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L-(+)-ascorbic acid

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1	Performance of alginate films for retention of L-(+)-ascorbic acid		
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ABSTRACT

In view of acting as controlled delivery systems for nutritional supplementation,			
therapy or antioxidant activity at interfaces, alginate films of different copolymer			
composition and glycerol plasticizer levels were developed in the presence of Ca2+ for			
achieving higher stability of L-(+)-ascorbic acid (AA). The ability of the alginate			
network to preserve AA from hydrolysis, tested by storage under vacuum at 25°C, only			
decreased with the relative humidity (RH) increase when alginates were mainly			
constituted by guluronic-guluronic acid blocks (GG), whereas also decreased with the			
glycerol level increase when mannuronic-mannuronic acid (MM) and/or alternating			
guluronic-mannuronic (GM+MG) flexible blocks were present in higher proportions.			
This result could be probably related to the lower capability of the latter alginate block			
compositions to immobilize water in the network as they are not able to constitute Ca ²⁺			
mediated junction zones where water molecules are highly retained. Films also studied			
under air storage showed that even at less favorable conditions of RH and glycerol			
levels, both GG or GM+MG enriched alginate networks in general preserved AA from			
oxidation. It also demonstrated that hydrolysis is the principal way by which AA is lost			
when supported in films.			

47 Keywords: alginate films, ascorbic acid hydrolysis, glycerol, biomolecule delivery,

48 antioxidant interface.

1. Introduction

Alginate is a biomaterial that has found numerous applications in biome	edical
science and engineering due to its favorable properties, including biocompatibility	y and
facility for gelation (Lee and Mooney, 2012). Alginate hydrogels have been particular	ılarly
attractive in wound healing, drug delivery, and tissue engineering applications, as	these
gels retain structural similarity to the extracellular matrices in tissues and ca	in be
manipulated to play several critical roles. Alginates are also very useful because of	their
utility in preparing hydrogels at mild pH and temperature conditions, suitable	e for
sensitive biomolecules (Pawar and Edgar, 2012). Alginic acid, a natural polysacch	aride
harvested from brown algae, is an unbranched binary copolymer constituted by ((1,4)-
linked β -D-mannuronic acid (M-block), α -L-guluronic acid (G-block) and sequence	es of
alternating β -D-mannuronic and α -L-guluronic acid (MG-block) (Jothisaraswathi α	et al.,
2006). Physical and mechanical properties as well as biocompatibility of alg	ginate
materials are highly dependent on the relative content of L-guluronic to D-mannu	ronic
acids (Klöck et al., 1997; Stabler et al., 2001). Calcium ions can replace in par	rt the
hydrogen bonding, zipping guluronate (but not mannuronate) chains tog	ether
stoichiometrically in an "egg-box" conformation. Guluronate chain pairing thr	rough
junction zones involves three components: uronate chains, calcium ions and	water
molecules. The antiparallel arrangement is the macromolecular interaction prol	bably
favored in the gel, showing a notable contribution of hydrogen bonds to gel stre	ngth.
Moreover, the antiparallel association of 2 ₁ helical chains is the arrangement fou	nd in
the solid state (Braccini and Pérez, 2001).	
Alginates of different monomeric composition can be assayed in their abil	ity to
form film matrices for compartmentalization of L-(+)-ascorbic acid (AA), also kn	nown
as vitamin C. Through a delivery film $\Delta \Delta$ could provide for example nutrit	tional

supplementation (Durschlag et al., 2007), selective killing of cancer cells or local treatment of infections where H₂O₂ (formed from AA) may be beneficial (Chen et al, 2005). AA is a water soluble reducing agent and a natural antioxidant which also can be used for pharmaceutical preservation. AA stability is affected by processing and storage conditions because it depends on a large number of factors such as temperature, equilibrium RH, oxygen partial pressure, light (Kitts, 1997). AA reacts with oxygen to produce L-dehydroascorbic acid (DHA) that also has vitamin C activity in vivo. Biological activity is irreversibly lost when DHA is hydrolyzed in the subsequent reaction. Furthermore, anaerobic degradation of AA through hydrolysis also occurs simultaneously to AA oxidation when oxygen is present, producing 2-keto-L-gulonic acid (Kurata and Sakurai, 1967). On the other hand, non enzymatic browning also proceeds with AA concentration decay since the products of the reactions that follow the first step of AA destruction are also part of the browning reaction chain (León and Rojas, 2007). Compartmentalization of AA into a film network could help achieve stabilization because it can preclude the AA interaction with oxygen, with other pharmaceutical preservatives or chemical components of the system where the film is applied, and films can constitute controlled delivery systems and provide localized antioxidant activity at interfaces. In order to evaluate the ability of alginate matrices to stabilize AA, the objective of the present work was to study the effect of alginate composition and level of glycerol (plasticizer) applied to film constitution as well as of the RH (33.3; 57.7, 75.2%) used for film storage (25°C) on the hydrolytic and oxidative stability of AA in these matrices.

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2.1. Chemicals

Manugel DM and Protanal LF240 alginates were a gift from FMC BioPolymer (Billingstad, Norway). Cargill (Mechelen, Belgium) and Sigma-Aldrich (herein called "VR") alginates were also used in this study. All other chemicals were of analytical grade from Merck (Argentina) or Sigma-Aldrich (St. Louis, MO, USA). Deionized water (Milli-Q, USA) was used.

2.2. Analyses of alginates

The diadic frequency composition of alginate (F_{GG} , F_{MM} and F_{GM+MG}) or block-proportions were determined by means of circular dichroism. Spectra of samples containing ≈ 0.8 mg/mL of alginate in deionized water were recorded on a Jasco J-810 (Japan) spectropolarimeter. Data in the far UV (195-250 nm) region was collected at 25°C using a 2 mm path length cuvette. A scan speed of 20 nm/min with a time constant of 1 s was used. Each spectrum was measured four times and the data was average to minimize noise. Deconvolution of experimental spectra was done according to the procedure described by Donati et al. (2003). Mollar ellipticity was calculated using a mean residue weight value of 176.14 (the molecular weight of the monomer minus one water molecule). The diadic composition calculations were performed according to Donati et al. (2003). Based on these results the four alginates above mentioned were then selected among others for film development.

Afterwards, these four alginates were submitted to chemicals assays to determine the total acid carbohydrate content according to the spectrophotometric method of Edstrom (1969), using 4,5,4',5'-dibenzo-3,3'-diethyl-9-methylthio-carbocyanine

bromide. The protein content was determined according to Lowry et al. (1951). Methanol and acetyl contents respectively derived from methoxyl esterification of carboxylate groups and acetate ether bonding to –OH groups of the acid polysaccharides (alginates), were determined according to Wood and Siddiqui (1971) and Naumenko and Phillipov (1992), respectively. The degree of methyl esterification (DM) and acetylation (DA) of the acid polysaccharides were then calculated as:

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$$DM = 100 \cdot \frac{moles_{CH_3OH}}{moles_{total} \ acid \ carbohydra \ tes}$$

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$$DA = 100 \cdot \frac{moles_{CH_3}COO}{moles_{total} \ acid \ carbohydra \ tes}$$

Molecular weight profile of alginates was determined through gel filtration using a Fast Protein Liquid Chromatograph (FPLC, Pharmacia, Sweden) with a Superose 12HR 10/30 column (Amersham Biosciences-GE Healthcare, USA). Each alginate sample was dissolved and also eluted by using 0.5 M of imidazole buffer (pH 7.0) (Mort et al., 1991) or deionized water, at 0.5 mL/min. Dextrans of 65,000 and 40,210 molecular weights as well as blue dextran, CoCl₂ and sucrose were used as standards for column calibration at both elution conditions. A pectin of known molecular weight was used as reference to control the column performance under both elution conditions. Total carbohydrate content was determined into each collected fraction by the phenolsulfuric acid spectrophotometric method (Dubois et al., 1956) when samples were collected with 0.5 M imidazole buffer (pH 7.0), and according to the method of Edstrom (1969) when samples were collected with deionized water. The former colorimetric technique underestimated the content of alginates in each fraction.

Iron and copper contents in the alginates were directly determined through
inductively coupled plasma atomic emission spectrometry (ICP-AES), using a Thermo
Jarrel Ash Atom Scan 25 (Thermo Jarrel, USA), according to Rubio et al. (2009).

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2.3. Film formation

For the purpose of this study, each film system was developed from one of the four alginates above mentioned. A 2% (w/w) alginate concentration was used for the film making solution, thus permitting to obtain plasticized films with the adequate handling resistance. The aqueous solution was continuously stirred under controlled high speed (1,400 rpm-constant) using a vertical stirrer (LH model, Velp Scientifica, Italy) in order to reach homogeneous hydration. While stirring, the obtained viscous, homogeneous and transparent system was then heated up to 85°C at a constant heating rate (5.3 °C/min) by means of a hot plate (Velp Scientifica, Italy) and with simultaneous recording of the temperature by using a thermocouple connected to a Consort millivoltmeter (P901, Belgium). The following substances were subsequently added: glycerol [26.7, 35.6 or 52.3 g per 100g of (polymer+glycerol)] for plasticization (Yang and Paulson, 2000), potassium sorbate (0.030% w/w) as antimicrobial agent and AA (0.100% w/w). Finally, 1.1×10⁻³ moles of Ca²⁺ (as CaCl₂.2 H₂O) were added for gelling after cooling. The hot solution was placed under vacuum for 20 s to remove air bubbles and then immediately poured onto horizontally leveled polystyrene plates. The solution dispensed into each identified plate was weighted in an analytical scale (0.0001 gprecision) in order to have constant thickness as well as a known initial content of AA into the subsequently generated film. The fractionated system was dried for 2.5 hours in a forced convection oven at 60 °C. Films were also weighted after drying, peeled from

the polystyrene plates and stored in light-protected desiccators over saturated solutions of known water activity (a_{W}^{o}) , in order to maintain a constant RH for film equilibration:

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$$a_W^o = \frac{RH\%}{100}$$

The salts used were $MgCl_2$ ($a_W^\circ = 0.333$), NaBr ($a_W^\circ = 0.577$) and NaCl ($a_W^\circ = 0.752$) at 25 °C (Greenspan, 1977). Equilibration was followed by the daily measurement of a_W in the film samples until attaining the final equilibrium. Afterwards, the sample thickness was measured at six different locations in each of ten specimens by using a digital micrometer (Mitutoyo, Kawasaki, Japan).

Three batches of films (replicates) were prepared as above described. The film samples obtained from each batch were identified and distributed among the light-protected desiccators with the different RHs (33.3; 57.7 or 75.2%) and stored at 25 °C in order to establish the influence of the film making in the following determinations. Storage was first performed under vacuum (P = 130 Pa) with controlled RH in order to ensure that AA degradation begins through the irreversible hydrolysis of its lactone ring as the first and limiting reaction step (León and Rojas, 2007). Hence, the specific influence of water in the AA stability could be analyzed. On the other hand, samples of the three batches of Cargill and Sigma (VR) alginate films made with 35.6 or 52.3% of glycerol were further stored under normal air conditions (P = 1.013×10⁵ Pa), protected from light, at 25°C and 57.7% or 75.2% RH, in order to also infer the specific influence of oxygen on the total kinetic of AA destruction.

The following analyses were performed on each film sample collected from the three batches at each corresponding time, glycerol level and RH of interest.

190 2.4. Water activity

191	To evaluate film equilibration, the true water activity (awo) was determined on
192	the film samples with a Decagon AquaLab (Series 3 Water activity meter, USA) at 25
193	°C, using a calibration curve made with the standard saturated salt solutions of MgCl ₂ ,
194	NaBr and NaCl mentioned before.
195	
196	2.5. Measurement of pH
197	This was performed on the gel-forming solutions as well as on films equilibrated
198	at the corresponding RH, using a bulb-combined glass electrode or a flat surface
199	electrode (Phoenix, AZ, USA) connected to a pH meter (Consort P901, Belgium). Film
200	pH was determined after a slight surface hydration with 20.0 μL deionized water (Joel
201	et al., 1972). Standard buffer solutions (pH 4.00 and 7.02) were used for calibration.
202	
203	2.6. Determination of L-(+)-ascorbic acid (AA)
204	A film sample taken from each of the three batches of films stored at each RH
205	was carefully cut into pieces smaller than 1-mm in size, weighed on an analytical scale
206	(0.0001 g), placed into a 25.00 ml-volumetric flask with a 1%(w/v)-oxalic acid solution
207	and submitted to magnetic stirring for 1.5 h at 5 °C to achieve the total extraction of AA
208	from the film sample. During this time, it was also submitted to vortexing (Velp, Italy)
209	for 90 s at 35 Hz, every 15 min. The suspension was finally centrifuged at 10,000 rpm
210	and 6 °C for 30 min (Eppendorf 5810R, USA). An aliquot was taken from the
211	supernatant and the AA concentration was determined by using the 2,6-dichloro phenol
212	indophenol (2,6-DPIP) spectrophotometric method (Rojas and Gerschenson, 1991)
213	though xylene was not used for extraction of the remaining 2.6-DPIP. The AA

concentration was determined in two different aliquots (duplicate) for each film sample.

215	The initial amount of AA into each identified film sample was known because			
216	the solution dispensed into each plate and the corresponding film obtained after drying			
217	were both weighted as indicated above. In a previous assay, the AA concentration was			
218	spectrophotometrically determined in 10 films of three different batches ($n=30$) which			
219	were processed as described, and it was compared with the expected concentration. The			
220	recovery of AA from the films assayed to determine the optimum experimental			
221	conditions for extraction ranges from 98.9 to 104.6%. Good interday (relative standard			
222	deviation, RSD \leq 2.84%) and intraday (RSD \leq 1.98%) precision was achieved.			
223	The procedure retains its accuracy up to 81% of AA degradation kinetics. The			
224	calibration curve was constructed with nine AA concentrations ranging between 0 and			
225	34 μg/mL every time the 2,6-DPIP solution was prepared. Regression analysis of Beer's			
226	plots showed good correlation in the 0 and 34 µg/mL concentration range, showing the			
227	same regression parameters [interception= 0.616 ± 0.001 ; slope= $-(725\pm3)\times10^{-5}$;			
228	residual standard error = 8.7×10^{-6} ; $R^2 = 0.9997$]. The limit of detection of the			
229	spectrophotometric method is 0.68 µg/mL.			
230				
231	2.7. Color			
232	Measurement of the film color was performed in each sample according to the			
233	ASTM E1925 (1995) employing a Minolta colorimeter (Minolta CM-508d) with an			
234	aperture of 1.5 cm-diameter (León and Rojas, 2007). Film samples for color			
235	measurement were taken from each of the three batches of films obtained in order to			
236	determine the kinetics of browning (yellowness index, YI %) increase. Also, L , a , and b			
237	(HunterLab) color parameters were measured, which ranged from $L=0$ (black) to $L=0$			
238	100 (white or maximum) for lightness (L); $-a$ (greenness) to $+a$ (redness), and $-b$			

239	(blueness) to $+b$ (yellowness). Standard values considered were those of the white
240	background.
241	
242	2.8. Moisture or water content
243	Films were sampled after equilibration at each RH, cut into pieces smaller than
244	1-mm size, weighed (0.0001 g) and placed into small, light glass containers. Samples
245	were dehydrated in a vacuum oven at 70°C until constant weight, which involved
246	approximately 22-30 days. Determinations were performed on six film specimens at
247	each evaluated condition. Moisture or water content was informed on dry basis.
248	
249	2.9. Glass transition temperature (T_g) .
250	Modulated differential scanning calorimetry (MDSC, TA Instruments, USA)
251	was used to determine the $T_{\rm g}$ (midpoint temperature) from the second scan performed
252	on an equilibrated film sample (10–15 mg) placed into an hermetically sealed 40 μL -
253	aluminium medium pressure pan. An empty pan served as reference. Temperature was
254	brought down to -140 °C (20 °C/min) followed by a 5 min-isotherm at -140 °C. A \pm
255	0.5°C every 40 s modulation was applied. A ramp was then performed up to 40°C
256	(10°C/min), followed by a second decrease in temperature to −140°C (20°C/min), and a
257	5 min-isotherm at -140°C. Afterwards, a second ramp was performed up to 200°C
258	(10°C/min), from which the $T_{\rm g}$ value was determined. MDSC was periodically
259	calibrated with a sapphire disk, in the full temperature range at which the equipment is
260	usually employed.
261	
262	2.10 Statistical analyses

The results are reported as the average and standard deviation. Rate constants of AA destruction (k_{AA} ' and k_{T}) were calculated by linear regression according to a first order reaction, where each experimental point corresponded to the ratio between the AA concentration remaining at a given storage time t (C_{AA}) and the initial (t = 0) concentration of AA (C_{AA} °):

$$C_{AA}(t) = \frac{weight_{AA}(t)}{weight_{film}}$$

wherein the "weight" is expressed in grams.

Browning rate constants (k_{YI}) were calculated from the slope of the linear regression of experimental data (YI% vs time). Analysis of covariance (ANCOVA) was applied for comparison of slopes, that is, of the rate constants (k_{AA} ' and k_{T} , or k_{YI}), as indicated by Sokal and Rohlf (2000). The statistical analyses of results were performed by applying ANOVA (α : 0.05), followed by pairwise multiple comparisons evaluated by Tukey's significant difference test. The GraphPad Prism software (version 5.00, 2007, GraphPad Software Inc., USA) was used for all analyses previously detailed.

The effect of two quantitative factors (RH and glycerol) on the calculated rate constants (k_{AA} ' and k_{YI}) were analyzed with a complete 3×3 experimental design at the three levels described before for both factors, coded as -1, 0, +1. This design was repeated for the four polymer tested. In the first model the polymer type was included as a categorical variable, but subsequently each polymer was analyzed separately. A regression model was applied as a function of the lineal and quadratic values of the quantitative factors and their interactions. This statistical analysis was performed with R (version 2.15: R Core Team, 2012).

3. Results and Discussion

3.1. Polymer characterization

The relevant molecular characteristics of the alginate polymers used in this work are listed in **Table 1**. Proteins were not detectable. Alginates showed an acidic polysaccharide content of $\approx 95\%$ (Edstrom, 1969) and they were no methoxyl-esterified. As expected from algal alginates, O-acetyl groups were absent (Davidson et al., 1977). Similar and low amounts of iron and copper were observed (**Table 1**). Molecular weights and their distributions were similar (≈ 876 kDa). This value corresponds to a high molecular weight alginate which is reported to be related to higher viscosity (Aoyama et al., 2007). Important biophysical properties of alginates are also related to the molecular weight (Kong et al., 2004).

The high selectivity of alginate binding towards calcium ions, which accounts for its capacity to form ionotropic gels, is determined by the polymer composition (Simpson et al., 2004). Furthermore, parameters such as the stability, strength and porosity of the obtained gels are influenced by the diadic frequency composition (F_{GG} , F_{GM+MG} and F_{MM}) of alginate (Donati et al., 2003). In order to study the influence of the macromolecule structure in the development of film networks able to stabilize AA, alginates with different monomeric composition were then used in this work. Alginate composition and block-proportions can be determined by the circular dichroism characteristics of alginate molecules (Morris et al., 1980; Klöck et al., 1997; Donati et al., 2003). Circular dichroism spectra are shown in **Fig. 1**. All polymers used showed the negative MG and GG diads bands. The circular dichroism spectra of Manugel and Cargill alginates were characterized by the minima at 210 nm (\approx –1330 and –1260 molar ellipticity, respectively), whereas VR and Protanal alginates show a shallower spectra with minima at 213 nm (\approx –1050 for both alginates). These features can account

312 for the different diadic composition. According to the procedure described by Donati et 313 al. (2003), deconvolution of experimental spectra (Fig. 1) allowed calculating the diadic 314 composition (F_{GG}, F_{MM} and F_{GM+MG}), and results are shown in **Table 1**. Manugel 315 alginate was mainly constituted by GG-blocks, with lower proportion of MM-blocks. 316 Cargill alginate showed lower proportion of GG- and MM-blocks than Manugel 317 alginate, but Cargill differs mainly in its higher proportion of flexible GM+MG-blocks. 318 On the other hand, Protanal and VR alginates showed similar composition, although 319 Protanal was characterized by a higher proportion of MM-blocks and a lower one of 320 GM+MG-blocks. Contrary to polymannuronates, a high affinity of polyguluronates to calcium 321 322 ions was determined by Kohn (1975). By studying the encapsulation of βTC3 cells, 323 Simpson et al. (2004) determined that alginate with high mannuronic acid content was 324 not affected by changes in CaCl₂ concentration due to the low percentage of consecutive guluronic acid residues. A cooperative effect in calcium binding is observed for 325 326 polyguluronic acid at chain lengths above a threshold of ≈ 20 residues (Braccini and Pérez, 2001; Fang et al., 2008). The alginate fragments with alternating sequence of D-327 328 mannuronic and L-guluronic acid units (GM+MG-blocks) exert only a low selectivity in 329 ion exchange reaction, whereas the affinity of the monomers (D-mannuronate, L-330 guluronate) to calcium ions was found to be virtually the same (Kohn, 1975). GG-331 blocks are the most inflexible ones in alginate macromolecules, whereas GM+MG-332 blocks are the most flexible. Chain breakage by oxidants was demonstrated to occur 333 mainly at the most flexible blocks of the alginate macromolecules, whereas the GGblock length largely determines the elastic modulus of calcium cross-linked gels (Kong 334 et al., 2004). In the present work, the amount of Ca²⁺ required for gelling of the film 335 336 making solutions was then calculated from the proportion of GG-blocks, being it

337	reported in Table 1 . For film formulations, 1.1×10° moles of Ca ² were then used in
338	order to satisfy a minimum requirement for all alginates. This content also permits to
339	obtain films with an adequate handling flexibility, especially at the lowest level of
340	glycerol used for plasticization.
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342	3.2. Film characteristics
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344	Homogeneous and flexible films plasticized by glycerol proportions of 26.7,
345	35.6 or 52.3%w/w were obtained after casting from each alginate solution. Films were
346	transparent, almost colorless or yellowish ($b = +6$ to $+9$; YI = 12-18%) and showed high
347	initial lightness (Table 2). The AA concentration initially determined (C_{AA}^{o}) was \approx
348	3.02×10 ⁻² g AA per g of film, which means that a 100% of AA recovery was achieved
349	after casting. Temperature should be as low as possible to get short periods of drying (≤
350	2.5 hours), which avoid AA losses through hydrolysis during this processing. Therefore,
351	films were finally dried at 60°C. Film samples attained equilibration at 20 hours of
352	vacuum storage at each RH, as determined by measurement of the film a_{W}^{o} (0.333,
353	0.577 and 0.752, respectively) at 25°C. Thickness measured after equilibration was \approx
354	0.12 mm (Table 2). There was not significant influence of RH and glycerol content on
355	film thickness. The film pH recorded along storage varied as indicated in Table 2.
356	Moisture contents increased with the RH of film equilibration. In general, the increase
357	in the glycerol level produced a significant increase in the moisture content only for
358	films equilibrated at 75.2% RH (Table 3).
359	At $\approx -38^{\circ}$ C and/or 0°C, MDSC scans did not show any endothermic peak that
360	could correspond to freezable bound and free water, respectively (Hatakeyama and
361	Hatakevama 1998) Therefore water gained from the storage environment was

362	adsorbed or retained by the polymeric network. The $T_{\rm g}$ values found for all equilibrated
363	films herein studied were lower than the storage temperature (25°C) (Table 3). Hence,
364	the equilibrated films were amorphous rubber materials at ambient temperature. Into
365	each type of alginate assayed, $T_{\rm g}$ values in general decreased significantly ($p < 0.05$)
366	with the increase in the glycerol proportion used as well as in the water content (Table
367	3). Hence, glycerol as well as the water captured during storage plasticized the film
368	networks. At each level of glycerol, Manugel alginate films showed, in general, the
369	highest values of $T_{\rm g}$ and, hence, the lowest macromolecular mobility. Probably, this
370	result may be associated with its higher proportion of inflexible GG-blocks and/or with
371	a very small proportion of flexible GM+MG-blocks (Table 1). According to Roger et
372	al. (2004), powder samples of alginate exhibited $T_{\rm g}$ ranging from 95°C to 136°C and no
373	significant effect on $T_{\rm g}$ was observed for different molecular weight samples. However,
374	an increase in $T_{\rm g}$ values with the G content was observed. This effect was attributed to
375	the presence of residual Ca2+ ions in the alginate powder, crosslinking oligomeric G-
376	rich chains.
377	Alginates are block copolymers and, hence, they can behave as two-phase
378	systems or physical blends. Each phase exhibits its own distinct $T_{\rm g}$ (Ferry, 1980). Only
379	one $T_{\rm g}$ was detected in thermograms of alginate films developed in the present work.
380	This could be attributable to a plasticization effect and/or to a probable random
381	alternating distribution of blocks in the alginate macromolecules.
382	
383	3.3. Stability of L-(+)-Ascorbic Acid to Chemical Hydrolysis in Films
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385	The study of AA stability by storage in the absence of air $(P = 130 \text{ Pa})$ allowed
386	to determine that the ratio between the remaining AA concentration $[C_{AA}(t)]$ and the

initial one $[C_{AAO}]$ statistically changed with the storage time (t) according to a pseudofirst order (p < 0.05) kinetic law (Leon and Rojas, 2007). The rate constants of AA hydrolysis (k_{AA}) were then calculated from the slope obtained after fitting a straight line to the data. On the other hand, browning development was measured as the increment of the YI with time, which statistically fitted (p < 0.05) to a pseudo-zero order reaction (Rojas and Gerschenson, 2001). Browning rate constant (k_{YI}) was then calculated from each slope obtained after linear regression fitting to the experimental data. The AA stability to hydrolysis (k_{AA}) values seemed to be mainly affected by the glycerol level as well as by the RH of film storage at 25°C, as shown in the example depicted in **Fig. 2**. Similar conclusions were drawn from comparison of k_{YI} values.

Collected $k_{\rm AA}$ ' data was analyzed by an experimental design of two quantitative factors (RH and glycerol) at the three levels described before, coded as -1, 0, +1. A regression model was applied to analyze $k_{\rm AA}$ ' as a function of the linear and quadratic values of the quantitative terms and their interactions. In a preliminary analysis, the type of polymer was considered as a third quantitative factor which differed in the frequency composition of each alginate ($F_{\rm GG}$, $F_{\rm MM}$ and $F_{\rm GM+MG}$) applied to film development. It was observed that AA hydrolysis was only affected by a significant interaction between Protanal (p < 0.001) or VR (p < 0.05) and the alginate diadic composition and glycerol levels. Hence, only RH and glycerol were considered finally as quantitative factors and separated models were built for each type of polymer.

The statistical results are reported in **Table 4**. The experimental design of RH and glycerol factors indicated that the rate constant of AA hydrolysis (k_{AA} ') significantly (p < 0.05) increased as a consequence of the separated increase in RH or glycerol content, when AA was compartmentalized in Protanal or VR alginate networks. It has been suggested that the previous presence of glycerol permits or

412 facilitates the penetration of water into the polymeric network during storage (Pérez et 413 al., 2009). On the other hand, k_{AA} only increased significantly (p < 0.05) with the RH 414 of film storage when AA was supported either in Manugel (p < 0.05) or Cargill (p < 0.05) 0.001) alginate network. The dependence was also significant (p < 0.05) for the 415 416 quadratic term of the RH factor for Cargill alginate films. The proportion of glycerol 417 used for plasticization did not affect the AA stability in Manugel or Cargill alginate 418 film. The highest proportion of GG-block in Manugel followed by Cargill alginate 419 produces ordered templates for polymer chain associations mediated by Ca²⁺ 420 crosslinking between neighboring macromolecules (Braccini and Pérez, 2001). 421 Chandrasekaran et al. (1988) indicated that glycerol can produce disturbance of filament 422 aggregation in the case of gellan polymer, which may also be extended to Manugel and 423 Cargill alginate films. However, zipping of GG-block chains together by calcium ions 424 may overcome the glycerol effect in these films. As previously mentioned, GG-block 425 length determines the elastic modulus of calcium cross-linked alginate gels (Kong et al., 426 2004). 427 A somewhat higher hydrolytic stability of AA supported in Manugel or Cargill 428 alginate films is observed by plotting the rate constants of AA hydrolysis (k_{AA}) versus 429 glycerol or RH linear factor (Fig. 3), especially by storage at 33.3% of RH but also at 430 75.2%. Hence, alginates with a predominant proportion of GG-blocks showed a higher 431 ability to stabilize AA against hydrolysis. This effect could be associated with their 432 higher capability to immobilize water by physical retention, as previously demonstrated 433 for gellan films (León and Rojas, 2007). As mentioned above, guluronate chain pairing 434 junction zones also involve water molecules (Braccini and Pérez, 2001), which 435 correspond to highly adsorbed or non-freezable bound water (Ping et al., 2001).

436 Water is responsible for hydrolysis and the irreversible opening of the lactone 437 ring of the AA molecule, producing 2-keto-L-gulonic acid (Kurata and Sakurai, 1967). 438 Hence, at constant temperature (25°C), k_{AA} ' depended on the RH factor because k_{AA} ' is 439 the product of the true second order rate constant for AA hydrolysis (k) and the 440 concentration of water available for reactions (C_{WATER}) (León and Rojas, 2007). This 441 kind of water is that loosely retained by the solid-like film network. As RH of film 442 equilibration increases, the polymeric network leaves higher proportion of loosely 443 adsorbed water, which is available for chemical reactions. This condition also promotes 444 the parallel development of browning reactions from 2-keto-L-gulonic acid. 445 The half-life times $(t_{1/2})$ of the AA supported in the alginate films were 446 calculated from the values for k_{AA} . In the most favorable condition of RH (33.3%), $t_{1/2}$ 447 values ranged between 10 and 16 months for AA supported in Manugel and Cargill 448 alginate films, a result not affected by the glycerol level, and between 3 and 11 months 449 in Protanal and VR alginates, for decreasing proportions of glycerol. At 57.7% RH, the 450 values of $t_{1/2}$ were in general no lower than 2 months. At 75.2% RH, the AA supported 451 in Manugel alginate films showed a $t_{1/2} \approx 27$ days for all glycerol levels, whereas in VR 452 and Protanal films, the $t_{1/2}$ decreased from 32 to 14 and from 32 to 9 days, respectively, 453 as glycerol level increased. 454 The rate constants of browning development (k_{YI}) were also analyzed through the experimental design applied to AA degradation kinetics, with the polymer type as a 455 456 categorical variable. The results indicated that k_{YI} significantly (p < 0.01) increased in a 457 linear trend with the RH of film storage and glycerol proportion for all polymers 458 assayed (Table 4), excepting for VR alginate films. In the latter system, browning 459 kinetic was only dependent (p < 0.01) on the RH of storage. Significant dependence of 460 $k_{\rm YI}$ on the RH in Cargill (p < 0.01) and Protanal (p < 0.05) alginate films was also

461	observed in a quadratic term. An interaction between RH and glycerol was also detected
462	for Protanal films. Response surfaces were then plotted (Fig. 4). They allowed us to find
463	the best conditions for minimal browning, which corresponded to a 41-44 % RH for
464	storage and 29% w/w of glycerol content for plasticization, whereas the highest values
465	of k_{YI} were observed at the highest RH of storage and glycerol content in films (Fig. 4).
466	By plotting the rate constants of browning (k_{YI}) versus glycerol or RH lineal
467	factor (Fig. 3), no clear tendencies towards slower browning were observed in film
468	systems. In general, lower $k_{\rm YI}$ values were obtained for Manugel alginate films at
469	increasing RH and glycerol levels.
470	Despite the different kinetic order, k_{YI} correlated significantly (Pearson's
471	correlation coefficient $r = 0.8731$; $p < 0.001$) with the k_{AA} ' values.
472	
473	3.4. Stability of L-(+)-Ascorbic Acid to Chemical Hydrolysis and Oxygen in Films
474	Films respectively made with Cargill or VR alginate using the two highest
475	glycerol proportions were also studied in their ability to stabilize AA in the presence of
476	oxygen. Storage was performed at 57.7 or 75.2% RH (25°C) under normal air pressure
477	(P=1.013×10 ⁵ Pa). Hence, the oxygen partial pressure (p_i) was 0.21 atm constant during
478	storage. Under these conditions, a pseudo-first order kinetics could be fitted to the
479	experimental data of AA concentration ($p < 0.05$) in a manner similar to that previously
480	observed in Fig. 2 for AA loss in alginate films stored under vacuum. AA destruction in
481	the presence of oxygen occurred simultaneously to the hydrolytic reaction previously
482	studied under vacuum storage of films (Kurata and Sakurai, 1967). It can be then
483	considered that at least two irreversible parallel or competitive reactions proceed: the
484	AA hydrolysis (k_{AA}) and the AA oxidation (k_{AA}) , which can be expressed as a

- differential kinetic equation written for the AA as the reagent, in the form of pseudo-
- 486 first-order rate reactions:

487
$$r_{AA} = -\frac{1}{v_{AA}} \frac{dC_{AA}}{dt} = k'_{AA} \cdot C_{AA}(t) + k^{OX}_{AA} \cdot C_{AA}(t)$$
 (1)

- wherein v_{AA} is the stoichiometric coefficient for AA hydrolytic reaction, r_{AA} is the AA-
- reaction rate/unit volume at a constant temperature, $C_{AA}(t)$ is the AA concentration
- 490 remaining at time t, k_{AA} ' is the rate constant of the pseudo first order kinetics for AA
- 491 hydrolysis, k_{AA}^{OX} is the oxidation rate constant of AA.
- 492 By integration ($v_{AA} = 1$), results:

493
$$C_{AA} = C_{AA}^{O} \cdot \exp[-(k'_{AA} + k_{AA}^{OX}) t]$$

- Hence, the slope calculated from the experimental data obtained after storage under air
- 495 give the total rate constant (k_T) :

$$k_T = k'_{AA} + k_{AA}^{OX}$$
 (2)

- 497 and the oxidation rate constant (k_{AA}^{OX}) can be obtained as the arithmetic difference. For
- oxygen partial pressures lower than 0.40 atm, the apparent rate constant (k_{AA}^{OX} ; eq. 1
- and 2) involved the product between the true kinetic rate constant of oxidation (only
- dependent on temperature) and the oxygen concentration, related to the p_i (Khan and
- 501 Martell, 1967).
- In general, film systems stored under air did not show significant differences
- between $k_{\rm T}$ and $k_{\rm AA}$ values (**Table 5**). Higher $k_{\rm T}$ values were only observed for Cargill
- alginate film formulated with 35.6% glycerol and stored at 57.7 or 75.2% RH. Even in
- film systems where a non significant difference between k_T and k_{AA} was observed,
- browning rate constants (k_{YI}) determined under air storage at 75.2% RH were in general
- 507 higher than the k_{YI} values found under vacuum (**Table 5**). It can be concluded that, in
- 508 general, the alginate film networks seemed to effectively preserve AA from oxidation.

4	$\boldsymbol{\alpha}$	1		
4.	Co	ncl	lusio	ns

Water is the factor responsible for AA hydrolysis, and glycerol may facilitate
water penetration from the environment into the polymeric network. In the presence of
Ca^{2+} , alginates with higher proportion of GG-blocks (F _{GG} = 0.66) and lower one of
MM- and, mainly, of GM+MG flexible blocks, generate film networks that immobilize
water sufficiently to reduce the degradation of hydro-sensitive biomolecules such as
AA. When comparing the hydrolytic with the total rate constant of AA destruction
under air, it was observed that even at less favorable conditions of RH and glycerol
levels, both GG and GM+MG enriched alginate networks in general preserve AA from
oxidation. It also demonstrated that hydrolysis is the principal way by which AA is lost
when supported in films and, hence, water immobilization is a key factor to be
controlled.

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674	Figure captions
675	
676	Fig. 1. Circular dichroism spectra recorded for (— black) Protanal, (— thick orange
677	line) VR, (—blue) Cargill and (—red) Manugel alginates.
678	
679	Fig. 2. Kinetics of AA hydrolysis determined in Cargill alginate films are shown for
680	two levels of glycerol and three levels of storage relative humidity (RH).
681	
682	Fig. 3. Rate constants of AA hydrolysis (k_{AA} ') are plotted against glycerol (glyc.n) (A)
683	or relative humidity (RH.n) (B) linear factor. Idem for the rate constants of browning
684	development (k_{YI}): (glyc.n) (C) and (RH.n) (D).
685	
686	Fig. 4. Relative humidity (RH) and glycerol content influences on the rate constant of
687	browning development ($k_{ m YI}$) are plotted as response surfaces for Protanal (A) and
688	Cargill (B) alginate films.
689	
690	

690 **Table 1**691 Chemical composition of the alginate polymers used for film development.

692

		Alg	inate	X
	Manugel	Cargill	VR	Protanal
Molecular weight ^a (kDa)	876 ± 180	876 ± 200	876 ± 140	876 ± 180
Protein content ^a (g / 100 g) ^f	0.10 ± 0.09	0.59 ± 0.08	0.5 ± 0.3	0.03 ± 0.06
Total acid carbohydrates ^a (g / 100 g) ^f	95.9 ± 0.8	93.0 ± 0.6	97.05 ± 0.07	95.6 ± 0.4
DM ^b (%)	0.10	0.10	0.08	0.5
DA ^c (%)	ND	ND	ND	ND
Iron ^a (mg/1000 g) ^f	45 ± 4	39 ± 6	34± 6	36 ± 5
Copper ^a (mg/1000 g) ^f	38 ± 7	42 ± 8	24 ± 7	29 ± 5
F _{GG} d	0.66	0.57	0.27	0.25
$F_{MM}^{}$	0.26	0.22	0.32	0.42
F_{GM+MG}^{d}	0.08	0.21	0.40	0.33
F_G^{d}	0.70	0.67	0.47	0.42
F_M^{d}	0.30	0.33	0.53	0.58
Ca ²⁺ required e (mol/100 g) f	2.85×10^{-3}	2.45×10^{-3}	1.18×10^{-3}	1.08×10^{-3}

^a Mean and standard deviation (n = 3) are shown.

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^b Degree of methyl esterification is expressed as 100 × moles of methoxyl group / moles of total acid carbohydrates.

^c Degree of acetylation is expressed as 100 × moles of acetyl group/moles of total acid carbohydrates.

d Diadic frequency composition of GG-, MM- and GM+MG-blocks [guluronic (G); mannuronic (M)] in alginates determined through circular dichroism (Donati et al., 2003). F_G and F_M are the total proportions of G and M monomers, respectively.

^e moles of Ca²⁺ required per 100 g of film making solution calculated from the respective F_{GG} value.

^f Expressed per 100 g or 1000 g of alginate.

ND: non detectable.

Table 2

Color parameters^{a,b} and thickness^{c,d} are reported as well as the pH^a variation recorded 704 during the complete period of film storage,. 705

706

703

Alginate	YI %	L %	+ <i>b</i>	Thickness (mm)	рН
Manugel	17 ± 1	82 ± 1	7.9 ± 0.3	0.100 ± 0.040	4.42 ± 0.07
Cargill	12 ± 2	85 ± 1	6.3 ± 0.5	0.100 ± 0.030	4.54 ± 0.08
VR	16 ± 2	80 ± 1	6.1 ± 0.6	0.140 ± 0.070	4.37 ± 0.03
Protanal	18 ± 3	80 ± 3	9 ± 1	0.110± 0.030	4.61 ± 0.03

^a Mean and standard deviation ($n \ge 27$) are shown. ^b Yellowness index (YI), lightness (L) and b (blue–yellow component) recorded initially.

^c Mean and standard deviation ($n \ge 11$) are shown.

^d It was measured after film equilibration at each relative humidity (HR) and 25°C.

715 Table 3 716 Moisture content and glass transition temperature (Tg) determined after film 717 equilibration at each relative humidity (RH) of storage (25°C).

Alginate	Glycerol	RH	Moisture content ^a	$T_{ m g}^{\ m b}$
1118111111	(% w/w)	(%)	(g water / g dm)	(C)
		33.3	14.3 ± 0.9	-40.06
Manugel	26.7	57.7	22.9 ± 0.1	-44.18
_		75.2	27.1 ± 0.1	-66.83
		33.3	17 ± 2	-59.05
Cargill	26.7	57.7	23.42 ± 0.09	-62.57
		75.2	29.8 ± 0.1	-64.37
		33.3	15.3 ± 0.4	-61.70
VR	26.7	57.7	21.4 ± 0.1	-70.57
		75.2	31.3 ± 0.4	-72.66
		33.3	17 ± 2	-63.82
Protanal	26.7	57.7	22.8 ± 0.7	−71.36
		75.2	27.9 ± 0.1	-72.46
		33.3	16.3 ± 0.2	-53.14
Manugel	35.6	57.7	23.9 ± 0.4	-65.72
		75.2	34.1 ± 0.4	−71.4
		33.3	16.0 ± 0.4	-58.00
Cargill	35.6	57.7	23.9 ± 0.4	-66.36
		75.2	28.77 ± 0.07	−72.95
		33.3	17.1 ± 0.5	-63.50
VR	35.6	57.7	23.7 ± 0.3	-71.83
		75.2	37.9 ± 0.4	-75.11
		33.3	15.8 ± 0.4	-63.81
Protanal	35.6	57.7	23.5 ± 0.9	-73.92
		75.2	33.1 ± 0.1	-75.66
		33.3	17 ± 1	-63.37
Manugel	52.3	57.7	25.1 ± 0.3	− 75.21
		75.2	35.9 ± 0.6	-84.14
		33.3	16.3 ± 0.7	-68.05
Cargill	52.3	57.7	24.71 ± 0.08	-75.21
		75.2	35.7 ± 0.3	-84.53
		33.3	16.8 ± 0.4	-73.62
VR	52.3	57.7	25.5 ± 0.7	-77.00
		75.2	38.3 ± 0.2	-89.71
		33.3	17.1 ± 0.2	-74.19
Protanal	52.3	57.7	24.2 ± 0.2	-76.25
-	ndard deviation (75.2	39 ± 3	-88.43

^aMean and standard deviation (n = 6) are shown. ^bMean is shown. SD is not reported because it is lower than 1% of the $T_{\rm g}$ value. dm: dry mass.

721 Table 4 722 Results of the statistical analysis are summarized for the rate constants of AA hydrolysis (k_{AA}) and subsequent browning development (k_{YI}) . 723

724

-	Manugel	Cargill	VR	Protanal				
-								
RH.n	0.00148	0.000516	0.00791	0.0104				
RH.n ²	0.46283	0.012284	0.23020	0.0726				
glyc.n	0.13367	0.763096	0.04400	0.0227				
glyc.n ²	0.12316	0.098638	0.63613	0.3497				
RH.n:glyc.na	0.28204	0.209220	0.10240	0.1099				
Residual standard error	1.655×10^{-6}	1.534×10^{-6}	3.583×10^{-6}	5.72×10^{-6}				
Multiple R ²	0.9789	0.9901	0.9520	0.9563				
F-test probability	0.0102	0.003335	0.03419	0.02985				
	$k_{ m YI}$							
RH.n	0.000963	0.0013	0.00727	0.000153				
RH.n ²	0.262219	0.0071	0.22641	0.003147				
glyc.n	0.016522	0.0343	0.43724	0.005224				
glyc.n ²	0.511434	0.7844	0.69191	0.474943				
RH.n:glyc.n	0.074762	0.0852	0.49912	0.011385				
Residual standard error	4.49×10^{-5}	3.142×10^{-5}	1.158×10^{-4}	4.195×10 ⁻⁵				
Multiple R ²	0.9856	0.9981	0.9396	0.9960				
F-test probability	0.005814	0.004774	0.04773	0.000849				

⁷²⁵ 726 In no case was the interaction between RH and glycerol level significant (p < 0.05) for AA hydrolytic rate constants.

727 728

Relative humidity (RH) or glycerol (glyc) linear (RH·n; glyc·n) and quadratic (RH·n²; glyc·n²) factors. Bold numbers highlight significance (p < 0.05).

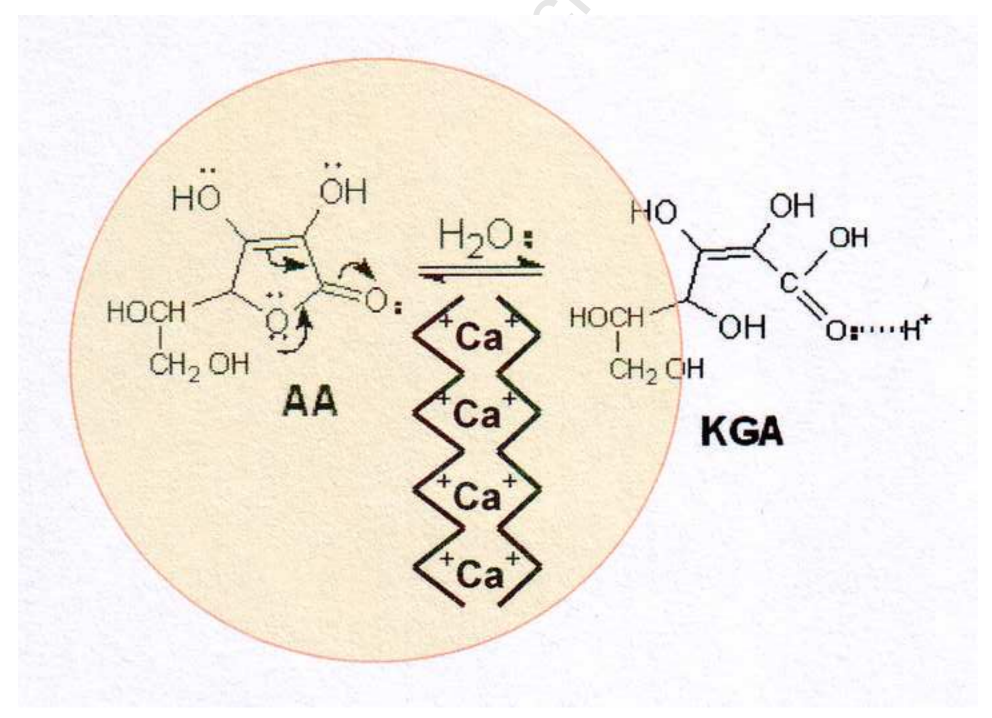
Table 5

Rate constants^a of AA hydrolysis (k_{AA}) or hydrolysis and oxidation (k_T)^b, as well as of

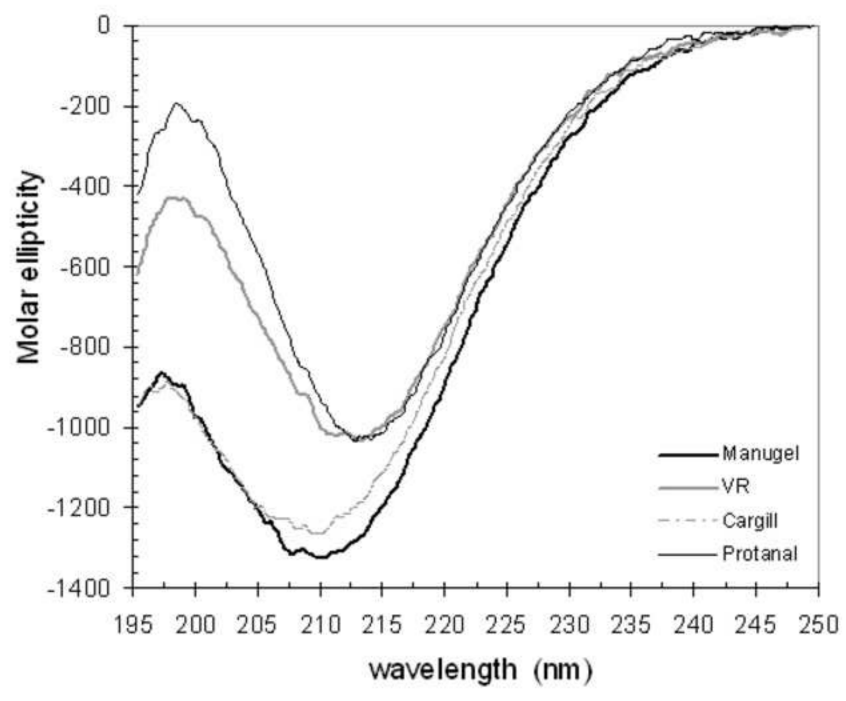
browning development (k_{YI}) at 25°C, are reported.

	Okasanal	Dalation	Storage without air		Storage under air	
Alginate	Glycerol		k_{AA} ' ×10 ⁵	$k_{\rm Yl} \times 10^4$	<i>k</i> _T ×10 ⁵	$k_{\rm YI} \times 10^4$
	(% w/w)	% w/w) humidity (%)	(min ⁻¹)	(YI%•min ⁻¹)	(min ⁻¹)	(YI%-min ⁻¹)
Cargill 35.6	35.6	57.7	0.32 ± 0.01	2.2 ± 0.2	1.07 ± 0.05	$3.33 \pm \ 0.09$
		75.2	1.97± 0.08	$10.3 \pm\ 0.5$	2.68 ± 0.03	$10.7 \pm\ 0.4$
VR	35.6	57.7	1.02 ± 0.07	3.3 ± 0.3	0.98 ± 0.08	3.6 ± 0.1
		75.2	2.7 ± 0.1	5.8 ± 0.4	3.1 ± 0.3	11.1 ± 0.9
Cargill	52.3	57.7	0.79 ± 0.04	2.9 ± 0.1	0.78 ± 0.06	3.0 ± 0.2
		75.2	2.8 ± 0.2	6.1 ± 0.5	2.86 ± 0.04	8.0 ± 0.3
VR	52.3	57.7	1.06 ± 0.07	3.4 ± 0.2	1.04 ± 0.08	3.1 ± 0.2
		75.2	3.4 ± 0.1	7.2 ± 0.3	3.7 ± 0.3	10.2 ± 1

^a Mean and standard deviation (n > 21) are shown. ^b $k_{\rm T}$ is the total rate constant of AA oxidation (eq. 2). 734



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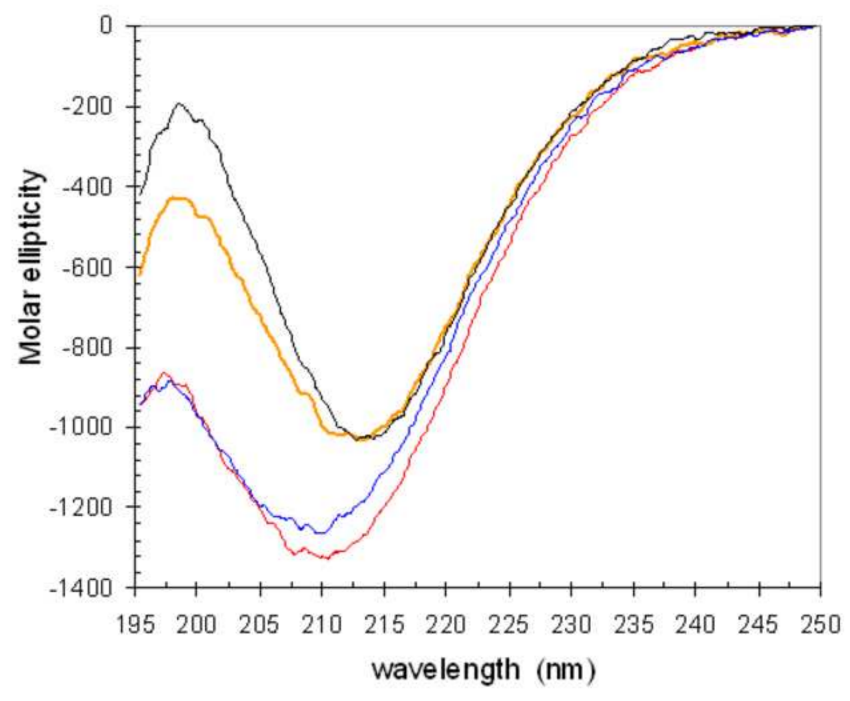


Fig. 2

