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1 **Release of glucose and maltodextrin DE 2 from gellan gum gels and the impacts of gel structure**

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6

7 **Abstract**

8 Structural influences of hydrocolloids gels on the release of carbohydrates was examined with a  
9 focus on structure-function relationships. This understanding will guide formulation of food gels for  
10 targeted sugar release in populations such as diabetics and athletes. Hydrocolloid gels with well  
11 characterised structures, with a focus on high acyl (HA) and low acyl (LA) gellan gum, were  
12 formulated with glucose, maltose, DE 10 maltodextrin (MD) and DE 2 MD. Gel structure did not  
13 significantly affect glucose release, but mixed gel type had a significant effect on MD availability. A  
14 DE 2 MD required amylase to release more than 10% of the carbohydrates but still had 38% retained  
15 in a gel formulated with 30% MD. Formulation with any non-melting gelling hydrocolloid decreased  
16 the amount of released MD and phase separated networks released more than interpenetrating  
17 networks. Differential scanning calorimetry was used to compare helix formation of MD gels and the  
18 number of helices was inversely correlated with carbohydrate release. These results demonstrated a  
19 range of sugar release profiles achievable from formulation from specific gelling agent structures  
20 and carbohydrates.

21

22 **Keywords:** sugar release, controlled release, maltodextrin, carbohydrate release, carbohydrate  
23 digestion, High acyl gellan gum

## 24 **1. Introduction**

25 Hydrocolloids can control the release, digestion, or adsorption of various nutrients ranging from  
26 starches to vitamins by controlling their network structure and response to known digestive stimuli  
27 (Norton et al., 2014; McClements, 2021). Controlling carbohydrate digestion is crucial in formulation  
28 of foods when considering specific populations which require a sustained energy release or low  
29 glycaemic index such as diabetics and athletes (Gidley, 2013; Norton et al., 2014). Carbohydrates, a  
30 major source of energy for humans, come in many sizes ranging from monomers (glucose) and  
31 dimers (maltose) and up to 30-150 saccharide units (maltodextrin (MD)) and 100-1,800 saccharide  
32 units (starches). Smaller molecules are able to diffuse through viscous solutions or gel networks  
33 (Mills et al., 2011). Larger molecules need to be broken into smaller units which can then move into  
34 the chyme where they can be absorbed (Tharakan et al., 2010; Gidley, 2013; Fabek et al., 2014). To  
35 formulate products with controlled sugar release, the relationships between carbohydrate size and  
36 hydrocolloid gel structure must be understood.

37 For most hydrocolloid gels, small molecules such as mono and disaccharides, salts, and artificial  
38 sweeteners, are all smaller than the pores of the gel network and are able to diffuse through the gel.  
39 Thus, the network mostly acts to prevent mixing (a faster mass transfer) and only a small deviation  
40 from diffusion coefficients has been measured (Jönsson et al., 1986; Lorén et al., 2009a; Lorén et al.,  
41 2009b). In one study, a 14% - 30% decrease in diffusion coefficient of salt was measured for  
42 hydrocolloids between 1 and 4% mass (Mills et al., 2011). Larger molecules (3 vs 8 nm) showed a  
43 greater decrease in diffusion coefficients with increasing polymer concentration (Lorén et al.,  
44 2009b). Differences between gelling agent and concentration were not significant with an exception  
45 major structural changes such as melting (Mills et al., 2011). It is well known that surface area of a  
46 gel has a large influence on the rate of release, so that brittleness or a tendency to fracture causes a  
47 quicker release (Morris, 1994; Mills et al., 2011). Texture, breakdown, and serum release of the gel

48 during mastication thus plays a major role in differentiation between hydrocolloids gels (Khin et al.,  
49 2021).

50 Release of larger carbohydrates (those not able to diffuse through the pores) is more complex. The  
51 digestive enzyme  $\alpha$ -amylase cleaves maltose units from starch (Butterworth et al., 2011; Dhital et al.,  
52 2017) which is then small enough to diffuse out of the gel. Small amounts of glucose, maltotriose,  
53 and dextrin are also created (Butterworth et al., 2011; Dhital et al., 2017). An increased viscosity or a  
54 gel network impedes the mass transfer of the enzyme and slows the rate of digestion (Tharakan et  
55 al., 2010; Gidley, 2013; Fabek et al., 2014). Gel surface area, packing density, and subsequent  
56 entrapment impact the ability of  $\alpha$ -amylase to reach the carbohydrate and thus the rate of digestion  
57 (Wee and Henry, 2020; McClements, 2021). In addition to the physical inhibition, chemical binding  
58 can also occur. For example, cellulose was found to inhibit  $\alpha$ -amylase activity by binding with the  
59 enzyme (Dhital et al., 2017). Studies examining the effects of gelling hydrocolloids on the digestion  
60 of starch have typically found a decreased rate of digestion and total digestion (Butler et al., 2008;  
61 Koh et al., 2009; Sasaki and Kohyama, 2011; Ramírez et al., 2015; Zhang et al., 2018; Srikaeo and  
62 Paphonyanyong, 2020). Gelatinization, retrogradation, and steric hindrance of starch are all  
63 expected to play a role making differentiation of the separate mechanisms impossible (Zhang et al.,  
64 2018). Comparisons to MD may give insight into the contribution of network effects on  
65 retrogradation because gelatinization does not occur.

66 An understanding of the effects of gel structure on carbohydrate release, as a function of molecular  
67 weight (MW), is important for the formulation of products with controlled carbohydrate release.  
68 Most work has focused on either high MW carbohydrates (specifically starches) (Koh et al., 2009;  
69 Sasaki and Kohyama, 2011; Ramírez et al., 2015; Zhang et al., 2018; Srikaeo and Paphonyanyong,  
70 2020) or very low MW, such as glucose, maltose, and sucrose (Morris, 1994; Wang et al., 2014; Yang  
71 et al., 2015; Nishinari and Fang, 2016; Khin et al., 2021), but left out intermediate MW  
72 carbohydrates like MD. The mechanism of release for the digestion of MDs has not yet been

73 determined and was thought to be slowed by incorporation of hydrocolloids. Gellan gum has been  
74 shown to form an interpenetrating network with MD (Clark et al., 1999; Kanyuck et al., 2021a) and  
75 high acyl (HA) gellan gum variant was capable of forming a wide variety of material properties  
76 (Kanyuck et al., 2021a). Therefore, this gelling agent has considerable potential for use in  
77 customizable carbohydrate release systems.

78 The objective of the present investigation was to examine the role of gelling agents on carbohydrate  
79 release by comparing carbohydrates of different MW trapped within hydrocolloid gels with well-  
80 characterized properties. It was hypothesized that the MW of the carbohydrate, the gel network  
81 structure, and the response to environmental conditions (a stimuli-response) can predict release  
82 behaviour from hydrocolloids gels. MW of a carbohydrate is known to determine the path of release  
83 and digestion from gels. Although there are certainly other structural factors such as branching and  
84 linkages in carbohydrates, molar mass is just as important and is sometimes overlooked (Nishinari  
85 and Fang, 2021). Some gel networks display a response to stimuli such as melting, dissolution, or  
86 swelling which typically have large impacts on release (McClements, 2021). After determining the  
87 pathway for MD release, a structural comparison will examine the influence of mixed gel network  
88 type and MD helix formation. Exploring these fundamental relationships between carbohydrate MW  
89 and gelling agent structure will facilitate strategic formulation of products to achieve desired release  
90 profiles.

## 91 **2. Materials and Methods**

### 92 **2.1 Materials.**

93 Both MDs were derived from potato and acquired from Avebe (Veendam, Netherlands) with a  
94 dextrose equivalent (DE) of 2 (Paselli SA 2, batch H3362903) and 10 (Paselli MD 10, batch  
95 H4852902). The HA (LT100) and LA (F) gellan gum were acquired from CP Kelco (Atlanta, USA). The  
96 following hydrocolloids were purchased from Sigma Aldrich (St. Louis, USA): Gelatin type A with a  
97 bloom strength of 300, kappa-carrageenan, iota-carrageenan, agarose type A9539, and sodium

98 alginate. Maltose, KCl, and CaCl<sub>2</sub> were also purchased from Sigma Aldrich. The α-amylase was from  
 99 *Aspergillus oryzae* (10065 Sigma-Aldrich) with an activity of 32 U/mg. Glucose was purchased from  
 100 Fisher Chemical (Loughborough, UK).

101 Table 1. A summary of the hydrocolloids used in formulation of the carbohydrate gels.

Gelling agent	Source	Conc.	Gel preparation
Agarose	Type A9539, Sigma Aldrich	2%	Powder was dispersed and heated in water at 90 °C for 30 minutes
Alginate	Sodium type, Sigma Aldrich	2%	Powder was dispersed and heated in water at 90 °C for 30 minutes and then gelled by diffusion method in a 91 mM CaCl <sub>2</sub> solution
Gelatin	Type A, Sigma Aldrich	2%	Powder was dispersed and heated in water at 50 °C for 30 minutes
High acyl gellan gum	CP Kelco LT100	0.25-3%	Powder was dispersed and heated in water at 90 °C for 2 hours
iota-carrageenan	Sigma Aldrich	2%	Powder was dispersed and heated in water at 90 °C for 30 minutes and then 2.68 mM KCl was added and immediately poured into moulds
kappa-carrageenan	Sigma Aldrich	2%	Powder was dispersed and heated in water at 90 °C for 30 minutes and then 2.68 mM KCl was added and immediately poured into moulds
Low acyl gellan gum	CP Kelco F	0.25-3%	Powder was dispersed and heated in water at 90 °C for 2 hours

102

## 103 2.2. Gel preparation

104 All gels were prepared by dispersing the hydrocolloids in heated deionized (DI) water with stirring to  
 105 fully hydrate the polymers individually in stock solutions. MD and the glucose or maltose solutions  
 106 were heated at 90 °C for 4 hours. Gellan gums were heated at 90 °C for 2 hours, the carrageenan,  
 107 agarose, and alginate heated at 90 °C for 30 minutes, and gelatin heated at 50 °C for 30 minutes.  
 108 After the individual hydrocolloids were hydrated, a hot solution (90 °C) of a stock concentration of  
 109 the carbohydrate (glucose, maltose, MD DE2, or MD DE10 as indicated) was mixed with the  
 110 hydrocolloids for 5 minutes to combine. These mixtures were poured into 20mm diameter  
 111 cylindrical plastic moulds and set at room temperature for at least 48 hours before analysis. Samples  
 112 with MD were analysed after 4 days to give sufficient time for helix aggregation (Kanyuck et al.,  
 113 2019). Kappa-carrageenan and iota-carrageenan contained an added 2.68 mM KCl to reach gelation.

114 Sodium alginate was gelled by the diffusion method (Draget, 2009) with 91 mM CaCl<sub>2</sub> and formulated  
115 at a higher glucose concentration (40%) to account for loss during the gelation preparation.

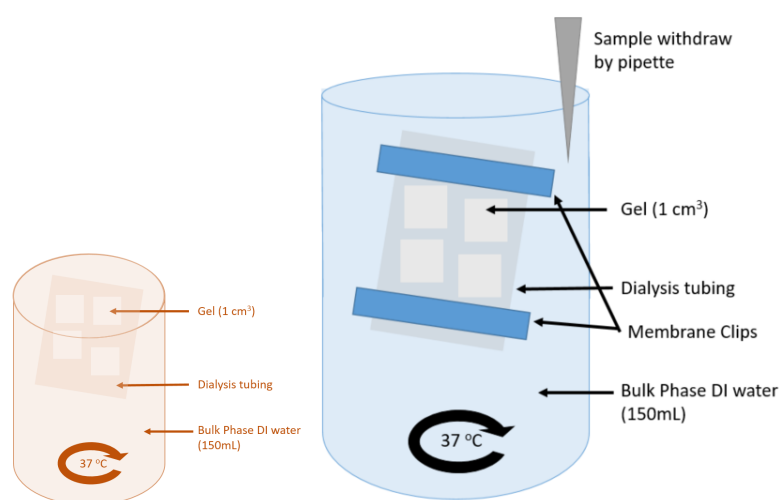
### 116 **2.3. Release measurements**

117 The method for measuring release of carbohydrates from gels followed the procedure by Koh et al.  
118 (2009) with some modification. Gels were cut into 4 pieces of ~1 cm<sup>3</sup> each (5 g ± 1 g). To prevent  
119 amylase from interfering with the refractive index measurement, gel pieces were placed within a  
120 dialysis tubing membrane of molecular weight cut-off 14 kDa (MEMBRA-CEL MD44-14). Amylase  
121 isolated from *Aspergillus oryzae* has been found to have molecular weights of 51 kDa (sedimentation  
122 and diffusion) and 49 kDa (gel filtration) and thus is too large to cross the membrane. Membrane  
123 clips were used to seal the gel sample and 5mL of amylase solution (or water when amylase was not  
124 used) within the dialysis tubing. The sample pouch was added to a volume of 150 mL of DI water  
125 pre-warmed to 37 °C inside plastic bottles with lids. The apparatus was held in a shaker (Sciquip,  
126 Newtown, UK) at 37 °C with rotation of 200 RPM for the duration of the experiment. A schematic of  
127 the experimental setup is shown in Figure 1. At each time point, 0.5 mL from the bulk was removed  
128 for measurement by refractive index (Rudolph research J357 automatic refractometer from  
129 Hackettstown, USA) and was returned to the bulk phase. Refractive index is a measure of the  
130 relative speed of light in a solution and is linearly related to sugar concentration. Calibration curves  
131 for glucose and maltose were used to calculate the sugar concentration in each sample by the  
132 refractive index measurement.

133 Measurements were normalized to the 'percent of total carbohydrates released' by dividing by the  
134 amount of carbohydrate known from the sample mass. A 'total release' value was measured after 48  
135 hours. Initial experiments showed the sugar concentration did not increase after the 48 hour time  
136 point. A 90 minute time point is compared between samples as an indication of the relative rate of  
137 digestion. This time point has also been shown to have the highest correlation with glycemic index  
138 ( $r=0.909$ ) (Goñi et al., 1997).

139 Samples containing MD utilized a triggered release by addition of  $\alpha$ -amylase to mimic human  
140 digestion. The enzyme  $\alpha$ -amylase cleaves linear carbohydrate chains into maltose units (Butterworth  
141 et al., 2011). A stock solution of 100 U/mL amylase was prepared by dispersing the powder in DI  
142 water at room temperature for 30 minutes. 5 mL of the amylase solution was added to the dialysis  
143 tubing to reach an activity of 500U. The dialysis tubing was then sealed with a clip and placed into  
144 the bulk water phase at 37 °C within 5 minutes. Amylase concentrations in human saliva have wide  
145 variability based on time of day, most recent meal, and also the individual. The value of 500 U was  
146 chosen because it is within the range of human salivary enzyme activity (Mandel et al., 2010) and  
147 similar to the concentration used by Koh et al. (2009) and (Janssen et al., 2009).

148 It should be acknowledged that *in vitro* tests such as this can only approximate differences between  
149 samples. Amylase sourced from porcine or *Aspergillus oryzae* have shown minor differences to  
150 human amylase, but their use allows for consistent comparison between experiments. Some  
151 deviations should be expected, so for true glyceic index human tests should be used. However,  
152 true human experiments also have natural variability in oral processing, enzyme concentrations,  
153 hormones, and residence times in the stomach and intestine (Dhital et al., 2017). For comparison  
154 between different samples, the use of any of the amylases has shown to be effective.



155

156 **Figure 1.** Diagram of release experimental setup showing the gels within dialysis tubing and inside of  
157 a larger bulk phase (150mL) which was shaken at 200 rpm at 37 °C. Method was adapted from Koh  
158 et al. (2009).



159 **2.4. Modelling of glucose release**

160 Glucose concentrations measured after 24 hours had reached the predicted value (with ± 5% error)  
161 and were thus normalized to account for natural sample variability with the equation:

162 
$$\text{Released Glucose} = M_t/M_\infty \tag{Eq 1}$$

163 Where  $M_t$  is the measured concentration at time 't' and  $M_\infty$  the final maximum concentration. The  
164 only sample with greater than 5% error was sodium alginate which was expected to lose some  
165 glucose during the gelation methodology (diffusion of calcium ions into the alginate solution along  
166 with diffusion of glucose out). The collected data was then fitted to a power law model and the  
167 Peppas-Sahlin Model. As the models are not able to account for the lowering difference in  
168 concentration gradient over time, data was only fitted below 60 % of the release (the model  
169 assumes steady-state). These were then compared to a COMSOL fit of Fickian diffusion within the gel  
170 which was able to account for changing concentrations.

171 **Power Law Model:** A simple exponential model was fit using Microsoft Excel:

172 
$$M_t/M_\infty = k*t^n \tag{Eq 2}$$

173 Where 'k' is the rate constant specific to the gel formula and 'n' the diffusional exponent (Siepmann  
174 and Peppas, 2011).

175 **Peppas-Sahlin Model:** Release curves were also fit to the model proposed by Peppas and Sahlin  
176 (1989). An equation with the following form was fit to the release profiles:

177 
$$M_t/M_\infty = k_1*t^{0.45} + k_2 *t^{0.9} \tag{Eq 3}$$

178 Where k is the rate constant where  $k_1*t^{0.45}$  represents the Fickian diffusion and  $k_2 *t^{0.9}$  the case II  
179 transport contributions for a cylindrical shape (Peppas and Sahlin, 1989; Siepmann and Peppas,  
180 2011). Fickian diffusional describes the release of an active caused by a concentration gradient while  
181 the case II transport mechanism is dictated by a transition of the polymer which changes the release

182 rate of the active (Peppas and Sahlin, 1989; Siepmann and Peppas, 2011). Similar to the single  
183 exponential model, the model is only valid for the initial 60% of glucose release to avoid the effects  
184 of lowering differences in concentration gradients. The biexponential regression was fit using  
185 SigmaPlot (Version 12.5 SYSTAT Software, USA). Proportional contributions were calculated using  
186 the equations proposed by Peppas and Sahlin (1989). In summary, the percent contribution was  
187 calculated by the ratio of each coefficient for each time point.

188 **COMSOL Model:** The engineering software COMSOL (COMSOL Inc. Burlington, MA, USA), was used  
189 to predict diffusion of glucose using the experimental dimensions and concentration gradients. The  
190 flux of glucose from within the gel (into the water) was calculated by Ficks' law of diffusion using the  
191 dimensions of the objects (shown in Figure 1) and an initial concentration of 2.38 M (2381 moles/m<sup>3</sup>)  
192 in the gel and 0 in the water. Gels were surrounded by a water region of 150 mL with a diffusivity of  
193 (1 m<sup>3</sup>/s) meaning practically that mixing was instantaneous. A thin mesh was drawn around the gel  
194 to ensure release only occurred at the surface of the gel and diffusion was modelled to the edges of  
195 the gel. The model was fit for a single cube of gel (1 mL) with the measured values adjusted by a  
196 factor of 0.25 for simplicity. A diffusion coefficient of glucose in water was 6.0 x 10<sup>-10</sup> m<sup>2</sup>/sec was  
197 obtained from literature (Stein and Litman, 2014). The model accounted for changes in flux with the  
198 changing concentration gradients (which the other models do not).

## 199 **2.5. Swelling**

200 Swelling of gellan gum gels was measured by increases in mass after soaking in aqueous solutions.  
201 Gels were cut into ~20 mm height pieces from the cylindrical moulds and the mass weighed to 7.5 ±  
202 1 g. The gel was then placed into 150 mL of DI water at room temperature. At each time point, the  
203 gel was removed using a strainer, patted dry to remove surface water, and weighed. The amount of  
204 swelling was determined from the ratio of initial mass to final mass by the equation:

$$205 \text{ Swelling Ratio} = M/M_0 \quad (\text{Eq 4})$$

206 where  $M$  is the measured sample mass after swelling and  $M_0$  is the initial mass.

## 207 **2.6. DSC**

208 Gelation of maltodextrin was studied by measuring the enthalpy and entropy using a  $\mu$ DSC3 evo  
209 (Setaram Instrumentation, France). Samples were added in the sol phase (hot) to the sample vessels  
210 and held for 4 days at room temperature prior to analysis to allow sufficient gelation of the MD  
211 component (Kanyuck et al., 2019). A heating and cooling cycle began with a hold at 5 °C for 10  
212 minutes and then increased at 1°C/min up to 95 °C. After a 10 minute hold at 95°C, the temperature  
213 was cooled at 1°C/min down to 5 °C.

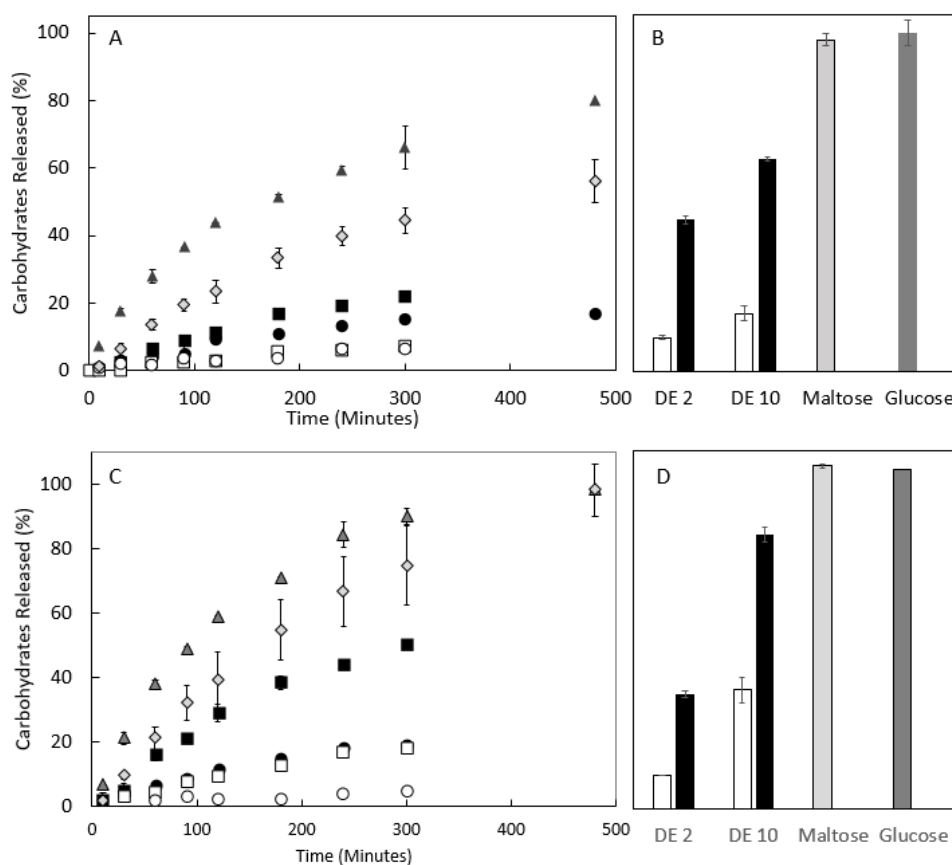
## 214 **2.7. Statistical Analysis**

215 All samples were measured in at least triplicate and data are presented as means  $\pm$  standard  
216 deviation. Release curves were repeated four times for each sample. Error bars show one standard  
217 deviation above and below the mean value. On the bar charts, different letters suggest significantly  
218 different mean values. A t-test with a p-value of 0.05 was used to determine which samples were  
219 significantly different.

## 220 **3. Results and Discussion**

221 Of the many factors to consider in predicting the release, the carbohydrate MW is of crucial  
222 importance (Nishinari and Fang, 2021). To examine this effect, carbohydrates of varying molecular  
223 weight (MW) were compared by release profiles from HA gellan gel (Figure 2A) and LA gellan gum  
224 (Figure 2C). Small molecules were represented by glucose (180 Da) and maltose (342 Da) and  
225 showed complete release from the gel within 48 hours. The rate of release was slower for maltose  
226 because it is a larger molecule than glucose. Larger molecules are expected to have slower diffusion  
227 coefficients due to the greater hydrodynamic radius (Nishinari and Fang, 2021). MDs are known to  
228 contain a wide range of different molecular sizes with a bimodal distribution and the distribution of  
229 the DE 2 MD is centred at 10,000 Da and 492,000 Da (Loret et al., 2004). Both MDs used (DE 2 and

230 DE 10) led to a slower and incomplete release of carbohydrates from the gels. Molecules smaller  
 231 than the pores are able to diffuse out of the gel network while larger molecules are trapped (Lin and  
 232 Metters, 2006; McClements, 2017). The small amount of carbohydrates measured without addition  
 233 of an enzyme (10% and 17% for DE 2 and 10, respectively) reflect the proportion of molecules which  
 234 were small enough to diffuse out of the gellan gum gel network. Addition of amylase, the enzyme  
 235 which cleaves maltose units from a larger carbohydrate chain, considerably increased the amount  
 236 (to 44% and 63% for DE 2 and 10, respectively). Based on the work of (Dhital et al., 2017), amylase  
 237 was thought to enter the gel network and break the MD into maltose molecules which were then  
 238 small enough to diffuse out of the gel. Starch, a much larger carbohydrate, is well known to need  
 239 this enzyme to break into saccharides that can be released from a gel (Koh et al., 2009; Butterworth  
 240 et al., 2011; Dhital et al., 2017).



241  
 242 **Figure 2.** Release of carbohydrates from 1% HA gellan (A) and 2% LA gellan (C) gels formulated with  
 243 30% glucose (▲), maltose (◇), and with MD DE 2 (●) and DE 10 (■) with amylase (black) and without  
 244 (white). Total release from the gels at 48 hours shown for 1% HA gellan (B) and 2% LA gellan (D).

245 Two clear pathways of release were established based on the size of the carbohydrate; diffusion  
246 based release of the small molecules, and an amylase-triggered release for large molecules that  
247 cannot diffuse out of the gel network. The following work will be split into subsequent sections to  
248 examine effects of gel network type on the release of small (3.1) and medium sized aggregated (3.3)  
249 carbohydrates. Glucose was selected to be representative of small carbohydrates and DE 2 MD  
250 chosen for the enzyme-triggered release.

### 251 **3.1. Diffusion based release of small MW carbohydrates**

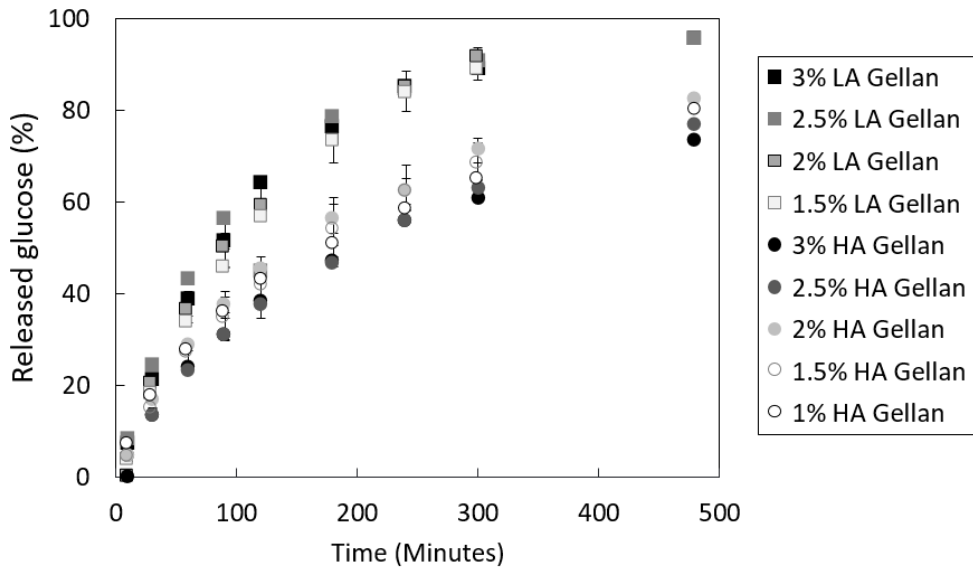
252 Glucose was chosen as a model for small molecule carbohydrates and the release from different gel  
253 structures (polymer types and concentration) were compared. All samples reached  $100 \pm 5\%$  after 24  
254 hours and are shown normalized in the graph to decrease the impact of variability in the gel  
255 formulation. Changes in gellan concentration led to significant differences in the release speed ( $p <$   
256  $0.05$ ), however the differences of a few percentage points had minimal practical differences (Figure  
257 3). Increases in concentration of polymer are known to decrease the release rates of small  
258 molecules, but this is typically quite a small shift (10-20%). This trend was observed for sucrose from  
259 agar gels (Wang et al., 2014; Yang et al., 2015), salt from LA gellan and gelatin (Mills et al., 2011), and  
260 dendrimers of 3 and 8 nm from kappa-carrageenan (Lorén et al., 2009b). Higher polymer  
261 concentrations are expected to decrease the pore size within gels and provides a greater physical  
262 barrier. For glucose this is minimally important because the pores are already much larger and the  
263 hydrocolloid such a small proportion of the mass (Mills et al., 2011). Larger actives (3 and 8 nm)  
264 showed progressively a greater slowing from a kappa-carrageenan gel network (Lorén et al., 2009b).

265 A comparison of LA and HA however shows a difference between these two polymer types,  
266 irrespective of the concentration (Figure 3). Both HA and LA gellan gum form physical gels by double  
267 helix formation upon cold-setting and do not melt at  $37\text{ }^{\circ}\text{C}$  or below (Morris et al., 2012). Removal of  
268 acyl groups for the LA gellan yields a completely different gel texture than HA gellan due to  
269 differences in helix aggregation which may have been a factor. The surface area for each gel was

270 matched in these experiments (controlled in the sample preparation). After 24 hours, all of the  
271 glucose (within a reasonable standard deviation of +/- 5%) had been released which suggested there  
272 was no significant binding between HA gellan and glucose to cause the lower diffusion rates and  
273 therefore the difference appeared to be kinetic in nature.

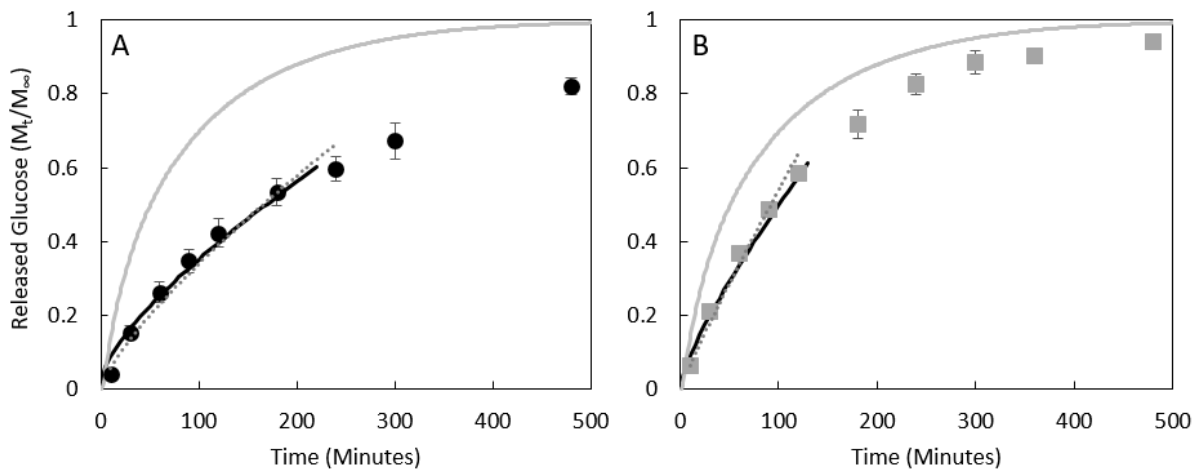
274 Mathematical modelling of release profiles has become a popular method for understanding the  
275 mechanisms of release from gels (Lin and Metters, 2006). Comparison of models for HA and LA  
276 gellan gum release were used to elucidate the origin or mechanism of the difference. Quality of fit  
277 for the models is shown in Figure 4 and the equations displayed in Table 2. The commonly used  
278 Peppas-Sahlin equation (Eq 3) models the release of an active as a summation of the Fickian  
279 diffusion ( $k_1 * t^{0.45}$ ) and case II release ( $k_2 * t^{0.9}$ ). As the model cannot account for changes in  
280 concentration, analysis should only be conducted on the initial 60% of the release profile. This was  
281 reflected in the curves of Figure 4 which end at the 60% release point.

282 Comparing the importance of each coefficient ( $k_1$  for the Fickian contribution and  $k_2$  for relaxational  
283 case II contribution) was used to give evidence of the type of release (Siepmann and Peppas, 2011).  
284 According to this model, the Fickian or case II contribution can be modelled over time to show any  
285 changes in type of release. Relative contributions of each type, and how that shifts over the release  
286 profile, are shown in (Supplemental Figure 1). For LA gellan gum, the release was suggested to be  
287 largely case II driven (Supplemental Figure 1) which could also be predicted from the diffusional  
288 exponent (n) value of the single power exponent of 0.92 which is near to that of 'pure relaxation' of  
289 a 0.9 value. Alternatively, the single n for HA gellan (0.76) was between that of Fickian and case II  
290 and was reflected in the relative greater Fickian contribution (Supplemental Figure 1). The Peppas-  
291 Sahlin model suggested the HA gellan release profile was more similar to a Fickian release while the  
292 LA gellan was predominately dictated by case II release. A greater similarity of HA gellan gum to  
293 Fickian diffusion may suggest that the LA gellan gum network relaxes to increase the release rate but  
294 was not sufficient to fully explain the difference between HA and LA gellan gum.



295

296 **Figure 3.** Concentration dependence of release profiles from LA gellan (squares) and HA gellan  
 297 (circles) formulated with 30% glucose.



298

299 **Figure 4.** Modelling glucose release for 2% HA gellan (A) and LA gellan (B) comparing the Peppas-  
 300 Sahlin Model (solid black line), single exponential (dashed grey line), and COMSOL mass transfer  
 301 model (grey line). Equations are shown in Table 2.

302 **Table 2.** Equations modelled to the release of glucose from high acyl (HA) and low acyl (LA) gellan  
 303 gum.

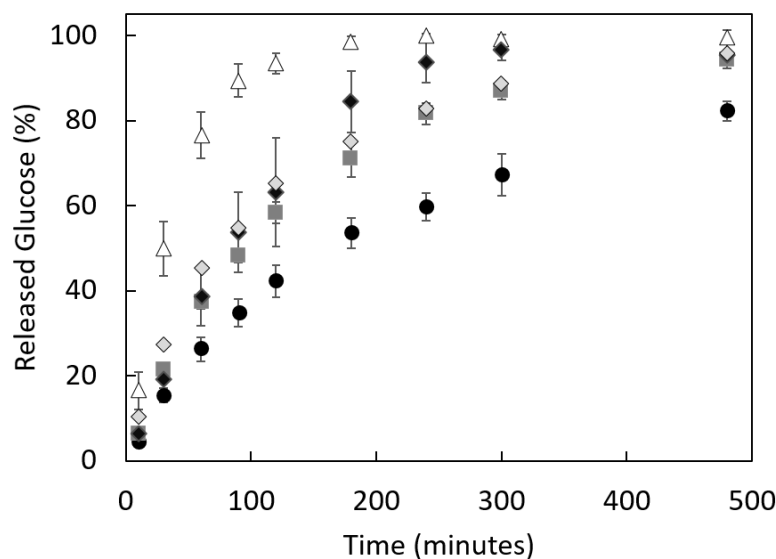
Model	HA Gellan	LA Gellan
Single Exponential	$M_t/M_\infty = 0.010 * t^{0.76}$ $R^2 = 0.98$	$M_t/M_\infty = 0.008 * t^{0.92}$ $R^2 = 0.99$
Peppas-Sahlin	$M_t/M_\infty = 0.028 * t^{0.45} + 0.002 * t^{0.9}$	$M_t/M_\infty = 0.017 * t^{0.45} + 0.007 * t^{0.9}$

304

305 Using a chemical engineering modelling software (COMSOL) and considering dimensions, initial  
306 concentrations, and changes during release a curve from 'pure diffusion' through the gel can be  
307 predicted for this specific system. The software was able to account for non-steady state behaviour  
308 and used the literature diffusion coefficient (D) of  $6.0 \times 10^{-10} \text{ m}^2/\text{s}$  (Stein and Litman, 2014). Higher  
309 similarity was observed between the expected pure diffusion and LA gellan, while the release from  
310 HA gellan was clearly slower. A marginal slower release from a hydrogel was expected due to the  
311 steric obstacle of the network by 14-30% (Mills et al., 2011). A shift farther from pure diffusion for  
312 the HA gellan gel suggested a stimuli-driven change to the gel was responsible for the slower release  
313 behaviour.

314 With evidence from modelling that LA gellan was closer to a 'typical' diffusion pattern, glucose  
315 release from other gelling agents were compared to give context to the different gel network  
316 structures. Release profiles from gelatin, alginate, and kappa-carrageenan are compared to that of  
317 HA and LA gellan gum in Figure 5. Release from alginate and kappa-carrageenan were similar to LA  
318 gellan gum. An alginate gel network is held together by chemical crosslinks (calcium bridges)  
319 between chains (Draget, 2009) while kappa-carrageenan forms a gel network through potassium  
320 induced aggregation of double helices (Morris et al., 1980). These three different gel structures did  
321 not appear to affect the release of glucose. At the measurement temperature (37 °C) the gelatin  
322 network melted and caused a quicker release profile than any of the other gelling agents. The other  
323 gelling agents did not melt. In comparison, the behaviour of HA gellan was unprecedentedly slower  
324 than any of the other gelling agents. Recent work has shown that swelling of HA gellan was  
325 responsible for the slower release of glucose compared to LA gellan (Kanyuck et al., 2021b). This  
326 stimuli-driven swelling was proposed to be the cause of the slower release from HA gellan gum and  
327 will be discussed in the following section (3.2). For diffusion-based release, network structure of the  
328 hydrocolloid was not important and differences were only observed from stimuli-driven changes to  
329 the gel, specifically melting and swelling.





330

331 **Figure 5.** Release of glucose from 2% HA gellan gum (●), LA gellan gum (■), alginate (◆), kappa-  
 332 carrageenan (◇), and gelatin (△) at 37 °C.

333

### 334 3.2. Impacts of stimuli-driven structural changes of gels on release

335 **Melting:** Stimuli from the environment which cause structural changes to a hydrocolloid gel, such as

336 swelling, dissolution, and erosion can modify the release profile (McClements, 2017). The quicker

337 release of glucose from gelatin was hypothesized to have been caused by the melting at the analysis

338 temperature of 37 °C. The experimental procedure was repeated at 25 °C which is below the melting

339 temperature of gelatin. When gelatin did not melt, the release was similar to LA gellan gum

340 (Supplemental Figure 2). Thus the inherent structure of gelatin did not distinguish from the other

341 gels but instead the temperature-driven structural change. Previous work has also confirmed that

342 environmental temperatures which cause melting of a gelatin gel showed much faster release of salt

343 than release at a temperature that did not cause melting (Mills et al., 2011). Melting of hydrocolloids

344 gels was a stimuli-driven structural change that impacts release from gelatin.

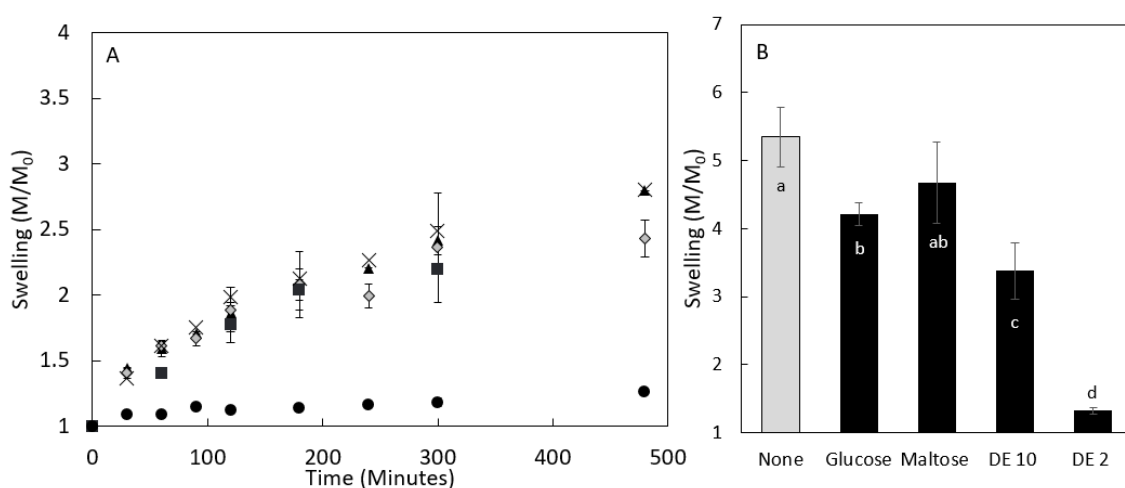
345 **Swelling:** In many cases, swelling of a polymer increases the release rate of a small molecule due to

346 the increased pore size of the hydrocolloid (McClements, 2017). In the case of HA gellan, swelling

347 actually slowed the release of glucose (Kanyuck et al., 2021b) and has the potential to impact larger

348 carbohydrates. Swelling kinetics of HA gellan is shown in Figure 6 for formulations with glucose,

349 maltose, DE 10 MD, and DE 2 MD. There was significantly less swelling with inclusion of glucose or  
 350 DE 10 MD, but the mass had still doubled after 180 minutes. The aggregates formed by the DE 10  
 351 MD decreased the swelling more than glucose or maltose (Figure 6). Very clearly the network  
 352 formed by DE 2 MD inhibited the swelling of the mixed gel. This MD (DE 2) is known to form large  
 353 and bulky aggregates within the HA gellan gum network (Kanyuck et al., 2021a). Not surprisingly,  
 354 these appeared to have prevented much of the typical swelling for HA gellan. Slower release of  
 355 glucose, maltose, and DE 10 MD was subsequently suspected for HA gellan due to a decreased mass  
 356 transfer caused by swelling (Kanyuck et al., 2021b). Swelling of a gel causes a greater volume and  
 357 larger dimensions, and subsequently the slower release was thought to have been caused by a lower  
 358 effective concentration inside the gel and a greater distance for the active to travel (Kanyuck et al.,  
 359 2021b). This effect was observed by comparing HA gellan and LA gellan gum release (Figure 2). A  
 360 slower release from these carbohydrate sources (glucose, maltose, and DE 10 MD) was measured for  
 361 HA gellan and emphasised the importance of this stimuli-response. Just as the environmental  
 362 temperature dictated melting of gelatin, the osmotic environment dictated HA gellan swelling  
 363 (Kanyuck et al., 2021b). These stimuli-responsive changes were shown to be crucial for predicting  
 364 release profiles and specific conditions were of critical importance.



365  
 366 **Figure 6.** Swelling of 1% HA gellan during the timeframe of release experiments (X) compared to  
 367 formulations with 30% glucose (▲), maltose (◇), and with MD DE 2 (●) and DE 10 (■). Part B displays  
 368 the swelling after 48 hours.

369

### 370 **3.3. Amylase-triggered release of MD**

371 Addition of the digestive enzyme amylase was essential for the release of MD from gellan gum gels.  
372 Only 10% of the DE 2 MD chains were small enough to diffuse out of the 1% HA gellan gum gel, while  
373 addition of the hydrolysing enzyme allowed 44% of the carbohydrates to be release from the gel  
374 (Figure 2). Similarly for 1% LA gellan gum, 40% was released with amylase but only 10% without. The  
375 ability of amylase to enter the gel network and reach the MD to begin cleavage was of chief  
376 importance (Dhital et al., 2017). However, even with addition of amylase more than half of the  
377 carbohydrate was resistant in the experiment. Aggregates of MD were hypothesized to be the  
378 source of enzyme resistance and will be explored. Impacts of hydrocolloid gel structure on the  
379 availability of these MD aggregates will then be explored with amylase-triggered release.

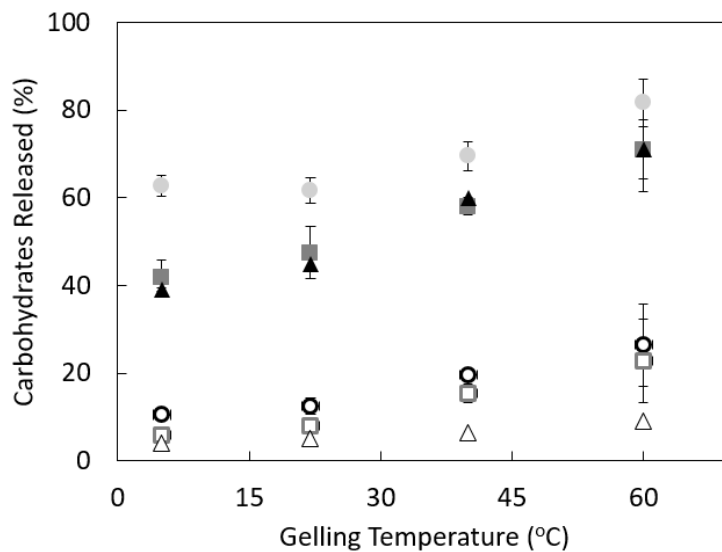
#### 380 **3.3.1. MD aggregation**

381 Self-aggregation of MD was hypothesized to play a role in the carbohydrate availability. This MD (DE  
382 2) is well characterized in literature and known to form aggregates of double helix that form a gel at  
383 high enough concentrations (15-20%) by connection of these dense aggregates (Kasapis et al.,  
384 1993a; Loret et al., 2004; Kanyuck et al., 2019). Holding temperature during gelation is known to  
385 affect the size and enthalpy of the aggregates formed (Kanyuck et al., 2019). Exploiting that  
386 knowledge, the impacts of MD aggregation on availability for amylase cleavage were determined by  
387 varying the gelation temperature. Higher temperatures formed fewer aggregates but at a higher  
388 entropy which was thought to be from the participation of longer chains in aggregate formation and  
389 connectivity (Kanyuck et al., 2019). Release of 30% and 40% MD gels formed at different  
390 temperatures is shown in Figure 7. For both concentrations of MD, lower carbohydrate release was  
391 measured for gels formed at lower temperatures. Correlation between enthalpy (Kanyuck et al.,  
392 2019) and carbohydrates released ( $R^2 = 0.82$ ) suggested the helices contributed to the enzyme  
393 resistance. Structural composition was the same between gels (linkages and branch points) and all  
394 release experiments were conducted at the same temperature (37 °C) so the differences showed

395 how aggregation impacted the accessibility of carbohydrates to amylase. Aggregation of MD was  
396 thought to function similarly to retrograded starch. Recrystallization and retrogradation of starch  
397 resulted in amorphous structures with inhibited enzyme affinity because of the irregular structure  
398 (Gidley et al., 1995; Butterworth et al., 2011; Dhital et al., 2017).

399 An aggregation effect was also seen in mixed gels of MD with HA gellan gum. Higher gelling  
400 temperatures resulted in greater percentages of released carbohydrates (Figure 7). The presence of  
401 the HA gellan gum network also decreased the amount of available carbohydrates, and at lower  
402 temperatures there was a greater inhibitory effect. At 60 °C the HA gellan network decreased  
403 availability by 11% while at 5 °C the difference was 24%. Based on the 90-minute values which were  
404 lower than either concentration of MD alone, the HA gellan network slowed amylase diffusion  
405 through the gel irrespective of the gelling temperature. At lower temperatures the gel network was  
406 more inhibitory and possibly due to a greater steric inhibition from more MD aggregates. The  
407 structure of this mixed gel consists of MD aggregates within pores of the HA gellan network  
408 (Kanyuck et al., 2021a). More MD aggregates would add considerable bulk within the HA gellan  
409 network that appeared to have blocked and prevented amylase from reaching as many aggregates in  
410 the mixed gel. These factors emphasise the contributions of MD aggregation and a gel network  
411 exclusion effect in the release of carbohydrates from mixed gel formulations.

412



414

415 **Figure 7.** Carbohydrate release by amylase hydrolysis compared by gelling temperature for DE 2 MD  
 416 at 30% (●), 40% (■), and 30% with 1% HA gellan gum (▲). Percentage released at 90 minutes shown  
 417 by open symbols and final release shown by filled symbols. Samples were held at the indicated  
 418 gelling temperature for 4 days prior to measurements and release experiments were all conducted  
 419 at the same temperature (37 °C).

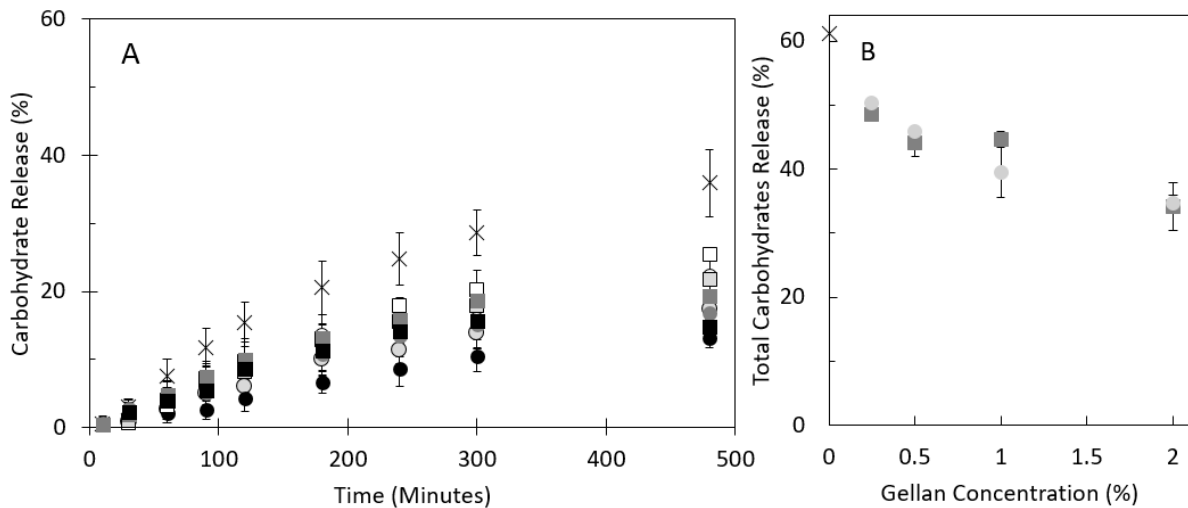
420

### 421 3.3.2. Gelling agents

422 MD gels were formulated with various gelling hydrocolloids to examine their structural effects on  
 423 MD availability. Different concentrations of HA and LA gellan gum at a constant 30% MD and the  
 424 release profiles are shown in Figure 8. Addition of either HA or LA gellan gum decreased the speed of  
 425 carbohydrate release as well as the total availability compared to a MD-only gel. Higher  
 426 concentrations also decreased the total carbohydrate availability (Figure 8B). Unlike the similarity  
 427 observed for glucose release, a smaller pore size from greater concentrations of polymer decreased  
 428 the speed of release. The slower release from HA gellan gum was likely due to swelling from the  
 429 greater distance amylase needed to travel into the gel (Figure 6). Additionally, a lower amount of  
 430 total carbohydrate was released with higher concentrations of gellan gum. Even at concentrations  
 431 below gelation of gellan (0.5% for LA and 0.25% for HA) the gel network caused a decrease in  
 432 availability compared to MD alone (Figure 8). Higher concentrations of gelling agents produce gels

433 which have a higher modulus, more helices, and a smaller pore size (Djabourov et al., 2013). As  
434 shown previously (section 3.1), this change in gel network density had no significant effect on the  
435 diffusional release of glucose because it was much smaller than the pores. However, the behaviour  
436 of MD was different. Release was likely prevented because of entrapment of MD aggregates within  
437 these pores and a network density that limited the accessibility of amylase to reach all parts of the  
438 gel. For gels that slow the release, typically the polymer slows the movement of critical lyzing  
439 enzymes into the gel (McClements and Xiao, 2014). A comparison to other gelling agents with  
440 differing network types and structural arrangements, was thought to also have an impact on MD  
441 availability.

442 The structural influence of gelling agents was hypothesized to be based on the type of mixed gel  
443 network. Both HA and LA gellan gum are known to form interpenetrating polymer networks (IPNs)  
444 with MD and the structures have been described as MD aggregates within pores of the gellan  
445 network (Clark et al., 1999; Kanyuck et al., 2021a). Phase separated networks are known to form  
446 with gelatin (Kasapis et al., 1993a), agarose (Loret et al., 2005), and carrageenan (Wang and Ziegler,  
447 2009; Gładkowska-Balewicz, 2017). Additionally, gelatin melted at the measurement temperature  
448 (37 °C). These network characteristics will be compared to explain the structural influences of the  
449 gels on release behaviour. MD with an IPN (HA and LA gellan gum) resulted in the slowest release  
450 and the lowest total release (Figure 9). Non-melting phase-separated gels (k-carrageenan, i-  
451 carrageenan, and agarose) resulted in greater total release than the IPNs but less than MD alone.  
452 The phase-separated melting gel (gelatin) released more than with no gelling agent.



453

454 **Figure 8.** Release profiles (A) of 30% DE 2 MD without any gelling agent (X), HA gellan gum at 0.25%  
 455 (○), 0.5% (●), 1% (●), and 2% (●), and with LA gellan gum at 0.25% (□), 0.5% (■), 1% (■), and 2% (■).  
 456 Part B displays the total release after 48 hours for only MD (X) and each concentration of HA gellan  
 457 (●) and LA gellan (■).  
 458

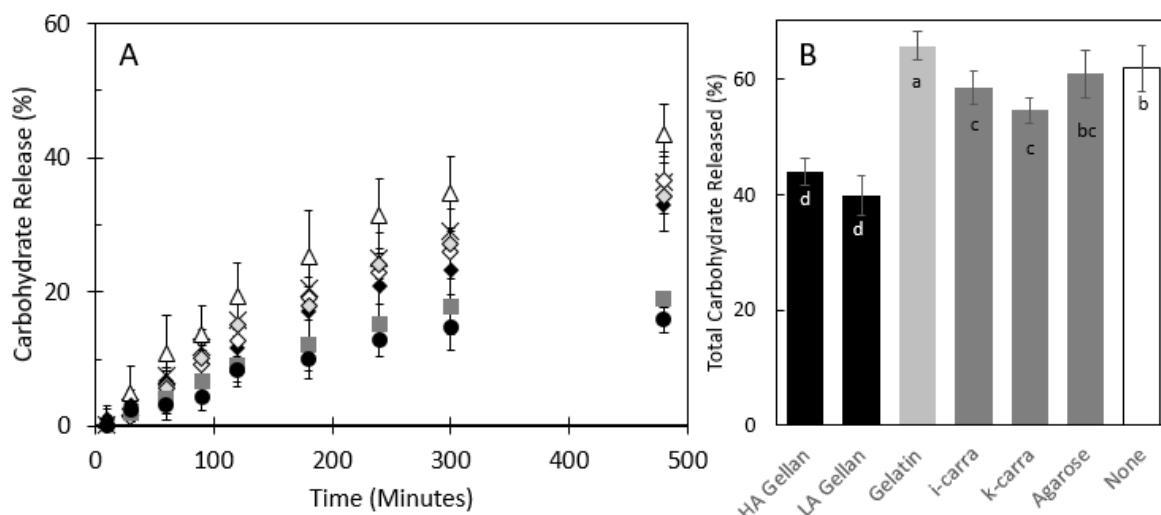
459 For the IPNs, the arrangement of MD aggregates within the pores of the network reasonably could  
 460 have inhibited amylase movement. Additionally, a more heterogeneous arrangement of the  
 461 aggregates and inhibition of the formation of large aggregates (Clark et al., 1999; Kanyuck et al.,  
 462 2021a) could have caused the lower release. Phase separated networks (all characterized as MD  
 463 continuous at these concentrations) have gelling agent rich domains dispersed amongst a  
 464 continuous MD phase where the gelling agents is not present (Kasapis 1993, Loret 2005). Separation  
 465 into these domains means the gelling agent would have had less potential to sterically block amylase  
 466 movement through the gel. Consequently, the release from phase separated gels was higher than  
 467 IPNs gels (Figure 9). Clustering by network type confirmed structure was an important factor for  
 468 comparing carbohydrate availability. In other work, hydrocolloids have been shown to decrease the  
 469 amylase digestion of retrograded starch, but differences between gelling agents has largely been  
 470 nominal (no intrinsic ordering or grouping) and the association with mixed gel network type may  
 471 carry over to starch applications.

472 Although gelatin forms a phase separated structure, the network also melted at the analysis  
 473 temperature and was thought to be the cause of the difference from the other phase separated

474 networks. This was confirmed by repeating the release measurement at 25 °C where the profile was  
475 less than MD alone and no longer significantly different from the other phase separated network  
476 (Supplemental Figure 3). Similarity to the other phase separated networks demonstrated the  
477 importance of melting in increasing the total amount released. Melting of a gel network would  
478 suggest it was no longer able to slow amylase from entering gel, but the greater total release from  
479 gelatin could not be explained as simply.

480 Fractionation (self-separation) of MD within phase separated biopolymer systems have been  
481 observed for agarose (Loret et al., 2005) and gelatin (Kasapis et al., 1993b). One phase contained the  
482 larger molecular weight fraction of MD and the other phase the gelling agent mixed with a fraction  
483 of the smaller molecular weight MD chains. It was thought that phase separation may have changed  
484 the structure of helices to increase the availability of carbohydrates (Kasapis et al., 1993b). Any  
485 changes in the MD aggregation and distribution from the gelling agent could have contributed to the  
486 enzyme accessibility. DSC was used to measure the melting temperatures and enthalpy of MD to  
487 detect any changes in the aggregation behaviour with gelatin (Table 3) with thermographs shown in  
488 Supplemental Figure 4. From Table 3 the network enthalpy of a mixed gel with gelatin was not  
489 significantly different than summation of the individual gels. Prior DSC analysis of mixed gels of  
490 gelatin and MD was not able to achieve this resolution (Kasapis et al., 1993a). Enthalpy of MD with  
491 HA gellan was demonstrated to be not significantly different than alone (Kanyuck et al., 2021a). No  
492 change in enthalpy or melting temperature (Table 3) suggested any fractionation of MD did not  
493 change the helix or aggregate formation. It was possible that the smaller MW chains that separate  
494 do not participate in helix formation and thus the change was not detectable. From these results,  
495 mixed gels did not cause a measureable change in the helix formation. This suggests the greater  
496 percentage of carbohydrates released from gelatin was caused by an organisational difference.  
497 Melting of gelatin regions may have allowed the amylase increased accessibility through the MD  
498 continuous network by liquefying the gelatin phase regions. These structural differences of network  
499 type with MD were shown to be predictive of the amount of carbohydrate release.





500

501 **Figure 9.** Release profiles from mixed gels of 30% MD comparing gelling agents forming and IPN (HA  
 502 gellan (●) and LA gellan (■)) and phase separated networks (i-carrageenan (◇), k-carrageenan (◆),  
 503 agarose (♦), and gelatin (Δ)) and no gelling agent (X). Gelatin melted at the release temperature.  
 504 Total release (B) is shown for IPN (black), phase separated networks (grey), and no gelling agent  
 505 (white).

506

507 **Table 3.** Peak melting temperatures and enthalpy from DSC heating thermographs for MD and  
 508 gelatin independently and the mixed gel of both hydrocolloids. Curves are shown in Supplemental  
 509 Figure 3.

	Total Enthalpy (J/g)	Gelatin		MD	
		Peak (°C)	Enthalpy (J/g)	Peak (°C)	Enthalpy (J/g)
MD (30%)	3.18 ± 0.2			70 ± 2	3.18 ± 0.2
Gelatin (2%)	0.59 ± 0.2	33 ± 0.3	0.59 ± 0.2		
<i>Summation</i>	<i>3.77 ± 0.2</i>				
MD (30%) with Gelatin (2%)	3.60 ± 0.3	33 ± 0.3		71 ± 1	

510

511 **4. Conclusion**

512 Carbohydrate size and a hydrocolloid's response to stimuli were shown to be important for all types  
 513 of release. Gel structure, specifically the network type, was influential for the larger aggregate-  
 514 forming MD but not for glucose. Glucose offered a simplified system to compare the effects of  
 515 responses to stimuli (melting and swelling) of hydrocolloid gels on release. The structuring of MD  
 516 introduced dependencies on amylase accessibility, self-aggregation, and the microstructure of the  
 517 system. Interestingly, because MD does not undergo gelatinization, the results may provide an

518 indication of the effects of starch retrogradation within hydrocolloid networks on the carbohydrate  
519 availability. The complex findings from this simplified two polymer system is another demonstration  
520 of the complexity of food digestion when dealing with multiple ingredients. The work presented  
521 here provides a framework for formulating and processing to achieve specific carbohydrate release  
522 profiles from gels. Future work, including evaluation of digestion characteristics in humans, is  
523 encouraged.

524

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530

531 **CRedit author statement.**

532 **Kelsey M. Kanyuck:** Conceptualization, Methodology, Formal analysis, Investigation, Writing -  
533 original draft. **Tom B. Mills:** Supervision, Writing - review & editing. **Ian T. Norton:** Supervision,  
534 Funding acquisition, Writing - review & editing. **Abigail B. Norton-Welch:** Supervision, Funding  
535 acquisition, Writing - review & editing.

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