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ON THE MECHANISMS OF CONSUMPTION OF CALCIUM LIGNOSULFONATE BY CEMENT PASTE

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7 ABSTRACT

8 The aim of this paper is to assess the mechanisms of consumption of softwood calcium lignosulfonate 9 (LSs) by cement paste. The LSs consumption by two different cements (CX and ANL) and two 10 reference materials (CaCO₃ and Ca(OH)₂) was investigated, either by adding the LSs immediately 11 with the mixing water (IA) or after 10 minutes of hydration (DA). For IA, the increase in LSs dosage 12 caused additional ettringite formation and an increase in particle surface area. This was not observed 13 for DA. Since no AFm phase could be detected, intercalation in AFm seemed not to occur for the 14 investigated materials. The main mechanism of LSs consumption for CX cement (both for IA and DA) 15 and for ANL cement (only for DA) appeared to be monolayer adsorption. For IA, the amount of 16 consumed LSs could not be ascribed exclusively to monolayer surface adsorption and other LSs 17 consumption mechanisms might play a role.

18 1. INTRODUCTION

Water-reducers, or plasticizers, are commonly used as admixture for concrete. Their addition to fresh concrete allows obtaining highly fluid concrete at low water-binder ratios, improving the mechanical properties of the hardened concrete. To optimize the polymer-cement combination and the amount of admixture needed to achieve the desired workability, it is important to understand the mechanisms of consumption of plasticizer by cement paste. 24 The plasticizer investigated in this paper is a low-sugar softwood calcium lignosulfonate (LSs), 25 commonly used in concrete in the dosage of 0.25-0.40 mass % of binder. Lignosulfonates are 26 polyelectrolytes derived from lignins from pulping industry. The lignins are fragmented and 27 sulfonated, thereby becoming water-soluble. Lignin can be derived both from softwood and hardwood trees, which results in lignosulfonates with different molecular weight and amount of molecular 28 29 functional groups (carboxyl groups, phenolic-OH, sulfonic groups). Lignosulfonate is known to have 30 medium retarding effect on cement hydration. The sugars naturally contained in lignin remain in 31 lignosulfonate after its production. These sugars contribute to longer setting times of cement, in 32 particular the hexoses, which can be removed by fermentation in low-sugar lignosulfonates. However, 33 studies in literature found that also sugar-free lignosulfonates exhibited pronounced retardation of 34 cement paste hydration, e.g. [1, 2].

Plasticizers interact with unhydrated and hydrated cement grains, as summarized in a recent literature review by Marchon and Flatt [3]. In this paper, the amount of polymer uptaken by the cement paste is defined as "consumed" as opposed to the free one dissolved in the pore solution. The mechanisms of polymer consumption will be separately discussed in this paragraph.

39 The dispersing effectiveness of a superplasticizer on cementitious materials is, amongst others, a 40 function of its degree of adsorption on the surface of cement grains and hydrates. The adsorbed 41 plasticizer layer renders the total particle surface negatively charged, i.e. with a negative zeta potential. 42 As negatively charged particles approach each other there will be an electrostatic repulsion preventing 43 them from forming agglomerates. Additionally, when two surfaces approach enough for their adsorbed 44 layers to overlap, a steric force develops. This will contribute in hindering particles to get close 45 enough to form agglomerates. The key parameters that govern the steric repulsion are the adsorption layer thickness and its conformation at the solid liquid interface [4]. 46

The polymer will not be adsorbed equally on the four main cement phases. According to Yoshioka et al. [5], much higher adsorption occurs on aluminate and ferrite than on the silicate phases. The amount of adsorbed polymer on ettringite was found to be the largest amongst the cement hydrates by Zingg et al. [6]. It must be noted that, in both the cited references, the results were reported by unit of mass andnot by unit of specific surface.

Polymer adsorption can also take place in multiple layers on cement particles and hydrates. After ideal monolayer coverage, the cement particles will have a negative surface charge. Ca^{2+} ions will then be electrostatically attracted to the negatively charged groups of the polymer and they will bond with them. This Ca^{2+} outer layer will allow the adsorption of a further layer of negatively charged polymer [7, 8], facilitating additional consumption of polymers.

57 Adsorption onto the surface of cement particles and hydrates is not the only potential consumption 58 mechanism taking place when a plasticizer is added to a cementitious system. Part of the water-59 reducing admixture might also be intercalated in the hydration products, mainly in the layered 60 structure of AFm, and part of the admixture will remain dissolved in the aqueous phase, according to, amongst others, [9-12]. When tricalcium aluminate (C_3A) enters in contact with water, it reacts 61 immediately forming, in absence of gypsum, the metastable layered phases C_4AH_{19} and C_2AH_8 . In 62 presence of gypsum, C₃A will react with water forming $C_6A\overline{S}_3H_{32}$ (ettringite) and $C_4A\overline{S}H_{12}$ 63 (monosulphate). C_4AH_{19} , C_2AH_8 and $C_4A\overline{S}H_{12}$ belong to the group of layered double hydroxides 64 (LDHs). Several anions and polyelectrolytes can intercalate in between the cationic layers of LDH 65 compounds by replacing their hydroxyl ions. According to Plank et al. [10], intercalation was found to 66 67 be possible for polymers with different structure, namely, linear, comb-like and polymer brushes with very long side chains. The polymer intercalated in LDHs will no longer be available for dispersing 68 69 cement particles; therefore a higher dosage of polymer will be necessary to reach the desired 70 workability.

Another possible mechanism of polymer-cement interaction is complexation between functional groups of the plasticizer and calcium ions dissolved in the pore solution, as observed in several studies [13-16]. Collins et al. [13] found that calcium ions dissolved in a calcium lignosulfonate solution with pH over 10-11 can hold together a matrix of lignosulfonate molecules, forming a gel. Other cations, e.g. aluminium and iron, could also tightly bind to LS molecules. As mentioned by Sowoidnich et al. 76 [15], the interaction between calcium ions and polymer functional groups (mainly sulfonic and 77 carboxyl groups) can be divided into complexation of calcium ions in aqueous solution, complexation 78 of calcium ions on particles surface (adsorption) and formation of polymer-containing clusters. 79 Formation of Ca-polymer complexes will increase the amount of consumed polymer if they are 80 precipitated or form colloids that are filtered away when collecting pore solution, and decrease the 81 amount of free Ca^{2+} ions in the pore solution, lowering the Ca-Si ratio of the pore solution, hence 82 possibly modifying the hydration reactions and the resulting hydrates, as stated by Yousuf et al. [17]. 83 The polymer molecules captured in the complexes with calcium ions might still have some free anionic functional groups on their outer regions. These anionic functional groups might as well be 84 85 attracted to the positive charged calcium ions adsorbed on the polymer layer over cement particles and hydrates. The calcium-polymer complexes might then bind to the cement particles already covered 86 with polymer, decreasing the amount of free LSs in the pore solution and forming multiple layers of 87 polymer adsorbed. For this reason the mechanisms of calcium complexation and multilayer adsorption 88 89 can be considered interrelated and, sometimes, undistinguishable from each other.

90 The subject of this paper is to investigate the mechanisms consuming a low-sugar softwood calcium 91 lignosulfonate (LSs) in paste of two Portland cements with different surface area and C₃A content. The 92 effects were studied both by adding the lignosulfonate immediately with the mixing water (IA) and by 93 adding it after 10 minutes of hydration (DA). The amount of polymer consumed by the cement paste 94 was determined by UV-spectrometry and adsorption isotherms were calculated. Comparison of 95 adsorption isotherms onto cement and reference materials representative of cement but which do not hydrate (CaCO₃ and Ca(OH)₂), allow a better elucidation of the LSs consumption mechanisms in 96 97 simpler systems than cement. Changes in the surface area of the hydrated cement particles were 98 investigated by BET. The changes in composition and amount of cement hydrates caused by the 99 addition of lignosulfonate were investigated with thermogravimetric analysis (TGA).

100 2. EXPERIMENTAL

101 2.1 Materials

102 The mechanisms of plasticizer consumption by cement paste were studied adding a low-sugar 103 softwood calcium lignosulfonate (LSs) to two Portland cements: a CEM I 52.5 N (ANL) and a CEM I 104 52.5 R (CX). The two cements were chosen because of their different surface area and C_3A content. 105 The content of the main clinker phases of the cements quantified by XRD Rietveld are given in Table 106 1. The chemical composition of the cements and the loss of ignition at 950 °C determined by XRF are 107 reported in Table 2. The particle size distribution (d_{10} , d_{50} , d_{90}), Blaine surface area and density, BET 108 surface area are given in Table 3.

109 A sugar-reduced softwood calcium lignosulfonate (LSs) was used as plasticizer. Its mass weighted 110 molecular weight (M_w), as measured with gel permeation chromatography (GPC), was 29000 g/mol 111 and the number weighted molecular weight (M_n) was 2100 g/mol, giving broad molar-mass dispersity 112 $(\mathfrak{D}_{\mathsf{M}})$ equal to 13.8. The molar-mass dispersity, also called polydispersity index, is defined as the ratio 113 between M_w and M_n [18]. Additional physical and chemical properties of the lignosulfonate are listed 114 in Table 4. For the lignosulfonate used in the present investigation, the sugars were removed from the 115 polymer molecule by fermentation and resulting alcohol by distillation. The LSs was dissolved in 116 deionised water to concentrations varying from 1 to 45 % to ease dosing, and the water content was 117 included in the calculation of the water-to-binder ratio (w/b).

118 CaCO₃ and technical-grade precipitated $Ca(OH)_2$ were also mixed with lignosulfonate in order to 119 study some simplified model systems. Their specific surface areas determined by BET are reported in 120 Table 5. In order to mimic the basic pH of cement paste, these samples were mixed with LSs solution 121 diluted in artificial pore water. The artificial pore water was a solution of NaOH and KOH with K/Na 122 molar ratio equal to 2 and measured pH of 12.9.

123 **2.2 Sample preparation**

124 **2.2.1 Portland cements**

125 Cement was mixed with distilled water and/or lignosulfonate solution in a high-shear mixer MR530 by 126 Braun at intensity 6 obtaining pastes with w/b = 0.4. About 200 ml of cement paste was mixed per 127 batch. In order to investigate the effect of the time of addition of lignosulfonate, two different mixing 128 procedures were compared: immediate addition of LSs with the mixing water (IA) and delayed 129 addition of LSs at 10 minutes of hydration (DA).

For IA, the binder was mixed with distilled water (and/or lignosulfonate diluted in distilled water or artificial pore water) according to the procedure used by Vikan [19]: 30 seconds mixing and scraping the walls of the mixer to homogenize the mix, 5 minutes resting and 1 minute mixing.

For DA, the binder and 85% of the needed water were mixed according to the following mixing procedure: 30 seconds mixing and scraping the mixer walls to homogenize the mix, 10 min resting (delay time chosen according several studies in literature [20-23]). LSs and the remaining 15% of the needed water were then added to the mix which was mixed for 1 minute.

After mixing, about 35 ml of paste was poured in 50 ml sealed plastic centrifuge tubes and let restuntil the chosen analysis time.

139 **2.2.2 Calcium carbonate**

140 Calcium carbonate has been shown to be a suitable model system for investigating stability and 141 rheology of cement paste. Mikanovic et al. [24] showed that calcium carbonate exhibits colloidal 142 properties very similar to those of cement paste at early ages (hydration < 1 hour). CaCO₃ exhibit 143 surface properties and flocculation behavior similar to that of cement paste, namely, an irregular 144 spheroid shape, and a low surface charge in water. In addition it has a very low solubility, also at high 145 pH. In addition, this material was expected not to noticeably react with water.

About 300 g of $CaCO_3$ was mixed with about 90 g of LSs solution dissolved in artificial pore water in a high-shear mixer MR530 by Braun at intensity 6. The mixing procedure was identical to the one used for neat cement with IA. The water-solid ratio by mass was 0.3. Several LSs dosages wereanalysed, as reported in Table 6. The samples were let to rest for 30 minutes prior to being analysed.

150 2.2.3 Calcium hydroxide

151 $Ca(OH)_2$ is one of the main cement hydrates and its solubility is about 100 times higher than the one of 152 $CaCO_3$, so it appears to be useful to investigate the possible interaction between the LSs and calcium 153 ions. About 5 g of Ca(OH)_2 was mixed with about 40 g of LSs solution dissolved in artificial pore 154 water in plastic centrifuge tubes and mixed by hand for 1 minute. The high fineness of the Ca(OH)_2 155 powder required a water-solid ratio by mass of 8.0. The mixing solution contained increasing amounts 156 of LSs, as reported in Table 6. All the samples were let to rest for 30 minutes prior to being analysed.

157 **2.3 Methods**

158 **2.3.1** Adsorption isotherms

Polymer adsorption by a solid is usually described through isotherms, in which the amount of polymer adsorbed is plotted against the total amount of polymer added to the system [25]. The shape of an isotherm is largely determined by the adsorption mechanism. In this study, the isotherms were drawn relating the amount of LSs consumed by the cement paste to the amount of total LSs added to the sample.

With the help of a calibration curve, achieved by measuring the UV absorbance of pure LSs solutions in artificial pore water at different concentrations, the amount of free plasticizer (g LS/100 g solution) was calculated. This amount was related to the amount of binder in the sample (g LS/100 g binder). The LSs consumed by the investigated systems was then calculated by subtracting the amount of free LSs from the total amount of LSs initially added to the sample, as displayed in the following equation:

$$169 \quad \text{consumed LSs} = \text{total LSs} - \text{free LSs} \tag{1}$$

170 The absorbance of the pore solution was measured with UV-spectroscopy. In order to confirm the 171 results obtained with UV-spectroscopy, the adsorption isotherm of ANL cement for IA was measured 172 also with total organic carbon analysis (TOC). Potential removal of polymer aggregates by filtration was eliminated as error source, by comparing TOC analysis of filtered and un-filtered samples whichwere found to be very similar

175 **2.3.1.1 UV-spectroscopy**

176 UV-spectrometry allowed measuring the absorbance of the pore solution at increasing LSs dosage.177 The LSs dosages tested are summarized in Table 6.

178 The pore solution was extracted from the cement paste by centrifuging the samples in a Heraeus 179 Megafuge 8 centrifuge by Thermo Scientific for 3 minutes at the speed of 4500 rpm. The supernatant 180 pore solution was extracted and filtered with 0.45 µm cellulose acetate syringe filters by VWR. The 181 amount of free LSs in the pore water was measured with UV-spectrometry with a Genesys 10S UV-182 spectrophotometer by Thermo Scientific. Several wavelengths have been reported in literature to study 183 the amount of lignosulfonate in pore solution: Perche [26] and Ratinac [27] used 280 nm, Uchikawa et 184 al. [28] and Houst at al. [4] used 284 nm, Vikan [19] used 283 nm. Samples diluted 1:100 with distilled water were scanned with different wavelengths from 190 to 300 nm using distilled water as 185 blank reference sample. For the plasticizer used in this study, 281 nm was chosen as the best 186 187 wavelength to measure the absorbance value.

The amount of plasticizer consumed by cement paste as a function of increasing hydration time was determined by centrifugation of pastes aged for different times (5-120 minutes). As displayed in Figure 1, it was found that at 10 minutes of hydration the LSs uptake reached an equilibrium value. All the samples were then analysed aftert 30 minutes of hydration.

192 **2.3.1.2 Total organic carbon (TOC)**

193 The concentration of free polymer in the pore solution extracted by ANL cement (0.2; 0.4; 0.8; 1.0; 194 1.5 mass % of binder LSs IA) was measured with the total organic carbon analysis (TOC). The TOC 195 analysis was performed using a Vario TOC Cube by Elementar. The extracted pore solution was 196 filtered with 0.20 μ m cellulose acetate filters. Part of the sample was acidified with 2 drops of 197 concentrated HCl to prevent any formation of precipitates in the solution. The amount of consumed 198 LSs was measured by TOC on the same sample before and after acidification. Acidification did not lead to any variation in the results. No notable difference in the results obtained with UV-spectroscopy and with TOC was displayed up to a LSs dosage of 0.6 mass % of binder. Over this dosage, the consumed LSs from TOC measurements was from 3 to 15 % higher than the one measured with UVspectroscopy. The difference might be due to the differences in sample preparation (different sample dilution, different filter used) and the measurement techniques. However, the results obtained with the two different techniques showed similar trends.

205 2.3.2 Solvent exchange

A solvent exchange procedure with isopropanol was used to stop the hydration of the cement paste after 30 minutes of hydration. The samples were then analysed with thermogravimetric analysis (TGA) and BET.

209 About 5 ml of cement paste was transferred in a 50 ml centrifuge tube and centrifuged for 1 minute at 210 2000 rpm. The supernatant water was removed. About 40 ml of isopropanol was poured in the 211 centrifuge tube. The tube was shaken for 30 seconds and let to rest for 5 minutes. The sample was 212 centrifuged again for 1 minute at 2000 rpm and the supernatant liquid was removed. The solvent 213 exchange procedure with isopropanol was repeated once, followed by a final solvent exchange with 10 214 ml of petroleum ether. The resulting paste was let to dry for 2 days in a desiccator over silica gel, and 215 soda lime to minimize carbonation. After drying, the samples were homogenized in a porcelain mortar 216 and stored in sealed containers in a desiccator over silica gel and soda lime until analysis.

217 **2.3.3 Thermogravimetric analysis (TGA)**

The thermogravimetric analysis (TGA) was performed with a Mettler Toledo TGA DSC3+ on hydrated cement paste after stopping the hydration using solvent exchange. Approximately 200 mg of cement paste powder was loaded in 600 μ l alumina crucibles. The samples were heated from 40 to 900°C at a rate of 10°C/min while purging with 50 ml/min N₂.

222 2.3.4 Characterization of surface area of hydrated cement pastes by BET

The BET measurements were performed using a Tristar II Plus by Micromeritics on samples of which the hydration was stopped with the solvent exchange procedure. Before the measurement, the samples were degassed for about 5 minutes at room temperature. The measurement was performed purging the samples with nitrogen at room temperature, which took about 10 minutes. The sample mass was about 2 g. The samples did not undergo any thermal treatment before the measurement to avoid any possible destruction of ettringite, as recommended by Mantellato et al. [29, 30].

229 3 RESULTS

230 **3.1 Adsorption isotherms**

The adsorption isotherms of ANL and CX cement pastes, CaCO3, and Ca(OH)2 were obtained by 231 232 measuring the amount of consumed polymer for an increasing dosage of polymer in the mix as described in Table 6. Higher LSs dosages were used for Ca(OH)₂ due to its very small particle size, 233 234 and thus larger specific surface. For CX cement it was not possible to extract pore water at LSs 235 dosages over 1.0 mass % of binder due to paste hardening after 30 minutes of hydration. The LSs was 236 added to the cement paste either immediately with the mixing water (IA), or after 10 minutes of 237 hydration (DA). The results are presented as consumed LS per mass % of binder in Figure 2a, and per m^2 of unhydrated substrate surface area available for adsorption in Figure 2b. The isotherms were 238 239 obtained by fitting the experimental data to the non-linear Langmuir model, according to the equation

240 described by Marchon et al. [31]:
$$m_{SA} = \frac{m_{SA}^{\infty} K c_A}{1+K c_A}$$
 (2)

241 Where m_{SA} is the adsorbed mass, m_{SA}^{∞} is the adsorbed mass at the plateau, K is a chemical equilibrium 242 constant, and c_A is the concentration in solution.

As reported in [31], the Langmuir model is not ideal for a system like cement paste, which surface area changes with hydration and where the polymer is not adsorbed equally on all cement phases. In addition, in the present paper, the Langmuir equation was used to fit results plotted as amount of consumed polymer versus dosage of added polymer, rather than versus the equilibrium concentration of polymer remaining in solution. Therefore, its K-value does not bear any physical meaning, and the fit can only be considered as a visual guide for the eye. The adsorption isotherm of $Ca(OH)_2$ is omitted in Figure 2 a because of the higher LSs dosages used. It must be noted that the water-solid ratio by mass was 0.3 for $CaCO_3$ and 8.0 for $Ca(OH)_2$, while it was 0.4 for the neat cements.

252 The results were also presented in Figure 3 as consumed LSs amount (% of the LSs amount added) 253 versus the LSs amount added. Figure 3 shows that all the adsorption isotherms for the cements 254 displayed similar LSs consumption at low LSs dosage (up to about 0.25 mass % of binder LSs). At 255 these low LSs dosages, about 75 % of the LSs added was consumed both for IA and for DA. At LSs 256 dosages higher than about 0.25 mass % of binder, the curves obtained for IA showed a LSs 257 consumption of about 70 % of the LSs added. The consumption kept constant with increasing LSs 258 dosages. For DA, the amount of LSs consumed decreased from about 75 % to about 30 % of the LSs 259 added when the LSs dosage increased from 0.25 to 1.5 mass % of binder.

As displayed in Figure 2, for IA, no adsorption plateau could be detected within the tested range, neither for ANL nor for CX cement. This has also been observed by others, e.g. by Vikan [19] and Ratinac et el. [27]. The isotherms' shape indicated a continuous LSs uptake when more LSs was added to the mix.

For DA, an adsorption plateau was found for both cements. The isotherms that reached an adsorption plateau also displayed a considerably lower amount of LSs consumed by the cement pastes compared to those that did not reach any plateau.

267 **3.2** Surface area of hydrated cement particles

The BET surface area was measured for ANL and CX cement pastes with increasing LSs amounts hydrated for 30 minutes. The hydration was stopped by solvent exchange after 30 minutes. The results and their linear fit are shown in Figure 5.

For both cements the surface area after 30 minutes of hydration was found to increase as the dosage of
plasticizer added to the cement paste increased. The increase in surface area was remarkably larger for
CX than for ANL cement, and for IA compared to DA.

274 New adsorption isotherms were calculated dividing the amount of consumed LSs by the actual surface 275 area of ANL and CX cements after 30 minutes of hydration as measured with BET for both IA and 276 DA. The isotherms were obtained by fitting the experimental data to the non-linear Langmuir model, 277 as described in paragraph 3.1. The results are displayed in Figure 6. For CX cement, the isotherms 278 obtained with IA and DA nearly coincide, both reaching an adsorption plateau. On the contrary, for 279 ANL cement, even when expressed relative to the hydrated surface area, the adsorption isotherms 280 remain qualitatively similar to those obtained considering the unhydrated surface area.

3.3 281

LSs molecular footprint

282 As displayed in Figure 6, for ANL cement, the adsorption plateau was reached for a total LSs amount 283 between 0.8 and 1.2 mass % of binder, which the authors assume to be due to the achievement of 284 monolayer surface coverage. The amount of LSs consumed at the achievement of the plateau was considered equal to the value m_{SA}^{∞} obtained with a non-linear regression analysis based on equation 2. 285

286 Knowing the molecular weight of the LSs molecule (29000 g/mol) and the specific surface area of hydrated ANL cement as measured with BET (174 m²/100g cement with 0.8 mass % of binder LSs, 287 and 184 m²/100g cement with 1.2 mass % of binder LSs), it is possible to calculate the LSs "molecular 288 289 footprint". The error connected to the molecular footprint was calculated considering an estimated 290 error or 30% for the measurement of the surface area with BET and of the LSs molecular weight, and the standard error of m_{SA}^{∞} . The LSs molecular footprint resulted to be about 20 ± 10 nm² for ANL 291 292 cement for both LSs dosages, which mostly agrees to the data given by the producer for the same LSs 293 $(50 \pm 30 \text{ nm}^2)$. The producer calculated this value from the plateau values of the adsorption isotherms 294 of LSs on MgO at alkaline pH.

295 For CX cement, the adsorption plateau was reached for a total LSs amount between 1.2 and 1.5 mass % of binder. Knowing the specific surface area of hydrated CX cement as measured with BET (286 296 $m^2/100g$ cement with 1.2 mass % of binder LSs, and 301 $m^2/100g$ cement with 1.5 mass % of binder 297 LSs), the LSs molecular footprint was calculated as about $20 \pm 10 \text{ nm}^2$ for both LSs dosages. This 298 299 value is equal to the one obtained for ANL cement.

300 In order to get a deeper understanding of the actual plasticizer consumption mechanisms by cement 301 paste, the adsorption isotherms determined with ANL or CX cements were compared to those obtained 302 for simplified model systems, CaCO₃ and Ca(OH)₂.

The CaCO₃ isotherm is displayed in Figure 2 a per mass % of dry powder. The CaCO₃ isotherm reached an adsorption plateau for a total LSs amount between 0.2 and 0.4 mass % of binder. At surface saturation, about 0.06 g of admixture was adsorbed on 100 g of CaCO₃. Since the specific surface area of the unhydrated CaCO₃ was measured with BET as 57 m²/100g, the surface coverage can be calculated as about 920 m²/g _{LS}. Hence, the molecular "footprint" (coverage) of LSs was calculated as about 60± 30 nm² for both LSs dosages. This result partly agrees with the data given by the producer (50 ± 30 nm²).

310 The Ca(OH)₂ isotherm, omitted in the figures due to the high LSs dosages used, reached an adsorption 311 plateau for a total LSs amount between 8.0 and 12.0 mass % of binder. The high LSs dosages at which 312 the plateau is reached is most likely due to the high specific surface area of the Ca(OH)₂ particles. At 313 surface saturation, about 0.06 g of admixture was adsorbed on 100 g of Ca(OH)₂. Since the specific surface area of the unhydrated $Ca(OH)_2$ was measured with BET as 1666 m²/100g, the molecular 314 "footprint" of LSs was calculated as about 40 ± 20 nm² for both LSs dosages. This value resulted lower 315 316 than the one obtained for CaCO₃, still being included in the range given by the producer (50 ± 30) nm^2). 317

The consumed LSs at plateau achievement and the molecular footprint calculated for the different materials can be found in Table 7. It has to be kept in mind that the LSs has a broad polydispersity index, which leads to a wide error in the molecular footprint given by the producer. The results reported in Table 7 showed that a relatively similar footprint was obtained for the model materials, i.e. $CaCO_3$ and $Ca(OH)_2$, and the real cements.

324 **3.4 Hydrates characterization**

In order to investigate the effects of LSs on hydrates formation, thermogravimetric curves were measured on ANL and CX cement pastes with 1.5 mass % of binder LSs added with IA and DA. A reference sample of ANL and CX pastes without LSs was also measured. The hydration of the tested samples was stopped after 30 minutes with the solvent exchange procedure using isopropanol and petroleum ether, as described in paragraph 2.3.2. The results are displayed in Figure 4 a, b.

330 Several peaks could be observed. According to Lotenbach et al. [32], amongst others, the peak at about 120 °C and the smaller one at 240 °C indicate the presence of ettringite (ettr.). The peak around 331 332 160 °C represents the decomposition of sulfates. The sulfates will be most likely gypsum (x = 2) for ANL cement. CX cement originally contains anhydrite (x = 0), which will not show any peak in the 333 334 TGA curve, since it does not contain water, and hemihydrate (x = 0.5). After 30 minutes of hydration, 335 hemihydrate will be partly or completely converted to gypsum. The peak around 160 °C in CX cement will then be due to the decomposition of gypsum and/or hemihydrate. The peak around 420 °C shows 336 337 the presence of portlandite (CH); the ones around 610 °C and 780 °C represent the decomposition of 338 carbonates (CO₂). The peaks over 500 °C can be attributed both to the decomposition of limestone 339 included in the used cements and, for the samples containing LSs, to the decomposition of LSs. No 340 peak ascribable to AFm phases was detected.

The results show that, for IA, the addition of LSs led to additional formation of ettringite and reduced amount of gypsum. For DA, in presence of LSs, no remarkable changes in the ettringite or gypsum amounts were observed. The LSs addition caused an increase in the weight loss in the carbonate region both for IA and DA. Similar trends were observed for both cements, even though remarkably more ettringite and fewer sulfates were measured for CX cement, which might be linked to the larger amount of C_3A in CX cement than in ANL cement.

348 4 DISCUSSION

The subject of this paper is to investigate the mechanisms consuming LSs in paste for two Portland cements, both for immediate and delayed addition of plasticizer. From literature the potential main mechanisms are: polymer intercalation into early cement hydrates; surface multilayer adsorption / complexation between functional groups of the plasticizer and calcium ions dissolved in cement pore solution; and monolayer surface adsorption of LSs on cement particles and hydrates. Each mechanism will be discussed separately in the following section.

4.1 Mechanisms of polymer consumption by the cement paste

356 4.1.1 Intercalation

According to, amongst others, Flatt and Houst [9], intercalation in AFm is generally considered to be the main cause of the difference between the adsorption isotherms obtained for IA or for DA. According to the theory, from the moment cement enters in contact with water, calcium aluminates start being consumed in AFt formation. In order for polymer intercalation to take place, calcium aluminates and polymer must be simultaneously available in solution. In case of DA, most hydrated aluminates have already been consumed in AFt formation at the time the polymer is added, hence the aluminates will no longer be available to form intercalated AFm.

364 Zingg et al. [6] formulated an alternative explanation for the difference in polymer consumption 365 between IA and DA, which does not include intercalation. They hypothesized that, for IA, due to the 366 dispersive effect of the plasticizer, numerous fine ettringite particles are floating in the pore solution, 367 providing additional particle surface area for adsorption. On the contrary, with DA, the ettringite 368 particles have already precipitated on the C₃S surface and cannot be redispersed. Hence, there will be 369 no increase in surface area and, consequently, in polymer adsorption. However, it must be kept in 370 mind that Zingg et al. did not investigate the amount of surface of ettringite which is made available 371 through a change of the initial hydration reactions by the addition of the polymer.

The adsorption isotherms presented in Figure 2a,b displayed a remarkable difference in LSs consumption between the isotherm obtained for IA and the one obtained for DA. However, no peak

374 corresponding to AFm phase could be detected with TGA, as shown in Figure 4a,b. For the materials
375 investigated in this paper, intercalation in AFm seems less likely as a LSs consumption mechanism for
376 the investigated materials and dosages, while the theory of Zingg et al. [6] appears more feasible.

377 4.1.2 Calcium complexation / multilayer surface adsorption

In the case that LSs would be consumed in calcium complexes, either in solution or as multilayer surface adsorption, the adsorption isotherms would display an increase in LSs consumption with the amount of total LSs added, even for LSs dosages over the surface saturation value, as calcium is buffered by the cement hydration.

382 In this study, the adsorption isotherms of both $CaCO_3$ and $Ca(OH)_2$ reached an adsorption plateau, as 383 shown in Figure 2 b. According to Lide [33], CaCO₃ has a solubility of 0.0014 g/100g in cold water, 384 while $Ca(OH)_2$ of 0.185 g/100g. Because of the negligible release of calcium ions by $CaCO_3$, calcium 385 complexation or multilayer adsorption with LSs molecules would be limited. The solubility of 386 $Ca(OH)_2$ is about 100 times higher than the one of $CaCO_3$. However, also for $Ca(OH)_2$ no calcium 387 complexation or multilayer adsorption with LSs molecules seemed to take place. Therefore, the LSs 388 consumed by both CaCO₃ and Ca(OH)₂ was most likely entirely due to monolayer adsorption of LSs 389 molecules on the particles surface.

390 Concerning the two cements, as shown in Figure 2a,b, the isotherm for both ANL and CX cement 391 reached an adsorption plateau for DA. As for CaCO₃, the main LSs consumption seems to be 392 monolayer surface adsorption. For IA, on the contrary, the isotherm of both the cements did not reach 393 an adsorption plateau, indicating a continuous polymer uptake by the cement paste the more polymer 394 is added. However, the authors assume that, as calcium complexation/multilayer adsorption was minor 395 or not existing for both CaCO₃ and Ca(OH)₂, and for the cements in case of DA, also for the cements 396 in case of IA calcium complexation/multilayer adsorption should not be a major mechanism behind 397 the LSs consumption.

398 4.1.3 Monolayer surface adsorption

In the case that LSs would adsorb as a monolayer onto the cement particles, the adsorption isotherms would display a plateau when the entire surface is covered by the polymer, as displayed in several studies [5, 19, 34, 35]. This mechanism, monolayer adsorption, can be described by the Langmuir model [31]. Since such a plateau is reached by both ANL and CX cements when the polymer was added with DA, and considering the above conjectures, it is likely that monolayer surface adsorption is the main mechanism of LSs consumption for DA.

No adsorption plateau was reached when the polymer was added with IA. This might be due to different reasons. First, it is possible that the increase in LSs consumption solely occurs due to the increase in particle surface area available for adsorption due to cement hydration, as displayed in Figure 5. In this case, the only mechanism of LSs consumption would be monolayer surface adsorption. Another possibility is that other consumption mechanisms were acting in addition to surface adsorption. In order to examine these possibilities, it is first necessary to further investigate the effect of the increase of particle surface area with hydration on LSs consumption by the cement paste.

412 **4.2** The effect of LSs on the increase in particle surface area with hydration

As displayed in Figure 5, the BET measurements showed an increase in particle surface area with hydration, which was influenced both by the plasticizer dosage and by its addition time. The increase in surface area was larger for CX than for ANL cement, and greatly larger for IA than for DA for both cements. As shown in Figure 6, the influence of the different surface area of the two hydrated cements could be eliminated by expressing the results relative to the surface area of the hydrated substrate as measured with BET.

For CX cement, the isotherms obtained with IA and DA nearly coincide, both reaching an adsorption plateau. This indicates that the higher LSs consumption by CX cement paste measured with IA is mainly due to the larger increase in particle surface area that takes place with IA, which, on the contrary, does not take place with DA. Hence, for CX cement, the main mechanism for LSs consumption seems to be monolayer surface adsorption for both IA and DA. 424 Regarding ANL cement, even when expressed relative to the hydrated surface area, the adsorption 425 isotherms remain qualitatively similar to those obtained considering the unhydrated surface area. 426 Therefore, for ANL cement, monolayer surface adsorption does not seem to be the only LSs 427 consumption mechanism for IA.

Finally, as displayed in Figure 4, when LSs was added to the cement pastes with IA, a higher amount of ettringite was formed by both cements compared to the sample without LSs. Moreover, as shown in Figure 5, the LSs addition led to an increase in particle surface area. The increase was larger for CX cement, which also presented a larger amount of ettringite. Therefore, the increase in particle surface area seems to be directly correlated to the increased amount of ettringite produced by the cements in presence of LSs.

434 **5 SUMMARY AND CONCLUSIONS**

435 The aim of this paper is to obtain a better understanding of the mechanisms for lignosulfonate (LSs) consumption by cement paste. This is considered to be crucial to maximize the efficiency of the 436 437 plasticizer. According to literature, the mechanisms behind the consumption of a softwood low-sugar 438 calcium lignosulfonate can potentially be: monolayer adsorption, intercalation, and calcium 439 complexation/multilayer adsorption. The LSs consumption by two different cements at immediate (IA) 440 and delayed (DA) addition was investigated using UV-spectroscopy and adsorption isotherms were 441 calculated. The changes in particle surface area with hydration were measured with BET. The changes in hydrates due to the addition of LSs were examined with TGA. The results obtained for cement 442 pastes were compared to those obtained for reference materials (i.e. calcium carbonate, calcium 443 444 hydroxide). The following conclusions were drawn:

- 445 The presence of LSs in the cement paste led to an additional formation of ettringite for IA,
 446 while an increase was not detected for DA;
- An increase in LSs dosage led to an increase in particle surface area after 30 minutes of
 hydration. This increase was considerably higher for IA rather than for DA.

- 449 Neither AFm nor intercalated AFm were observed. Therefore, intercalation cannot explain the
 450 differences in LSs consumption observed between IA and DA.
- The mechanism of LSs consumption seems to be mainly monolayer surface adsorption for CX
 cement (both for IA and for DA), and for ANL cement when DA was applied. For ANL, other
 mechanisms in addition to monolayer surface adsorption appear to play a role in LSs
 consumption when IA is applied.

455 6 FUTURE RESEARCH

In a follow-up study, the effect of LSs on rheology, rate of hydration, and amount and morphology of ettringite in pastes of the same cements will be investigated both for IA and DA. A deeper investigation on the mechanisms behind LSs consumption in ANL cement for IA will also be considered for future research.

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553
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- 554 Notation
- 555 apw artificial pore water
- 556 ANL Anlegg cement
- 557 CX Cemex cement
- 558 IA immediate addition of plasticizer
- 559 DA delayed addition of plasticizer
- 560 LSs softwood low-sugar Ca-lignosulfonate
- 561 OPC ordinary Portland cement
- 562 w/b water-binder ratio
- 563 LDHs layered double hydroxides

⁵⁶⁵ List of tables

566	Table 1 – Main phases in cement ANL and CX from XRD-Rietveld analysis performed at École de	5
567	Mines d'Alès, France. ^a : results obtained with TGA analysis	23
568	Table 2 – Chemical composition of the raw materials given by the producers	23
569	Table 3 - Physical properties of ANL and CX cements performed at École des Mines d'Alès, France	ce,
570	or given by the producer (*)	24
571	Table 4 - Chemical and physical properties of the lignosulfonate plasticizer used	24
572	Table 5 – Surface area (from BET) of $CaCO_3$ and $Ca(OH)_2$	24
573	Table 6 – Tested samples to obtain adsorption isotherms	24
574	Table 7 – Molecular footprints calculated for the different materials. * Equal to m_{SA}^{∞} in equation	2 **
575	The surface area used in the calculations was the hydrated one for the two cements and the	
576	unhydrated one for the two model materials	25

578 List of figures

579	Figure 1 - Amount of consumed LSs as % of added LSs vs. hydration time of ANL and CX cement				
580	pastes with 0.40 mass % of binder LSs25				
581	Figure 2a, b – Amount of consumed LSs after 30 min. of hydration vs. amount of LSs added to neat				
582	ANL and CX cements (IA and DA), and to $CaCO_3^*$. The results are calculated per mass % of binder in				
583	fig. 2 a and per unit of surface area available for adsorption of unhydrated particles in fig. 2 b26				
584	Figure 4 a, b - Thermogravimetric curves and their derivatives for ANL (a) and CX (b) cement paste				
585	without LSs (black) and with 1.5 mass % of binder LSs (gray)(full line for IA and dotted line for DA)				
586	for which hydration was stopped after 30 minutes. The peaks corresponding to the decomposition of				
587	ettringite (ettr.), hemihydrate (CaSO4·0.5H2O) or gypsum (CaSO4·2H2O), portlandite (CH) and				
588	carbonates (CO2)27				
589	Figure 5 – Surface area of ANL and CX cement particles hydrated for 30 min. both for IA and DA vs.				
590	the total dosage of LSs added (mass % of				
591	binder)				
592	Figure 6 - Amount of consumed LSs at 30 min. hydration (calculated as unit of surface area of				
593	hydrated substrate) vs. amount of LSs added to neat ANL and CX cement (IA and DA) (calculated as				
594	mass % of binder). For CX cement, the data points for 0.2; 0.4; 1.2 mass % of binder LSs (IA/DA)				
595	were calculated with interpolation and were not experimentally measured				

597 Table 1 – Main phases in cement ANL and CX from XRD-Rietveld analysis performed at École des

598 Mines d'Alès, France.^a: results obtained with TGA analysis

Phase composition (%wt)	ANL	СХ
Alite	60.5	54.3
Belite	14.2	18.8
Aluminate cubic	1.3	4.7
Aluminate ortho.	0.9	2.4
Ferrite	14.0	6.5
periclase	0.4	1.1
quartz	0.3	-
calcite	3.2/ 3.8 ^a	3.6/ 3.7 ^a
portlandite	1.1/ 1.4 ^a	2.6/ 2.5 ^a
anhydrite	-	2.1
hemihydrate	2.6	1.8
gypsum	1.0	-
arcanite		0.6
aphthitalite	0.4	0.7
thenardite	-	0.8

599

600 Table 2 – Chemical composition of the raw materials given by the producers

Chemical compound (%wt)	ANL	СХ
Fe ₂ O ₃	3.50	2.60
TiO ₂	0.22	0.25
CaO	62.70	64.00
K ₂ O	0.40	1.00
P_2O_5	0.15	0.23
SiO ₂	20.60	20.00
Al ₂ O ₃	4.40	4.60
MgO	1.60	2.40
Na ₂ O	0.30	0.20
SO ₃	3.30	3.60
LOI (%) 1000 °C	1.6	1.7
Sum	97.17	98.88

602 Table 3 - Physical properties of ANL and CX cements performed at École des Mines d'Alès, France,

ANL	СХ
890	1326
360	540
3.13	3.09
2.0	2.0
12.0	10.0
34.0	26.0
	890 360 3.13 2.0 12.0

603 *or given by the producer (*).*

604

605 Table 4 - Chemical and physical properties of the lignosulfonate plasticizer used

Plasticizer type	Mw	Org S (∝ SO ₃)	SO ₄ ²⁻ (mass %)	Ca ²⁺	Na	СООН	φ-OH	Total sugar (%)
LSs	29000	4.6	0.9	4.6	0.9	7.1	1.4	8.3

606

607 Table 5 – Surface area (from BET) of $CaCO_3$ and $Ca(OH)_2$

	CaCO ₃	Ca(OH) ₂
Surface area (BET) (m²/kg)	570	16661

608

609 Table 6 – *Tested samples to obtain adsorption isotherms*

Material	LSs addition procedure	LSs dosage tested (mass % solid)
ANI comont	IA	0.1; 0.2; 0.4; 0.6; 0.8; 1.2; 1.5
ANL cement	DA	0.05; 0.1; 0.25; 0.4; 0.8; 1.2; 1.5
	IA	0.1; 0.2; 0.4; 0.6; 0.8; 1.0
CX cement	DA	0.05; 0.1; 0.2; 0.4; 0.8; 1.2; 1.5
CaCO ₃	IA	0.05; 0.1; 0.2; 0.4; 0.8; 1.0; 1.2; 1.5

Ca(OH) ₂	IA	1.0; 2.0; 5.0; 8.0; 12.0; 22.0
610		

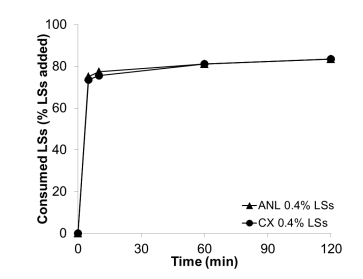
611 Table 7 – Molecular footprints calculated for the different materials. * Equal to m_{SA}^{∞} in equation 2 **

612 The surface area used in the calculations was the hydrated one for the two cements and the

613 *unhydrated one for the two model materials*

Material Added LSs at platea		Consumed LSs at plateau *	Molecular footprint **	
	(mass % of binder)	(mass % of binder)	(nm ²)	
ANL cement	0.8 - 1.2	0.42 ± 0.05	20 ± 10	
CX cement	1.2 - 1.5	0.68 ± 0.03	20 ± 10	
CaCO ₃	0.2 - 0.4	0.045 ± 0.009	60 ± 30	
Ca(OH) ₂	8.0 - 12.0	1.99 ± 0.31	40 ± 20	

614



616 Figure 1 - Amount of consumed LSs as % of added LSs vs. hydration time of ANL and CX cement

617 pastes with 0.40 mass % of binder LSs

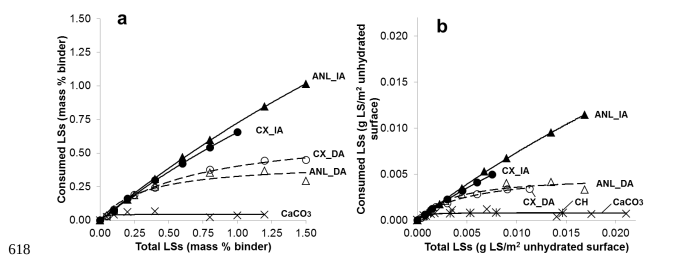
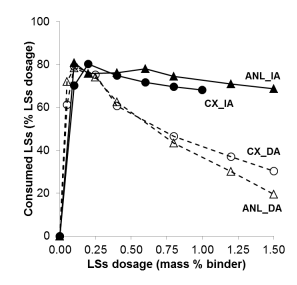


Figure 2a, b – Amount of consumed LSs after 30 min. of hydration vs. amount of LSs added to neat
ANL and CX cements (IA and DA), and to CaCO₃*. The results are calculated per mass % of binder in
fig. 2 a and per unit of surface area available for adsorption of unhydrated particles in fig. 2 b. * In
Figure 2a, the isotherms of CH is omitted due to the higher LSs dosage used for this sample.



624 Figure 3 – Consumed LSs (% LSs dosage) vs. LSs dosage (mass % of binder) for pastes of ANL and

625 CX cements where LSs was added both with IA and DA

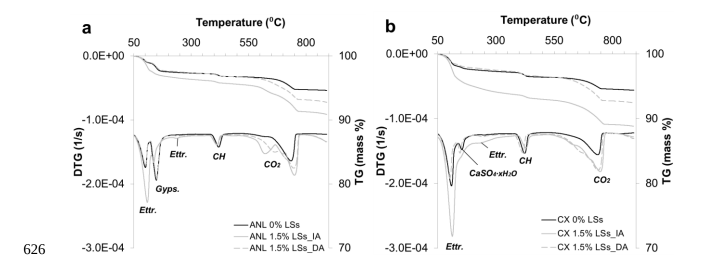
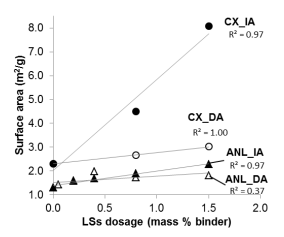


Figure 4 a, b - Thermogravimetric curves and their derivatives for ANL (a) and CX (b) cement paste without LSs (black) and with 1.5 mass % of binder LSs (gray)(full line for IA and dotted line for DA) for which hydration was stopped after 30 minutes. The peaks corresponding to the decomposition of ettringite (ettr.), hemihydrate (CaSO₄·0.5H₂O) or gypsum (CaSO₄·2H₂O), portlandite (CH) and carbonates (CO₂)



633 Figure 5 – Surface area of ANL and CX cement particles hydrated for 30 min. both for IA and DA vs.

634 the total dosage of LSs added (mass % of binder)

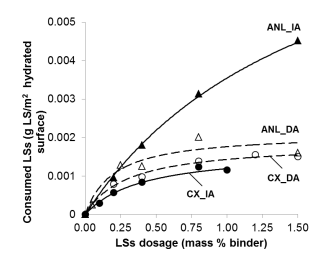


Figure 6 - Amount of consumed LSs at 30 min. hydration (calculated as unit of surface area of
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- 638 mass % of binder). For CX cement, the data points for 0.2; 0.4; 1.2 mass % of binder LSs (IA/DA)
- 639 were calculated with interpolation and were not experimentally measured