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Exploitation of κ -carrageenan aerogels as template for edible oleogel preparation.

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Abstract

In the current research, oleogels were prepared by using κ-carrageenan aerogels as template. In particular, hydrogels containing increasing concentration (0.4, 1.0, and 2.0% w/w) of κ-carrageenan were firstly converted into alcoholgel and subsequently dried by using supercritical CO₂ to obtain aerogels. The latter were porous and structurally stable materials with high mechanical strength. The polymer content affected the aerogel structure: increasing the initial κ-carrageenan concentration a coarser structure with larger polymer aggregates was obtained. However, the aerogel obtained at intermediate polymer concentration resulted the firmest one, probably due to the formation of a less aerated and more isotropic structure. Aerogels demonstrated a reduced capacity of water vapor sorption, remaining glassy and porous at room temperature at relative humidity lower than 60%. Aerogels showed a good capacity of oil absorption. The maximum oil loading capacity (about 80 %) was obtained for aerogel containing the highest κ-carrageenan content. Thus, it can be concluded that aerogels based on the structuring of water soluble polymers have potential as material for oil absorption and delivery.

Keywords	oleogel; hydrogel; κ-carrageenan; structure; supercritical CO ₂ drying; sorption kinetics
Taxonomy	Carrageenans, Aerogels
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Dear Editor,

I would like to submit to your attention the manuscript entitled “Exploitation of κ -carrageenan aerogels as template for edible oleogel preparation” (Lara Manzocco, Fabio Valoppi, Sonia Calligaris, Francesco Andreatta, Sara Spilimbergo, Maria Cristina Nicoli for consideration for publication on Food Hydrocolloids.

Oleogels result from liquid oil entrapment in a three-dimensional network without modifying the chemical characteristics of the oil. Although oleogelation is a recent research topic, the possibility to structure oil into self-standing structured solids has received considerable attention in the last decade due to their high potential number of applications of food area.

In the current research, oleogels were prepared by using κ -carrageenan aerogels as template. In particular, hydrogels containing increasing concentration of κ -carrageenan were firstly converted into alcoholgel and subsequently dried by using supercritical CO₂ to obtain aerogels. The latter were porous and structurally stable materials with high mechanical strength. Aerogels showed a good capacity of oil absorption. The maximum oil loading capacity (about 80 %) was obtained for aerogel containing the highest κ -carrageenan content. Thus, it can be concluded that aerogels based on the structuring of water soluble polymers have potential as material for oil absorption and delivery.

We would greatly appreciate your comments on the paper.

Best regards
Sonia Calligaris

Dear Editor,

Please find the revised version of our manuscript (FOODHYD_2017_271). We have endeavoured to take into account or to respond to the Reviewer's comments as indicated below.

We hope that this response is satisfactory and that the manuscript will be suitable for publication in Food Hydrocolloids.

Best regards,
Sonia Calligaris

Reviewer 1

This work is interesting and worthy of publication but the absence of key references makes this reviewer concerned about the work. Firstly, the foam-templating approach was first reported by Patel et al. in 2013, are referenced by the authors. The approach was established there and shown to work. The authors extend the approach to Xanthan gum here. However, for some reason, the authors fail to quote key recent references on foam-templated cellulosic xerogels used to bind oil and stabilize peanut butter and cookie creams, both published in Food Hydrocolloids in 2016. Copies are enclosed with this review.

We thank the reviewer for his/her appreciation of the topic. We also agree with the reviewer that the reference from Tanti et al. (2016) should have been properly quoted. For this reason we added it in the text (lines 59-61). It is our feeling that the proposed approach is different from that reported in the literature since we considered oil absorption by aerogels and not by xerogels, as performed by Patel et al. 2013 and the two papers of Tanti et al., 2016. Please note that Xanthan gum was not used in our experiments, which were developed by using k-carrageenan.

Moreover, the authors also discuss oil binding as a two-step process, without quoting another key reference which has previously used this approach to model oil binding in oleogel systems (Blake et al, 2014).

Blake et al. (2014) was actually cited in the R&D session (line 404). As suggested by the reviewer, mention to this paper was also added in the M&M section (line 267).

Reviewer 2

I congratulate for this very new approach of oleogel preparation. Although it seems hard to apply for actual productions, it is quite novel an approach. Furthermore, the study was well planned and carried out. It is worth to be published.

We thank the reviewer for his/her comments. As the reviewer pointed out, more work needs to be done to apply this approach for an actual production. Mention to this need was reported in the conclusion section of the manuscript (lines 449-451).

Reviewer 3

The paper contains a sufficient work in term of analysis of the k-carrageenan aerogels and the subsequent conversion into oleogels. The authors applied supercritical carbon dioxide drying to obtain the aerogels. They converted the hydrogels with increasing concentration (0.4, 1.0, and 2.0% w/w) of k-carrageenan into alcoholgel and afterwards they dried them.

In the introduction the authors stated that supercritical carbon dioxide drying shown several advantages in terms of product quality compared to air and freeze drying. These assumptions could be accepted considering that several studies have been published so far claiming the potential of supercritical carbon dioxide drying. However, it is completely unacceptable the lack of the experimental design presented in this study. The authors just applied one pressure and one temperature (11 MPa and 45 °C) to obtain aerogels at different concentrations of k-carrageenan. How did they choose this conditions? As the process is so innovative, I suggest to test several conditions of pressure and temperature to investigate the effect of the process on the microstructure and oil absorption of the product.

In materials and methods, the authors wrote that the drying was performed “....at 3.5 NL/min using a micrometric valve. After 3 h of drying, the outlet flow rate was increased to 5.0 NL/min for additional 4 h, maintaining the same pressure. Finally, flow rate was increased to 6.0 NL/min for 1 h.” Why did they use this drying procedure? Did they find references supporting it? Probably no differences (or significant differences) could be observed drying at 3.5 NL/min or 6.0 NL/min or combining several carbon dioxide flow rates. But the authors needed to prove it.

We thank the reviewer for his/her suggestions and comments. The conditions adopted in the experiments were selected based on different considerations:

Pressure and temperature above the critical point of carbon dioxide were selected based on pilot plant performance. Flow rate program was selected on the basis of preliminary trials. The latter were performed in a wide range of flow rate from 2.0 to 8.0 NL/min. The adopted program was selected since associated with short drying time while guarantee the structural integrity of the sample. Excessive flow rates in the initial part of the drying process actually produced large cracks on the sample surface and were discarded. By contrast, initial flow rate was kept low and progressively increased up to the final steps of drying. This information was added in the manuscript (lines 140-143 and lines 151-156)

In materials and methods, Table 1 could be erased and included in the text (paragraph 2.2 hydrogel preparation).

We agree with the reviewer and text was modified accordingly (lines 123-125).

In results and discussion, it was stated that “The removal of ethanol resulted complete after 6, 7, and 8 h of drying for samples containing 0.4, 1.0 and 2.0 % (w/w) k-C.”

How did the authors assume that drying was complete after 6, 7, and 8 h? What did they measure? I suggest to include the drying kinetics to show the efficiency of the process at different conditions of pressure and temperature. An optimization of the process parameters will highly increase the scientific value of the manuscript.

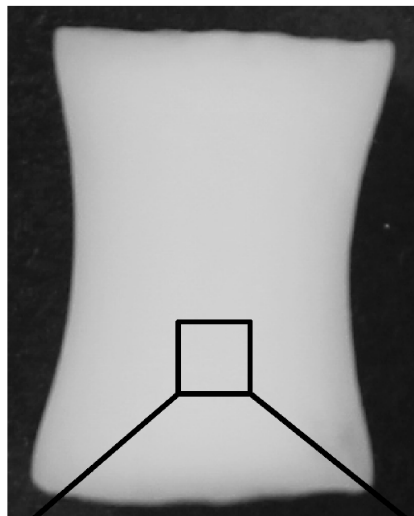
The drying kinetics were assessed recording the evolution of ethanol concentration over time using a digital alcoholmeter (see Material and Method section, lines 160-161). Drying kinetics were added as supplementary information (Figure S1).

Hydrogel



Solvent substitution
+
Supercritical
CO₂ drying

Aerogel

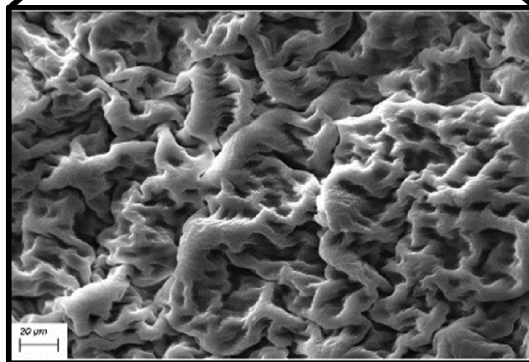


Oil
absorption

Oleogel



Oil loading
capacity up to
80% (w/w)



1 **Exploitation of κ -carrageenan aerogels as template for edible oleogel**
2 **preparation**

3

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25 **Abstract**

26 In the current research, oleogels were prepared by using k-carrageenan aerogels as template. In
27 particular, hydrogels containing increasing concentration (0.4, 1.0, and 2.0% w/w) of k-
28 carrageenan were firstly converted into alcoholgel and subsequently dried by using supercritical
29 CO₂ to obtain aerogels. The latter were porous and structurally stable materials with high
30 mechanical strength. The polymer content affected the aerogel structure: increasing the initial
31 k-carrageenan concentration a coarser structure with larger polymer aggregates was obtained.
32 However, the aerogel obtained at intermediate polymer concentration resulted the firmest one,
33 probably due to the formation of a less aerated and more isotropic structure. Aerogels
34 demonstrated a reduced capacity of water vapor sorption, remaining glassy and porous at room
35 temperature at relative humidity lower than 60%. Aerogels showed a good capacity of oil
36 absorption. The maximum oil loading capacity (about 80 %) was obtained for aerogel
37 containing the highest k-carrageenan content. Thus, it can be concluded that aerogels based on
38 the structuring of water soluble polymers have potential as material for oil absorption and
39 delivery.

40

41

42 **Keywords:** oleogel; hydrogel; κ-carrageenan; structure; supercritical CO₂ drying; sorption
43 kinetics

44

45 **1. Introduction**

46 Oleogels result from liquid oil entrapment in a three-dimensional network without modifying
47 the chemical characteristics of the oil. Although oleogelation is a recent research topic, the
48 possibility to structure oil into self-standing structured solids has received considerable
49 attention in the last decade since they have been proposed as hydrogenated/saturated fat
50 replacers, oil migration inhibitors, oil binders, and oxidation protective systems (Da Pieve,
51 Calligaris, Panozzo, Arrighetti, & Nicoli, 2011; Patel et al., 2014; Stortz & Marangoni, 2013;
52 Yilmaz & Ogutcu, 2015; Zetzl, Marangoni, & Barbut, 2012; Zulim Botega, Marangoni, Smith,
53 & Goff, 2013).

54 The simplest approach to oil gelation is based on the formation of crystalline networks by self-
55 assemble lipid additives (Co & Marangoni, 2012) or by networking of chemically modified
56 biopolymers such as ethyl cellulose and hydrolyzed chitin (Co & Marangoni, 2012; Laredo,
57 Barbut, & Marangoni, 2011; Nikiforidis & Scholten, 2015). However, oleogels could also be
58 generated by absorption of liquid oil into a porous template made of a dried polymeric network
59 of gelatin, xanthan gum, methylcellulose and hydroxypropyl methylcellulose (Patel &
60 Dewettinck, 2016; Patel, Schatteman, Lesaffer, & Dewettinck, 2013; Tanti, Barbut, &
61 Marangoni, 2016a, 2016b). To this aim, the polymer is pre-hydrated to form a hydrogel. The
62 latter is then dried to block the polymer network and obtain a porous material that can uptake
63 oil. However, this procedure is hardly applied due to structural collapse during hydrogel drying.
64 Traditional air drying is actually unable to prevent hydrogel collapse due to the formation of
65 liquid-vapor menisci in the gel pores. This produces a capillary pressure gradient that causes
66 pore collapse, leading to xerogel materials with limited oil sorption capacity (Scherer & Smith,
67 1995). Similarly, freeze-drying of hydrogels causes intense network stress due to formation of
68 crystals before drying. As a result, cryogels undergo internal breakage of polymer network and
69 surface cracking (Garcia-Gonzalez, Camino-Rey, Alnaief, Zetzl, & Smirnova, 2012). To
70 prevent pore collapse phenomena and maintain as much as possible the hydrogel network

71 architecture, a two-step procedure may be exploited: firstly, solvent exchange is carried out so
72 that water in the hydrogel is replaced by ethanol to obtain an alcoholgel; secondly, ethanol is
73 extracted from the alcoholgel by supercritical carbon dioxide drying to obtain an aerogel
74 (Garcia-Gonzalez et al., 2012). Supercritical drying prevents structure collapse since it does not
75 involve vapor transitions nor intense surface tensions in the pores. The resulting aerogels are
76 thus low density and highly porous materials (Gesser & Goswami, 1989; Hrubesh & Poco,
77 1995).

78 Most aerogels are inorganic, being often made of silica, metal oxides or polystyrenes (Du, Zhou,
79 Zhang, & Shen, 2013; Gesser & Goswami, 1989; Pierre & Pajonk, 2002). They are lightweight
80 materials with high mechanic strength and excellent thermal insulation and dielectric properties
81 (Pierre & Pajonk, 2002). However, according to Pierre & Pajonk (2002), not only inorganic
82 polymerizing agents but all organic biopolymers are potential candidates to form aerogels. To
83 this regard, the preparation of aerogels from different polysaccharides, including starch,
84 cellulose, pectin, and carrageenan, have been recently reviewed by Mikkonen, Parikka, Ghafar,
85 & Tenkanen (2013) and Ivanovic, Milovanovic, & Zizovic (2016). These materials have been
86 proposed for packaging purposes but also for encapsulation and controlled release of drugs,
87 aroma or antioxidants. They have also been shown to quickly absorb aqueous solutions and
88 surfactants by capillary forces, due to the open pore structure and large surface area (Escudero,
89 Robitzer, Di Renzo, & Quignard, 2009; Mallepally, Bernard, Marin, Ward, & McHugh, 2013).

90 Recently, aerogels have been proposed also as oil carrier. Comin, Temelli, & Saldana (2012)
91 studied the oil impregnation capacity of β -glucan aerogels. In this case, the highest
92 impregnation capacity was about 65%. Similarly, Ahmadi, Madadlou, & Saboury (2016)
93 proposed aerogels made of whey proteins and crystalline cellulose. The latter presented a
94 maximum oil loading capacity of about 70%.

95 Based on this information, the possibility to obtain food-grade aerogels with high oil loading
96 capacity could open new opportunities in the exploitation of aerogels for novel food
97 applications.

98 This work represents a first attempt to develop food-grade oleogels by oil sorption into aerogels
99 by using κ -carrageenan as structuring biopolymer. This widely used food additive was chosen
100 because, in the presence of K^+ , it forms hydrogels with a tubular architecture, which could be
101 particularly interesting for oil sorption (Dunstan et al., 2001). κ -carrageenan hydrogels with
102 different concentration were converted to alcoholgels by a solvent exchange procedure. Ethanol
103 was then removed from the alcoholgel by supercritical carbon dioxide drying to obtain the
104 aerogels. The supercritical drying has been indicated as the most promising drying methodology
105 to obtain aerogels mainly because it prevents the gel structure from pore physical collapse
106 phenomenon and shrinkage upon solvent removal (Ivanovic et al., 2016). κ -carrageenan based
107 aerogel were characterized for appearance, network density, firmness, microstructure, water
108 vapor adsorption and glass transition. Finally, the capability of aerogels to absorb sunflower oil
109 and form oleogels was evaluated.

110

111 **2. Materials and methods**

112 **2.1 Materials**

113 κ -carrageenan (κ -C) was purchased from Sigma-Aldrich (Milan, Italy); lithium chloride (LiCl),
114 calcium chloride hexahydrate ($CaCl_2 \cdot 6H_2O$), potassium carbonate (K_2CO_3), sodium chloride
115 (NaCl), potassium acetate (CH_3COOK), potassium chloride (KCl), and potassium sulfate
116 (K_2SO_4) were purchased from Carlo Erba Reagents (Milan, Italy); absolute ethanol was
117 purchased from J.T. Baker (Griesheim, Germany); phosphorus pentoxide (P_2O_5) was purchased
118 from Chem-Lab NV (Zedelgem, Belgium); sunflower oil was purchased in a local market. All
119 solutions were prepared using milli-Q water.

120

121 **2.2 Hydrogel preparation**

122 Aqueous suspensions containing 0.4, 1.0, or 2.0% (w/w) κ -C and 1.0, 1.0 or 2.0 % (w/w) KCl,
123 respectively, were prepared. In particular, κ -C was slowly added to the KCl aqueous solution
124 at 90 °C under stirring. The homogeneous κ -C suspension was then poured into cylindrical
125 molds of 2.9 cm diameter and 12 cm height. Samples were cooled in an ice bath and stored for
126 1 day at 4 °C before analysis or further processing.

127

128 **2.3 Hydrogel to alcoholgel conversion by solvent substitution**

129 κ -C hydrogels were cut in cylinders with a height of about 4.5 cm and diameter of 2.9 cm and
130 were maintained for 1 day into aqueous solutions of ethanol with increasing concentrations (25,
131 50, 75% v/v). Finally, samples were introduced into absolute ethanol twice (the first time for 8
132 h and the second one for 1 day) in order to remove residual water. The ratio between hydrogel
133 and ethanol solutions was 1:8 (v/v). Conversion was carried out at room temperature (about 22
134 °C).

135

136 **2.4 Alcoholgel to aerogel conversion by supercritical CO₂ drying**

137 Alcoholgels were converted to aerogel by supercritical CO₂ drying using the apparatus (Figure
138 1) developed at the Department of Agricultural, Food, Environmental and Animal Sciences of
139 the University of Udine. Preliminary tests were carried out to define supercritical CO₂ drying
140 conditions to obtain aerogels in the available equipment. Based on these preliminary results,
141 aerogels were produced after their maintenance in a continuous flow of supercritical CO₂ at 11
142 \pm 1 MPa and 45 °C. Liquid carbon dioxide (purity 99.995%, Sapio, Monza, Italy) was cooled
143 to 4 °C using a F34-ED chiller (C; Julabo, Milano, Italy) after been filtered with a 15 μ m filter
144 (B₁; Ham-Let, Milano, Italy). Subsequently, CO₂ was pressurized at 11 \pm 1 MPa with an Orlita
145 MhS35/10 diaphragm pump (D; ProMinent Italiana S.r.l., Bolzano, Italy) and heated to 45 °C
146 using a water bath connected to a CB8 – 30e thermostatic bath (G; Heto, Allerød, Denmark).

147 Before pressurization, alcoholgel sample was placed inside the stainless steel cylindrical reactor
148 (E, volume ~265 mL) with two screwed caps, each one equipped with a sintered stainless steel
149 filter that allowed a uniform distribution of the CO₂ during drying. Different combinations of
150 supercritical CO₂ flows in the range from 2.0 to 8.0 NL/min were initially tested. The
151 combination allowing drying time to be minimized while maintaining the structural integrity of
152 the material were selected by visual assessment of the absence of surface cracks on the samples.
153 The adopted conditions were: the outlet flow through the reactor was 3.5 NL/min for 3 h; 5.0
154 NL/min for subsequent 4 h and 6.0 NL/min for subsequent 1 h. Finally, a slow decompression
155 from 11 MPa to atmospheric pressure was carried out at 6.0 NL/min in 30 min. The outlet flow
156 was set by a micrometric valve (V₄) and controlled with a RAGK41 rotameter (H; Rota
157 Yokogawa, Milan, Italy). To avoid malfunctioning of the rotameter, CO₂ was filtered with a 40
158 µm filter (B₂; Ham-Let, Milano, Italy). Ethanol content in the gaseous outlet was measured
159 using a AL9000L digital alcoholmeter (L; Alcoscan, Milan, Italy) every 60 min. In order to
160 carefully control temperature and pressure during experiments, a thermocouple (TT) connected
161 to a digital data logger (F) and a manometer (PT2) were used. The valves V₃ and V₄ were heated
162 in a water bath connected to a thermostatic bath (G) to prevent freezing during decompression.
163 In order to assure an adequate heat exchange in the water bath, a small water pump (P) was
164 used.

165 Aerogels were stored in a desiccator containing P₂O₅ at room temperature until use.

166

167 **2.5 Aerogel to oleogel conversion by oil absorption**

168 Aerogel samples were introduced into 250 mL beakers previously filled with 125 mL of
169 sunflower oil. At defined time intervals during conversion from aerogel to oleogel, samples
170 were withdrawn, wiped with absorbing paper and weighted. Absorbed oil was expressed as the
171 ratio between weight gain at time t and the initial weight of the aerogel sample. The immersion

172 of aerogel into oil was prolonged until a constant weight after two consequent readings was
173 reached.

174

175

176 **2.6 Analytical determinations**

177 ***2.6.1 Volume and network density***

178 Sample volume was calculated as the volume of the cylinder whose diameter and height were
179 measured by a CD-15APXR digital caliper (Absolute AOS Digimatic, Mitutoyo Corporation,
180 Kanagawa, Japan). Volume changes following conversion of hydrogel to alcoholgel and
181 aerogel were expressed as the percentage ratio between the variation of sample volume and
182 volume of the corresponding hydrogel. Network density was then calculated as the ratio
183 between aerogel sample weight and volume of hydro-, alcohol-, aero- or oleogel samples.

184

185 ***2.6.2 Firmness***

186 Firmness was measured by uniaxial compression test using an Instron 4301 (Instron LTD., High
187 Wycombe, UK). The instrumental settings and operations were accomplished using the
188 software Automated Materials Testing System (version 5, Series IX, Instron LTD., High
189 Wycombe, UK). In particular, hydrogel and alcoholgel samples (about 2.9 cm diameter and 1.5
190 cm height) were tested using a 6.2 mm diameter cylindrical probe mounted on a 100 N
191 compression head at a 25 mm/min crosshead speed. Force–distance curves were obtained from
192 the compression tests and firmness was taken as the maximum force (N) required to penetrated
193 the sample for 5 mm. Aerogel and oleogel samples (about 1 cm diameter and 3 mm height)
194 were tested using a 12.7 mm diameter cylindrical probe mounted on a 1000 N compression
195 head at a 25 mm/min crosshead speed. Force–distance curves were obtained from the
196 compression tests and firmness was taken as the maximum force (N) required to compress the
197 sample by 1 mm. The analyses were repeated at least 3 times for each sample.

198

199 **2.6.3 Image acquisition**

200 Sample images were acquired using an image acquisition cabinet (Immagini & Computer,
201 Bareggio, Italy) equipped with a digital camera (EOS 550D, Canon, Milano, Italy). In
202 particular, the digital camera was placed on an adjustable stand positioned 45 cm above a black
203 or white cardboard base where the samples were placed. Light was provided by 4 100 W frosted
204 photographic floodlights, in a position allowing minimum shadow and glare. Images were
205 saved in *jpeg* format resulting in 3456×2304 pixels.

206

207 **2.6.4 Scanning Electron Microscopy (SEM)**

208 Aerogel samples were mounted on aluminum sample holders and sputter coated with 10 nm of
209 gold using a Sputter Coater 108 auto (Cressington Scientific Instruments, Watford, United
210 Kingdom). The aluminum holder was transferred to the SEM unit (EVO 40XVP, Carl Zeiss,
211 Milan, Italy), which was at ambient temperature and under vacuum. Samples were imaged using
212 an acceleration voltage of 20 kV and SmartSEM v. 5.09 (Carl Zeiss, Milan, Italy) application
213 software was used to capture images of the samples. Images were saved in *tiff* format resulting
214 in 1696×2048 pixels.

215

216 **2.6.5 Water vapor sorption**

217 Aerogel samples were weighted and transferred into a dried weighting bottle. The latter was
218 then transferred into desiccators containing LiCl, CH₃COOK, CaCl₂, K₂CO₃, NaCl, KCl, and
219 K₂SO₄ saturated solutions with equilibrium relative humidity (ERH%) values of 11, 25, 31, 43,
220 75, 86, and 96%, respectively. Samples were kept inside desiccators until constant weight was
221 reached. The Brunauer-Emmet-Teller (BET) sorption isotherm model (eq. 1) was fitted into
222 water sorption data (Brunauer, Emmett, & Teller, 1938).

$$223 \frac{a_w}{m \cdot (1 - a_w)} = \frac{1}{m_0 \cdot c} + \frac{c - 1}{m_0 \cdot c} \cdot a_w \quad (1)$$

224 where a_w is the water activity, m is the moisture of the sample expressed as ratio between the
225 weight (g) of absorbed water and the weight (g) of dry matter, m_0 is the moisture of the water
226 monolayer, and c is an experimental constant.

227

228 **2.6.6 Differential Scanning Calorimetry (DSC)**

229 DSC analysis was carried out using a TA4000 differential scanning calorimeter (Mettler-
230 Toledo, Greifensee, Swiss) connected to a GraphWare software TAT72.2/5 (Mettler-Toledo).
231 Heat flow calibration was achieved using indium (heat of fusion 28.45 J/g). Temperature
232 calibration was carried out using hexane (m.p. -93.5 °C), water (m.p. 0.0 °C) and indium (m.p.
233 156.6 °C). Samples were prepared by carefully weighing around 10 mg of hydrogel or aerogel
234 in 160 mL aluminum DSC pans, closed with hermetic sealing. An empty pan was used as a
235 reference in the DSC cell.

236 Aerogel samples equilibrated at different a_w values were heated from -150 to 250 °C. The scan
237 speed was set at 10 °C/min and samples were analyzed under nitrogen flow (20 mL/min). The
238 start of melting transition was taken as on-set (T_{on}) point of transition, that is the point at which
239 the extrapolated baseline intersects the extrapolated tangent of the calorimetric peak in the
240 transition state. Total peak enthalpy (ΔH_m) was obtained by integration of the melting curve. T_g
241 was determined from the on-set temperature of the glass transition of aerogels. The machine
242 equipment program STARe ver. 8.10 (Mettler-Toledo, Greifensee, Switzerland) was used to
243 plot and analyze the thermal data.

244 The amount of frozen water was then calculated as the ratio between aerogel ΔH_m and pure ice
245 ΔH_m (333.5 J/g). The concentration of the maximally cryo-concentrated solution (c'_g) was
246 calculated from the amount of unfrozen water and total solids.

247

248

249

250 **2.6.7 State diagram and modified state diagram**

251 Aerogel state diagrams were obtained plotting the T_g values for samples equilibrated at different
252 ERH% as a function of mass fraction of the sample. The obtained curve was fitted using the
253 Gordon-Taylor equation (eq. 2) (Gordon & Taylor, 1952).

$$254 \quad T_g = \frac{w_1 T_{g1} + k w_2 T_{g2}}{w_1 + k w_2} \quad (2)$$

255 where T_{g1} is the glass transition temperature of the amorphous solute, T_{g2} is the glass transition
256 temperature of the solvent (-137.5°C), w_1 and w_2 are the mass fraction of the solute and the
257 solvent, respectively, and k is an experimental constant.

258 The modified state diagram was then obtained plotting the T_g values for samples equilibrated
259 at different ERH% as a function of their a_w values.

260

261 **2.6.8 Oil absorption kinetics**

262 Oil content of the oleogel was defined as the percentage ratio between the maximum amount
263 of absorbed oil and the weight of the oleogel.

264 Oil absorption capacity was calculated as the ratio between weight gain at time t and aerogel
265 network density. The kinetics of oil absorption were then elaborated by fitting a two-phase
266 exponential decay model (eq. 3) to absorption data (Blake, Co, & Marangoni, 2014).

$$267 \quad y = y_{fast} \left(1 - e^{(-k_{fast}t)}\right) + y_{slow} \left(1 - e^{(-k_{slow}t)}\right) \quad (3)$$

$$268 \quad y_{max} = y_{fast} + y_{slow} \quad (4)$$

269 where y_{fast} and y_{slow} are the asymptote values of the fast- and slow-decaying components,
270 respectively, k_{fast} and k_{slow} are the rate constants for the fast- and slow-decaying component,
271 respectively, and y_{max} is the maximum amount of absorbed oil when time t tends to infinite and
272 is the sum of y_{fast} and y_{slow} (eq. 4). y_{max} can also be considered the theoretical plateau value.

273

274 **2.6.9 Oil holding capacity (OHC)**

275 Around 100 - 200 mg of oleogel was placed into 1.5 mL microtubes between two pieces of
276 absorbing paper. Samples were centrifuged at 13,000 rpm (15,871 x g) for 30 min using a
277 microcentrifuge (Mikro 120, Hettich Zentrifugen, Andreas Hettich GmbH and Co, Tuttlingen,
278 Germany). Oil holding capacity (OHC) was computed as the percentage ratio among the weight
279 of oil retained in the oleogel after centrifugation and total weight of oil in the sample.

280

281 ***2.6.10 Data analysis***

282 All determinations were expressed as the mean \pm standard error (SE) of at least two
283 measurements from two experiment replicates ($n \geq 4$), if not otherwise specified. Statistical
284 analysis was performed by using R v. 3.0.2 (The R foundation for Statistical Computing).
285 Bartlett's test was used to check the homogeneity of variance, one way ANOVA was carried
286 out and Tukey-test was used as post-hoc test to determine statistical significant differences
287 among means ($p < 0.05$). Linear regression analysis by least squares minimization was
288 performed using GraphPad Prism v.5.03 (GraphPad Software, San Diego, USA). The goodness
289 of fit was evaluated on the basis of statistical parameters of fitting (R^2 , p, standard error) and
290 the residual analysis. Non-linear regression analysis of T_g values as a function of aerogel mass
291 fraction was performed on TableCurve 2D software (Jandel Scientific, ver. 5.01). Levenberg–
292 Marquardt algorithm was used to perform least squares function minimization and the goodness
293 of fit was evaluated on the basis of statistical parameters of fitting (R^2 , p, standard error) and
294 the residual analysis.

295

296 **3. Results and discussion**

297 ***3.1 From hydrogel to aerogel***

298 κ -carrageenan (κ -C) hydrogels were used as template to obtain aerogels by applying a solvent
299 exchange procedure. Hydrogels were initially formed thanks to the well-known ability of κ -C
300 random coils to transit to a double helix conformation. The double helices, in the presence of

301 monovalent ions, such as potassium (K^+), aggregated in water forming a gelled system
302 (Rinaudo, 2008; Rochas & Rinaudo, 1984). Hydrogels containing different concentration of k-
303 C appeared as self-standing materials with a network density and firmness that linearly
304 increased as the concentration of the structuring polymer increased ($R^2 > 0.99$) (Table 1). This
305 was due, as known, to the formation of a higher number of junction zones among the double
306 helices (Rinaudo, 2008). The hydrogels were firstly converted to alcoholgels by substituting
307 the water solvent with ethanol. Ethanol was then removed from the alcoholgel by a continuous
308 flow of supercritical CO_2 , obtaining the aerogels. The removal of ethanol resulted complete
309 after 6, 7, and 8 h of drying for samples containing 0.4, 1.0 and 2.0 % (w/w) k-C (see Figure
310 S1 in supplementary material).

311 Figure 2 shows the visual appearance of hydrogels, alcoholgels and aerogels. It is evident that
312 gel characteristics changed upon solvent exchange. Aerogels appeared completely opaque,
313 differently from hydrogel and alcoholgel, suggesting the aerogel can be regarded as a porous
314 material. Porosity would favor intense light scattering, providing a dense and white appearance.
315 Turning hydrogels into alcoholgels and then aerogels also promoted an intense shrinkage
316 (Figure 2). The latter can be probably attributable to the different structural organization of the
317 gel network depending on the solvent nature. During the first solvent substitution, ethanol is
318 forced to diffuse through the k-C network even if this polymer is insoluble in ethanol
319 (Therkelsen, 1993). For this reason, ethanol difficulty interacts with k-C and is unable to fill all
320 the space previously occupied by water. The interactions among k-C chains became thus
321 stronger, leading to gel shrinkage. The alcohol removal caused a further shrinkage, probably
322 indicating a collapse of the structure upon drying (Figure 2). k-C concentration in the hydrogels
323 negatively affected the level of shrinkage, so that volume contraction progressively decreased
324 as k-C concentration increase. This suggests that samples richer in structured polymer chains
325 were less prone to shrinkage and begot a more porous aerogel structure.

326 As a consequence of shrinkage, the gel network density progressively increased moving from
327 hydrogel to aerogel, at all k-C concentrations (Table 1). This result confirms the hypothesis that
328 solvent substitution led to a reduction of the distance among polymer network chains. It is
329 interesting to note that the network density of aerogels decreased as k-C concentration
330 increased, differently from hydrogels and alcoholgels. This suggests that the higher the polymer
331 content, the more porous the aerogel structure. A maximum firmness value was observed for
332 sample at intermediate k-C concentration. It could be inferred that the higher resistance to
333 mechanical deformation of this sample could be the result of less aerated and/or more isotropic
334 aerogels. To confirm this hypothesis, SEM analysis of aerogels was performed. Images in
335 Figure 2 revealed the presence of a compact matrix of k-C with superficial pores in all samples,
336 even if with some morphological differences. Sample at the lowest k-C concentration was
337 characterized by a compact structure embedding restricted porous areas. The latter showed a
338 fine-grain and appeared evenly distributed in the aerogel with intermediate k-C content. Finally,
339 sample with the highest k-C content showed a coarse structure with larger polymer aggregates
340 as well as cracks and microchannels onto the surface. Results clearly indicate that aerated
341 structures were achieved when hydrogel structural collapse was hindered by increasing its
342 initial polymer concentration.

343 In order to investigate the properties of the developed aerogels, their capacity of absorbing
344 water vapor was evaluated. Samples were thus equilibrated at constant temperature and
345 different relative humidity to obtain their sorption isotherms. Moisture content data were then
346 modelled as a function of a_w , using the procedure proposed by Brunauer et al. (1938).
347 Regression analysis showed good determination coefficients (> 0.85) and statistically
348 significant model parameters ($p < 0.05$). The monolayer water content (m_0) and the BET
349 constant (c) were thus estimated (Table 2).

350 The m_0 parameter showed comparable values for the three aerogels. The constant c showed
351 values between 2 and 50 revealing the presence of a type II isotherm (Al-Muhtaseb, McMinn,

352 & Magee, 2002; Brunauer, Deming, Deming, & Teller, 1940). This means that the aerogels
353 were characterized by a poor capacity of water vapor sorption since an increase in the relative
354 humidity was reflected into a great a_w increase. In other words, water vapor difficulty interacted
355 with the porous aerogel structure. Its swelling and solvation only occurred when direct
356 hydration of the aerogels was carried out by water immersion (data not shown).

357 To study the physical stability of the aerogels, DSC analysis was performed. Aerogels were
358 characterized by a glass transition temperature of 180 ± 1 °C, indicating that they were in the
359 glassy state at room temperature. The effect of equilibration at different ERH% on aerogel glass
360 transition temperature was then studied. Glass transition temperature data were modeled as a
361 function of mass fraction using the approach proposed by Gordon & Taylor (1952). Non-linear
362 regression analysis showed good determination coefficients (> 0.93) and significant ($p < 0.001$)
363 values of the model experimental constant (k) (Table 2). Also in this case, no differences among
364 samples were detected.

365 The modified state diagrams of the aerogels were thus obtained combining the water vapor
366 sorption curve with the glass transition temperature one, and resulted comparable for the three
367 aerogels. This suggests that the ERH% dependence of aerogel physical stability is mainly
368 governed by the intrinsic properties of k-C rather than by the structure of the aerogel. As an
369 example, Figure 3 shows the modified state diagram of the aerogel obtained from a hydrogel
370 containing 1% k-C.

371 At room temperature, the system was below the glass transition temperature up to an a_w value
372 of 0.6. However, when an amount of water equal to the 10% of the aerogel sample mass was
373 absorbed, the system decreases its glass transition temperature below 20 °C and a transition to
374 the rubber state was observed. This transition led to a structural collapse and the system became
375 thermodynamically unstable. Based on these data, the aerogels here developed would remain
376 glassy and porous at room temperature if maintained at ERH lower than 60%. For these reasons,

377 aerogels can be easily stored for prolonged time if protected from atmospheric moisture through
378 appropriate packaging.

379

380 ***3.2 From aerogels to oleogels***

381 Based on their physical properties, aerogels could be exploited to entrap liquid oil, potentially
382 leading to oleogels. The capacity of aerogels to absorb oil was thus evaluated (Figure 4).

383 Oil absorption progressively increased during immersion in oil and was considered complete
384 after the plateau value was reached (Figure 4A). The rate of oil absorption was also greatly
385 affected by the aerogel structure, so that the maximum amount of absorbed oil was reached
386 after 3, 24 and 48 h for samples containing 0.4, 1.0 and 2.0% k-C, respectively. Data in Figure
387 4A were further elaborated to evidence the effect of the network density on oil absorption. Oil
388 absorption capacity of the aerogel was computed as the ratio between absorbed oil and network
389 density. Normalization of absorbed oil based on network density (Figure 4B) clearly shows that
390 the capacity of the aerogel to absorb oil progressively increased in the order $0.4 < 1.0 < 2.0\%$
391 k-C. This suggests that the aerogels, which had experienced a lower level of structural collapse
392 (lower shrinkage), also showed a higher capacity of oil absorption, regardless the network
393 density. In other words, the capacity of oil to be entrapped in the aerogel depends not only on
394 the density of the polymer network but also on its architecture. Liquid absorption by a porous
395 material is actually affected by several factors such as number, dimension and size distribution
396 of pores, pores tortuosity and internal surface (i.e. roughness) (Bear, 1972; Khosravi & Azizian,
397 2016). The diameter of the pores is known to steer the rate of oil absorption while the number
398 and length of pores affect the amount of absorbed oil. To further investigate these aspects,
399 kinetics of aerogel oil absorption were analyzed. In particular, data shown in Figure 4B were
400 elaborated by fitting a two-phase exponential decay model (eq. 3) (Blake et al., 2014). This
401 model was chosen since it describes oil absorption kinetics as a result of two different
402 components. The fast component, which is related to pore diameter, and the slow component,

403 which accounts for pore number and length. Non-linear regression analysis showed good
404 determination coefficient (> 0.99) and significant ($p < 0.001$) model parameters (Table 3).
405 The rate constant for the fast-decaying component (k_{fast}) resulted always higher than the slow
406 one, indicating that the limiting factor of the initial phase of oil absorption was pore size. The
407 value of k_{fast} decreased by increasing the aerogel network density. As discussed by Khosravi &
408 Azizian (2016), lower value of k_{fast} could be related to the presence of pores with larger
409 diameter, which are known to be less effective in initial oil uptake. These larger pores were
410 probably more numerous and longer according to the sample order $2.0 > 1.0 > 0.4\%$ (w/w) κ -
411 C in the initial hydrogel. This result is in agreement with the microscopic structure of the
412 aerogels (Figure 2) and gives reason for the increasing overall absorption of oil, as indicated by
413 the higher value of y_{max} (Table 3).

414 Samples after absorption of the maximum amount of oil can be regarded as oleogels. Table 4
415 shows their visual appearance, composition, firmness and oil holding capacity.

416 Oleogels obtained from hydrogels containing 0.4 and 1.0% (w/w) κ -C were able to entrap
417 around 2.5 times their initial weight, whereas the aerogel obtained from 2.0% κ -C hydrogel
418 held *circa* 4.5 times its initial weight (Figure 4A). Firmness of oleogels showed the same trend
419 observed for aerogels (Table 1) with a maximum value for sample containing 1.0% (w/w) κ -C.

420 The maximum loading capacity resulted about 81% (w/w) that is higher than that reported for
421 aerogels containing other food-grade biopolymers, such as β -glucans (Comin et al., 2012) and
422 whey proteins (Ahmadi et al., 2016). Finally, the capability of the oleogels in retaining absorbed
423 oil was finally assessed by an accelerated oil release test based on centrifugation. The highest
424 values of oil holding capacity (OHC) were recorded for samples containing 0.4 and 1.0% (w/w)
425 κ -C in the initial hydrogel. By contrast, aerogels from 2.0% (w/w) κ -C hydrogel showed a lower
426 ability to retain oil. In other words, this sample, which was characterized by a higher number
427 of longer pores absorbed the highest amounts of oil (Figure 4) that can be easily released upon
428 centrifugation. This suggest that oil is physically entrapped in the system cavities.

429

430 **Conclusions**

431 κ -carrageenan aerogels resulted to be highly porous and structurally stable materials with high
432 mechanic strength. Similarly to other organic aerogels, they were made from renewable sources
433 and were completely biodegradable. Given these properties, they could be used for a number
434 of different applications, including thermal and electric insulation but also development of
435 novel packaging materials and selective carriers for drugs, nutrients, aroma compounds or
436 additives. In the present work, a novel application of κ -carrageenan aerogels was studied. The
437 latter were actually demonstrated to uptake large amounts of oil without compromising their
438 structural integrity and leading to stiff oleogels. Oil content and retention depended on the
439 aerogel architectural organization, as described by the pore number, size and length. These
440 results suggest that κ -carrageenan based aerogels could be used to absorb lipophilic molecules,
441 including unintentionally discharged oil spills. Reversely, κ -carrageenan oleogels could be
442 exploited in the food, pharmaceutical or cosmetic sectors for pioneering applications. The
443 results acquired were relevant to κ -carrageenan oleogels but the methodology here developed
444 could be definitely extended to other biopolymers. Further research is thus needed to explore
445 this possibility and obtain food grade oleogels with tailored characteristics.

446

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452

453

454

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458

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545 Zulim Botega, D. C., Marangoni, A. G., Smith, A. K., & Goff, H. D. (2013). The potential
546 application of rice bran wax oleogel to replace solid fat and enhance unsaturated fat
547 content in ice cream. *Journal of Food Science*, *78*, C1334-1339.

548

549 Table 1. Network density and firmness of hydrogel containing increasing κ -carrageenan
 550 concentration, and of the derived alcoholgel and aerogel.

κ -carrageenan in hydrogel (% w/w)	Network density (g _{d.m.} /cm ³)			Firmness (N)		
	Hydrogel	Alcoholgel	Aerogel	Hydrogel	Alcoholgel	Aerogel
0.4	0.004 ±	0.009 ±	0.237 ±	0.84	8.00 ±	114.00 ±
	0.001 ^c	0.001 ^b	0.026 ^a	± 0.06 ^c	0.66 ^c	7.21 ^a
1	0.008 ±	0.014 ±	0.180 ±	3.37 ±	18.92 ±	165.67 ±
	0.001 ^b	0.002 ^b	0.014 ^{ab}	0.05 ^b	0.94 ^b	7.22 ^b
2	0.016 ±	0.026 ±	0.129 ±	7.37	45.22 ±	136.67 ±
	0.001 ^a	0.003 ^a	0.001 ^b	± 0.70 ^a	1.50 ^a	2.03 ^a

551 ^{a, b, c}: means with different letters in the same column are significantly different ($p < 0.05$).

552

553 Table 2. Experimental regression coefficients estimates m_0 ($\text{g}_{\text{H}_2\text{O}}/\text{g}_{\text{d.m.}}$) and c for BET equation
 554 ($R^2 > 0.85$; $p < 0.05$), and k for Gordon-Taylor equation ($R^2 > 0.93$; $p < 0.001$) of aerogels
 555 obtained from hydrogels containing increasing κ -carrageenan concentration. Standard error of
 556 fitting is also reported.

κ -carrageenan in hydrogel (% w/w)	BET equation		Gordon-Taylor equation		
	$m_0 \pm \text{SE}$	$c \pm \text{SE}$	R^2	$k \pm \text{SE}$	R^2
0.4	0.04 ± 0.01	9.23 ± 6.16	0.95	10.33 ± 1.55	0.97
1.0	0.05 ± 0.01	43.88 ± 13.91	0.85	9.79 ± 1.27	0.93
2.0	0.04 ± 0.01	4.86 ± 0.52	0.99	9.92 ± 0.94	0.98

557

558



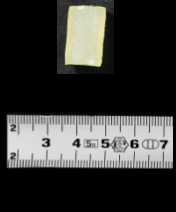
559 Table 3. Experimental regression coefficients estimates k_{fast} , k_{slow} , and y_{max} for the two-phase
 560 model of oil absorption in aerogels obtained from hydrogels containing increasing κ -
 561 carrageenan concentration. Standard error of fitting is also reported.

	κ -carrageenan in hydrogel		
	(% w/w)		
	0.4	1.0	2.0
k_{fast} (h ⁻¹)	3.825 ± 0.320	2.080 ± 0.153	1.734 ± 0.198
k_{slow} (h ⁻¹)	0.602 ± 0.058	0.313 ± 0.017	0.139 ± 0.006
y_{max} (g oil/g aerogel/cm ³ aerogel)	1.077 ± 0.049	3.195 ± 0.112	13.845 ± 0.310

562

563

564 Table 4. Visual appearance, firmness, oil content, and oil holding capacity (OHC) of oleogels
 565 obtained from hydrogels containing increasing κ -carrageenan concentration.

κ -carrageenan in hydrogel (% w/w)	Visual appearance	κ -carrageenan in oleogel (% w/w)	Oil in oleogel (% w/w)	Firmness (N)	OHC (% w/w)
0.4		27.58 ± 0.83^a	72.42 ± 0.83^a	158.33 ± 9.16^c	83.44 ± 1.35^a
1.0		26.76 ± 0.33^a	73.24 ± 0.33^a	311.70 ± 11.78^a	82.18 ± 1.11^a
2.0		18.72 ± 0.31^b	81.28 ± 0.31^b	216.40 ± 6.79^b	62.21 ± 1.31^b

566 ^{a, b, c}: means with different letters in the same column are significantly different ($p < 0.05$)

567

568 **Figure captions**

569 Figure 1. Schematic representation of supercritical CO₂ drying apparatus.

570

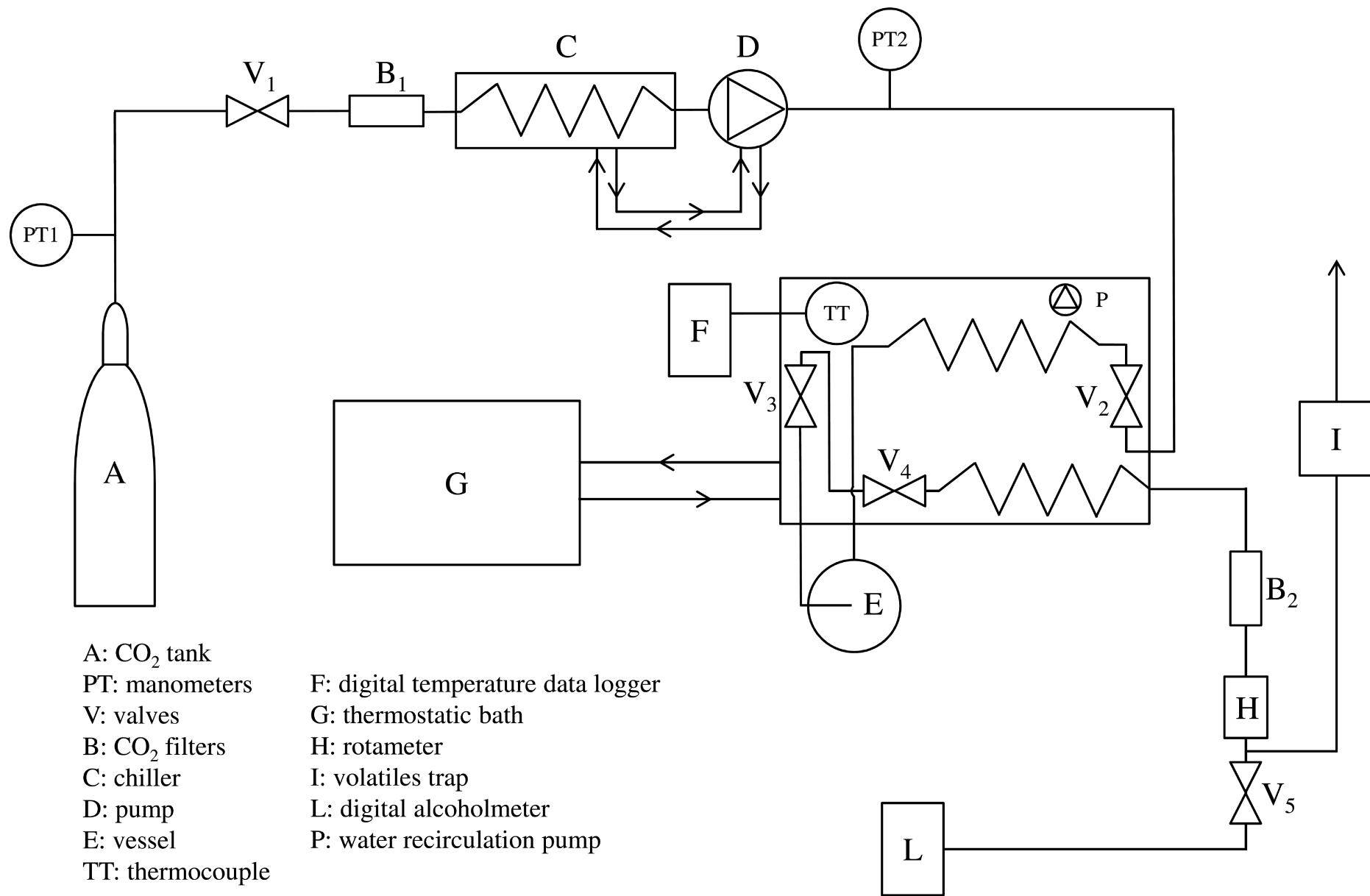
571 Figure 2. Visual appearance of hydrogel containing increasing κ -carrageenan concentration,
572 and of the derived alcoholgel and aerogel. A ruler in cm is also reported as reference. Scanning
573 electron microscopy images of aerogels obtained from hydrogels containing increasing κ -
574 carrageenan concentration.

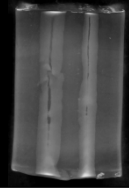
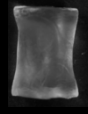

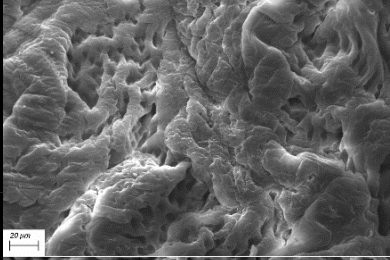

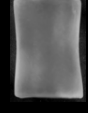

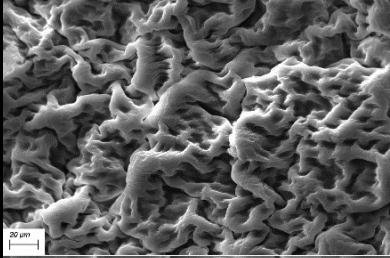
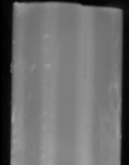


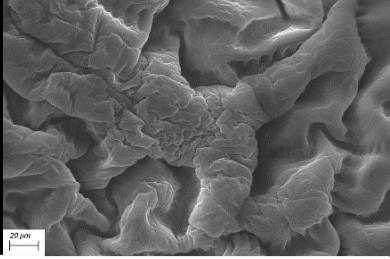
575

576 Figure 3. Modified state diagram of aerogel obtained from hydrogel containing 1% (w/w) κ -
577 carrageenan.

578

579 Figure 4. Absorbed oil (A) and oil absorption capacity (B) of aerogels obtained from hydrogels
580 containing increasing κ -carrageenan concentration as a function of time.



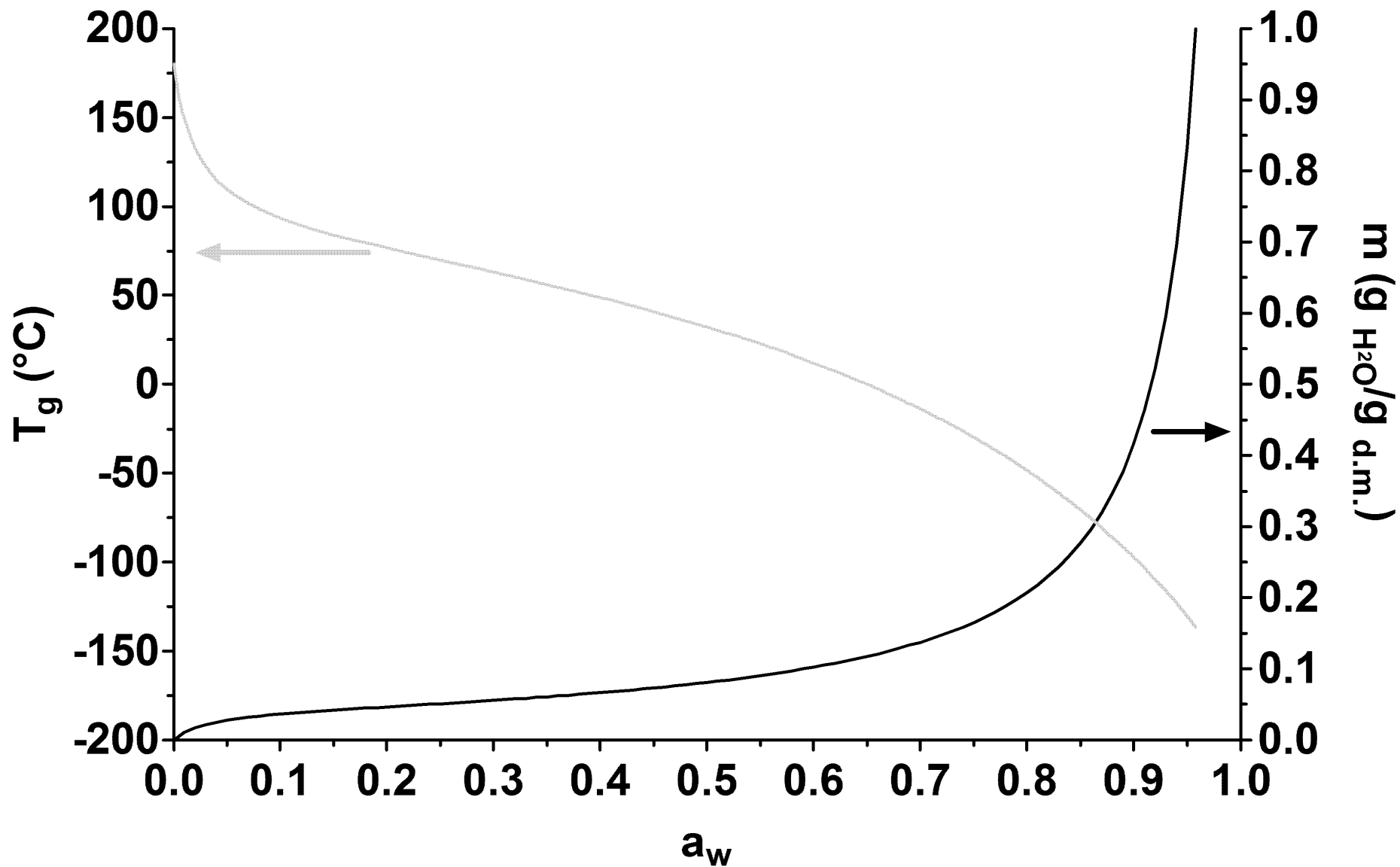
κ -carrageenan in hydrogel (% w/w)	Hydrogel	Alcoholgel	Aerogel	Aerogel SEM images
0.4				
1.0				
2.0				

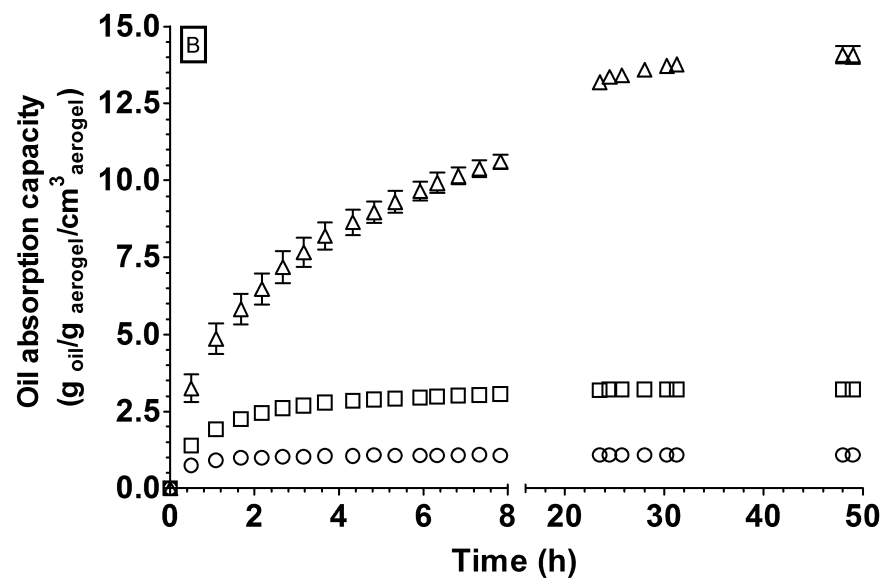
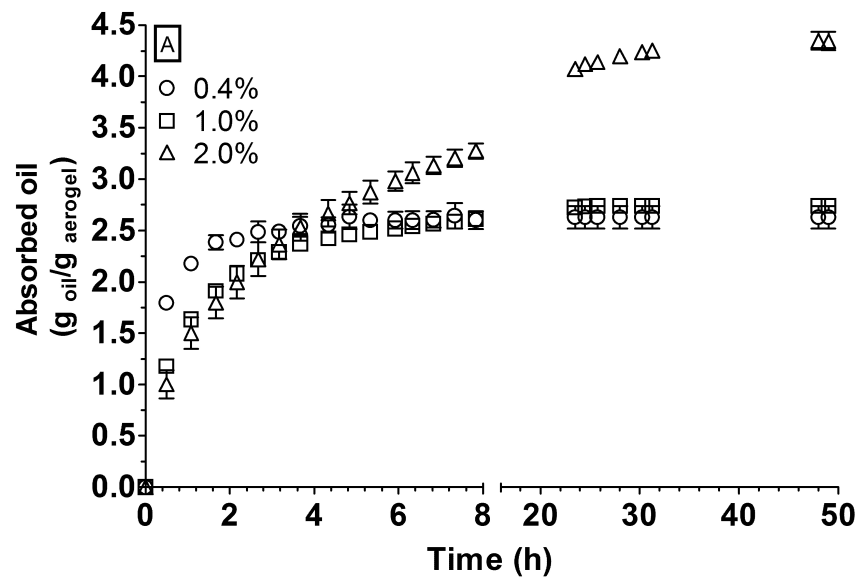


20 μ m

20 μ m

20 μ m





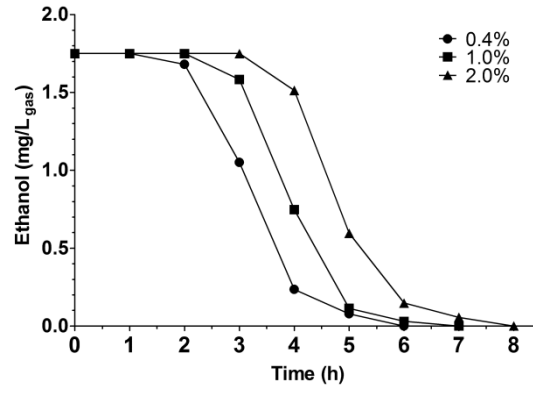


Figure S1. Ethanol concentration as a function of time during supercritical CO₂ drying of alcoholgels obtained from hydrogels containing increasing κ -carrageenan concentration.

Highlights

k-carrageenan aerogels with tailored properties can be obtained by supercritical CO₂ drying;

k-carrageenan aerogels showed high oil loaded capacity;

k-carrageenan oleogels could have interesting potential application in food area.