days |
| Untreated | 15.5 | 18.2 | 15 | 0.44 | 0.46 |
| 0.01% AS | 15.0 | 18.3 | 19 | 0.40 | 0.41 |
| 0.002% ES | 14.5 | 18.6 | 14 | 0.53 | 0.54 |
| 0.5% LS | 15.0 | 18.4 | 13 | 0.36 | 0.40 |
| 2.0% LS | 13.5 | 18.5 | 14 | 0.45 | 0.49 |
| 5.0% LS | 14.0 | 18.3 | 5 | 0.22 | 0.24 |

Tab. 5 Deformation modulus at 50% of maximal strength

| | | | | Standard |
|--------------------|-------|-----------------------------------|-----------------------|-----------|
| Treatment | w (%) | γ_{d} (kN/m ³) | E ₅₀ (MPa) | deviation |
| | | | | (MPa) |
| | | | | |
| L Instructure of a | 15.0 | 18.2 | 13 | 2 |
| Unirealed | | | | |
| | 12.5 | 18.4 | 56 | 7 |
| | 14.0 | 18.5 | 15 | 1 |
| 0.002% ES | | | | |
| | 12.5 | 18.3 | 50 | 4 |
| | | | | |
| 2.0% LS | 12.7 | 18.5 | 22 | 2 |

- Tab. 6 Total void ratio for the compacted samples and calculated mercury investigated
 maximum void ratio during the MIP tests

| Treatment | e _{tot} | einv | γ _d (kN/m³) |
|-----------|------------------|------|------------------------|
| Untreated | 0.47 | 0.44 | 18.3 |
| 0.01% AS | 0.46 | 0.44 | 18.3 |
| 0.002% ES | 0.43 | 0.42 | 18.7 |
| 2.0% LS | 0.44 | 0.41 | 18.6 |

Tab. 7 Matrix suction of treated soil

| Treatment | w (%) | γ _d (kN/m³) | Matrix suction (kPa) |
|-----------|-------|------------------------|-------------------------|
| Untreated | 14.7 | 18.0 | 56 |
| 0.01% AS | 15.4 | 18.0 | 54 |
| 0.002% ES | 14.7 | 18.2 | 58 |
| 2.0% LS | 15.2 | 18.2 | 225 |



2 Fig. 1 Compaction and IPI curves for the untreated, 0.01% AS and 0.002% ES treated soil



7 Fig. 2 Compaction and IPI curves for the LS treatment at 0.5, 2.0 and 5.0%



Fig. 3 Compaction and IPI curves of the ES treatment at 75 and 100% of the standard
Proctor energy



Fig. 4 Compaction and IPI curves of the 2.0% LS treatment at 75% and 100% of thestandard Proctor energy



Fig. 5 Unconfined compressive strength at 7 and 28 days for the studied non-traditional
 treatments compacted at their respective optimum Proctor



Fig. 6 One-dimensional free swelling potential for the ES and LS treatments at different initialcompaction states



Fig. 7 Hydraulic conductivity coefficient for the ES and LS treatments in different initial
 compaction states



40 Fig. 8 SEM image of the untreated soil



- **Fig. 9** SEM image of the 0.01% AS treated soil



- 49 Fig. 10 SEM image of the 0.002% ES treated soil



55 Fig. 11 SEM image of the 2% LS treated soil







Fig. 13 Pore size distribution for the untreated, 0.01% AS, 0.002% ES and 2.0% LS treated

65 soil



69 Fig. 14 Surface tension of the three non-traditional additives and SDS in aqueous solution



74 Fig. 15 Compaction and IPI curves for 0.01% SDS and 0.002% ES treated soil