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Mignon, Arn; DEVISSCHER, Dries; GRAULUS, Geert-Jan; Stubbe, Birgit; Martins, Jose; Dubruel, Peter; De Belie, Nele & Van Vlierberghe, Sandra (2017) Combinatory approach of methacrylated alginate and acid monomers for concrete applications. In: CARBOHYDRATE POLYMERS, 155, p. 448-455.

DOI: 10.1016/j.carbpol.2016.08.102 Handle: http://hdl.handle.net/1942/22766

1	Combinatory approach of methacrylated alginate and acid monomers			
2		for concrete applications		
3				
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31	Abstract:			
32		ally alginate, can be useful for self-healing of cracks in concrete. Instead		
33		present within calcium alginate, covalent bonds, by methacrylation of		
34		esult in mechanically stronger superabsorbent polymers (SAPs). These		
35		ins as backbone are combined with two acrylic monomers in a varying		
36		show a moisture uptake capacity up to 110% their own weight at a		
37		vith a negligible hysteresis. The swelling capacity increased (up to 246		
38	times its own weight) with	a decreasing acrylic acid/2 acrylamido-2-methylpropane sulfonic acid		
39	ratio. The SAPs also showe	ed a thermal stability up to 200°C. Interestingly, the SAP composed of		
40	alginate and acrylic acid ex	erted a very limited decrease in compressive strength (up to 7% with		
41	addition of 1 wt% SAP) rend	ering this material interesting for the envisaged self-healing application.		
42				
43	Keywords:			

44 polysaccharide; alginate; self-healing; concrete; swelling potential; compressive strength

45

46 1. Introduction

47

Reports have been presented in literature covering synthetic superabsorbent polymers (SAPs) used 48 49 in concrete for various applications such as internal curing (Mechtcherine et al., 2014; Viktor 50 Mechtcherine, 2006), frost resistance (Jensen, 2013; Laustsen, Hasholt & Jensen, 2013) and self-51 sealing and -healing of cracks (Mignon et al., 2014; Snoeck, Van Tittelboom, De Belie, Steuperaert & 52 Dubruel, 2012; Snoeck, Van Tittelboom, Steuperaert, Dubruel & De Belie, 2014). The use of synthetic 53 SAPs has been proven to be promising as a smart internal solution for this latter application (Mignon et al., 2014; Mignon et al., 2015b; Snoeck, Van Tittelboom, Steuperaert, Dubruel & De Belie, 2014). 54 55 However, these synthetic materials often compromise the material strength, which is undesired in 56 the construction industry. On the one hand, the SAP particles will cause internal curing by releasing 57 their entrained mixing water, stimulating the densification and further hydration of the cementitious 58 matrix and reducing autogenous shrinkage and hence, the risk at early age cracking. These 59 consequences can lead to an increase of the overall material strength. Conversely, after the release 60 of the entrained water by the SAPs, air-filled macropores will remain present in the matrix, which 61 generally leads to a decrease of the overall concrete strength (Hasholt, Jensen, Kovler & Zhutovsky, 62 2012; Laustsen, Hasholt & Jensen, 2013; Snoeck, Schaubroeck, Dubruel & De Belie, 2014). As for self-63 sealing (Snoeck, Steuperaert, Van Tittelboom, Dubruel & De Belie, 2012) and self-healing applications 64 (Snoeck, Schaubroeck, Dubruel & De Belie, 2014; Snoeck, Van Tittelboom, Steuperaert, Dubruel & De 65 Belie, 2014) high SAP amounts are required (up to 1 % relative to cement mass), this macropore 66 formation becomes more critical especially when high amounts of additional water are used to 67 compensate for the loss in workability. Previous research has proven that polysaccharides and 68 especially alginate, are very promising as starting materials for self-sealing and -healing of cracks. 69 However, instead of the weak electrostatic bonds present within calcium alginate which are prone to 70 disintegration in the presence of monovalent cations (Bajpai & Sharma, 2004), covalent linkages 71 generally result in mechanically superior SAPs (Povh & Rosina, 2005). One way to create covalent 72 bonds in a SAP network based on polysaccharides is by first modifying them with methacrylic anhydride (MAAH) (Chou, Akintoye & Nicoll, 2009; Chou & Nicoll, 2009). MAAH can be used to 73 74 enable cross-linking of the polysaccharides. In the case of alginate, the hydroxyl groups will react 75 with the anhydride resulting in methacrylated alginate. These methacrylate functions can be used to 76 execute a free radical polymerization in the presence of acrylic monomers.

77

78 Acrylic acid (AA) has already been proven to be a widespread used monomer for the development of 79 synthetic SAPs, often in combination with other acrylic monomers such as acrylamide (Ding, Xiao, An 80 & Jia, 2006; Mohammad J. Zohuriaan-Mehr 2008; Nesrinne & Djamel, 2013; Zhang, Cui, Yin, Li, Liao & 81 Cai, 2011; Zhou et al., 2003). Another interesting monomer is 2-acrylamido-2-methylpropanesulfonic 82 acid (AMPS) as this hydrophilic, sulfonic acid possesses a large swelling capacity and has already been 83 used in a variety of applications going from water treatment (Kang & Cao, 2012; Yu et al., 2011) and 84 drug delivery (Pourjavadi, Barzegar & Zeidabadi, 2007) towards other biomedical applications 85 (Anirudhan & Sandeep, 2011; Keogh, 1995; Keogh, Hobot, Eaton, Jevne & Bergan, 1995) and personal 86 care (Abdel-Azim, Farahat, Atta & Abdel-Fattah, 1998; Cahalan & Coury, 1986; Lundmark, Melby & 87 Chun, 1978).

88

89 The present manuscript reports on the development and characterization of SAPs based on 90 methacrylated alginate as backbone combined with a varying molar fraction of a carboxylic acid (AA) 91 and/or a sulfonic acid (AMPS). The degree of methacrylation after modification will be determined by 92 proton nuclear magnetic resonance spectroscopy. After polymerization, the gel fraction and particle 93 size distribution will be identified. The chemical structure of the synthesized SAPs will be verified 94 through attenuated total reflectance-infrared (ATR-IR) spectroscopy. High resolution magic-angle 95 spinning (HR-MAS) ¹H-NMR spectroscopy will be used to identify the polymerization efficiency. In 96 addition, the sorption and desorption of moisture at different relative humidities will be measured by 97 dynamic vapor sorption (DVS) measurements. It is also important to identify the swelling degree in 98 aqueous (ultrapure water and demineralized water) and cement filtrate solutions. Additionally, the 99 thermal stability is evaluated using thermogravimetric analysis (TGA). Finally, the influence of these 100 SAPs on the strength of mortar is also determined.

101

2. Materials and methods

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2.1. Materials

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106 2-Acrylamido-2-methylpropane sulfonic acid (AMPS), ammonium persulfate (APS), sodium alginate 107 (NaAlg, M/G ratio of 2.3, Mw of 76 kDa and a Đ of 4.6) and methacrylic anhydride (MAAH) have been 108 bought from Sigma-Aldrich (Bornem, Belgium). Acrylic acid (AA) and N,N,N',N'- tetramethylethylene-109 diamine (TEMED) originated from Acros Organics (Geel, Belgium). N,N'-methylene bisacrylamide 110 (MBA) came from Merck (Nottingham, UK). The paper filters (retention of 8 – 12 μ m) were 111 purchased at Munktell filters (Bärenstein, Germany).

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2.2. Modification of alginate backbone with methacrylate moieties

115 A derivatization of alginate with methacrylic anhydride (MAAH), based on a reaction of the hydroxyl 116 moieties of the polysaccharide backbone with the anhydride, was carried out to incorporate 117 methacrylate groups for network formation. A 2 wt% sodium alginate solution was prepared in 118 demineralized water using a mechanical stirrer. Subsequently, MAAH was added dropwise to the 119 solution. The added amount corresponded with 2 equivalents of MAAH with respect to the hydroxyl 120 groups from alginate. During the reaction, methacrylic acid was released which lowers the pH of the 121 reaction. However, the acidity of the mixture was constantly monitored and the pH was increased to 122 8 by adding small amounts of a 5 M sodium hydroxide (NaOH) solution. Conversely, the pH should 123 not be too high to avoid hydrolysis of the ester in the reaction product. The mixture was reacted at 124 room temperature for 24 hours after the addition of MAAH. Afterwards, dialysis (12 – 14 kDa) was 125 performed for 72 hours while changing the dialysis water twice a day to remove unreacted agents 126 and formed methacrylic acid. The resulting solution afforded modified alginate (algMOD) by 127 lyophilization.

- 128
- 129 130
- 2.3. Detailed characterization of the synthesized SAPs
- 131 After polymerization, unreacted particles were removed from the end product via dialysis. By 132 measuring the dry weight of the sample before and after purification during 24 hours, the gel 133 fraction could be determined using the following equation:

134	$G[\%] = \frac{W}{W_0}$ (1)
135	W = mass of the dry insoluble part of the sample
135	W_0 = initial dry mass of the sample
137	
138	To examine the resulting particle size diameters of all grinded materials (grinded with an A11 basic
139	Analytical Mill), a Zeiss Axiotech optical microscope was used together with the digital image
140	capturing software ZEN core and the analysing software ImageJ. A sample population > 100 has been
141	used. The statistical analysis was performed by creating a size distribution curve.
142	
143	The SAP powder was subsequently characterized using attenuated total reflectance infrared (ATR-IR)
144	spectroscopy. A BioRad FTS 575C combined with a MKII Golden Gate setup equipped with a diamond
145	crystal from Specac was used for all experiments. These results were analysed with the Bio-Rad Win-
146	IR Pro software.
147	
148	The 1 H-NMR spectra were obtained by dissolving the samples in D ₂ O (10 mg/ml) and recording the
149	samples at room temperature on a Varian Inova 400 MHz spectrometer using a 5 mm four-nucleus
150	PFG probe with water suppression.
151	
152	High Resolution Magic-Angle Spinning NMR (HR-MAS ¹ H-NMR) spectroscopy analysis of the
153	developed SAPs was performed on a Bruker Avance II 700 spectrometer (700.13 MHz) using a HR-
154	MAS probe equipped with a ¹ H, ¹³ C, ¹¹⁹ Sn and gradient channel. The spinning rate was set to 6 kHz.
155	Samples were prepared by placing a small amount of freeze-dried material inside a 4 mm zirconium
156	oxide MAS rotor (80 μ L). 30 μ L D ₂ O was added to the rotor, allowing the samples to swell. The
157	samples were homogenized by manual stirring prior to analysis. A teflon [®] coated cap was used to
158 159	close the rotor.
160	Dynamic vapor sorption (DVS) experiments were applied as described earlier in (Mignon et al.,
161	2015a) with systematic steps of 20, 40, 60, 80, 90 and 95% RH.
162	
163	Swelling tests performed in ultrapure and demineralized water
164	The swelling capacity of the SAPs was determined as the mass change between the freeze-dried and
165	the swollen (cfr. saturated) state. A mass of 0.15 - 0.20 g polymer was incubated in 100 mL of
166	aqueous solution. After 3h incubation, a funnel and a filter were used to capture the water that was
167	not absorbed by the SAPs. By calculating the difference between the initially added and the filtered

not absorbed by the SAPs. By calculating the difference between the initially added and the filtered water, the residual water inside the material could be determined together with the swelling capacity of the material using Equation (5):

170

Swelling capacity =
$${(m_0 - m_{filter})}/{m_{SAP}}$$
 (5)

172where m_0 is the initially added water mass [g], m_{filter} represents the mass of the water going through173the filter [g] and m_{SAP} is the added mass of dried SAP (i.e. 0.20 g). The filtration paper was typically174saturated prior to the filtration to exclude its influence on the mass of the filtered water.

- 175
- 176 <u>Swelling tests in cement filtrate solutions</u>

177 Cement filtrate (CF) was made by mixing 10 g ordinary Portland cement (OPC) and 100 mL demineralized water for three hours with a mechanical stirrer, followed by filtration to remove the 178 179 cement particles and collecting the solution. The same conditions as for aqueous solutions were used for further analysis. All swelling tests were performed in triplicate. 180

181

182 The thermal properties were analyzed using thermogravimetric analysis (TGA) using a TA-183 instruments Q-50 Thermogravimetric Analyzer as described earlier in (Mignon et al., 2015b) with a maximal temperature of 600°C and a starting equilibrium temperature of 30°C. 184

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2.4. Bending and compression strength measurements

188 Mortar samples were manufactured by a standard mortar mixing procedure, as described in detail in 189 EN 196-1 to investigate the influence of the addition of SAPs on the flexural and compression 190 strength of mortar samples. Flexural and compressive strength were measured at the age of 28 days 191 by means of a three-point-bending test followed by a compression test on the resulting halves 192 following the Standard NBN EN 196-1 and described in detail in (Mignon et al., 2015b).

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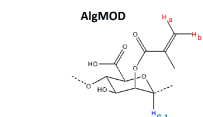
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 - 3. Results and discussion
 - 3.1. Methacrylation of alginate through modification with methacrylic anhydride

199 In a first step, alginate was methacrylated yielding algMOD. The degree of methacrylation (DM, or 200 the degree of substitution, DS) (i.e. the amount of double bonds incorporated on alginate) has a 201 strong influence on the physical properties as well as the swelling capacity of the formed SAP. For 202 that reason, it is important to calculate the DS using ¹H NMR spectroscopy (as described in section 203 2.6).



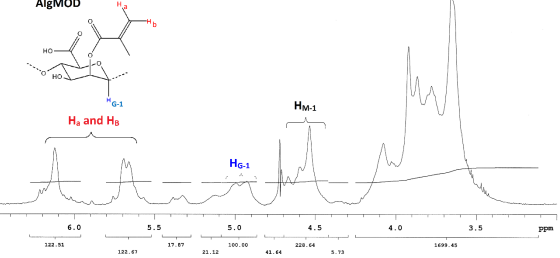


Figure 1: ¹H NMR spectrum of algMOD with annotation of the relevant peaks for the calculation of 206

207 the DS (H_a and H_b as protons connected to the C=C bond, a proton on the α -l-guluronic acid structure

208 (H_{G-1}) , and a reference proton on the β -d-mannuronic acid (H_{M-1}) , not shown on molecular structure). 209

210 The obtained spectra were analyzed by comparing the characteristic peaks of the methacrylate 211 moieties corresponding to the vinyl protons at 5.73 and 6.16 ppm with the reference peak from the alginate backbone at 4.97 ppm. As the latter corresponds with the anomeric carbon of the guluronic 212 213 acid block (G-units), the G-value had to be calculated and incorporated as a correction factor due to 214 the presence of mannuronic acid (M-units). Therefore, the relative proportion of the G-units 215 compared to the total amount of G- and M-units had to be determined by using Equation 2. 216 Subsequently, the average of the peaks of the methacrylate groups was calculated and used to 217 compute the degree of substitution (DS) per repeating unit by applying Equation 3. To obtain the DS 218 with respect to the hydroxyl groups, this value had to be divided by two as the considered repeating 219 unit (G-unit) possessed two hydroxyl groups.

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- 222

$$G\left[\%\right] = \frac{H_{G-1}}{H_{M-1} + H_{G-1}} = \frac{I_{4.97ppm}}{I_{4.58ppm} + I_{4.97ppm}} * 100\%$$
(2)

 $DS [\%] = G * \frac{\frac{H_a + H_b}{2}}{H_{G-1}} = G * \frac{\frac{I_{5.73ppm} + I_{6.16ppm}}{2}}{I_{4.97ppm}}$ (3)

224

225 The protons H_{G-1} , H_{M-1} , H_a and H_b are related to the intensity of their related peaks on the NMR 226 spectrum, seen further in Figure 1. Using Equations 2 and 3, a relative concentration of 227 guluronic acid units or G-content of 30.5% and a DS of 18.7% as a function of the present hydroxyl groups were determined. The latter implies that almost one out of five functional 228 229 hydroxyl groups or one of every three repeating units (37.4%) have been modified.

230

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3.2. SAP development and characterization

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233 After modification, methacrylated alginate (algMOD) was combined with the monomers (AA and 234 AMPS) to create a cross-linked network through free radical polymerization. AMPS was selected as second monomer to compare the effect of a carboxylic acid versus the incorporation of a sulfonic 235 236 acid with respect to the swelling capacity as well as the influence on the mortar strength. AlgMOD 237 combined with AA forms a strong and transparent solid gel while a decrease in AA/AMPS ratio leads 238 to a more brittle gel. The composition and nomenclature of the SAPs is given in Table 1.

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Table 1: Overview of theoretical chemical composition of the developed SAPs and the gel fraction. 241 The algMOD fraction was added in addition to the total monomer amount.

Sample	AlgMOD AA + AMPS g/g	AA mol%	AMPS mol%	Gel fraction (%)
p(alg(1)_AA ₁₀₀ /AMPS ₀ (7))	1/7	100	0	85.3 ± 1.2
p(alg(1)_AA ₇₅ /AMPS ₂₅ (7))	1/7	75	25	80.9 ± 2.6
p(alg(1)_AA ₅₀ /AMPS ₅₀ (7))	1/7	50	50	61.3 ± 2.2
p(alg(1)_AA ₂₅ /AMPS ₇₅ (7))	1/7	25	75	58.9 ± 1.3
p(alg(1)_AA ₀ /AMPS ₁₀₀ (7))	1/7	0	100	42.7 ± 6.2

Table 1 also shows that a lower AA/AMPS ratio leads to a significant decrease in the gel fraction. On the one hand, the materials containing more AMPS were more brittle and thus more prone to become damaged during purification. In addition, the higher polarity of AMPS could have resulted in an increased repulsion between negatively charged carboxylate groups on the alginate backbone and sulfate moieties on the AMPS monomer, thereby hindering the crosslinking reaction.

248

249 Smaller particles have a higher surface area which can have an influence on a.o. the moisture uptake 250 capacity. In addition, upon incorporation in mortar, these smaller particles will lead to more smaller 251 pores compared to larger SAPs. All materials showed a similar particle size for d_{10} and d_{50} (see Table

252 2). In the table, d_x indicates the diameter where x percentage of the particles is smaller or equal to. 253 Approximately half of the particles were characterized by a diameter of 20 μ m or lower. However,

- the d_{q_0} values show that there was a small variation between the SAPs, which can be related to the
- 255 duration of the grinding. However, the difference was quite limited and all diameters ranged 256 between $50 - 80 \mu m$.
- 257
- 255

Table 2: Particle size ranges of the algMOD - AA/AMPS SAPs where d_{10} , d_{50} and d_{90} indicate the percentage of particles (10%, 50% and 90%) with a diameter smaller or equal to x μ m.

Material	d _{_10} [μm]	d _{_50} [μm]	d _{_90} [μm]
p(alg(1)_AA ₁₀₀ /AMPS ₀ (7))	9-10	20-21	80
p(alg(1)_AA ₇₅ /AMPS ₂₅ (7))	8-9	20-21	47-48
p(alg(1)_AA ₅₀ /AMPS ₅₀ (7))	8-9	18-19	48-49
p(alg(1)_AA ₂₅ /AMPS ₇₅ (7))	10-11	21-22	75-76
p(alg(1)_AA ₀ /AMPS ₁₀₀ (7))	10-11	20-21	67-68

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- 261

262 3.3. Chemical structure confirmation and cross-linking efficiency assessment

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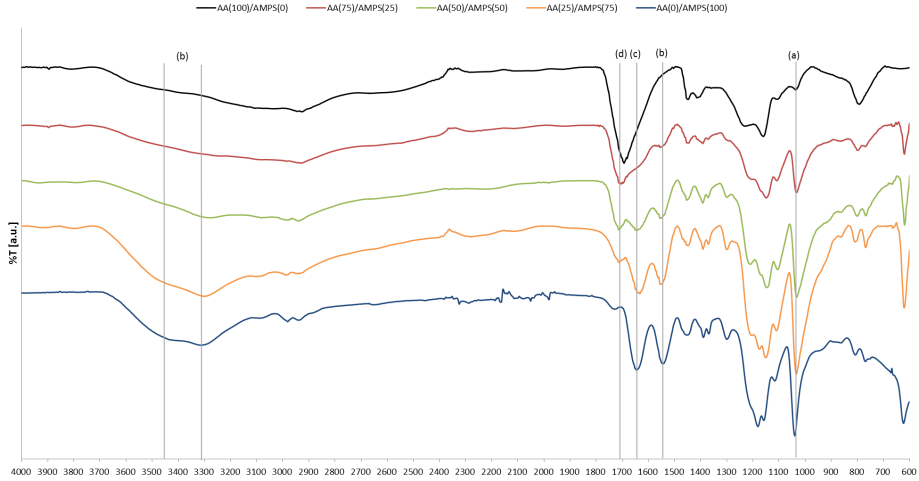
ATR-IR spectroscopy has been performed on the p(alg_AA/AMPS) SAPs (see Figure 2). The S=O stretch at 1040 cm⁻¹ of the sulfonic acid ((a) in Figure 2), the N-H stretch of the secondary amides between 3500 and 3100 cm⁻¹ and at 1550 cm⁻¹(b) and the C=O stretch of the amides at 1650 cm⁻¹ (c) are clearly stronger present with an increasing amount of AMPS.

The C=O stretch of the acid moiety at 1700 cm⁻¹ (d) is on the other hand strong for p(alg(1)_AA₁₀₀/AMPS₀(7)) due to the high amount of AA. These peaks are found by comparing to earlier performed research (Athawale & Lele, 1998; Athawale & Lele, 2000; Najjar, Yunus, Ahmad & Rahman, 2000; Rosa, Bordado & Casquilho, 2003).

272 These results give a qualitative indication of the presence of the polysaccharide and the monomers

as well as an increase or decrease of the intensity of certain peaks which is related to the variation of

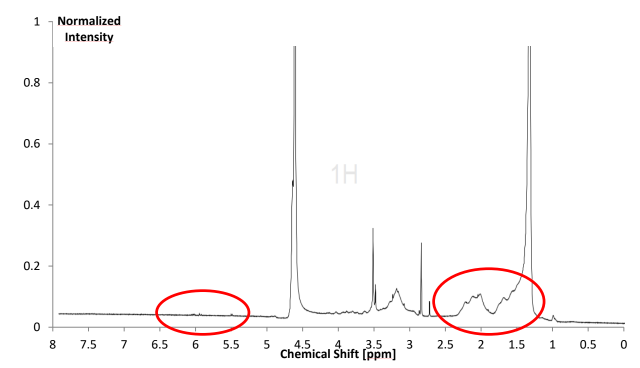
the AA/AMPS ratio.



v (cm-1)

Figure 2: FTIR spectra of the algMOD - AA/AMPS SAPs with annotation of the most relevant peaks. The samples are abbreviated by the molar ratio of the incorporated monomers according to: AA(x) /AMPS(y).

1 HR-MAS ¹H-NMR spectroscopy was used as a technique to confirm the polymerization (cross-linking) 2 efficiency. The peaks corresponding to protons from the C=C double bonds from either the 3 methacrylate functionalities introduced in algMOD or from the monomers are apparent in the range 4 of 5.5 - 6.5 ppm (as indicated in Figure 1 for algMOD). After polymerization, these signals have 5 completely disappeared, as shown in Figure 3 (i.e. p(alg(1)_AA₅₀/AMPS₅₀(7)) as an example). These 6 peaks have shifted to the right (1.5 - 2.5 ppm) by conversion of the double bond protons into alkane 7 protons, adjacent to an electronegative group.



9

8

Figure 3: HR-MAS ¹H NMR spectrum of p(alg(1)_AA₅₀/AMPS₅₀(7)). The absence of peaks
 corresponding to double bond protons (situated between 5.5 and 6.5 ppm) from either algMOD, AA
 or AMPS confirms the success of the cross-linking reaction.

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3.4. Determination of the moisture uptake capacity of the SAPs via dynamic vapor sorption

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16 The moisture uptake capacity of the SAPs has been assessed to identify their behavior in mortar or 17 concrete, when no direct ingress of water or rain is possible upon crack formation. If these SAPs can 18 significantly absorb moisture in humid environments, the cracks may be partially sealed. The results 19 (see Figure 4 (a)) show that with a decrease of the AA/AMPS ratio, the moisture uptake capacity 20 increases, especially at high relative humidity (RH). The values range between 53.6 and 109.8% at a 21 RH of 95%. Materials with \geq 50 mol% AMPS can even take up more than their original weight in 22 moisture at a RH of 95%. In addition, these materials show a negligible degree of hysteresis. As a 23 result, all moisture taken up can also be completely desorbed again. Indeed, concrete constructions 24 that are not exposed to humid environments can thus still retain a certain amount of moisture from 25 air at the crack faces. In a subsequent stage, the SAPs are likely to completely deliver this moisture to 26 remaining unreacted cement particles, which can lead to formation of new cement hydration 27 products and deposition of CaCO₃ from dissolved Ca(OH)₂ and CO₂, thereby contributing to the 28 development of self-healing applications (Snoeck, Van Tittelboom, Steuperaert, Dubruel & De Belie, 29 2014).

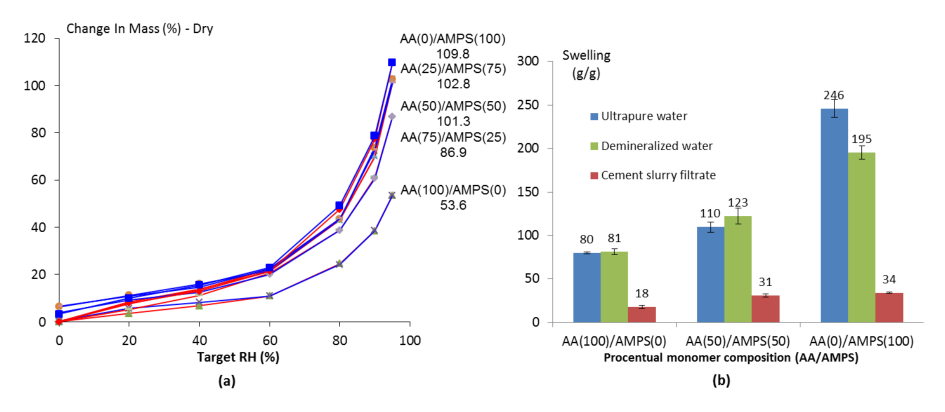
The reason for the higher moisture uptake capacity upon decreasing the AA/AMPS ratio can be attributed to the higher polarity of the sulfonic acid compared to the carboxylic acid. On the other hand, the increase can also be related to the higher amount of sulfonic acid groups being ionized at the same pH. Although this effect will be rather limited for DVS measurements, it will clearly be more pronounced in the swelling capacity.

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3.5. Swelling capacity measurements on the synthesized SAPs

38 The swelling potential of the SAPs was determined in three swelling media (i.e. ultrapure water, 39 demineralized water and cement filtrate solution). As such, a difference in the ionic concentration 40 will result in a variation of the swelling capacity. On the other hand, the monomer ratio (AA/AMPS) 41 will also have a strong influence on the swelling degree. Therefore, three samples with varying 42 monomer ratio were selected for further testing including $p(alg(1) AA_{100}/AMPS_0(7))$, 43 $p(alg(1)_{AA_{50}}/AMPS_{50}(7))$ and $p(alg(1)_{AA_0}/AMPS_{100}(7))$. The results show (see Figure 4 (b)) that the 44 swelling potential in cement filtrate is significantly (p < 0.05) lower, as anticipated due to the 45 presence of Ca²⁺, Mg²⁺ and other ions (Mignon et al., 2015b) as well as the lower osmotic pressure in a solution with a high ion concentration (Horkay, Tasaki & Basser, 2000). No significant differences 46 47 could be observed between ultrapure water and demineralized water except for 48 $p(alg(1)_{AA_0}/AMPS_{100}(7))$. Due to the high swelling of the latter, the relatively higher ion 49 concentration of demineralized water resulted in a swelling reduction. An observed trend for the 50 AA/AMPS ratio is that for all solutions, a decrease of the ratio leads to an increase of the swelling 51 capacity, which is in excellent agreement with the obtained DVS results.

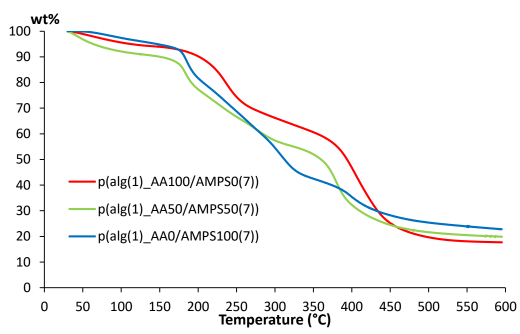


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Figure 4: (a) Sorption and desorption isotherms of p(algMOD _AA/AMPS) materials measured by a Dynamic Vapor Sorption apparatus. The values underneed the sample names describe the maximal moisture uptake capacity (at 95% RH). (b) Swelling potential of the synthesized SAPs in ultrapure water, demineralized water and cement filtrate solution with a pH of 12.6. The samples are abbreviated by the molar ratio of the incorporated monomers according to: AA(x) /AMPS(y).

58 3.6. Thermal stability of the SAPs assessed by thermogravimetric analysis

The production of the SAPs occurs at 45°C while the temperature of concrete can rise up to 50 - 60°C 60 61 during curing (Snoeck, 2015). Ideally, the polymers should not show thermal degradation at these 62 temperatures. In order to study potential thermally induced degradation, TGA measurements were performed (see Figure 5). The results showed that at 100°C, over 90% of the material was 63 64 maintained, while this loss can be attributed to residual water being present in the SAP as they were 65 kept in the lab and no freeze drying step was performed right before the TGA measurements. Subsequently, the step at 200 - 250°C can be related to either the decarboxylation of the polymer 66 67 and when AMPS is present the decomposition of sulfate groups. The last step around 360 – 450°C is 68 the main chain C-C fission (Abdelaal, Makki & Sobahi, 2012; Diao et al., 2010).



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Figure 5: TGA plots displayed as the percentual weight as a function of temperature for $p(alg(1)_AA_{100}/AMPS_0(7)), p(alg(1)_AA_{50}/AMPS_{50}(7)) and p(alg(1)_AA_0/AMPS_{100}(7)).$

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3.7. Effect of SAPs on the flexural and compressive strength of mortar samples

As the envisaged application includes their use in concrete, the SAPs were incorporated in mortar to investigate their effect on the flexural and compressive strength of the mortar samples. The composition of the mortar mixture is described in the materials and methods (chapter 2.11.)

78 Ideally, the SAPs should only exhibit a limited or no influence on the strength upon incorporating 79 them in mortar. Interestingly (Table 3), p(alg(1)_AA₁₀₀/AMPS₀(7)) only results in a limited 80 compressive strength reduction up to 7% upon addition of 1 wt% compared to the added amount of 81 cement. This material can thus be considered extremely promising. The data indicate that SAPs with 82 a higher molar fraction of AMPS lead to severely weaker mortar compared to the reference for both 83 the bending and compressive strength, up to the point that they are not useful anymore for the 84 envisaged application. Upon addition of 1 wt% $p(alg(1)_AA_0/AMPS_{100}(7))$ with respect to the added 85 amount of cement, the matrix collapsed and was too weak to assess bending and compressive 86 strength.

Table 3: Information on the additional water added on top of the 225g water initially, as well as the results of the three point bending and compression tests performed on the mortar samples

Sample description	SAP concentration [w%]	Additional water [ml]	Bending strength [MPa]	Compressive strength [MPa]
Reference	0	0	7.6 ± 0.5	73.1 ± 1.6
p(alg(1)_AA ₁₀₀ /AMPS ₀ (7))	0.5	15	6.8 ± 0.4 (-11%)	68.9 ± 1.6 (-6%)
p(aig(1)_AA100/Aivir 30(7))	1	30	7.6 ± 0.3 (/)	68.3 ± 0.6 (-7%)
p(alg(1)_AA ₅₀ /AM ₅₀ (7))	0.5	80	6.3 ± 0.4 (-17%)	42.1 ± 1.4 (-42%)
p(aig(1)_AA50/AW150(7))	1	150	5.1 ± 0.6 (-33%)	26.9 ± 0.2 (-63%)
p(alg(1)_AA ₀ /AMPS ₁₀₀ (7))	0.5	100	5.9 ± 0.3 (-22%)	36.6 ± 0.9 (-50%)

89 containing 0.5 or 1 wt% $p(alg(1)_AA_x/AMPS_y(7))$.

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4. Conclusions and future perspectives

93 Alginate was successfully modified using methacrylic anhydride to create methacrylated alginate (algMOD) with a DS of 19% with respect to the hydroxyl groups present. ATR-IR and HR-MAS ¹H-NMR 94 95 spectroscopy enabled to confirm the SAP structure as well as the cross-linking efficiency. The 96 materials possessed gel fractions ranging between 43 and 85%, which decreased upon decreasing 97 the AA/AMPS molar ratio. The SAPs showed a moisture uptake capacity going from 54 to 110% their 98 own weight at a RH of 95% upon increasing the AMPS ratio. Interestingly, all materials showed a 99 negligible hysteresis which implies that they can be used as a reservoir and all moisture taken up will 100 also be completely desorbed again. When investigating the swelling capacity, it was observed that an 101 increase of the AMPS content led to an increased swelling, independent of the used solution, up to a 102 maximal swelling potential of 246 times its own weight for $p(alg(1)_AA_0/AMPS_{100}(7))$ in ultrapure 103 water. Additionally, these polymers showed a thermal stability above 100°C, indicating that they will 104 not degrade during the curing of the concrete matrix. After incorporation of the SAPs in mortar, it 105 was determined that a decrease of the AA/AMPS ratio resulted in a severe decrease of the 106 compressive strength of mortar. Interestingly, $p(alg(1)_AA_{100}/AMPS_0(7))$ showed only a very limited 107 decrease in compressive strength (up to 7% decrease upon addition of 1 wt% SAP). Despite the 108 rather limited moisture uptake and swelling capacity, this latter material showed the highest gel 109 fractions and very limited effect on the compressive strength, rendering this material very interesting 110 for the envisaged application. This material will thus be further tested for its self-sealing and -healing 111 potential. The latter will be the topic of a forthcoming paper.

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113 Acknowledgement

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The authors would like to thank the FWO (Research Foundation Flanders) for project funding (3G019012, Effect of tunable hydrogels on concrete microstructure, moisture properties, sealing and self-healing of cracks). The authors would like to acknowledge the group of Peter Adriaensens and especially Gunter Reekmans for the calculation of the degree of methacrylation of algMOD (Organic

- and Bio-polymer Chemistry Department, UHasselt).
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121 References:

- Abdel-Azim, A. A. A., Farahat, M. S., Atta, A. M., & Abdel-Fattah, A. A. (1998). Preparation and
 properties of two-component hydrogels based on 2-acrylamido-2-methylpropane sulphonic acid. *Polymers for Advanced Technologies*, 9(5), 282-289.
- Abdelaal, M. Y., Makki, M. S. I., & Sobahi, T. R. A. (2012). Modification and characterization of
- polyacrylic acid for metal ion recovery. *American Journal of Polymer Science*, 2(4), 73-78.
- Anirudhan, T. S., & Sandeep, S. (2011). Synthesis and characterization of molecularly imprinted polymer of N-maleoylchitosan-grafted-2-acrylamido-2-methylpropanesulfonic acid and its controlled
- delivery and recognition of bovine serum albumin. *Polymer Chemistry*, *2*(9), 2052-2061.
- Bajpai, S. K., & Sharma, S. (2004). Investigation of swelling/degradation behaviour of alginate beads crosslinked with Ca2+ and Ba2+ ions. *Reactive and Functional Polymers*, *59*(2), 129-140.
- 132 Cahalan, P. T., & Coury, A. J. (1986). Method of preparing tape electrode. Google Patents.
- 133 Chou, A. I., Akintoye, S. O., & Nicoll, S. B. (2009). Photo-crosslinked alginate hydrogels support 134 enhanced matrix accumulation by nucleus pulposus cells in vivo. *Osteoarthritis and Cartilage, 17*(10), 135 1377-1384.
- 136 Chou, A. I., & Nicoll, S. B. (2009). Characterization of photocrosslinked alginate hydrogels for nucleus
- 137 pulposus cell encapsulation. *Journal of Biomedical Materials Research Part A, 91*(1), 187-194.
- 138 Diao, H., Yan, F., Qiu, L., Lu, J., Lu, X., Lin, B., Li, Q., Shang, S., Liu, W., & Liu, J. (2010). High 139 Performance Cross-Linked Poly(2-acrylamido-2-methylpropanesulfonic acid)-Based Proton Exchange
- 140 Membranes for Fuel Cells. *Macromolecules*, 43(15), 6398-6405.
- 141 Ding, Y., Xiao, C., An, S., & Jia, G. (2006). Water-absorptive blend fibers of copoly(acrylic acid– 142 acrylamide) and poly(vinyl alcohol). *Journal of Applied Polymer Science*, *100*(4), 3353-3357.
- Hasholt, M. T., Jensen, O. M., Kovler, K., & Zhutovsky, S. (2012). Can superabsorbent polymers
 mitigate autogenous shrinkage of internally cured concrete without compromising the strength? *Construction and Building Materials, 31*, 226-230.
- 146 Horkay, F., Tasaki, I., & Basser, P. J. (2000). Osmotic swelling of polyacrylate hydrogels in 147 physiological salt solutions. *Biomacromolecules*, 1(1), 84-90.
- Jensen, O. M. (2013). Use of superabsorbent polymers in concrete. *Concrete International, 35*(1), 4852.
- Kang, G.-d., & Cao, Y.-m. (2012). Development of antifouling reverse osmosis membranes for water
 treatment: a review. *Water research*, 46(3), 584-600.
- Keogh, J. R. (1995). Contacting blood with medical equipment having polymerized 2-acrylamido-2-methylpropanesulfonic acid on exposed surface. Google Patents.
- 154 Keogh, J. R., Hobot, C. M., Eaton, J. W., Jevne, A. H., & Bergan, M. A. (1995). Made from monomers
- such as N-(3-aminopropyl) methacrylamide hydrochloride, 2-acrylamido-2-methylpropanesulfonic
 acid, acrylamide and acrylic acid bonded to polymeric substrate surface; useful in medical devices.
- 157 Google Patents.
- Laustsen, S., Hasholt, M., & Jensen, O. (2013). Void structure of concrete with superabsorbent polymers and its relation to frost resistance of concrete. *Materials and Structures, 48*(1-2), 357-368.
- Lundmark, L. D., Melby, A., & Chun, H.-m. (1978). Method of imparting lubricity to keratinoussubstrates and mucous membranes. Google Patents.
- 162 Mechtcherine, V., Gorges, M., Schroefl, C., Assmann, A., Brameshuber, W., Ribeiro, A., Cusson, D.,
- 163 Custódio, J., da Silva, E., Ichimiya, K., Igarashi, S.-i., Klemm, A., Kovler, K., de Mendonça Lopes, A.,
- Lura, P., Nguyen, V., Reinhardt, H.-W., Filho, R., Weiss, J., Wyrzykowski, M., Ye, G., & Zhutovsky, S.
- 165 (2014). Effect of internal curing by using superabsorbent polymers (SAP) on autogenous shrinkage
- and other properties of a high-performance fine-grained concrete: results of a RILEM round-robin
 test. *Materials and Structures*, 47(3), 541-562.
 - 168 Mignon, A., Graulus, G.-J., Snoeck, D., Martins, J., De Belie, N., Dubruel, P., & Van Vlierberghe, S.
 - 169 (2014). pH-sensitive superabsorbent polymers: a potential candidate material for self-healing
 - 170 concrete. Journal of Materials Science, 50(2), 970-979.

- 171 Mignon, A., Graulus, G.-J., Snoeck, D., Martins, J., De Belie, N., Dubruel, P., & Van Vlierberghe, S. 172 (2015a). pH-sensitive superabsorbent polymers: a potential candidate material for self-healing 173 concrete. *Journal of Materials Science*, *50*(2), 970-979.
- 174 Mignon, A., Snoeck, D., Schaubroeck, D., Luickx, N., Dubruel, P., Van Vlierberghe, S., & De Belie, N.
- 175 (2015b). pH-responsive superabsorbent polymers: A pathway to self-healing of mortar. *Reactive and*
- 176 Functional Polymers, 93(0), 68-76.
- 177 Mohammad J. Zohuriaan-Mehr , K. K. (2008). Superabsorbent Polymer Materials: A Review. *Iranian* 178 *Polymer Journal,* 17((6)), 451-477.
- Nesrinne, S., & Djamel, A. (2013). Synthesis, characterization and rheological behavior of pH sensitive
 poly(acrylamide-co-acrylic acid) hydrogels. *Arabian Journal of Chemistry*(0).
- Pourjavadi, A., Barzegar, S., & Zeidabadi, F. (2007). Synthesis and properties of biodegradable
 hydrogels of κ-carrageenan grafted acrylic acid-co-2-acrylamido-2-methylpropanesulfonic acid as
 candidates for drug delivery systems. *Reactive and Functional Polymers*, *67*(7), 644-654.
- Povh, B., & Rosina, M. (2005). *Scattering and structures: essentials and analogies in quantum physics*.
 Springer Science & Business Media.
- 186 Snoeck, D. (2015). Self-healing and microstructure of cementitious materials with microfibres and187 superabsorbent polymers. Ghent University.
- 188 Snoeck, D., Schaubroeck, D., Dubruel, P., & De Belie, N. (2014). Effect of high amounts of 189 superabsorbent polymers and additional water on the workability, microstructure and strength of 190 mortars with a water-to-cement ratio of 0.50. *Construction and Building Materials*, *72*(0), 148-157.
- 191 Snoeck, D., Steuperaert, S., Van Tittelboom, K., Dubruel, P., & De Belie, N. (2012). Visualization of 192 water penetration in cementitious materials with superabsorbent polymers by means of neutron
- 193 radiography. *Cement and Concrete Research, 42*(8), 1113-1121.
- 194 Snoeck, D., Van Tittelboom, K., De Belie, N., Steuperaert, S., & Dubruel, P. (2012). The use of
- superabsorbent polymers as a crack sealing and crack healing mechanism in cementitious materials.
- 196 Concrete Repair, Rehabilitation and Retrofitting III: 3rd International Conference on Concrete Repair,
- Rehabilitation and Retrofitting, ICCRRR-3, 3-5 September 2012, Cape Town, South Africa (p. 58): CRC
 Press.
- Snoeck, D., Van Tittelboom, K., Steuperaert, S., Dubruel, P., & De Belie, N. (2014). Self-healing
 cementitious materials by the combination of microfibres and superabsorbent polymers. *Journal of Intelligent Material Systems and Structures*, 25(1), 13-24.
- Viktor Mechtcherine, L. D., Joachim Schulze. (2006). Internal curing by super absorbent polymers
 (SAP) effects on material properties of self-compacting fibre-reinforced high performance concrete.
 In P. L. O. M. Jensen, K. Kovler (Ed.) (pp. 87 96): RILEM Publications SARL.
- Yu, X.-I., Liu, X.-h., Xia, Z.-r., Cheng, D.-b., Xiao, S., Hu, Z.-j., & Yu, X.-m. (2011). Synthesis and Waterretention Property of the PAA-AM-AMPS and PAA-AM Superabsorbent Polymer. *Fine Chemicals*, *5*, 004.
- Zhang, B., Cui, Y., Yin, G., Li, X., Liao, L., & Cai, X. (2011). Synthesis and swelling properties of proteinpoly(acrylic acid-co-acrylamide) superabsorbent composite. *Polymer Composites, 32*(5), 683-691.
- 210 Zhou, X., Weng, L., Chen, Q., Zhang, J., Shen, D., Li, Z., Shao, M., & Xu, J. (2003). Investigation of pH
- sensitivity of poly(acrylic acid-co-acrylamide) hydrogel. *Polymer International*, *52*(7), 1153-1157.

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