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1	Physicochemical functionality of chimeric isomaltomegalosaccharides with α -(1 \rightarrow 4)-glucosidic
2	segments of various lengths
3	
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13	
14	Abbreviations: DDase, dextran dextrinase; DP, average value of degree of polymerization; Gn,
15	maltodextrin with $DP = n$; IMS, isomaltomegalosaccharide; IMS- p/q (e.g., IMS-15/9), IMS with DPs
16	of p and q for α -(1 \rightarrow 6)- and α -(1 \rightarrow 4)-segments, respectively; K _c , stability constant; K _s , equilibrium
17	constant; MWCO-2000 and MWCO-3500, dialysis tubes with molecular weight cutoffs of 2,000 and
18	3,500, respectively; QC, quercetin; Q3G, quercetin-3- <i>O</i> -β-glucoside; TDT; thermal decomposition
19	temperature; TGA, thermogravimetric analysis; TNS, 2-p-toluidinylnaphthalene-6-sulfonate.
20	
21	Abstract
22	Isomaltomegalosaccharide (IMS) is a long chimeric glucosaccharide composed of α -(1 \rightarrow 6)-
23	and α -(1 \rightarrow 4)-linked segments at nonreducing and reducing ends, respectively; the hydrophilicity and
24	hydrophobicity of these segments are expected to lead to bifunctionality. We enzymatically
25	synthesized IMS with average degrees of polymerization (DPs) of 15.8, 19.3, and 23.5, where α -
26	$(1\rightarrow 4)$ -segments had DPs of 3, 6, and 9, respectively. IMS exhibited considerably higher water
27	solubility than maltodextrin because of the α -(1 \rightarrow 6)-segment and an identical resistance to thermal

28	degradation as short dextran. Interaction of IMS with a fluorescent probe of 2-p-
29	toluidinylnaphthalene-6-sulfonate demonstrated that IMS was more hydrophobic than maltodextrin,
30	where the degree of hydrophobicity increased as DP of α -(1→4)-segment increased (9 > 6 > 3).
31	Fluorescent pyrene-estimating polarity of IMS was found to be similar to that of methanol or 1-
32	butanol. The bifunctional IMS enhanced the water solubility of quercetin-3-O-glucoside and quercetin:
33	the solubilization of less-soluble bioactive substances is beneficial in carbohydrate industry.
34	
35	Keywords: dextran dextrinase, isomaltomegalosaccharide, chimeric structure, bifunctionality,
36	quercetin flavonoid, water solubility enhancement
37	
38	1. Introduction
39	Glucosaccharides, such as soluble starch and dextran, are hydrophilic due to the presence of
40	numerous surface hydroxyl groups that promote water solubility. Monosaccharide glucose containing
41	methine groups has a negligibly weak hydrophobicity. However, α -(1 \rightarrow 4)-glucosaccharides (i.e.,
42	maltodextrin and cyclodextrin) exhibit high hydrophobicity arising from stereochemical constrained
43	chains with helical structures (Balasubramanian et al., 1993), whereby the internal surface is relatively
44	apolar because all the methine groups point inwards. The degree of apolarity of the sugar chain can be
45	evaluated by hydrophobic interaction with fluorophore probe 2 - p -toluidinylnaphthalene-6-sulfonate
46	(TNS) (Das et al., 1995) because the emission intensity of TNS is higher in a nonpolar state (i.e., in
47	sugar chain) than in an aqueous state. TNS interacts with the helical chain of maltodextrin where this
48	α -(1 \rightarrow 4)-glucosaccharide forms a helical conformation for degrees of polymerization (DPs) greater

than 5 and an incipient helix-type structure for lower DPs (Sundari et al., 1991; Aoyama et al., 1992).

50 By contrast, dextran composed of sole α -(1 \rightarrow 6)-glucosyl units possesses no hydrophobicity since this

51 α -(1 \rightarrow 6)-glucosaccharide does not affect the fluorescence of TNS (Sundari & Balasubramanian, 1997),

52 indicating that dextran is a hydrophilic saccharide. Functional properties are affected by the type and

53 length of glucosaccharide chains. In this study, we hypothesize that a glucosaccharide containing both

54 α -(1 \rightarrow 4)- and α -(1 \rightarrow 6)-glucosidic segments ["chimeric" saccharide; i.e., isomaltomegalosaccharide

(IMS)] (Lang et al., 2022) could exhibit the bifunctionality, that is, both hydrophobicity and
hydrophilicity.

57 Dextran dextrinase (DDase; EC 2.4.1.2) is a glucosyltransferase that catalyzes a two-substrate 58 reaction with maltodextrins as the substrates. DDase can cleave an α -(1 \rightarrow 4)-linkage at the 59 nonreducing end of a donor substrate and transfer the glucosyl unit to the nonreducing end of an 60 acceptor substrate to generate an α -(1 \rightarrow 6)- or α -(1 \rightarrow 4)-linkage (Yamamoto et al., 1992). Consecutive 61 transfers from the α -(1 \rightarrow 4)-linkage result in the extension of the α -(1 \rightarrow 6)-linked segment, whereby 62 the enzyme finally synthesizes a macromolecule of viscous dextran (Yamamoto et al., 1993). 63 Theoretically, the dextran produced by DDase is not an α -(1 \rightarrow 6)-glucan because the reducing terminal 64 region is composed of α -(1 \rightarrow 4)-linkages originating from maltodextrin of the acceptor substrate, 65 implying the product is a "chimeric" polysaccharide with α -(1 \rightarrow 6)- and α -(1 \rightarrow 4)-linked segments. 66 Recently, we found that intermediate-sized saccharides could be effectively produced by DDase 67 from Gluconobacter oxydans ATCC 11894 under optimized conditions (Lang et al., 2022). The 68 product is called an IMS based on the definition of a megalosaccharide (Thoma et al., 1959), for which 69 the DP ranges between that of an oligosaccharide (DP = 2-9) and a polysaccharide. The DP of a 70 polysaccharide is not well-defined, but the properties of polysaccharides emerge for DPs of more than 71 100 or 200 (Kitamura S, personal information); thus, the DP of a megalosaccharide is considered to be 72 10–100 (or 100–200). The IMS is a chimeric saccharide comprised of α -(1 \rightarrow 6)- and α -(1 \rightarrow 4)-73 segments, where the α -(1 \rightarrow 4)-segment is derived from a maltodextrin substrate. The IMS is expected 74 to possess a low viscosity and a high water solubility because the linear α -(1 \rightarrow 6)-segment promotes 75 hydrophilicity, as suggested by Sundari et al. (1991) and Balasubramanian et al. (1993). An 76 in vivo study was performed to investigate the functionality of IMS. The IMS with short α -(1 \rightarrow 4)-77 segment enhanced the absorption of quercetin-3-O- β -glucoside (Q3G) in the rat small intestine 78 (Shinoki et al., 2013). In addition, higher enzymatic degradation of an azobenzene dye (ethyl red, an 79 environmental pollutant) by azoreductase occurred in the presence of an IMS than in the presence of 80 β-cyclodextrin possibly because of weak hydrophobic interaction between the IMS and ethyl red (Lang et al., 2014). Strong interaction of β -cyclodextrin–ethyl red complex inhibited enzymic 81 degradation by its steric hindrance. Recently, an IMS was reported to enhance the barrier function of 82

83 intestinal epithelial tight junctions, which could facilitate inflammation suppression and reduce the
84 risks of chronic diseases (Hara et al., 2017).

85 In this study, we revealed the bifunctionality of the chimeric IMS with α -(1 \rightarrow 4)-segments of various lengths: i.e., three IMSs with DPs of 15.8, 19.3, and 23.5 (IMS-13/3, IMS-13/6, and IMS-15/9, 86 87 respectively) (Fig. 1). Fluorescence studies were performed to determine the hydrophobic 88 characteristics of the IMS substrates: TNS was used to determine the stability constants (K_c s), and the 89 sense of polarity was determined using polycyclic aromatic pyrene; to the best of our knowledge, the 90 sense of polarity was determined for the first time for carbohydrates in this study. The IMS-induced 91 water solubilization of Q3G and quercetin (QC) (which are bioactive flavonoids that are widely used 92 as health supplements) was also evaluated using a phase solubility diagram. The hydrophobicity of 93 IMS facilitates the aqueous solubilization of these flavonoids. Properties related to the α -(1 \rightarrow 6)-94 segment (water solubility and thermal stability) were analyzed.



96 Fig. 1. Proposed structure of isomaltomegalosaccharide (IMS); a, IMS-13/3; b, IMS-13/6; c, IMS-15/9; 97 numbers, DPs (average values) of α -(1 \rightarrow 6)- and α -(1 \rightarrow 4)-segments.

98

99 2. Materials and methods

100 **2.1. Materials**

101 Q3G and QC hydrate were purchased from Extrasynthese (Cedex, France) and Tokyo

102 Chemical Industry (Tokyo, Japan), respectively. Dextran T1 (DP = 7), T1.5 (DP = 10), and T3.5 (DP

103 = 20) were obtained from Pharmacosmos (Holbaek, Denmark). Dextran T10 (DP = 60) and T40 (DP = 104 250) were purchased from Amersham Biosciences (Uppsala, Sweden). Amylose (DP = 28 and DP = 105 600; synthetic products) was purchased from Glico Nutrition (Osaka, Japan). Amylose (DP = 18; corn) 106 was purchased from Hayashibara (Okayama, Japan). TNS was obtained from Sigma-Aldrich 107 (Shinagawa, Japan). Pyrene, polar protic solvents, and polar aprotic solvents were purchased 108 from Nacalai Tesque (Kyoto, Japan). Cellulose dialysis membranes [with molecular weight cutoffs 109 of 2,000 (MWCO-2000) and 3,500 (MWCO-3500)] were purchased from Spectrum (Rancho 110 Dominguez, CA, USA). Maltotriose (G3) and G6/G7 (maltohexaose and maltoheptaose; commercial product Fugioligo) were donated by Nihon Shokuhin Kako (Tokyo, Japan). Maltodextrin with average 111 112 DP = 11.8 (G12), of which DP ranged in 11.2–13.6 (see Fig. 2C), was prepared by the hydrolysis of short amylose (DP = 16.5–19.5; Carbosynth, Berkshire, UK) as follows (Fig. S1): short amylose (100 113 114 mg/mL) was suspended in a 50 mM sodium acetate buffer (pH 5.0) containing 0.02% sodium azide 115 and treated with recombinant pullulanase (5 µL/mL; Sigma–Aldrich, St. Louis, MO, USA) at 37 °C 116 for 2 d to cleave the α -(1 \rightarrow 6)-linkage; the G12 fraction was then isolated by precipitation with 75% (v/v) methanol and dried in vacuo (45% yield). The ¹H-NMR analysis of isolated G12 is shown in Fig. 117 118 S2A and S2B.

119

120 **2.2. Preparation of freeze-dried DDase**

121 Cultivation of G. oxydans, extracellular DDase preparation, and enzyme activity assays were 122 performed according to methods previously used (Lang et al., 2022). Suitable quantities of cells were 123 suspended in a 25 mM sodium acetate buffer (pH 4.2, 50 mL) containing 1% G3 and incubated at 124 30 °C for 3 h under agitation at 200 rpm (rev/min). During incubation of the cells with G3, DDase was 125 secreted extracellularly and produced dextran from G3, which was tightly bound to DDase (dextran-126 bound DDase) (Lang et al., 2022). The resultant dextran-bound enzyme was dialyzed against water 127 containing 0.02% sodium azide at 4 °C for 2 d and then frozen at -80 °C. Dried DDase was obtained 128 by freeze-drying for 2 d using a lyophilizer FDU-1200 (Eyela, Tokyo, Japan).

129

130 **2.3. Production and structural analysis of IMS**

131 As previously reported (Lang et al., 2022), increasing the substrate concentration promoted 132 IMS formation and depressed dextran formation, and increasing the agitation efficiency increased the 133 α -(1 \rightarrow 6)-linkage content of IMS and decreased dextran formation. Freeze-dried DDase (0.1 U/mL) 134 was mixed with 200 mM G3, G4, G6/G7 or G12 containing a 50 mM sodium acetate buffer (pH 4.2), 135 where the reaction volume was 40 mL, and incubated at 50 °C for 96 h in a 200-mL baffled flask 136 under agitation at 100 rpm. An appreciable quantity of the mixture (100–200 μ L) was recovered at the 137 designated time. The reaction was terminated by heating at 100 °C for 20 min, followed by 138 centrifugation at 12,000× g for 10 min at 4 °C to remove the denatured enzyme. The progress of the 139 formation of the α -(1 \rightarrow 6)-linkage of saccharides contained in the reaction mixture was monitored 140 using ¹H-NMR according to a method we have previously reported (Lang et al., 2022). 141 The reaction was allowed to proceed for 96 h, after which IMS and dextran were separated by 142 methanol fractionation based on a procedure we have previously reported (Lang et al., 2022), except 143 that 50% (v/v) methanol was used to precipitate the dextran generated from G3 or G4. The IMSs 144 obtained from G6/G7 and G12 were further purified by dialysis with MWCO-2000 and MWCO-3500, respectively, for 18 h at 4 °C. The IMSs were desalted using an ion exchange resin (Lang et al., 2014). 145 146 The IMS obtained from G6/G7 was treated with porcine pancreatic α -amylase (Sigma–Aldrich) to 147 shorten the length of the reducing terminal α -(1 \rightarrow 4)-segment (Lang et al., 2022). The carbohydrate 148 concentration was measured by the phenol-sulfuric acid method (Dubois et al., 1959) using a glucose 149 standard. Both the average value of DP and type of α -glucosyl linkage of the IMSs were analyzed by 150 ¹H-NMR (Lang et al., 2014) (Fig. S2C and S2D). The saccharide size distribution was estimated using 151 gel permeation HPLC (Lang et al., 2022). According to our previous method (Lang et al., 2022), the 152 length of α -(1 \rightarrow 4)-chain of IMS was investigated by digestion with dextran glucosidase, which 153 catalyzes the exo-wise hydrolysis of α -(1 \rightarrow 6)-glucosidic linkage of substrate at nonreducing end and 154 no attack on α -(1 \rightarrow 4)-glucosidic linkage (Saburi et al., 2006), and the formed maltodextrins were 155 analyzed. Water content of IMS is shown in Table S1.

156

157 **2.4.** Thermal properties and water solubility

158The thermal decomposition of the IMSs was measured with a thermogravimetric analysis159(TGA) apparatus (TG 8120; Rigaku, Tokyo, Japan). An aluminum crucible was used to hold 10 mg of160an oven-dried sample at 40 °C for 1 d. TGA was performed by elevating the temperature from 50 to161500 °C at a heating rate of 10 °C/min under an N2 atmosphere with a flow rate of 50 mL/min. The data162were analyzed using the Thermo plus EVO version 2.060-1 software program (Rigaku).163Water solubility was determined by dissolving the IMS sample (40 mg) in 0.1 mL of water164using a 1.5-mL microcentrifuge tube. The liquid in the tube was mixed thoroughly and left to stand at

165 25 °C for 5 h. The suspension was centrifuged at $12,000 \times$ g at 25 °C for 20 min. The supernatant (50 166 µL) was lyophilized and weighed, and the results were used to calculate the dissolved quantity of the 167 sample in 1 mL of water. Commercial samples of three amyloses and four dextrans were analyzed in 168 the same way.

169

170 **2.5. TNS binding analysis**

Fluorescence titration was conducted to investigate the formation of complexes between 0-10mM IMS and 10 μ M TNS in water on a Hitachi F-4500 spectrometer (Tokyo, Japan) at an excitation wavelength of 360 nm according to a previously reported method (Buranaboripan et al., 2014). K_c was determined from Benesi–Hildebrand plots [Eq. (1)] at an emission intensity of 447 nm for IMS-13/3 and IMS-13/6 and 467 nm for IMS-15/9.

176 $1/(I - I_0) = 1/(I' - I_0) + 1/\{K_c(I' - I_0)[H]\}$ (1)

In the equation above, *I*₀, *I* and *I'* are the initial fluorescence intensities of TNS without saccharide,
with saccharide of different concentrations ([H]), and with saccharide at the maximum concentration,
respectively.

180

181 **2.6. Polarity estimated by pyrene**

182 Ten μ L pyrene in ethanol was added to saccharide in water (final volume, 1 mL), and then the 183 fluorescence spectra were immediately monitored at 25 °C on a Hitachi F-4500 spectrometer at an 184 excitation wavelength of 335 nm, where the ratio of the emission intensity at 375 nm (I₁) to that at 384 185 nm (I₃) was estimated. Final concentrations of pyrene, saccharide, and ethanol were 0.5 μ M, 20–100

187	The same approach was applied to estimate the I_1/I_3 values of polar protic solvents (e.g., methanol)
188	and polar aprotic solvents (e.g., dimethyl sulfoxide). The determined I_1/I_3 values and dielectric
189	constants for the solvents (https://macro.lsu.edu/howto/solvents/Dielectric%20Constant%20.htm) were
190	used to generate calibration curves (see Fig. 3D) to evaluate the polarities of the IMSs and dextrans.
191	
192	2.7. Phase solubility of flavonoids
193	An excess quantity of Q3G or QC (1.0 mg) was mixed with 100 μ L of an aqueous solution
194	containing 0-40 mM saccharide. The suspension was mixed frequently, left to stand at 25 °C for 6 h,
195	and a solution saturated with Q3G or QC was recovered using a previously reported approach (Lang et
196	al., 2014). A portion of the saturated solution was diluted with dimethyl sulfoxide and used to quantify
197	the flavonoid by measuring the absorbance at 360 nm. The apparent equilibrium constant (K_s) was
198	estimated from the phase solubility diagram (Higuchi and Connors, 1965) according to a method we
199	have previously reported (Lang et al., 2014).
200	
201	3. Results and discussion
202	3.1 Production of IMSs with α -(1 \rightarrow 4)-segments of various lengths at the reducing end
203	G. oxydans DDase, a cell surface protein, is secreted from cells when starch hydrolysate
204	(maltodextrin or glucose) is present in the medium (Suzuki et al., 1999; Sadahiro et al., 2015; Lang et
205	al., 2022). Cultivation of cells with maltodextrin (except G2) results in the extracellular generation of
206	dextran-bound DDase, which is more thermostable than the dextran-free enzyme (Lang et al., 2022). It
207	is also known that coexistence with dextran stabilizes the activities of freeze-dried enzymes (Allison et
208	al., 1998; Anchordoquy et al., 2001), suggesting that dextran-bound DDase maintains its activity after
209	freeze-drying and can be stored for long periods. Dextran-bound DDase (0.127 U/mL) was obtained
210	from the medium in which cells (corresponding to 31.5 mg dry cells) were treated with 1.0% G3. The
211	enzyme maintained 94% of its original activity after lyophilization, which makes extended storage of
212	dextran-bound DDase feasible.

mM, and 1% (v/v), respectively. The $I_{\rm l}/I_{\rm 3}$ value of 1% (v/v) ethanol solution was measured to be 1.4.

A 200 mM substrate (G3, G4, G6/G7, or G12) was reacted with 0.1 U/mL freeze-dried DDase for 96 h according to our previous approach (Lang et al., 2022), except that the incubation temperature was increased from 45 °C to 50 °C, without affecting the IMS production yield. Incubation at 50 °C decreased the viscosity of the reaction mixture containing the highly concentrated (200 mM) substrate and encumbered the retrogradation of G12. Long maltodextrin G12 immediately thickened to form a gel at a lower temperature.

219 We recovered the reaction mixture at the indicated time, and all the saccharides contained in 220 the reaction mixture (remaining substrate and formed products) were subjected to ¹H-NMR monitoring 221 (Fig. 2A). The degree of formation of the α -(1 \rightarrow 6)-linkage increased in a sigmoidal manner due to α -(1 \rightarrow 4)-linkage formation by DDase (Lang et al., 2022). Table 1 shows that L_{max} [the maximum α -222 $(1\rightarrow 6)$ -linkage content of all the saccharides] increased from 56.7 to 66.3% for G3–G6/G7 as the DP 223 224 of the substrate increased. The $L_{\rm max}$ value of G12 (62.7%) is an exception of this manner since highly 225 concentrated G12 (200 mM) gradually retrograded during the long reaction, even at 50 °C. The reaction time (T_{50}) at which half of L_{max} was obtained increased in the order G12 > G6/G7 > G4 > G3 226 227 (Table 1), indicating that α -(1 \rightarrow 6)-linkage formation was slower for a longer substrate. 228



229

Fig. 2. Production of isomaltomegalosaccharide (IMS). A, Increase in the α -(1 \rightarrow 6)-linkage content of all saccharides in the reaction mixture. \circ , G3; \bullet , G4; \blacktriangle , G6/G7; \Box , G12. B, Yield of dextran (black) and IMS (red) isolated from the reaction mixture for a 96-h reaction time. Yield (%; w/w), amount of each product per that of substrate used. C, Size distribution of IMS (ii, iii, or v) and substrate (i or iv). i, G6/G7; ii, IMS-13/3; iii, IMS-13/6; iv, G12; v, IMS-15/9.

Table 1. Formation of α -(1 \rightarrow 6)-linkages in saccharides in the reaction mixture and characterization of

200 mM Substrate	All saccharides in reaction mixture		Isolated IMS				
	L_{\max}^{a}	L_{\max}^{a} T_{50}^{b}		α -(1 \rightarrow 6)-content	DP of two chains ^c		
	(%)	(h)		(%)			
G3	56.7	5.9	NO	NO	NO		
G4	61.1	9.8	13.2	70.3	4: 9		
G6/G7	66.3	14.3	19.3	69.3	6: 13		
G12	62.6	18.0	23.5	63.2	9: 15		

237	the isomaltomegalosaccharide	(IMS)	fraction.
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238 ^a Maximal α -(1 \rightarrow 6)-linkage content of all saccharides.

239 ^b Reaction time at which half of L_{max} was obtained.

^c The first and second values denote the DPs of the α -(1 \rightarrow 4)- and α -(1 \rightarrow 6)-segments, respectively.

241 NO was not obtained because of a low IMS yield from G3.

242

The subsequent DDase reaction at 96 h showed that an adequate content of α -(1 \rightarrow 6)-segment 243 244 was generated for each substrate (Fig. 2A); therefore, the product was separated into IMS and dextran 245 by methanol fractionation. The IMSs from G6/G7 and G12 were subjected to further purification by dialysis using MWCO-2000 and MWCO-3500, respectively, to remove saccharide components with 246 247 DPs below 12 and 22, respectively. Fig. 2B shows the yield of purified IMS or dextran (w/w, as a percentage of the initial quantity of substrate), demonstrating that the production of both IMS and 248 249 dextran increases with the DP of the maltodextrin substrates. IMSs were obtained in a high yield from the reaction of G6/G7 and G12 (36 and 49%, respectively) but in a very low yield from the reaction of 250 251 G3 and G4 (0.04 and 0.21%, respectively), where the main products were oligosaccharides. Even for a small yield, the IMS from G4 had a DP of 13.2, as analyzed by ¹H-NMR. When freeze-dried DDase 252 253 was reacted with highly concentrated G3 (500 mM) for 96 h, the IMS yield increased to 15.8%, but 254 the DP of the α -(1 \rightarrow 6)-segment was low [DP = 12.0 with nine α -(1 \rightarrow 6)-glucosyl units]. 255 The IMSs produced from G6/G7 and G12 (corresponding to IMS-13/6 and IMS-15/9, 256 respectively) had the estimated DPs of 19.3 and 23.5, respectively (Table 1), and contained α -(1 \rightarrow 4)-257 glucosyl units with average DPs of 6 and 9 at the reducing ends, respectively (Fig. 1). We used IMS-

238	15/6 as a substrate, for which the α -(1 \rightarrow 4)-segment was selectively cleaved by α -amylase (Lang et al.,
259	2020) and converted into a maltodextrin moiety (average $DP = 3$) to generate an additional IMS (IMS-
260	13/3) with $DP = 15.8$. The size distributions of the three IMSs were compared with that of the
261	substrate used (Fig. 2C). Chain length of α -(1 \rightarrow 4)-segment of each IMS was analyzed by dextran
262	glucosidase digestion (Lang et al., 2020), and maltodextrins corresponding to α -(1 \rightarrow 4)-segment were
263	formed (Fig. S3). The IMS structures are shown in Fig. 1, where IMS-13/3 and IMS-13/6 have the
264	same α -(1 \rightarrow 6)-segment structure with average DP = 13, and IMS-15/9 has a longer α -(1 \rightarrow 6)-segment
265	with average DP = 15. The α -(1 \rightarrow 6)-segment contents were 83.3, 69.3, and 63.2% for IMS-13/3, IMS-
266	13/6, and IMS-15/9, respectively.

250

268 **3.2 Thermal decomposition and water solubility of IMS**

269 The activation energy for amylopectin is higher compared with amylose (Pigłowska et al., 270 2020). This phenomenon can be explained by its high molecular weight and α -(1 \rightarrow 6)-segment content (Liu et al., 2010), suggesting that the α -(1 \rightarrow 6)-linked saccharide is more stable than the α -(1 \rightarrow 4)-271 272 linked saccharide. Therefore, an IMS containing these two segments should be more stable than 273 amylose with a similar size. To investigate the IMS chimeric property, we used TGA to measure the 274 apparent thermal decomposition temperature (TDT) of the IMSs, with amylose and dextran as control 275 saccharides. Under heating in a TGA apparatus in an N2 atmosphere, IMS-15/9 remained unchanged 276 below 260 °C except for moisture loss, followed by thermal decomposition corresponding to a total 277 weight loss of 89.13% at 500 °C (Fig. 3A). Similar curves were observed for the other samples. The 278 decomposition peak in the differential thermal analysis curve (Fig. 3A) was used to estimate the TDT of 279 each saccharide (Table 2). The TDT of amylose increased with DP, whereas that of dextran did not 280 [Table 2 shows that dextran T10 (DP = 60) and T40 (DP = 250) have lower TDTs than T1 (DP = 7)]. 281 This result is probably due to the low quantities of other glucosidic linkages, such as α -(1 \rightarrow 2)- or α -282 $(1 \rightarrow 3)$ -bonds, in the chains of the dextrans T10 and T40 (commercially available samples). A comparison of the TDTs of saccharides with nearly identical DPs [i.e., dextran T3.5 (DP = 20; TDT = 283 284 304.7 °C), IMS-13/6 (DP = 19.3; TDT = 304.7 °C), and amylose (DP = 20; TDT = 297.6 °C

285 (Saavedra-Leos et al., 2015)] shows that the thermal stability of IMS is similar to that of dextran and



higher than that of amylose.



288 Fig. 3. Characterization of isomaltomegalosaccharide (IMS). A, Thermal decomposition of IMS-15/9; DTA, differential thermal analysis. **B**, Emission spectra of 10 µM TNS in water (thin black line) and in 289 290 10 mM glucosaccharide; green, IMS-13/3; blue, IMS-13/6; thick black, IMS-15/9; red, G7. C, 291 Normalized pyrene emission spectra; black, water containing 1 % (v/v) ethanol (blank); blue, IMS-292 13/3; green, IMS-13/6; red, IMS-15/9. **D**, Calibration curves for I_1/I_3 versus the apparent dielectric 293 constant: polar protic solvents (•, in the direction of the black arrow, methanol, ethanol, 2-propanol, 1butanol, and 1-hexanol; Y = 0.015X + 0.647, $R^2 = 0.922$: Y and X represent I_1/I_3 and apparent 294 295 dielectric constant, respectively) and polar aprotic solvents (\circ , in the direction of the black arrow, dimethyl sulfoxide, acetonitrile, acetone, dichloromethane, and tetrahydrofuran; Y = 0.011X + 1.047, 296 297 $R^2 = 0.979$). Red arrows from left to right, IMS-15/9, IMS-13/6, and IMS-13/3 (100 mM for each). 298 Ethanol solution of pyrene was used, and the final concentration of ethanol was 1% (v/v).

Glucosaccharide	Decomposition temperature	Water solubility	$K_{\rm c}$ by TNS	I_1/I_3 determined by pyrene ^a (dielectric			<i>K</i> _s for Q3G	<i>K</i> _s for QC
				constant ^b)				
(DP)	(°C)	at 25 °C (mg/mL)	(M^{-1})	100 mM	40 mM	20 mM	(M^{-1})	(M^{-1})
IMS								
IMS-13/3 (15.8)	297.1	> 400	14.8	1.12 (31.3)	1.24 (39.5)	1.39 (49.5)	9.0	1.4
IMS-13/6 (19.3)	304.7	> 400	55.2	0.88 (15.8)	0.92 (18.2)	0.95 (20.2)	12.5	1.5
IMS-15/9 (23.5)	305.7	> 400	114	0.87 (15.1)	0.89 (16.2)	0.93 (18.9)	56.3	3.7
Amylose								
G5 (5)			1.9°					
G6 (6)			8.8 ^c					
G7 (7)			27°				6.01	0
Corn (18)		15.5 ± 2.1						
Amylose (20)	279.6 ^d							
Synthetic (28)	304.2	10.5 ± 1.5						
Synthetic (600)	309.8	0.52 ± 0.02						
Dextran								
T1 (7)	302.4	> 400		1.29 (42.9)	1.34 (46.2)	1.38 (48.9)	5.73	0
T1.5 (10)				1.25 (40.1)	1.25 (40.2)	1.29 (42.9)	5.86	0
T3.5 (20)	304.7	> 400		1.27 (41.2)	1.28 (42.2)	1.28 (42.2)		
T10 (60)	297.9	> 400		1.31 (44.5)	1.33 (45.5)	1.32 (44.9)		
T40 (250)	299.9	> 400 (swelling)						

Table 2. Physical properties of isomaltomegalosaccharide (IMS), amylose, and dextran.

^a Pyrene in ethanol solution was added to saccharide sample, where the final concentration of ethanol was 1% (v/v).

- ^b Estimated using the standard curve presented in Fig. 3D.
- ^c Aoyama et al., 1992.
- 303 ^d Saavedra-Leos et al., 2015.

304	IMS possessed a high water solubility identical to that of short dextran (T3.5; $DP = 20$), i.e., >
305	400 mg/mL at 25 °C (Table 2). Large dextran (T40; DP = 250) maintained a high water solubility and
306	became swollen and very viscous. By contrast, amylose with $DP = 18-28$ exhibited a low solubility of
307	15–10 mg/mL (Table 2). Long amyloses typically have poor water solubility (Mukerjea & Robyt,
308	2010) because the crystalline packing of double helices in A- and B-type conformations reduces the
309	binding capacity of water (Naknean & Meenune, 2010). The results obtained indicate that the high
310	aqueous solubility of IMS derives from the α -(1 \rightarrow 6)-segment. Hence, our findings elucidate the effect
311	of the α -(1 \rightarrow 6)-segment on thermal stability and water solubility, indicating the advantage offered by
312	the chimeric IMS structure.

314 **3.3 Using fluorescent TNS and pyrene to monitor the hydrophobicity of IMS**

315 The hydrophobic interaction between TNS and dextran was reported as undetectable, because 316 the methine groups in a flexible twofold crankshaft-like conformation of dextran are turned toward the bulk water medium (Sundari & Balasubramanian, 1997). However, the α -(1 \rightarrow 4)-segment of amylose 317 318 considerably increases the degree of hydrophobicity, the extent of which depends on the chain length, 319 which generates curved nonpolar surfaces or several helical forms. The hydrophobic interaction is 320 weak between TNS and G3 and not very selective at binding, but it is detectable for maltodextrin with 321 DP of 5 or higher. Perhaps maltodextrins with $DP \ge 5$ can undergo an induced-fit adjustment to 322 interact with hydrophobic ligands (e.g., TNS), similar to the hydrophobic cavity of cyclodextrin 323 (Aoyama et al., 1992). The K_{cs} of G5 (maltopentaose), G6, and G7 have been estimated using TNS to 324 be 1.9, 8.8, and 27 M⁻¹, respectively (Aoyama et al., 1992). These results suggest that IMS possesses 325 hydrophobicity that mainly originates from the α -(1 \rightarrow 4)-segment. The K_cs of IMS-13/3, IMS-13/6, 326 and IMS-15/9 were evaluated to be 14.8, 55.2, and 114 M⁻¹, respectively (Fig. S4), as expected (Table 2), and increase in the order of increasing length of the α -(1 \rightarrow 4)-segment (average DP = 9 > 6 > 3). 327 328 The fluorescence intensity of IMS-15/9 was significantly increased, and the maximum wavelength 329 shifted to 467 nm from 447 nm for IMS-13/3 and IMS-13/6 (Fig. 3B), suggesting that the hydrophobic 330 area of IMS-15/9 occupies a more polar environment than of IMS-13/3 and IMS-13/6 (Barel, 1975). The same phenomenon was also observed for G7, where the maximum wavelength was 467 nm (Fig. 331

332 3B), which further supports that the high hydrophobicity of IMS-15/9 originates from the α -(1 \rightarrow 4)-333 segment.

334 Furthermore, the polarity of the IMS hydrophobicity was examined by monitoring the 335 fluorescence spectrum of the complex formed between 0.5 μ M pyrene and 100 mM IMS (Fig. 3C), 336 which was used to estimate I_1/I_3 (Table 2). A high I_1/I_3 indicates a high polarity. The I_1/I_3 of IMS-13/3, 337 IMS-13/6, and IMS-15/9 showed that the I_1/I_3 decreased as the length of the α -(1 \rightarrow 4)-segment of IMS 338 increased, indicating that the IMS hydrophobicity derives from the α -(1 \rightarrow 4)-segment. The α -(1 \rightarrow 6)-339 saccharide dextran (DP = 7–60; 100 mM) possessed larger I_1/I_3 values than 100 mM IMS (Table 2). 340 We measured I_1/I_3 for polar protic and polar aprotic solvents. $I_1/I_3 = 0.81 - 1.12$ was measured for polar 341 protic solvents, which contain a hydroxy group (and/or other polar protic groups) that can donate a 342 proton, whereas $I_1/I_3 = 1.12 - 1.57$ was measured for polar aprotic solvents that do not contain such groups. The following solvents were investigated: polar protic solvents, methanol ($I_1/I_3 = 1.12$; 343 dielectric constant = 32.7), ethanol (1.04; 24.6), 2-propanol (0.97; 19.9), 1-butanol (0.91; 17.5), and 1-344 345 hexanol (0.81; 13.3); polar aprotic solvents, dimethyl sulfoxide (1.57; 46.7), acetonitrile (1.45; 37.5), 346 acetone (1.33; 20.7), dichloromethane (1.13; 8.93), and tetrahydrofuran (1.12; 7.58). Fig. 3D shows 347 the calibration curves (I_1/I_3 versus the dielectric constant) of the polar protic and aprotic solvents, and 348 the calibration curves of the polar protic solvents was used to estimate the IMS dielectric constants 349 (Table 2), because carbohydrates have polar hydroxyl groups. These dielectric constants indicate that 350 the polarities of IMS-13/3, IMS-13/6, and IMS-15/9 are comparable to methanol, 1-butanol, and 1-351 butanol, respectively. Table 2 shows that the I_1/I_3 s of 20 and 40 mM IMS are lower than those of the 352 20 and 40 mM dextrans, demonstrating that IMS hydrophobicity is maintained at low concentrations. 353 Fluorescence studies using TNS and pyrene indicate that the IMS hydrophobicity mainly originates 354 from the α -(1 \rightarrow 4)-segment.

355

356 **3.4 Solubility enhancement of QC and Q3C by IMS**

357 QC flavonoids are known to have poor water solubility: $4.7 \pm 0.4 \mu$ M for QC and 156.3 ± 5.5

358 μM for Q3G at 25 °C in water (both showed pH 6.80). This characteristic derives from the methyl

359 groups and *iso*-pentyl groups in these structures that increase lipophilicity (Crozier et al., 2009), which





375 Fig. 4. Phase solubility diagram for solubilization of quercetin-3-O-β-glucoside (Q3G) and quercetin 376 (QC) by saccharides. A, Q3G; B, QC; □, IMS-15/9; △, IMS-13/6; ■, IMS-13/3; ○, G7; •, T1; ▲, T1.5. 377 The hydrophobicity of IMS mainly originates from its α -(1 \rightarrow 4)-segment. However, both the 378 $K_{\rm c}$ for TNS and $K_{\rm s}$ for QC flavonoids are considerably higher than those of maltodextrins (Table 2). 379 We cannot completely explain these phenomena, but a ¹H-NMR analysis revealed that ethyl red interacted with the anomeric proton of the α -(1 \rightarrow 6)-segment in IMS-8/3 (Lang et al., 2014). These 380 results suggest that the high K_c and K_s derive from the α -(1 \rightarrow 6)-segment connecting the α -(1 \rightarrow 4)- and 381 382 α -(1 \rightarrow 6)-segments.

384 4. Conclusions

In the present study, we enzymatically synthesized IMSs containing α -(1 \rightarrow 4)-segments of various lengths from maltodextrins and investigated the IMS functions. The α -(1 \rightarrow 6)-segment of an IMS is hydrophilic and contributes to high water solubility and resistance to thermal decomposition. The α -(1 \rightarrow 4)-segment of an IMS can interact with hydrophobic ligands, and interaction depends on the length of α -(1 \rightarrow 4)-chain. The IMS chimeric nature, that is, bifunctionality of hydrophilicity and hydrophobicity, enhances the aqueous solubility of compounds with low water solubility (e.g., QC flavonoids), suggesting that IMSs could be extensively applied to industrial fields.

392

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402 CRediT authorship contribution statement

Weeranuch Lang: Conceptualization, Investigation, Formal analysis, Writing - original draft. Yuya
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Supervision. Atsuo Kimura: Writing - review & editing, Resources, Supervision, Funding acquisition,
Project administration.

409

410 **Declaration of Competing Interest**

- 411 The authors have no conflicts of interest to declare.
- 412

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