Preparation, structure-property relationships and applications of different emulsion gels: Bulk emulsion gels, emulsion gel particles, and fluid emulsion gels

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1	Preparation, structure-property relationships and applications of different
2	emulsion gels: bulk emulsion gels, emulsion gel particles, and fluid
3	emulsion gels
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### 12 Abstract

Background: In recent years, there has been increasing interest in emulsion gels, due to their
better stability during storage and potential for prolonged intestinal drug release compared to
emulsions. There are three kinds of emulsion gels, classified according to their morphological
properties: bulk emulsion gels, emulsion gel particles and liquid emulsion gels.

Scope and approach: This paper provides a comprehensive review of the mechanisms and
procedures of different methods for preparing different emulsion gels and relationships
between structures and properties of emulsion gels. The applications of emulsion gels in the
food industry are finally discussed.

Key findings and conclusions: Different emulsion gels result from different preparation 21 methods, and have various structure-property relationships and applications. Many methods 22 can be used to prepare bulk emulsion gels, involving different matrix materials, processing 23 techniques, and purposes. This can result in different structures of gel matrixes and emulsion 24 droplets and interactions between them, which can influence the structures of bulk emulsion 25 gels and then their mechanical and release properties. On the other hand, extrusion and 26 impinging aerosol methods are two methods for preparing emulsion gel particles, while liquid 27 28 emulsion gels can be prepared by Pickering emulsions and disrupted gel systems. Rheological, syneresis and swelling properties are critical for gel particle suspensions, while 29 flow behaviour and release properties are important to liquid emulsion gels. In addition, fat 30 replacements and delivery systems are main applications of emulsion gels in the food 31 32 industry. However, current research has mainly focused on bulk emulsion gels, so further studies on emulsion gel particles and liquid emulsion gels are required. 33

34 **Keywords:** emulsion gel; preparation; interaction; structure; property; fat replacer; delivery.

### 35 **1. Introduction**

36 Emulsion gels, also known as emulgels or gelled emulsions (Balakrishnan, Nguyen, Schmitt, Nicolai, & Chassenieux, 2017), are complex colloidal materials in which both emulsion 37 38 droplets and gels exist (Dickinson, 2012). According to the state of emulsion droplets in gels, structures of emulsion gels can be divided into two categories: emulsion droplet-filled gels 39 and emulsion droplet-aggregated gels (Fig. 1). In emulsion droplet-filled gels, the continuous 40 phase (e.g., protein- and polysaccharide-based gels) forms a continuous gel matrix which can 41 be defined as the support of emulsion gels, and emulsion droplets are embedded in this gel 42 matrix. In emulsion droplet-aggregated gels, emulsion droplets aggregate together and form a 43 44 network structure, such that the gel matrix is disrupted by the aggregated emulsion droplets. In most cases, the structural state of emulsion droplets is a combination of these two different 45 structures (i.e., partially droplets filled in the gel matrix and partially aggregated droplets), 46 47 probably owing to the inhomogeneous distribution of emulsion droplets. Moreover, according to the interactions between emulsion droplets and the gel matrix, emulsion droplets can be 48 49 divided into active and inactive fillers (Fig. 2). Active fillers are mechanically connected to 50 the gel network through emulsifiers by noncovalent and/or covalent bonds, especially when emulsifiers are natural molecules (e.g., proteins, egg lecithin, and soy lecithin); in contrast, 51 inactive fillers have little chemical or physical affinity with the molecules of gel matrix, 52 especially when low molecular weight (LMW) emulsifiers or no emulsifiers are used and 53 matrix materials have weak emulsifying properties (Van Vliet, 1988). 54

Preparing emulsion gels normally includes two steps: first preparing emulsions and then turning emulsions into gels. During the last decade, emulsion gels have received growing interest, due to their advantages compared to emulsions, such as higher stability during storage, owing to decreased oil movement and oxygen diffusion within the systems (Cofrades, Bou, Flaiz, Garcimartin, Benedi, Mateos, et al., 2017; Corstens, Berton-Carabin,

60	Elichiry-Ortiz, Hol, Troost, Masclee, et al., 2017; Lim, Kim, Choi, & Moon, 2015; Ma, Wan,
61	& Yang, 2017; Sato, Moraes, & Cunha, 2014), controlled and prolonged gastric and/or
62	intestinal drug release because of the protection by the gel matrix (Corstens, et al., 2017;
63	Guo, Bellissimo, & Rousseau, 2017), and practical applications, including overcoming the
64	textural problems caused by lipid particles in food products and mimicking the effect of fat
65	on hardness and water-holding capacity of meat products (Alejandre, Poyato, Ansorena, &
66	Astiasaran, 2016; Brito-Oliveira, Bispo, Moraes, Campanella, & Pinho, 2017).
67	Bulk emulsion gels, emulsion gel particles and liquid emulsion gels are three kinds of
68	emulsion gels (Fig. 3), which exhibit their own particular properties, due to their different
69	morphological properties. The size and shape of bulk emulsion gels are determined by the
70	emulsion volume and emulsion containers of different shapes and sizes used during the
71	emulsion gel preparation, and bulk emulsion gels can also be broke into smaller pieces with
72	different sizes and shapes. Therefore, mechanical properties (including viscoelastic and
73	textural properties) of bulk emulsion gels are important. Emulsion gel particles are normally
74	spherical with sizes (diameters) from nano to macro (Ching, Bansal, & Bhandari, 2017).
75	Thus, mechanical properties are also important for the macrogel particles. However, emulsion
76	gel particles can be dispersed in aqueous media, and gel particles allow swelling or de-
77	swelling as a function of environmental conditions, allowing turning of their size and/or
78	physicochemical properties (Torres, Tena, Murray, & Sarkar, 2017). There are two types of
79	fluid emulsion gels: gel-like Pickering emulsions and disrupted emulsion gel systems. Fluid
80	emulsion gels do not have solid shapes, but they have higher viscoelasticity than conventional
81	emulsions. Gel-like Pickering emulsions are similar to bulk emulsion gels, exhibiting a solid
82	state (Zou, Guo, Yin, Wang, & Yang, 2015), while disrupted emulsion gels normally exhibit
83	fluid characteristics (Soukoulis, Cambier, Hoffmann, & Bohn, 2016). This paper provides an

84 overview of the current knowledge of preparation methods, structure-property relationships,

85 and applications of different emulsion gel systems.

## 86 2. Preparation of different emulsion gels

87 2.1. Bulk emulsion gels

As shown in Table 1, proteins (e.g., myofibrillar protein, whey protein, soy protein, gelatin, 88 bovine serum albumin, sodium caseinate, and casein), polysaccharides (e.g., carrageenan, 89 gellan gum, agar, alginate, and inulin), and LMW compounds (e.g., sapoin glycyrrhizic acid 90 and the mixture of  $\beta$ -sitosterol and  $\gamma$ -oryzanol) are normally used as matrix materials in bulk 91 emulsion gels. According to the gelation process, preparation methods for bulk emulsion gels 92 include heat-set and cold-set (including one-step cold-set, cold-set after heat treatment, 93 enzyme treatment, acidification treatment, addition of ions, and self-assembly) gelation 94 methods. Choosing an appropriate method depends on the matrix materials (i.e., proteins, 95 polysaccharides or LMW compounds) and applications of prepared emulsion gels such as 96 mimicking food processing (i.e., heating process of meat), protecting encapsulated nutrients, 97 controlled release of encapsulated nutrients, or obtaining better mechanical properties. For 98 heat-set, one-step cold-set, cold-set after heating, and self-assembled gelation methods, the 99 concentration of proteins, polysaccharides and LMW compounds in the water phase should 100 be higher than the critical gelation concentration to guarantee the gelation. However, for 101 gelation methods based on enzyme treatment, acidification treatment, or addition of ions, the 102 103 concentration of matrix molecules can be below the critical gelation concentration, especially 104 to avoid gelation during the pre-heating process (Ye & Taylor, 2009).

105 2.1.1. Protein-based bulk emulsion gels

106 Several methods have been studied for preparing protein-based bulk emulsion gels: heat treatment, cold-set after pre-heating, acidification, addition of ions, and enzyme treatment, 107 depending on the gelation properties of proteins (Farjami & Madadlou, 2019). 108 Heat treatment can denature proteins, and denatured protein molecules can aggregate and 109 form three-dimensional structures through chemical forces (i.e., disulfide bonds, electrostatic 110 interactions, hydrophobic interactions, hydrogen bonds, and ionic bonds) under appropriate 111 conditions (e.g., protein concentration, pH, and ionic strength) (Tolano-Villaverde, Torres-112 Arreola, Ocaño-Higuera, & Marquez-Rios, 2015). Proteins (e.g., myofibrillar protein (MP), 113 whey protein isolate (WPI), and soy protein isolate (SPI)), which undergo heat-induced 114 gelation, can be used as matrix materials to prepare heat-induced bulk emulsion gels. 115 However, recent studies have focused on MP- and WPI-based bulk emulsion gels (Guo, Ye, 116 Lad, Dalgleish, & Singh, 2013; Wang, Zhang, Chen, Xu, Zhou, Li, et al., 2018). Studying 117 118 heat-induced MP-based bulk emulsion gels is important to develop high quality processed meat products such as sausages and surimi, because the interactions between MPs and fat 119 120 globules or oil droplets play an important role in textual properties and stability of meat products. In addition, heat treatment is the most common method for producing WPI-based 121 bulk emulsion gels in order to investigate the interactions between emulsifiers and the WPI-122 based gel matrix (Chen, Dickinson, Langton, & Hermansson, 2000). 123 One-step cold-set or cold-set after heat treatment is normally used for preparing gelatin-based 124

emulsion gels. The gelation mechanism of gelatin is that, when the gelatin solution is cooled

below 30°C, a self-assembly process of gelatin occurs and helices are created (Gómez-

127 Guillén, Giménez, López-Caballero, & Montero, 2011). Heat treatment (above 40°C) is

128 normally used to increase the solubility of gelatin before cold-set treatment. However, for

129 cold-soluble gelatin, the thermal process is not necessary (Pintado, Ruiz-Capillas, Jimenez-

130 Colmenero, Carmona, & Herrero, 2015). In addition, ethanol has been used to denature

proteins and produce cold-set whey protein emulsion gels (Xi, Liu, McClements, & Zou,2019).

The mechanism of acid-induced protein gelation is that the acidification, usually carried out 133 by adding glucono- $\delta$ -lactone (GDL), decreases the pH and neutralize the surface charges of 134 protein aggregates and a gel network then forms by hydrophobic interactions and Van der 135 Waals forces (Ringgenberg, Alexander, & Corredig, 2013). Before acidification. heat 136 treatment is normally used to denature proteins and form protein aggregates. In such cases, 137 two different processes can be used to produce acid-induced protein-based emulsion gels: 138 using pre-heated-induced protein aggregates to form emulsions (Lu, Mao, Zheng, Chen, & 139 Gao, 2020) or heating native protein-stabilized emulsion to form protein aggregates (Ye & 140 Taylor, 2009) before acidification. However, heating native protein-stabilized emulsions may 141 lead to droplet aggregation, which limits the application of emulsion gels for encapsulation of 142 heat-sensitive compounds (Mao, Roos, & Miao, 2014). 143

Addition of ions (normally  $Ca^{2+}$  in the form of  $CaCl_2$ ) can promote soluble protein aggregates 144 to form a gel network by ionic crosslinks (Wang, Luo, Liu, Adhikari, & Chen, 2019). It has 145 been reported that structures of CaCl<sub>2</sub>-induced SPI emulsion gels were mainly composed of 146 147 particulate protein-coated and were different from filamentous gel networks formed by MTGase and GDL (Tang, Chen, & Foegeding, 2011). In addition, the concentration of Ca<sup>2+</sup> 148 149 can affect the structures of protein-based emulsion gels; Sok Line, Remondetto, & Subirade (2005) found that low calcium concentrations (e.g.,  $11.7 \text{ mM Ca}^{2+}$ ) induced emulsion gels 150 with a fine-stranded structure, while high calcium concentrations (e.g., 40 mM or 68 mM 151  $Ca^{2+}$ ) led to random aggregates. Therefore,  $CaCl_2$  at a concentration of 8–20 mM is normally 152 used to produce Ca<sup>2+</sup>-induced emulsion gels (Liang, Leung Sok Line, Remondetto, & 153 Subirade, 2010; Tang, Chen, & Foegeding, 2011; Ye & Taylor, 2009). 154

155 Microbial transglutaminase (MTGase) can be used to promote cross-links between protein molecules and improve the properties of protein-based emulsion gels (Gaspar & de Goes-156 Favoni, 2015). Compared to other methods, enzyme treatment is a safe method to produce 157 protein-based emulsion gels with high quality under mild process conditions (35–37°C) and 158 without producing any side-products (Tang, Luo, Liu, & Chen, 2013). It was found that the 159 gel strength of MTGase-induced SPI-based emulsion gels was much higher than that of 160 GDL- or CaCl<sub>2</sub>-induced emulsion gels (Tang, Chen, & Foegeding, 2011). Two points should 161 be highlighted when enzyme treatment is used to prepare protein-based emulsion gels. Firstly, 162 the order of adding enzyme into emulsions may influence the properties of emulsion gels. 163 Tang, Yang, Liu, & Chen (2013) found that adding enzyme prior to emulsification required 164 less enzyme, but induced emulsion gels with higher stiffness compared to adding enzyme 165 after emulsification. Secondly, although the formation of protein aggregates is not necessary 166 for producing enzyme-induced gels, unfolding the compact structures of globular proteins 167 (e.g., SPI, WPI, and MP) can provide more target glutamine and lysine residues for the 168 MTGase treatment. For example, pre-incubation of SPI and egg white protein (Alavi, Emam-169 Djomeh, Salami, & Mohammadian, 2020; Tang, Yang, Liu, & Chen, 2013), pre-oxidation 170 treatment of MP (Wang, Xiong, & Sato, 2017), and breaking down disulfide bonds in bovine 171 serum albumin (Kang, Kim, Shin, Woo, & Moon, 2003) can improve gelation by MTGase. 172 However, it has been found that heated SPI-stabilized emulsions after emulsification could 173 174 not form gels following enzymatic treatment (Tang, Chen, & Foegeding, 2011).

175 2.1.2. Polysaccharide-based bulk emulsion gels

176 Several methods have been studied for preparing polysaccharide-based bulk emulsion gels,

such as heat-set, cold-set after pre-heating, addition of ions, and self-assembly

178 (crystallisation), depending on the gelation properties of polysaccharides.

179 Curdlan is a water-soluble  $\beta$ -(1,3)-glucan extracted from *Alcaligenes* faecalis, and curdlanbased emulsion gels can be obtained after heating emulsions, while cold-set after pre-heating 180 is normally used to prepare carrageenan-, agar-, and gellan gum-based emulsion gels (Jiang, 181 et al., 2019). The gelation mechanism involves forming double helices and cross-linking 182 helical domains to create a three-dimensional structure during cooling (Nishinari & 183 Takahashi, 2003). These are all cold-set and thermo-reversible gels. For producing cold-set 184 185 emulsion gels, polysaccharides should be dissolved at a high temperature (normally more than 70°C), and/or emulsions should be prepared at a medium temperature (normally between 186 187 45°C and 70°C), after which emulsion gels are formed at a low temperature (normally less than 25°C). 188

189 The addition of ions is normally used to produce alginate-based emulsion gels. Alginate, a

190 linear unbranched natural polysaccharide, is derived from brown seaweed extracts

191 (*Phaeophyceae*) (King, 1983). Sodium alginate has the ability to form 'egg-box' shaped gels

when the sodium ions are replaced by divalent cations (mostly calcium in the food industry)

193 (Ching, Bansal, & Bhandari, 2017). Two different methods can be used to prepare alginate-

based emulsion gels. Pintado, Ruiz-Capillas, Jimenez-Colmenero, Carmona, & Herrero

195 (2015) added  $CaSO_4$  into an alginate-based emulsion to produce an alginate-based emulsion

196 gel directly. Sato, Moraes, & Cunha (2014) used a different method to produce emulsion gels,

in which CaEDTA was added to the alginate-based emulsion first, after which the acid was

198 then introduced to liberate calcium ions.

Inulin is an oligosaccharide which includes 2 to 60 fructose molecules connected by  $\beta$ -(2 $\rightarrow$ 1) glycoside bonds (Glibowski & Pikus, 2011). Inulin with a crystal structure can disperse in an aqueous environment and form a suspension in which most of the crystals do not change their structures, except some of smallest crystals dissolving in water. Amorphous inulin can change its structure to crystallite in water (Glibowski & Pikus, 2011). Then, small crystallites can

aggregate to form larger clusters, which ultimately interact to form a gel (Bot, Erle, Vreeker,
& Agterof, 2004). Paradiso, Giarnetti, Summo, Pasqualone, Minervini, & Caponio (2015)
compared three different homogenization technologies (i.e., mechanical, ultrasonic and cold
ultrasonic homogenization) to prepare inulin-based emulsion gels, and found that ultrasonic
homogenization is a suitable method to prepare emulsion gels with better textural properties
compared to the other two homogenization technologies.

210 2.1.3. Self-assembly of low molecular weight compound-based bulk emulsion gels

211 Many LMW organic compounds, such as glycyrrhizic acid and a combination of  $\beta$ -sitosterol 212 and  $\gamma$ -oryzanol, can be used as oil-structuring agents, due to their self-assembly, to replace 213 solid fats and provide required sensory and flavor properties in food products (Pernetti, van 214 Malssen, Flöter, & Bot, 2007; Wan, Sun, Ma, Yang, Guo, & Yin, 2017). These organic 215 compounds, when in a water or oil phase, can form soft solid-like structured gels, which are 216 known as oleogels or organogels (Co & Marangoni, 2012), and they can be also used to 217 produce emulsion gels.

Saponin glycyrrhizic acid (GA) is a monodesmosidic saponin which is comprised of a 218 hydrophobic triterpenoid aglycon moiety (18  $\beta$ -glycyrrhetinic acid) attached to a hydrophilic 219 diglucuronic unit. GA molecules have both gelation and emulsifying properties, owing to 220 their self-assembly ability and amphiphilic structures. GA cannot structure vegetable oil 221 directly because of its low solubility in oil. However, GA molecules can self-assemble into 222 long nanofibrils in water, and nanofibrils not only absorb at the oil-water interface but also 223 224 further assemble and entangle to create a supramolecular hydrogel in water phase. Wan, Sun, Ma, Yang, Guo, & Yin (2017) investigated GA-based O/W emulsion gels and found that, for 225 more polar oils, GA fibrils had a higher affinity to the oil-water interface, leading to the 226 formation of a lot of fine multilayer emulsion droplets with smaller droplet size. Ma, Wan, & 227

228	Yang (2017) used GA to produce GA-based water-in-oil-in-water ( $W_1/O/W_2$ ) emulsion gels;
229	a $W_1$ /O emulsion was prepared first, before being mixed with GA solution at 80°C, and GA-
230	based $W_1/O/W_2$ emulsion gels were formed at room temperature by the self-assembly of GA.
231	The combination of $\beta$ -sitosterol and $\gamma$ -oryzanol can self-assemble in an oil phase to form a
232	helical ribbon, and then these tubules can aggregate and form networks, which are known as
233	oleogels or organogels. Thus, the combination of $\beta$ -sitosterol and $\gamma$ -oryzanol can be used to
234	prepare gelled W/O emulsions. However, the oil phase should be prepared at high
235	temperature (~100°C) to dissolve $\beta$ -sitosterol and $\gamma$ -oryzanol, and W/O emulsions should
236	also be prepared at 90°C to prevent the gelation of oil phase during emulsification. It has been
237	reported that, when a mixture of $\beta$ -sitosterol and $\gamma$ -oryzanol was used to prepare W/O
238	emulsion gels, the presence of water weakened the tubules and reduced the firmness of gelled
239	emulsions, due to the hydration of $\beta$ -sitosterol and the transition of crystals from anhydrous
240	and hemihydrate into monohydrate forms (Bot, den Adel, Regkos, Sawalha, Venema, &
241	Flöter, 2011). On the other hand, it was found that reducing the water activity and using oils
242	with low polarity could promote the formation of tubular microstructures of oryzanol and
243	sitosterol in emulsions (Sawalha, den Adel, Venema, Bot, Floter, & van der Linden, 2012).

244 2.2. Emulsion gel particles

Gel particles or gel beads can be divided into three categories according to their size: macrogel particles (> 1 mm), microgel particles (0.2–1000  $\mu$ m), and nanogel particles (< 0.2  $\mu$ m) (Ching, Bansal, & Bhandari, 2017). In the food area, studies have mainly focused on macrogel and microgel particles, and alginate was the matrix material most frequently used to produce gel particles. Ching, Bansal, & Bhandari (2017) reviewed current technologies for producing alginate hydrogel particles (e.g., simple dripping, jet back up extrusion, spinning disk, atomization, impinging aerosol method, emulsification technique, microfluidics, and

templating method), but studies on producing emulsion gel particles have rarely been
reported. As shown in Table 2, methods used to prepare emulsion gel particles include simple
dripping, electrostatic extrusion, and the impinging aerosol method.

Lević, Pajić Lijaković, Đorđević, Rac, Rakić, et al. (2015) used electrostatic extrusion 255 technique to prepare alginate-based emulsion beads with diameters in the range from 960 to 256 1650 µm, in which a syringe pump and electrostatic immobilization unit (at a voltage of 6.5 257 kV) were used to extrude an alginate-based emulsion through a needle (22 gauge) into a 258 collecting solution (0.015 g/ml of CaCl<sub>2</sub> solution). The reason for using an electrostatic 259 immobilization unit is that electrostatic forces can disrupt the liquid filament at the tip of the 260 needle and create a charged stream of small droplets. However, bigger beads were formed 261 with the diameters in the range from 2100 to 2350 µm without applying voltage, i.e., 262 extrusion by syringe or simple dripping, which is thus a simple method to produce emulsion 263 264 gel particles, but this method usually leads to large particle sizes. Ching, Bansal, & Bhandari (2016) developed a spray aerosol method to prepare alginate-based emulsion microgel 265 particles with the size of 36.2 to 57.8 µm. A fine aerosol mist of alginate-based emulsion and 266 an aerosol mist of 0.5 M calcium chloride solutions are created at the top and bottom of the 267 chamber, respectively, using an air atomizing nozzle. Two mists combine in the chamber, and 268 emulsion gel particles form in the chamber and are collected at the base of chamber. This is 269 an effective and continuous method to produce emulsion gel particles with small size, but this 270 method needs a special spray aerosol system. 271

272 2.3. Fluid emulsion gels

Apart from bulk emulsion gels and emulsion gel particles, fluid emulsion gels are the third
type of emulsion gels. Fluid emulsion gels are different from bulk gels and gel particles with
solid shapes, but they have higher viscoelastic properties than conventional emulsions. Fluid

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emulsion gels mainly include two types according to their preparation methods: gel-likeemulsions and disrupted emulsion gel systems (Table 3).

### 278 2.3.1. Gel-like emulsions

Pickering emulsions are a kind of emulsions which are stabilized by amphiphilic solid 279 particles, and can be divided into three categories: polysaccharide particle-, protein particle-280 281 and mixture-stabilized Pickering emulsions. Pickering emulsions are considered as better delivery systems than conventional emulsions, owing to their enhanced storage stability 282 against oxidation and coalescence and lower susceptibility to lipolysis. Pickering emulsions 283 can turn into gel-like emulsions under appropriate conditions (e.g., proper solid particle type, 284 solid particle concentration, oil phase concentration, pH, and ionic strength). It has been 285 reported that gel-like emulsions could be formed with 6 wt% preheated soy globulins at high 286 glycinin contents (> 75%) with soy oil at a oil volume fraction ( $\phi$ ) of 0.3, and that G' and G'' 287 values of gel-like emulsions increased as the increase of glycinin contents (from 75% to 288 289 100%), while neither unheated soy globulins nor preheated soy globulins with low glycinin contents could form gel-like emulsions (Luo, Liu, & Tang, 2013). This was probably because 290 the formation of a gel-like network was largely attributed to hydrophobic interactions 291 292 between denatured glycine molecules absorbed at the interface of oil droplets. However, Xu, Liu, & Tang (2019) found that, with increasing oil fractions ( $\varphi = 0.1$  to 0.88), a 0.5 wt% soy 293  $\beta$ -conglycinin-stabilized Pickering emulsion could turn into a gel-like emulsion at an oil 294 fraction of 0.7. It was also found that, with increasing wheat gluten level (emulsifier in oil-in-295 glycerol emulsions, 0.25–1.0 wt%), gel-like emulsions could be formed at high wheat gluten 296 contents (>= 0.5 wt%) (Liu, Chen, Guo, Yin, & Yang, 2016). Shao & Tang (2016) found that, 297 with increasing oil fraction (0.2 to 0.6), pea protein-based Pickering emulsions changed from 298 liquid to a gel-like state, while Zou, Guo, Yin, Wang, & Yang (2015) found that zein/tannic 299 acid complex-stabilized Pickering emulsion gels with high oil volume fraction ( $\phi > 0.5$ ) 300

301	could be successfully produced. Therefore, the oil phase and emulsifier contents should be
302	high enough to assure that solid particles absorbed at the surface of neighboring oil droplets
303	can connect and/or react with each other (Wouters & Delcour, 2019).
304	2.3.2. Disrupted gel systems
305	Fluid emulsion gels can also be prepared by breaking down bulk emulsion gels (Leon,
306	Medina, Park, & Aguilera, 2018). Soukoulis, Cambier, Hoffmann, & Bohn (2016)
307	investigated so-called sheared oil-in-gel (o/g) emulsions prepared by stirring an alginate-
308	based emulsion gel system at 1000 rpm for 6 h during the gelation process. Torres, Tena,
309	Murray, & Sarkar (2017) developed a method to produce starch-based gel emulsions by
310	homogenizing the bulk emulsion gels. This is a simple method to produce fluid emulsion gels
311	with small dispersed gel particles (5–50 $\mu$ m in diameter), but the gel matrix-covered structure
312	may be destroyed, leading to separation of the gel matrix and oil droplets during
313	homogenization, which may influence the stability of oil droplets and/or encapsulated
314	nutrients during storage.

## 315 **3. Structure-property relationships of different emulsion gels**

316 3.1. Bulk emulsion gels

Some properties of bulk emulsion gels are emphasized in the food industry, such as
mechanical properties (e.g., rheological, and textural perception), and release properties
(including stability during storage and targeted-release in digestion). Many factors (e.g.,
structures of the gel matrix, structures of emulsion droplets, and interactions between the gel
matrix and droplets) can influence the structures of bulk emulsion gels and then their
mechanical and release properties.

323	Common food emulsions include single emulsions (O/W and W/O emulsions) and multiple
324	emulsions ( $W_1/O/W_2$ and $O_1/W/O_2$ emulsions). After turning emulsions into bulk emulsion
325	gels, their structures usually do not change. Thus, the structures of emulsion gels also include
326	single structures (i.e., O/W and W/O) and multiple structures (i.e., $W_1/O/W_2$ and $O_1/W/O_2$ ).
327	The matrix materials of O/W and $W_1/O/W_2$ emulsion gels are protein-, polysaccharide-, or
328	organic compound-based hydrogels, while the matrix materials of W/O and $O_1/W/O_2$
329	emulsion gels are organic compound-based oleogels (also known as organogels or structured
330	oil). Moreover, properties of O/W and $W_1/O/W_2$ emulsion gels and W/O and $O_1/W/O_2$
331	emulsion gels differ, because the properties of emulsion gels mainly depend on the properties
332	of matrix materials (i.e., protein-, polysaccharide-, or organic compound-based gels),
333	although the properties of emulsion droplets and the interactions between the gel matrix and
334	droplets also influence the properties of emulsion gels. However, O/W emulsion gels have
335	been studied more widely than W/O, $W_1/O/W_2$ and $O_1/W/O_2$ emulsion gels, so the following
336	discussions in this review will focus on O/W emulsion gels unless other structures are
337	emphasized.

## 338 *3.1.1. The structure-mechanical property relationships of bulk emulsion gels*

Mechanical properties of bulk emulsion gels are closely associated with other properties (e.g., storage stability, oral perception, and controlled release) and their applications. The most common mechanical properties of bulk emulsion gels are dynamic modulus (i.e., storage and loss modulus), Young's modulus, fracture strength (i.e., strain and stress), yield strength, and hardness. There are many ways or tools to measure mechanical properties of bulk emulsion gels, such as rheometry, dynamic mechanical analysis (DMA), and textural analysis (Anseth, Bowman, & Brannon-Peppas, 1996).

### 346 *3.1.1.1. Matrix structures*

For protein-based bulk emulsion gels, use of different proteins and methods can lead to
different protein matrix structures and mechanical properties, owing to different gelation
mechanisms and resultant different molecular forces between protein molecules in the gel
matrix. Globular proteins (e.g., SPI, WPI, and MP) and non-globular proteins (e.g., gelatin,
casein, and sodium caseinate) have been widely used as matrix materials in producing bulk
emulsion gels.

The heat-set gelation method has been most used to prepare globular protein-based emulsion 353 gels, but globular protein-based emulsion gels can be also prepared through acidification 354 treatment, addition of ions, enzyme treatment, and MDA modification. For heat-induced 355 emulsion gels, noncovalent cross-links (i.e., electrostatic interactions, hydrophobic 356 interactions, and hydrogen bonds) and intermolecular disulfide bonds are the main forces 357 between globular protein molecules (Wu, Xiong, & Chen, 2011). The main linking forces in 358 359 the glucono-δ-lactone (GDL)-induced emulsion gels are hydrophobic interactions and Van der Waals forces, while salt-bridges are the main linking forces in salt-induced emulsion gels, 360 361 and TGase-induced emulsion gels involve more covalent cross-links (i.e.,  $\varepsilon$ -( $\gamma$ -glutamyl)lysine (G-L) cross-links). Therefore, different preparation methods may lead to different 362 mechanical properties of globular protein-based emulsion gels (Liang, et al., 2020; Wang, 363 Xiong, & Sato, 2017; Ye & Taylor, 2009); for example, it was found that CaSO<sub>4</sub>-induced SPI-364 based emulsion gels were stiffer with higher rigidity than MTGase-induced gels which 365 performed better elasticity (Wang, Luo, Liu, Zeng, Adhikari, He, et al., 2018). 366 Gelatin can form gels under one-step cold treatment or cold treatment after pre-heating as 367 descried in section 2.1.1.2. Cold-set gelatin gels, a kind of elastic polymer gel, are formed 368 with flexible and random-coil protein chains. Therefore, gelatin-based emulsion gels are 369 similar to gels with active-fillers (bound droplets) in which stress concentration phenomena 370 371 play a larger role compared to friction phenomena (Sala, van Vliet, Cohen Stuart, Aken, &

van de Velde, 2009). Other non-globular protein-based emulsion gels are normally prepared
with enzyme treatment and acidification treatment. For example, although the main linking
forces in acid-induced casein gels are also noncovalent cross-links, the firmness of acidinduced sodium caseinate gels was lower than that of acid-induced WPC gels, probably due
to their differences in gelation mechanism (Kiokias & Bot, 2005).

Overall, contributions to the connectivity of a three-dimensional protein network arise from 377 four different kinds of molecular forces: covalent bonds, electrostatic interactions, hydrogen 378 bonding and hydrophobic interactions. The presence of covalent bonds leads to permanent 379 'chemical' cross-links within the network, whereas the other three types of weaker 'physical' 380 forces contribute to a complex set of more temperature-dependent interactions (Chen & 381 Dickinson, 1999b). In addition, process parameters (e.g., temperature, protein content, ionic 382 strength, pH, the presence of other components, ultrasound pretreatment, and high-pressure 383 384 homogenization) also can influence the structures and mechanical properties of protein-based bulk emulsion gels (Bi, et al., 2020; Chen & Dickinson, 2000; Cheng, et al., 2019). 385 Firstly, temperature can influence the degree of denaturation of proteins, and thus affect the 386 stability of protein-stabilized emulsions and mechanical properties of emulsion gels. 387 388 Generally, a high degree of denaturation of proteins results in low stability of proteinstabilized emulsions but better mechanical properties of emulsion gels (Kiokias & Bot, 2005; 389 390 Ye & Taylor, 2009). Chen & Dickinson (2000) also found that gelation temperature could influence the rate of gelation and the dynamic modulus of acid-induced sodium caseinate-391

based emulsion gels by changing the strength of physical bonding rather than the networkstructures.

Secondly, the influence of protein content on the mechanical properties of emulsion gelsdepends on the state of emulsion droplets. The mechanical properties of droplet-filled gels

396 and inactive droplet-aggregated gels mainly depend on the gel strength of gel matrix structures, while interactions between the gel matrix (i.e., protein and polysaccharide) and 397 lipid droplets contribute more to the active droplet-aggregated gels (Pintado, Ruiz-Capillas, 398 399 Jimenez-Colmenero, Carmona, & Herrero, 2015). Therefore, increasing protein content can increase the gel strength of both kinds of emulsion gels but for different reasons (i.e., 400 increased gel strength of protein matrix for droplet-filled gels and inactive droplet-aggregated 401 gels, but strengthened interactions between the gel matrix and droplets and increased gel 402 strength of protein matrix for active droplet-aggregated gels). For example, it has been 403 404 reported that increasing the concentration of sodium caseinate can decrease the gelation time (Tgel) of sodium caseinate/sunflower oil emulsion-based gels (Montes de Oca-Ávalos, Huck-405 Iriart, Candal, & Herrera, 2016). 406

Thirdly, ionic strength and pH can influence intermolecular repulsion and gel structures in 407 408 emulsion gels. For example, at low ionic strength (< 50 mM NaCl) and pH values (below 4 or above 6) far away from pI of whey proteins, a fine-stranded network consisting of whey 409 protein strains with a length of ~50 nm and a diameter of ~10 nm is formed; at high ionic 410 411 strength (> 150 mM NaCl) and pH values near the pI, the strains with weak intermolecular repulsion can accumulate and form a particulate network structure (Chen, Dickinson, 412 Langton, & Hermansson, 2000; Guo, Bellissimo, & Rousseau, 2017; Langton & 413 Hermansson, 1992). However, both fine-stranded and particulate gels exhibit high gel 414 strength (Guo, Bellissimo, & Rousseau, 2017; Tang, Chen, & Foegeding, 2011). It was found 415 that fine-stranded whey protein gels prepared at low ionic strength (10 or 25 mM NaCl) were 416 rubbery and soft, but that particulate whey protein gels prepared at high ionic strength (100 or 417 200 mM NaCl) were hard and brittle (Guo, Ye, Lad, Dalgleish, & Singh, 2013). 418

419 Fourthly, the presence of other components (e.g., sucrose, glucose, hydroxytyrosol,

420 rosmarinic acid, genipin, sodium pyrophosphate, insoluble dietary fiber, and EGCG) also can

421 influence the structures and mechanical properties of emulsion gels (Chen, Ren, Zhang, Qu, Hu, & Yan, 2019; Feng, Chen, Lei, Wang, Xu, Zhou, et al., 2017; Freire, Bou, Cofrades, & 422 Jimenez-Colmenero, 2017; Montes de Oca-Ávalos, Huck-Iriart, Candal, & Herrera, 2016; 423 Wang, et al., 2018; Wang, Jiang, & Xiong, 2019; Zhuang, et al., 2019). Generally, if 424 components can strengthen protein-protein interactions and/or reduce droplet size, they can 425 increase gel strength of emulsion gels. However, if these components can interact with 426 protein molecules and disturb the interactions between protein molecules, they can weaken 427 the gel strength of emulsion gels, and these effects are normally dose-dependent. Overall, 428 429 preparation methods can affect linking forces between protein molecules, and protein type and processing parameters can influence the network structures of the gel matrix, both of 430 which can affect the mechanical properties of emulsion gels. 431

In terms of polysaccharide-based emulsion gels, polysaccharide type, preparation methods, 432 and processing parameters can influence the structures of the polysaccharide-based gel 433 matrix. Cold-set gellan gun-, agar-, and k-carrageenan-based emulsion gels are a kind of 434 435 polymer gels with strand-based structures (Kim, Gohtani, Matsuno, & Yamano, 1999; Wang, 436 Neves, Kobayashi, Uemura, & Nakajima, 2013). They normally show a predominantly elastic behavior, which resemble gelatin-based emulsion gels but differ from WPI-based emulsion 437 gels with particulate structures (Sala, van Vliet, Cohen Stuart, Aken, & van de Velde, 2009). 438 The network structures of alginate gels are in the shape of 'egg-box', in which sodium ion is 439 replaced by a divalent cation, and each cation can bind with four G residues to form a three-440 dimensional network structure (Ching, Bansal, & Bhandari, 2017), which can be affected by 441 freeze-thawing treatment (Li, Gong, Hou, Yang, & Guo, 2020). Inulin gels are formed by 442 connection of microcrystals, and their rheological properties resemble that of fat crystal-443 based networks in oil (Nourbehesht, Shekarchizadeh, & Soltanizadeh, 2018). However, there 444

445 are no studies on comparing mechanical properties of emulsion gels formed by different446 kinds of polysaccharides.

In addition, the influence of polysaccharide content on the mechanical properties of emulsion 447 gels depends on emulsifier type and gel structures. Most natural polysaccharides, except gum 448 Arabic and some kinds of pectin, have weak emulsifying abilities compared to proteins and 449 synthetic emulsifiers (Charoen, Jangchud, Jangchud, Harnsilawat, Naivikul, & McClements, 450 2011). Hence, the interactions between the gel matrix and emulsion droplets in 451 polysaccharide-based emulsion gels with/without synthetic emulsifiers are normally weak, 452 and increasing polysaccharide content can increase their gel strength, mainly due to the 453 decreased void spaces and increased gel strength of the gel matrix (Kim, Gohtani, Matsuno, 454 & Yamano, 1999). However, when proteins are used as emulsifiers, increasing polysaccharide 455 content can increase the gel strength of emulsion, mainly due to increased interactions 456 457 between polysaccharide molecules and droplets and/or the gel strength of polysaccharide gels. Although studies on the effects of ionic strength and pH on the mechanical properties of 458 459 polysaccharide-based emulsion gels have rarely been reported, Ozturk, Argin, Ozilgen, & 460 McClements (2015) found that ionic strength and pH did not have significant influences on the stability of a gum Arabic-stabilized emulsion, which was different from a WPI-stabilized 461 emulsion because of their different emulsification mechanisms (i.e., electrostatic repulsion for 462 WPI and steric repulsion for gum Arabic). Therefore, it is proposed that the influence of ionic 463 strength and pH on the structure and mechanical properties of polysaccharide-based emulsion 464 gels differs from that on protein-based emulsion gels. 465

For LMW organic compound-based emulsion gels, saponin glycyrrhizic acid (GA) and the

467 combination of  $\beta$ -sitosterol and  $\gamma$ -oryzanol have been investigated to prepare emulsion gels

468 by self-assembly. GA,  $\beta$ -sitosterol and  $\gamma$ -oryzanol have physical properties, so they have been

used to prepare different types of emulsion gels. GA can dissolve in water, and GA molecules

470	can self-assemble to form long nanofibrils and gels in water phase, and so can be used to
471	prepare emulsion gels with O/W or $W_1/O/W_2$ structures. The combination of $\beta$ -sitosterol and
472	$\gamma$ -oryzanol can self-assemble in an oil phase to form a helical ribbon, then these tubules can
473	aggregate and form a network, and so can be used to prepare gelled W/O emulsions.
474	Processing parameters (e.g., organic compound content and solvent type) also can influence
475	the structure and mechanical properties of organic compound-based emulsion gels. Ma, Wan,
476	& Yang (2017) found that an emulsion stabilized by GA at a low concentration (0.5 wt%)
477	could not form a gel, but self-standing emulsion gels could be formed and the viscoelastic
478	modulus also significantly increased with increasing GA concentration (1-4 wt%). It was also
479	found that no tubules were formed but only sitosterol and oryzanol crystals were present in
480	emulsion gels at 16% total sterol concentration, while there were tubules next to the crystals
481	at 32% total sterol concentration (Bot, den Adel, Regkos, Sawalha, Venema, & Flöter, 2011).
482	In addition, the polarity of solvents (i.e., oil in W/O emulsions) can influence the water
483	activity of W/O emulsions and structures of the oil phase. It has been reported that more
484	water molecules bind to the $\beta$ -sitosterol molecules and formed monohydrate crystals in
485	higher polarity oils (e.g., eugenol and castor oil), which hindered the formation of tubules and
486	resulted in weaker emulsion gels compared to less polar oils (e.g., decane and limonene)
487	(Sawalha, den Adel, Venema, Bot, Floter, & van der Linden, 2012). However, studies on
488	comparing structures and mechanical properties of emulsion gels prepared with different
489	kinds of organic compounds have rarely been reported. Over all, many factors (e.g., gel
490	matrix type, preparation method, and process parameters) can affect the gel structures of bulk
491	emulsion gels and thus their mechanical properties.

*3.1.1.2. Structures of emulsion droplets* 

493 The structure of emulsion droplets can influence the mechanical properties of bulk emulsion gels as well. Structures of emulsion droplets are normally influenced by oil phase (e.g., oil 494 type, oil content, and droplet size), and emulsifier type (e.g., low molecular weight 495 496 emulsifiers or proteins). In the food industry, emulsifiers mainly include two categories: low molecular synthetics (e.g., Span 80, Tween 80, and monoglycerides) and natural molecules 497 (e.g., proteins, egg lecithin, and soy lecithin) (Chen, Mao, Hou, Yuan, & Gao, 2020). 498 Emulsifiers can not only decrease the interfacial tension and thereby increase the stability of 499 emulsions but also affect the interactions between droplets and the gel matrix leading to 500 501 active or inactive fillers (Van Vliet, 1988). Therefore, the effect of emulsion droplets on the mechanical properties of emulsion gels depends on not only emulsion droplets (i.e., oil type, 502 oil content, and droplet size) but also the interactions between droplets and the gel matrix 503 504 (Farjami & Madadlou, 2019).

The effect of active fillers on the rheological properties of emulsion gels mainly depends on
the stiffness of the oil droplets and the droplet volume fraction (Sala, van Vliet, Cohen Stuart,
Aken, & van de Velde, 2009). The Kerner model can explain the effect of active fillers on the
mechanical properties of emulsion droplet-filled gels (Kerner, 1956):

509 
$$\frac{G'_{gel}}{G'_{matrix}} = \frac{15(1-v_m)(M-1)\phi_f}{(8-10v_m)M+7-5v_m-(8-10v_m)(M-1)\phi_f} + 1$$
(1)

where  $M = \frac{G'_{filler}}{G'_{matrix}}$ , and  $G'_{gel}$ ,  $G'_{filler}$ , and  $G'_{matrix}$  are the shear modulus of the overall gel, the filler droplets and the gel matrix, respectively,  $\phi_f$  is the actual droplet volume fraction, and  $v_m$ is the Poisson's ratio of the gel matrix. In addition, the Kerner model modified by Lewis and Nielsen can be used to explain the effect of active fillers on the mechanical properties of emulsion droplet-aggregated gels (Lewis & Nielsen, 1970):

515 
$$\frac{G'_{gel}}{G'_{matrix}} = \frac{15(1-v_m)(M-1)\psi\phi_f}{(8-10v_m)M+7-5v_m-(8-10v_m)(M-1)\psi\phi_f} + 1$$
(2)

where  $\psi \phi_f$  is the effective volume fraction of fillers, which takes into account the crowding effect of fillers and can be expressed as follows (Lewis & Nielsen, 1970):

518 
$$\psi \phi_f = \left[ 1 + \left( \frac{1 - \phi_{max}}{\phi_{max}^2} \right) \phi_f \right] \phi_f \tag{3}$$

where  $\phi_{\text{max}}$  is the maximum volume fraction of the fillers. According to Eq. (2), increasing the 519 shear modulus and the effective volume fraction  $(\psi \phi_f)$  or actual volume fraction  $(\phi_f)$  of fillers 520 can increase the mechanical properties of emulsion gels, which has been supported by many 521 studies (Gwartney, Larick, & Foegeding, 2004; Li, Kong, Zhang, & Hua, 2012; Oliver, 522 Scholten, & van Aken, 2015; Oliver, Wieck, & Scholten, 2016; Tang, Yang, Liu, & Chen, 523 2013). However, the Kerner model and the modified Kerner model are used under the 524 assumption that M or G'<sub>matrix</sub> do not change with changes in other factors (e.g.,  $\phi_f$  and G'<sub>filler</sub>) 525 (Chen & Dickinson, 1998a; Oliver, Berndsen, van Aken, & Scholten, 2015), especially at oil 526 volume fractions ( $\phi$ ) below 0.2 and protein (i.e., gel matrix) contents above 6 wt% (Guo, 527 Bellissimo, & Rousseau, 2017). However, the shear modulus of filler droplets ( $G'_{filler} = 4\gamma / d$ , 528 where  $\gamma$  is surface tension and d is the average diameter of the oil droplets ) is influenced by 529 oil type, oil content, droplet size, emulsifier type, emulsifier content, and process parameters 530 (Farjami & Madadlou, 2019; Sala, van Vliet, Cohen Stuart, Aken, & van de Velde, 2009; Van 531 Vliet, 1988). The shear modulus of the gel matrix  $(G'_{matrix})$  is influenced by droplet size, oil 532 533 content, gel matrix type, preparation method, and process parameters (Sato, Moraes, & Cunha, 2014). Therefore, when taking those factors (e.g., droplet size, process parameters, 534 and high oil content), which can affect the mechanical properties of both filler droplets and 535 the gel matrix, into account, the Kerner model and the modified Kerner model cannot be 536 applied. For instance, it has been reported that increasing the size of olive oil droplets in a 537 gelatin-based emulsion gel led to a weaker gel strength, probably due to the increase in 538 interfacial area, a higher amount of gelatin adsorbed to the interface, and a lower quantity of 539

540 protein available in the continuous phase (Sato, Moraes, & Cunha, 2014); however, it was found that increasing the size distribution of dispersed vegetable fat in a WPI-based emulsion 541 gel led to an increase in firmness, probably because of a larger number of contacts between 542 droplets (Kiokias & Bot, 2006). Oliver, Wieck, & Scholten (2016) found that increasing the 543 casein content (from 4% to 9%) could decrease the relative Young's modulus of emulsion 544 gels at high oil volume fractions ( $\phi_f > 0.15$ ), probably owing to the higher inhomogeneity of 545 casein-based gel matrix and increased effective volume fraction of droplets at lower casein 546 concentration; this indicated that the effective volume fraction ( $\psi \phi_f$ ) plays a more important 547 role than G'<sub>matrix</sub> in affecting the mechanical properties of emulsion gels with high matrix 548 inhomogeneity and at high oil volume fractions. 549

The effect of inactive fillers on the rheological properties of emulsion gels depends on the 550 properties and concentrations of LMW emulsifiers, droplet size, and oil content, although 551 552 there have been few studies on modelling the effect of inactive fillers on the rheological properties of emulsion gels. Chen & Dickinson (1999a) investigated the effect of LMW 553 554 emulsifiers on the viscoelastic properties of heat-set whey protein-based emulsion gels, and found that the elastic modulus of heat-set whey protein-based emulsion gels decreased after 555 adding a low level of diglycerol monolaurate (DGML, the surfactant/protein molar ratio (R) = 556 4) and diglycerol monooleate (DGMO, R = 4-32), while high levels of emulsifiers (R = 32) 557 for DGML, and R = 64 for DGMO) could recover the storage and loss modulus of emulsion 558 gels, probably due to depletion flocculation of the emulsion prior to heat-treatment. However, 559 it has been reported that Tween 20 (R = 0.25-8) always decreased the mechanical properties 560 of emulsion gels, and a high addition level (R = 8) could even break down the network 561 structure of proteins and lead to a liquid-like emulsion (Chen & Dickinson, 1998b). It has 562 been found that increasing oil content decreased fracture stress and stress intensity factor of 563 agar gels and  $\kappa$ -carra-geenan-locust bean gum gels (Koç, Drake, Vinyard, Essick, van de 564

Velde, & Foegeding, 2019). It has also been found that increasing solid lipid content could
increase the gel strength of emulsion gels at an emulsifier content of 4 g/100 g, but decreased
the gel strength at an emulsifier content of 2 g/100 g (Geremias-Andrade, Souki, Moraes, &
Pinho, 2017).

569 3.1.2. The structure-release property relationships of bulk emulsion gels

570 Bulk emulsion gels, especially O/W emulsion gels, are often used for the delivery and release of oil-soluble bioactive compounds and nutrients, such as  $\alpha$ -tocopherol (Liang, Leung Sok 571 Line, Remondetto, & Subirade, 2010) and β-carotene (Soukoulis, Tsevdou, Andre, Cambier, 572 Yonekura, Taoukis, et al., 2017). Compared to emulsions, emulsion gels can provide better 573 protection for encapsulated compounds and show slower release behavior (Cofrades, et al., 574 2017). Many studies have focused on the matrix erosion, lipid digestion and controlled 575 release of encapsulated compounds during digestion of emulsion gels. The digestion 576 behaviors of protein- and polysaccharide-based emulsion gels differ in the gastrointestinal 577 578 tract because of different digestion processes of proteins and polysaccharides. For proteinbased emulsion gels, Liang, Leung Sok Line, Remondetto, & Subirade (2010) found that gel 579 loss (i.e., matrix erosion owing to protein degradation) and release of  $\alpha$ -tocopherol occurred 580 in both simulated gastric fluid (SGF) and simulated intestinal fluid (SIF), respectively, which 581 indicated that release of  $\alpha$ -tocopherol was controlled mainly by matrix erosion because of 582 protein degradation. However, under simulated gastrointestinal (GI) conditions (0.5 h SGF 583 followed by 6 h SIF), gel loss and release of  $\alpha$ -tocopherol only occurred in the SGF step, 584 probably due to the formation of a viscous layer at the surface of gels. Moreover, gel rigidity 585 of protein-based emulsions is an important factor affecting the lipid digestion in GI digestion. 586 It has been reported that gastric digesta of a soft gel, prepared with 10 or 20 mM NaCl, 587 mainly consisted of individual oil droplets and small gel particles (~10 mm), while gastric 588 digesta of a hard gel, prepared with 100 or 200 mM NaCl, mainly consisted of small gel 589

590 particles (~10 mm) after 240 min gastric digestion, and the remaining network structure of gel particles hindered further breakdown during intestinal digestion (Guo, Ye, Lad, Dalgleish, 591 & Singh, 2016). It was also found that digestion of emulsion gels in the intestinal step was 592 593 delayed by denser, more spatially heterogeneous protein matrixes (Guo, Bellissimo, & Rousseau, 2017). In terms of polysaccharide-based emulsion gels, although there are fewer 594 reports about their digestion, it was found that oil droplets could be released from agar-based 595 emulsion gels during GI digestion in both SGF and SIF steps (2.0 h SGF followed by 4-14 h 596 SIF), while emulsifier type (glycerol monolaurate with different degrees of polymerization) 597 598 affected the size distribution of released oil droplets (Wang, Neves, Kobayashi, Uemura, & Nakajima, 2013). 599

Bulk emulsion gels are also used for the delivery and release of volatile flavor compounds, 600 such as ethyl butyrate, ethyl hexanoate, ethyl octanoate, propanol, diacetyl, pentanone, 601 602 hexanal, and heptanone (Hou, Guo, Wang, & Yang, 2016; Mao, Roos, & Miao, 2014). The release of volatile compounds in the oral cavity is normally measured by a simulated nose 603 604 breath device (Hou, Guo, Wang, & Yang, 2016) or gas chromatography (GC) headspace 605 analysis (Mao, Roos, & Miao, 2014). The release rate of volatile compounds depends on the gel matrix structure, oil content, the nature of volatile compounds, and the interactions 606 between flavor compounds and food ingredients (particularly oils in O/W emulsion gels) 607 (Boland, Delahunty, & van Ruth, 2006; Guichard, 2002). It has been reported that the release 608 rate of ethyl butyrate was significantly lower in a SPI/sugar beet pectin (SBP) complex-based 609 emulsion gel with a compact network than SPI- or SBP-based emulsion gels, but the release 610 rate of aroma compounds with higher hydrophobicity was not significantly influenced by the 611 structures of emulsion gels, probably because of their high affinity for the lipid phase rather 612 than interacting with proteins and/or polysaccharides (Hou, Guo, Wang, & Yang, 2016). Mao, 613 Roos, & Miao (2014) also found that emulsion gels with higher storage modulus at a low oil 614

- 615 content (20%) had lower release rates and partition coefficients of the volatiles, and that
- 616 increasing oil contents (from 5% to 20%) significantly decreased the release rate of
- 617 heptanone, probably owing to its highly lipophilic characteristics.
- 618 3.2. The structure-property relationships of emulsion gel particles
- 619 Although emulsion gel particles and bulk emulsion gels have similar structures (i.e., active
- 620 fillers, inactive fillers, emulsion droplet-filled gels, and emulsion droplet-aggregated gels)

and structure-property relationships, their physical characteristics and length scales differ

622 (Ching, Bansal, & Bhandari, 2016).

Firstly, the rheological behavior of gel particles differs to that of bulk gels, because the microgel particle system is a suspension (usually gel particles in water). The rheological properties of microgel particle suspensions are influenced by three parameters: volume fraction ( $\phi$ ), particle modulus (modulus of particles that make up the suspension) and interaction potential (Ching, Bansal, & Bhandari, 2016). The volume fraction ( $\phi$ ) can be determined using the equation below (Ching, Bansal, & Bhandari, 2016):

$$629 \qquad \phi = \frac{\frac{m}{\rho}}{\frac{m}{\rho} + \nu} \tag{4}$$

where  $\phi$  = final microgel suspension volume fraction, m = mass of microgel concentrate,  $\rho$  = 630 density of microgel concentrate measured with a 50 mL calibrated pycnometer, and v =631 volume of water added to microgel concentrate. Eq. (4) was modified by the equations 632 developed by Suzawa & Kaneda (2010), who calculated the volume fraction by the weight 633 and density of emulsions but did not consider the weight loss (normally water loss) of gel 634 particles during gelation. At low volume fraction, the flow behaviour is determined by the 635 continuous phase; at higher volume fraction, softer microgels will exhibit a lower storage 636 modulus compared to hard microgels (Adams, Frith, & Stokes, 2004). Ching, Bansal, & 637

638 Bhandari (2016) found that, at the same volume fraction, suspensions with more deformable alginate-based micorgels exhibited a lower bulk modulus. However, it is technically difficult 639 to investigate the rheological properties of macrogel particles, although their mechanical 640 properties could be investigated by a texture analyser. It has been reported that, with 641 increasing oil contents in alginate-based macrogels, the elastic modulus of particles 642 decreased, which indicates that oil droplets in alginate-based emulsion gel particles without 643 emulsifiers were inactive fillers (Ching, Bansal, & Bhandari, 2016). 644 Secondly, syneresis and swelling properties are important properties of gel particles (Ching, 645 Bansal, & Bhandari, 2017). It was found that alginate-based emulsion gel particles shrank 646

less if they had higher oil content, and that the swelling was more pronounced for smaller
particles, probably owing to the larger contact surface, but was less pronounced at increased
oil contents, probably because of droplets acting as physical barriers for water transport
(Lević, et al., 2015).

651 Thirdly, encapsulation efficiency (EE), loading capacity (LC) and encapsulation yield, important parameters in encapsulation processes of emulsion gel particles, are affected by 652 properties and contents of matrix material, emulsifier, and oil. It has been reported that 653 increasing alginate contents in the water phase could increase the oil EE in lupin protein 654 isolate (LPI)-stabilized emulsion gel particles, probably due to the formation of a stronger gel 655 matrix and better crosslinking on the external surfaces of particles (Piornos, Burgos-Díaz, 656 Morales, Rubilar, & Acevedo, 2017). However, when the protein content was higher than the 657 saturation concentration, or the oil content was very low, in which case excessive free protein 658 molecules existed in the water phase, the aggregation of non-adsorbed protein molecules 659 could lead to lower emulsion stability and lower EE (Guzey & McClements, 2006). In 660 addition, Ruffin, Schmit, Lafitte, Dollat, & Chambin (2014) found that, compared to native 661 662 WPI, using pre-heated WPI at 80°C for 30 min as emulsifier in pectin-based emulsion gel

particles slightly improved the yield and stability of encapsulated vitamin A, because of the
increased viscosity of denatured WPI dispersions and the decreased particle size of
emulsions.

666 3.3. The structure-property relationships of fluid emulsion gels

667 3.3.1. Gel-like emulsions

The oil content, particle content, and surface charge of particles can affect the rheological 668 properties of gel-like Pickering emulsions and release behavior of encapsulated compounds 669 from such emulsions (Shao & Tang, 2016; Xu, Liu, & Tang, 2019). For the effect of oil 670 content, Dai, Sun, Wei, Mao, & Gao (2018) found that zein/gum arabic complex-stabilized 671 Pickering emulsion gels solidified at high oil volume fractions in emulsions ( $\phi \ge 0.5$ ), and 672 increasing oil volume fractions ( $\varphi = 0.5-0.7$ ) increased the G' and G'' of gel-like emulsions, 673 probably due to more interactions between emulsion droplets (Xiao, Wang, Gonzalez, & 674 Huang, 2016). It was also reported that a gel-like emulsion at  $\varphi = 0.6$  exhibited much lower 675 release rate of  $\beta$ -carotene but higher stability during digestion than a Pickering emulsion at  $\varphi$ 676 = 0.3 (Shao & Tang, 2016). In terms of the effect of particle content, Xu, Liu, & Tang (2019) 677 found that increasing soy  $\beta$ -conglycinin contents from 0.2 to 1.0 wt% led to a progressive 678 decrease in droplet size, but a progressive increase in stiffness of the gel-like emulsions at  $\varphi =$ 679 0.8. Liu, Gao, McClements, Zhou, Wu, & Zou (2019) also found that increasing pre-heated 680 WPI contents from 2.5 to 10 wt% led to a progressive increase in gel strength, hardness, 681 WHC, and stability of the gel-like emulsions at 75 vol% oil; they also found that increasing 682 protein contents could increase the bioaccessibility of β-carotene because of the reduced 683 aggregation of the oil droplets and retarded degradation of  $\beta$ -carotene during digestion, owing 684 to a dense WPI-based gel structure around droplets. In addition, the surface charge of 685 (nano)particles can affect their emulsification and interfacial behavior (Larson-Smith, 686

687	Jackson, & Pozzo, 2012). It has been reported that electrostatic screening by adding NaCl
688	could improve the performance of soy glycine nanoparticles in forming gel-like emulsions
689	and increase stiffness of the resultant gel-like emulsions, due to enhanced diffusion and
690	adsorption of solid particles at the interface (Liu & Tang, 2016).
691	3.3.2. Disrupted gel systems
692	Although there are few studies on the structure-property relationships of disrupted gel
693	systems, Torres, Tena, Murray, & Sarkar (2017) found that increasing starch contents (from
694	15 to 20 wt%) and oil fractions (from 0 to 20 wt%) could improve the elastic modulus of
695	starch-based disrupted gels stabilized by octenyl succinin anhydride (OSA) modified starch,
696	which fitted the Kerner model. It has been reported that, compared to alginate-based
697	emulsions and bulk emulsion gels, sheared oil-in-gel (o/g) emulsions exhibited higher
698	bioaccessibility of encapsulated $\beta$ -carotene after <i>in vitro</i> digestion, due to the lower unbound
699	calcium content and higher colloidal stability throughout gastrointestinal passage, whereas
700	encapsulated $\beta$ -carotene in the bulk emulsion gels exhibited highest chemical stability

701 (Soukoulis, Cambier, Hoffmann, & Bohn, 2016).

## 702 **4.** Applications of emulsion gels in the food industry

4.1. Use of emulsion gels as fat replacers in meat products

Emulsion gels formed by myofibrillar proteins (MPs), water and lipid not only contribute to
the sensory properties (appearance and flavor) but also relate to the textural properties (waterand oil-holding, and cooking losses) of meat products (Wang, et al., 2018; Zhao, Zou, Shao,
Chen, Han, & Xu, 2017). Additives, such as extracts from herbs and spices, polyphenols, and
NaCl, can influence structures of emulsion gels and the properties of meat products (Wang, et al., 2018; Zhao, Zou, Shao, Chen, Han, & Xu, 2017). Wang et al. (2018) found that a low

710	level of rosmarinic acid (RA) (12 $\mu M/g$ protein) could protect thiol and $\epsilon\text{-NH}_3$ groups in MP-
711	based emulsion gels from oxidation, and thus improve the structure and water- and oil-
712	holding abilities of emulsion gels; however, a high level of RA (300 $\mu M/g$ protein) could
713	induce interactions between RA and MPs, which led to aggregation of MPs and a poor
714	emulsion gel network, while a high level of NaCl (0.6 M) could promote these interactions.
715	However, while health concerns around some meat products containing high fat content (over
716	27%) have increased in recent years, reducing fat content usually negatively influences
717	consumer acceptance and textural properties of final products (Oliver, Scholten, & van Aken,
718	2015). In order to avoid undesirable textural changes and improve the nutritional value of
719	meat products (e.g., sausages and patties), promising methods have been studied, such as
720	replacing fat with unsaturated oil (Oliver, Scholten, & van Aken, 2015) or structured oil (e.g.,
721	olive, linseed, fish, perilla, and sunflower seed oil encapsulated in emulsion gels formed with
722	SPI, WPI, sodium caseinate, carrageenan, gelatin, alginate, chia flour, oat bran, or inulin)
723	(Alejandre, Poyato, Ansorena, & Astiasaran, 2016; de Souza Paglarini, de Figueiredo
724	Furtado, Biachi, Vidal, Martini, Forte, et al., 2018; de Souza Paglarinia, Martinib, & Pollonio,
725	2019; Freire, Cofrades, Perez-Jimenez, Gomez-Estaca, Jimenez-Colmenero, & Bou, 2018;
726	Freire, Cofrades, Serrano-Casas, Pintado, Jimenez, & Jimenez-Colmenero, 2017; Glisic, et
727	al., 2019; Pintado, Herrero, Jimenez-Colmenero, Pasqualin Cavalheiro, & Ruiz-Capillas,
728	2018; Poyato, Astiasarán, Barriuso, & Ansorena, 2015; Serdaroglu, Nacak, & Karabiyikoglu,
729	2017). However, these methods may lead to undesirable sensory quality changes (e.g., color
730	parameters and sensory acceptability) (Serdaroğlu & Öztürk, 2017). Oliver, Scholten, & van
731	Aken (2015) found that physical properties of fat or oil and structural properties of the gel
732	matrix could influence the rheological properties of fat-filled emulsion gels or oil-filled
733	emulsion gels. Hence, the properties of fat in meat products should be considered, and the
734	gelling agent and oil should be chosen carefully when emulsion gels are used as a fat replacer

735 (Freire, Cofrades, Serrano-Casas, Pintado, Jimenez, & Jimenez-Colmenero, 2017). It has 736 been reported that combining emulsion gels and animal fat could be a good method to produce healthier meat products with acceptable sensory properties (de Souza Paglarini, et 737 738 al., 2019). In addition, emulsion gels help to control sodium availability and perception by changing sodium mobility and binding behavior, and can thus allow reduction of the salt 739 content in meat products (Okada & Lee, 2017). However, most studies have focused on bulk 740 emulsion gels and their uses in solid foods, and so more studies on emulsion gel particles and 741 their uses in liquid foods are needed. 742

4.2. Emulsion gels used as delivery systems to encapsulate and release food nutriments

Absorption of encapsulated lipophilic food nutrients (e.g., β-carotene, curcumin, n-3 fatty 744 745 acid, vitamin A, and  $\alpha$ -tocopherol) in emulsion gels include several steps: release from the gel matrix as the result of mechanical, chemical and enzymatic processes throughout the oral 746 processing and gastrointestinal passage, incorporation in the co-digested lipid droplets, 747 748 interaction with endogenous lipid surface active compounds (mainly bile salts and phospholipids) promoting the formation of mixed micelles, and eventual transportation of the 749 mixed micelles to the small intestinal epithelium (Soukoulis, Cambier, Hoffmann, & Bohn, 750 751 2016; Yonekura & Nagao, 2007). Polysaccharides (e.g., alginate, κ-carrageenan, and starch) and proteins (e.g., gelatin and WPI) are normally used as gelation materials in producing 752 emulsion gels encapsulating lipophilic food nutrients, but their digestion behaviors differ. 753 Protein-based emulsion gels are mainly disrupted in gastric digestion as the result of 754 enzymatic hydrolysis by pepsin, and the remaining protein-based network structures can 755 756 hinder further breakdown during intestinal digestion (Guo, Ye, Lad, Dalgleish, & Singh, 2016; Liang, Leung Sok Line, Remondetto, & Subirade, 2010). On the other hand, 757 polysaccharide-based (especially alginate-based) emulsion gels are less sensitive to gastric 758 759 fluid than protein-based emulsion gels, and may protect the encapsulated nutriments from

760 harsh gastric environment, and the remaining gel structures can be further disrupted during intestinal digestion (Wang, Neves, Kobayashi, Uemura, & Nakajima, 2013; Xu, et al., 2019). 761 However, emulsion gels normally give low effective bioavailability of encapsulated lipophilic 762 763 compounds, due to insufficient digestion of the gel matrix and resulting unreleased and undigested lipid phase (Liang, Leung Sok Line, Remondetto, & Subirade, 2010; Zhang, et al., 764 2016). Therefore, it is important to choose appropriate materials for different nutrients, which 765 can protect encapsulated nutrients and control their release, and also do not inhibit release in 766 the targeted gastrointestinal tract (Zhang, et al., 2016). Although emulsion gels may not 767 improve the final bioaccessibility of encapsulated food nutrients, they can improve emulsion 768 structures and stability of nutrients during storage, and exhibit slow release effects in the 769 gastrointestinal passage compared to emulsions (Brito-Oliveira, Bispo, Moraes, Campanella, 770 & Pinho, 2017; Ma, Wan, & Yang, 2017; Soukoulis, Cambier, Hoffmann, & Bohn, 2016; 771 Zhang, et al., 2016). 772

### 773 5. Conclusions

Various preparation methods of emulsion gels are available for different matrix materials 774 (e.g., heat treatment, enzyme treatment, acidification treatment, and addition of ions for 775 776 protein-based emulsion gels, cold-set and addition of ions for polysaccharide-based emulsion gels, and self-assembly for LMW compound-based emulsion gels), purposes (e.g., cold 777 778 treatment for protecting encapsulated nutrients and better mechanical properties), and emulsion gel types (e.g., internal gelation for bulk emulsion gels, external gelation for 779 780 emulsion gel particles, self-assembly for gel-like Pickering emulsions, and mechanical stir for 781 disrupted emulsion gels). Due to differences in the morphological properties among different emulsion gels, different physical properties are emphasized, such as the importance of 782 mechanical and release properties for bulk emulsion gels, syneresis and swelling properties 783 784 for emulsion gel particles, rheological properties for microgel particle suspensions, and flow

785 behaviour and release property for fluid emulsion gels. In terms of bulk emulsion gels, many factors (e.g., structures of gel matrix and emulsion droplets and interactions between them) 786 can influence their structures and thus mechanical and release properties. Structures of the gel 787 matrix in bulk emulsion gels are affected by matrix material, preparation method, and process 788 parameters, while structures of emulsion droplets are affected by oil type, oil content, droplet 789 size, and emulsifier type. In terms of emulsion gel particles, oil content and particle size can 790 influence their syneresis and swelling properties. The rheological properties of microgel 791 particle suspensions are influenced by volume fraction, particle modulus, and interaction 792 793 potential. In terms of gel-like Pickering emulsions, their rheological and release properties also are influenced by many factors (e.g., oil content, particle content, and surface charge of 794 particles). Finally, two main applications of emulsion gels in the food industry are as fat 795 replacers in meat products and delivery systems for food nutrients. 796

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## 800 **References**

- Adams, S., Frith, W. J., & Stokes, J. R. (2004). Influence of particle modulus on the rheological
   properties of agar microgel suspensions. *Journal of Rheology*, 48, 1195–1213.
- Alavi, F., Emam-Djomeh, Z., Salami, M., & Mohammadian, M. (2020). Effect of microbial
   transglutaminase on the mechanical properties and microstructure of acid-induced gels and
   emulsion gels produced from thermal denatured egg white proteins. *International Journal of Biological Macromolecules*, 153, 523–532.
- Alejandre, M., Poyato, C., Ansorena, D., & Astiasaran, I. (2016). Linseed oil gelled emulsion: A
   successful fat replacer in dry fermented sausages. *Meat Science*, *121*, 107–113.
- Anseth, K. S., Bowman, C. N., & Brannon-Peppas, L. (1996). Mechanical properties of hydrogels and
   their experimental determination. *Biomaterials*, 17, 1647–1657.
- Balakrishnan, G., Nguyen, B. T., Schmitt, C., Nicolai, T., & Chassenieux, C. (2017). Heat-set
  emulsion gels of casein micelles in mixtures with whey protein isolate. *Food Hydrocolloids*,
  73, 213–221.
- Benavides, S., Cortes, P., Parada, J., & Franco, W. (2016). Development of alginate microspheres
   containing thyme essential oil using ionic gelation. *Food Chemistry*, 204, 77–83.
- Bi, C. H., Wang, P. L., Sun, D. Y., Yan, Z. M., Liu, Y., Huang, Z. G., & Gao, F. (2020). Effect of
  high-pressure homogenization on gelling and rheological properties of soybean protein isolate
  emulsion gel. *Journal of Food Engineering*, 277, 109923.

819	Boland, A. B., Delahunty, C. M., & van Ruth, S. M. (2006). Influence of the texture of gelatin gels
820	and pectin gels on strawberry flavour release and perception. Food Chemistry, 96, 452–460.
821	Bot, A., den Adel, R., Regkos, C., Sawalha, H., Venema, P., & Flöter, E. (2011). Structuring in β-
822	sitosterol+ $\gamma$ -oryzanol-based emulsion gels during various stages of a temperature cycle. Food
823	Hydrocolloids, 25, 639–646.
824	Bot, A., Erle, U., Vreeker, R., & Agterof, W. G. M. (2004). Influence of crystallisation conditions on
825	the large deformation rheology of inulin gels. <i>Food Hydrocolloids</i> , 18, 547–556.
826	Brito-Oliveira, T. C., Bispo, M., Moraes, I. C. F., Campanella, O. H., & Pinho, S. C. (2017). Stability
827	of curcumin encapsulated in solid lipid microparticles incorporated in cold-set emulsion filled
828	gels of sov protein isolate and xanthan gum. Food Research International, 102, 759–767.
829	Charoen, R., Jangchud, A., Jangchud, K., Harnsilawat, T., Naivikul, O., & McClements, D. J. (2011).
830	Influence of biopolymer emulsifier type on formation and stability of rice bran oil in water
831	emulsions: whey protein gum arabic and modified starch <i>Journal of Food Science</i> 76
832	F165_F172
833	Chen H. Mao I. Hou Z. Yuan F. & Gao Y. (2020). Roles of additional emulsifiers in the
834	structures of emulsion gels and stability of vitamin F. Food Hydrocolloids 99, 105372
835	Chen I & Dickinson E (1998a) Viscoelastic properties of heat-set whey protein emulsion gels
836	Lournal of Texture Studies 20, 285–304
030	Chan I & Dickinson E (1008b) Viscoalestic Properties of Protein Stabilized Emulsions Effect of
027	Distribution Surfactant Interactions, Eard Chamistry 46, 01, 07
020	Chen L & Dialtingon E (1000a). Effect of monoglycorides and diglycoral actors on viscoelecticity of
039	heat act where protein ampleion cole. International Journal of Food Science & Technology
040	102 102 102 102 102 102 102 102 102 102
841	53,495-501.
842	Chen, J., & Dickinson, E. (1999b). Interfacial ageing effect on the meology of a heat-set protein
843	emulsion gel. Food Hydrocollolds, 13, 363–369.
844	Chen, J., & Dickinson, E. (2000). On the temperature reversibility of the viscoelasticity of acid-
845	induced sodium caseinate emulsion gels. International Dairy Journal, 10, 541–549.
846	Chen, J., Dickinson, E., Langton, M., & Hermansson, AM. (2000). Mechanical properties and
847	microstructure of heat-set whey protein emulsion gels: Effect of emulsifiers. LWT - Food
848	Science and Technology, 33, 299–307.
849	Chen, J., Ren, Y., Zhang, K., Qu, J., Hu, F., & Yan, Y. (2019). Phosphorylation modification of
850	myofibrillar proteins by sodium pyrophosphate affects emulsion gel formation and oxidative
851	stability under different pH conditions. <i>Food &amp; Function</i> , 10, 6568–6581.
852	Cheng, Y., Donkor, P. O., Ren, X., Wu, J., Agyemang, K., Ayim, I., & Ma, H. (2019). Effect of
853	ultrasound pretreatment with mono-frequency and simultaneous dual frequency on the
854	mechanical properties and microstructure of whey protein emulsion gels. Food
855	Hydrocolloids, 89, 434–442.
856	Ching, S. H., Bansal, N., & Bhandari, B. (2016). Rheology of emulsion-filled alginate microgel
857	suspensions. Food Research International, 80, 50-60.
858	Ching, S. H., Bansal, N., & Bhandari, B. (2017). Alginate gel particles-A review of production
859	techniques and physical properties. Critical Reviews in Food Science & Nutrition, 57, 1133-
860	1152.
861	Co, E. D., & Marangoni, A. G. (2012). Organogels: An alternative edible oil structuring method.
862	Journal of the American Oil Chemists' Society, 89, 749–780.
863	Cofrades, S., Bou, R., Flaiz, L., Garcimartin, A., Benedi, J., Mateos, R., Sanchez-Muniz, F. J.,
864	Olivero-David, R., & Jimenez-Colmenero, F. (2017). Bioaccessibility of hydroxytyrosol and
865	n-3 fatty acids as affected by the delivery system: simple, double and gelled double
866	emulsions. Journal of Food Science Technol, 54, 1785–1793.
867	Corstens, M. N., Berton-Carabin, C. C., Elichiry-Ortiz, P. T., Hol, K., Troost, F. J., Masclee, A. A.
868	M., & Schroën, K. (2017). Emulsion-alginate beads designed to control <i>in vitro</i> intestinal
869	lipolysis: Towards appetite control. Journal of Functional Foods. 34, 319–328.
870	Dai, L., Sun, C., Wei, Y., Mao, L., & Gao, Y. (2018). Characterization of Pickering emulsion gels
871	stabilized by zein/gum arabic complex colloidal nanoparticles. <i>Food Hydrocolloids</i> 74 239–
872	248.

873	de Souza Paglarini, C., de Figueiredo Furtado, G., Biachi, J. P., Vidal, V. A. S., Martini, S., Forte, M.
874	B. S., Cunha, R. L., & Pollonio, M. A. R. (2018). Functional emulsion gels with potential
875	application in meat products. Journal of Food Engineering, 222, 29–37.
876	de Souza Paglarini, C., de Figueiredo Furtado, G., Honório, A. R., Mokarzel, L., da Silva Vidal, V.
877	A., Ribeiro, A. P. B., Cunha, R. L., Pollonio, M. A. R. (2019). Functional emulsion gels as
878	pork back fat replacers in Bologna sausage. Food Structure, 20, 100105.
879	de Souza Paglarinia, C., Martinib, S., & Pollonio, M. A. R. (2019). Using emulsion gels made with
880	sonicated soy protein isolate dispersions to replace fat in frankfurters. LWT - Food Science
881	and Technology, 99, 453–459.
882	Devezeaux de Lavergne, M., Tournier, C., Bertrand, D., Salles, C., van de Velde, F., & Stieger, M.
883	(2016). Dynamic texture perception, oral processing behaviour and bolus properties of
884	emulsion-filled gels with and without contrasting mechanical properties. Food Hydrocolloids,
885	52, 648–660.
886	Dickinson, E. (2012). Emulsion gels: The structuring of soft solids with protein-stabilized oil droplets.
887	Food Hydrocolloids, 28, 224–241.
888	Dickinson, E., & Merino, L. M. (2002). Effect of sugars on the rheological properties of acid
889	caseinate-stabilized emulsion gels. Food Hydrocolloids, 16, 321-331.
890	Farjami, T., & Madadlou, A. (2019). An overview on preparation of emulsion-filled gels and
891	emulsion particulate gels. Trends in Food Science & Technology, 86, 85–94.
892	Feng, L., Jia, X., Zhu, Q., Liu, Y., Li, J., & Yin, L. (2019). Investigation of the mechanical,
893	rheological and microstructural properties of sugar beet pectin/soy protein isolate-based
894	emulsion-filled gels. Food Hydrocolloids, 89, 813–820.
895	Feng, W., Yue, C., Wusigale, Ni, Y., & Liang, L. (2018). Preparation and characterization of
896	emulsion-filled gel beads for the encapsulation and protection of resveratrol and alpha-
897	tocopherol. Food Research International, 108, 161–171.
898	Feng, X., Chen, L., Lei, N., Wang, S., Xu, X., Zhou, G., & Li, Z. (2017). Emulsifying properties of
899	oxidatively stressed myofibrillar protein emulsion gels prepared with (-)-epigallocatechin-3-
900	gallate and NaCl. Journal of Agricultural and Food Chemistryistry, 65, 2816–2826.
901	Flaiz, L., Freire, M., Cofrades, S., Mateos, R., Weiss, J., Jimenez-Colmenero, F., & Bou, R. (2016).
902	Comparison of simple, double and gelled double emulsions as hydroxytyrosol and n-3 fatty
903	acid delivery systems. Food Chemistry, 213, 49-57.
904	Freire, M., Bou, R., Cofrades, S., & Jimenez-Colmenero, F. (2017). Technological characteristics of
905	cold-set gelled double emulsion enriched with n-3 fatty acids: Effect of hydroxytyrosol
906	addition and chilling storage. Food Research International, 100(Pt 2), 298-305.
907	Freire, M., Cofrades, S., Perez-Jimenez, J., Gomez-Estaca, J., Jimenez-Colmenero, F., & Bou, R.
908	(2018). Emulsion gels containing n-3 fatty acids and condensed tannins designed as
909	functional fat replacers. Food Research International, 113, 465–473.
910	Freire, M., Cofrades, S., Serrano-Casas, V., Pintado, T., Jimenez, M. J., & Jimenez-Colmenero, F.
911	(2017). Gelled double emulsions as delivery systems for hydroxytyrosol and n-3 fatty acids in
912	healthy pork patties. Journal of Food Science Technol, 54, 3959-3968.
913	Gaspar, A. L., & de Goes-Favoni, S. P. (2015). Action of microbial transglutaminase (MTGase) in the
914	modification of food proteins: a review. Food Chemistry, 171, 315-322.
915	Geremias-Andrade, I. M., Souki, N. P. D. B. G., Moraes, I. C. F., & Pinho, S. C. (2017). Rheological
916	and mechanical characterization of curcumin-loaded emulsion-filled gels produced with whey
917	protein isolate and xanthan gum. LWT - Food Science and Technology, 86, 166–173.
918	Glibowski, P., & Pikus, S. (2011). Amorphous and crystal inulin behavior in a water environment.
919	Carbohydrate Polymers, 83, 635–639.
920	Glisic, M., Baltic, M., Glisic, M., Trbovic, D., Jokanovic, M., Parunovic, N., Vasilev, D. (2019).
921	Inulin based emulsion filled gel as a fat replacer in prebiotic and PUFA enriched dry
922	fermented sausages. International Journal of Food Science & Technology, 54, 787–797.
923	Glusac, J., Davidesko-Vardi, I., Isaschar-Ovdat, S., Kukavica, B., & Fishman, A. (2018). Gel-like
924	emulsions stabilized by tyrosinase-crosslinked potato and zein proteins. <i>Food Hydrocolloids</i> ,
925	82, 53-63.

Gómez-Guillén, M. C., Giménez, B., López-Caballero, M. E., & Montero, M. P. (2011). Functional 926 and bioactive properties of collagen and gelatin from alternative sources: A review. Food 927 Hydrocolloids, 25, 1813–1827. 928 929 Guichard, E. (2002). Interactions between flavor compounds and food ingredients and their influence on flavor perception. Food Reviews International, 18, 49-70. 930 931 Guo, Q., Bellissimo, N., & Rousseau, D. (2017). Role of gel structure in controlling in vitro intestinal 932 lipid digestion in whey protein emulsion gels. Food Hydrocolloids, 69, 264-272. 933 Guo, Q., Ye, A., Lad, M., Dalgleish, D., & Singh, H. (2013). The breakdown properties of heat-set 934 whey protein emulsion gels in the human mouth. Food Hydrocolloids, 33(, 215–224. 935 Guo, Q., Ye, A., Lad, M., Dalgleish, D., & Singh, H. (2016). Impact of colloidal structure of gastric digesta on in-vitro intestinal digestion of whey protein emulsion gels. Food Hydrocolloids, 936 937 54, 255-265. Guzey, D., & McClements, D. J. (2006). Formation, stability and properties of multilayer emulsions 938 939 for application in the food industry. Advances in Colloid and Interface Science, 128, 227-248. 940 Gwartney, E. A., Larick, D. K., & Foegeding, E. A. (2004). Sensory texture and mechanical 941 properties of stranded and particulate whey protein emulsion gels. Journal of Food Science, 942 69, S333-S339. Hemung, B.-O., Benjakul, S., & Yongsawatdigul, J. (2013). pH-dependent characteristics of gel-like 943 944 emulsion stabilized by threadfin bream sarcoplasmic proteins. Food Hydrocolloids, 30, 315-945 322. Herrero, A. M., Ruiz-Capillas, C., Pintado, T., Carmona, P., & Jiménez-Colmenero, F. (2018). 946 947 Elucidation of lipid structural characteristics of chia oil emulsion gels by Raman spectroscopy 948 and their relationship with technological properties. Food Hydrocolloids, 77, 212–219. Hou, J. J., Guo, J., Wang, J. M., & Yang, X. Q. (2016). Effect of interfacial composition and 949 950 crumbliness on aroma release in soy protein-sugar beet pectin mixed emulsion gels. Journal of the Science of Food and Agriculture, 96, 4449–4456. 951 952 Jiang, J., & Xiong, Y. L. (2013). Extreme pH treatments enhance the structure-reinforcement role of 953 soy protein isolate and its emulsions in pork myofibrillar protein gels in the presence of 954 microbial transglutaminase. Meat Science, 93, 469-476. 955 Jiang, Y., Liu, L., Wang, B., Yang, X., Chen, Z., Zhong, Y., Zhang, L., Mao, Z., Xu, H., Sui, X. 956 (2019). Polysaccharide-based edible emulsion gel stabilized by regenerated cellulose. Food 957 Hydrocolloids, 91, 232–237. Kang, Y. N., Kim, H., Shin, W. S., Woo, G., & Moon, T. W. (2003). Effect of disulfide bond 958 959 reduction on bovine serum albumin-stabilized emulsion gel formed by microbial 960 transglutaminase. Journal of Food Science, 68, 2215-2220. Kerner, E. (1956). The elastic and thermo-elastic properties of composite media. Proceedings of the 961 physical society. Section B, 69, 808. 962 Khalesi, H., Emadzadeh, B., Kadkhodaee, R., & Fang, Y. (2019). Effect of Persian gum on whey 963 964 protein concentrate cold-set emulsion gel: Structure and rheology study. International Journal 965 of Biological Macromolecules, 125, 17-26. 966 Kim, K., Gohtani, S., Matsuno, R., & Yamano, Y. (1999). Effects of oil droplet and agar concentration on gel strength and microstructure of o-w emulsion gel. Journal of Texture 967 Studies, 30, 319-335. 968 969 King, A. (1983). Brown seaweed extracts (alginates). Food Hydrocolloids, 2, 115–188. 970 Kiokias, S., & Bot, A. (2005). Effect of denaturation on temperature cycling stability of heated 971 acidified protein-stabilised o/w emulsion gels. Food Hydrocolloids, 19, 493-501. 972 Kiokias, S., & Bot, A. (2006). Temperature cycling stability of pre-heated acidified whey protein-973 stabilised o/w emulsion gels in relation to the internal surface area of the emulsion. Food 974 Hydrocolloids, 20, 245-252. Koc, H., Drake, M., Vinyard, C. J., Essick, G., van de Velde, F., & Foegeding, E. A. (2019). 975 Emulsion filled polysaccharide gels: Filler particle effects on material properties, oral 976 977 processing, and sensory texture. Food Hydrocolloids, 94, 311-325. 978 Lam, R. S. H., & Nickerson, M. T. (2014). The properties of whey protein-carrageenan mixtures 979 during the formation of electrostatic coupled biopolymer and emulsion gels. Food Research 980 International, 66, 140-149.

981	Langton, M., & Hermansson, AM. (1992). Fine-stranded and particulate gels of β-lactoglobulin and
982	whey protein at varying pH. Food Hydrocolloids, 5, 523–539.
983	Larson-Smith, K., Jackson, A., & Pozzo, D. C. (2012). SANS and SAXS analysis of charged
984	nanoparticle adsorption at oil-water interfaces. Langmuir, 28, 2493-2501.
985	Leon, A. M., Medina, W. T., Park, D. J., & Aguilera, J. M. (2018). Properties of microparticles from a
986	whey protein isolate/alginate emulsion gel. Food Science and Technology International, 24,
987	414–423.
988	Lević, S., Pajić Lijaković, I., Đorđević, V., Rac, V., Rakić, V., Šolević Knudsen, T., Pavlović, V.,
989	Bugarski, B., & Nedović, V. (2015). Characterization of sodium alginate/d-limonene
990	emulsions and respective calcium alginate/d-limonene beads produced by electrostatic
991	extrusion. Food Hydrocolloids, 45, 111–123.
992	Lewis, T. B., & Nielsen, L. E. (1970). Dynamic mechanical properties of particulate [filled]
993	composites. Journal of Applied Polymer Science, 14, 1449–1471.
994	Li, A., Gong, T., Hou, Y., Yang, X., & Guo, Y. (2020). Alginate-stabilized thixotropic emulsion gels
995	and their applications in fabrication of low-fat mayonnaise alternatives. International Journal
996	of Biological Macromolecules, 146, 821–831.
997	Li, F., & Hua, Y. (2013). Rheological properties of acid-induced soy protein-stabilized emulsion gels
998	in the absence and presence of N-ethylmaleimide. Food Hydrocolloids, 30, 641-646.
999	Li, F., Kong, X., Zhang, C., & Hua, Y. (2012). Gelation behaviour and rheological properties of acid-
1000	induced soy protein-stabilized emulsion gels. Food Hydrocolloids, 29, 347-355.
1001	Li, S., Zhang, B., Li, C., Fu, X., & Huang, Q. (2020). Pickering emulsion gel stabilized by
1002	octenylsuccinate quinoa starch granule as lutein carrier: Role of the gel network. Food
1003	Chemistry, 305, 125476.
1004	Liang, L., Leung Sok Line, V., Remondetto, G. E., & Subirade, M. (2010). In vitro release of a-
1005	tocopherol from emulsion-loaded β-lactoglobulin gels. International Dairy Journal, 20, 176–
1006	181.
1007	Liang, X., Ma, C., Yan, X., Zeng, H., McClements, D. J., Liu, X., & Liu, F. (2020). Structure,
1008	rheology and functionality of whey protein emulsion gels: Effects of double cross-linking
1009	with transglutaminase and calcium ions. Food Hydrocolloids, 102, 105569.
1010	Lim, S. H., Kim, H. R., Choi, S. J., & Moon, T. W. (2015). Lipid oxidation of sodium caseinate-
1011	stabilized emulsion-gels prepared using microbial transglutaminase. Food Science and
1012	Biotechnology, 24, 2023–2026.
1013	Liu, F., & Tang, CH. (2016). Soy glycinin as food-grade Pickering stabilizers: Part. II. Improvement
1014	of emulsification and interfacial adsorption by electrostatic screening. <i>Food Hydrocolloids</i> ,
1015	60, 620–630.
1016	Liu, W., Gao, H. X., McClements, D. J., Zhou, L., Wu, J., & Zou, L. Q. (2019). Stability, rheology,
1017	and beta-carotene bioaccessibility of high internal phase emulsion gels. <i>Food Hydrocolloids</i> ,
1018	88, 210–217.
1019	Liu, X., Chen, XW., Guo, J., Yin, SW., & Yang, XQ. (2016). Wheat gluten based percolating
1020	emulsion gels as simple strategy for structuring liquid oil. <i>Food Hydrocolloids</i> , 61, 747–755.
1021	Lorenzo, G., Zaritzky, N., & Califano, A. (2013). Rheological analysis of emulsion-filled gels based
1022	on high acyl gellan gum. Food Hydrocolloids, 30, 672–680.
1023	Lu, Y., Mao, L., Zheng, H., Chen, H., & Gao, Y. (2020). Characterization of β-carotene loaded
1024	emulsion gels containing denatured and native whey protein. <i>Food Hydrocolloids</i> , 102,
1025	
1026	Luo, LJ., Liu, F., & Tang, CH. (2013). The role of glycinin in the formation of gel-like soy protein-
1027	stabilized emulsions. Food Hydrocolloids, 32, 97–105.
1028	Ma, L., Wan, Z., & Yang, X. (2017). Multiple Water-in-Oil-in-Water Emulsion Gels Based on Self-
1029	Assembled Saponin Fibriliar Network for Photosensitive Cargo Protection. Journal of
1030	Agricultural and Food Chemistryistry, 65, 9/35–9/43.
1031	Mao, L., Koos, Y. H., & Miao, S. (2014). Study on the rheological properties and volatile release of cold set emulsion filled protein gals. <i>Journal of Agricultural and Ecod Chamisturistics</i> , 62
1032	con-set emuision-med protein geis. Journal of Agricultural and Food Chemistryistry, 02,

1033 11420–11428.

1034	Montes de Oca-Avalos, J. M., Huck-Iriart, C., Candal, R. J., & Herrera, M. L. (2016). Sodium
1035	caseinate/sunflower oil emulsion-based gels for structuring food. Food and Bioprocess
1036	<i>Technology</i> , 9, 981–992.
1037	Nishinari, K., & Takahashi, R. (2003). Interaction in polysaccharide solutions and gels. Current
1038	Opinion in Colloid & Interface Science, 8, 396–400.
1039	Nourbehesht, N., Shekarchizadeh, H., & Soltanizadeh, N. (2018). Investigation of stability,
1040	consistency, and oil oxidation of emulsion filled gel prepared by inulin and rice bran oil using
1041	ultrasonic radiation <i>Ultrason Sonochem</i> 42 585–593
1042	Okada K S & Lee Y (2017) Characterization of sodium mobility and binding by 23Na NMR
10/12	spectroscopy in a model lipoproteic emulsion gel for sodium reduction. <i>Journal of Food</i>
1045	Science, 82, 1563–1568.
1045	Oliver, L., Berndsen, L., van Aken, G. A., & Scholten, E. (2015). Influence of droplet clustering on
1046	the rheological properties of emulsion-filled gels. <i>Food Hydrocolloids</i> , 50, 74–83.
1047	Oliver L. Scholten E. & van Aken G. A. (2015) Effect of fat hardness on large deformation
1048	rheology of emulsion-filled gels. Food Hydrocolloids 43, 299–310
1049	Oliver I Wieck I & Scholten F (2016) Influence of matrix inhomogeneity on the rheological
1050	properties of emulsion-filled gels <i>Eood Hydrocolloids</i> 52 116–125
1050	Ozturk B Argin S Ozilgen M & McClements D I (2015) Formation and stabilization of
1051	nanoamulsion based vitamin E delivery systems using natural biopolymers: Whey protein
1052	isolate and gum arabic. <i>Ecod Chamistry</i> 188, 256, 263
1055	Deredice V. M. Cierretti M. Summe C. Descuelone A. Minervini E. & Canonio E. (2015)
1054	Production and characterization of amulaian filled cale based on invite and artra virgin alive
1055	Production and characterization of emulsion fined gets based on multif and extra virgin onve
1050	011. Food Hydrocollolds, 43, 30–40.
1057	Pernetti, M., van Maissen, K. F., Floter, E., & Bot, A. (2007). Structuring of edible oils by alternatives
1058	to crystalline fat. Current Opinion in Colloid & Interface Science, 12, 221–231.
1059	Pintado, T., Herrero, A. M., Jimenez-Colmenero, F., Pasqualin Cavalheiro, C., & Ruiz-Capillas, C.
1060	(2018). Chia and oat emulsion gels as new animal fat replacers and healthy bioactive sources
1061	in fresh sausage formulation. <i>Meat Science</i> , 135, 6–13.
1062	Pintado, T., Ruiz-Capillas, C., Jimenez-Colmenero, F., Carmona, P., & Herrero, A. M. (2015). Oil-in-
1063	water emulsion gels stabilized with chia (Salvia hispanica L.) and cold gelling agents:
1064	Technological and infrared spectroscopic characterization. <i>Food Chemistry</i> , 185, 470–478.
1065	Piornos, J. A., Burgos-Díaz, C., Morales, E., Rubilar, M., & Acevedo, F. (2017). Highly efficient
1066	encapsulation of linseed oil into alginate/lupin protein beads: Optimization of the emulsion
1067	formulation. Food Hydrocolloids, 63, 139–148.
1068	Poyato, C., Astiasarán, I., Barriuso, B., & Ansorena, D. (2015). A new polyunsaturated gelled
1069	emulsion as replacer of pork back-fat in burger patties: Effect on lipid composition, oxidative
1070	stability and sensory acceptability. LWT - Food Science and Technology, 62, 1069–1075.
1071	Ringgenberg, E., Alexander, M., & Corredig, M. (2013). Effect of concentration and incubation
1072	temperature on the acid induced aggregation of soymilk. Food Hydrocolloids, 30, 463-469.
1073	Ruffin, E., Schmit, T., Lafitte, G., Dollat, J. M., & Chambin, O. (2014). The impact of whey protein
1074	preheating on the properties of emulsion gel bead. Food Chemistry, 151, 324–332.
1075	Sala, G., van Vliet, T., Cohen Stuart, M. A., Aken, G. A. v., & van de Velde, F. (2009). Deformation
1076	and fracture of emulsion-filled gels: Effect of oil content and deformation speed. Food
1077	Hydrocolloids, 23, 1381–1393.
1078	Sato, A. C. K., Moraes, K. E. F. P., & Cunha, R. L. (2014). Development of gelled emulsions with
1079	improved oxidative and pH stability. <i>Food Hydrocolloids</i> , 34, 184–192.
1080	Sawalha, H., den Adel, R., Venema, P., Bot, A., Floter, E., & van der Linden E. (2012). Organogel-
1081	emulsions with mixtures of beta-sitosterol and gamma-oryzanol: influence of water activity
1082	and type of oil phase on gelling capability. <i>Journal of Agricultural and Food Chemistryistry</i>
1083	60 3462–3470
108/	Serdarogly M Nacak B & Karabivikogly M (2017) Effects of beef fat replacement with gelled
1004	Solution in the second se

- 1085 emulsion prepared with olive oil on quality parameters of chicken patties. *Korean Journal of* 1086 *Food Science Anim Resour, 37*, 376–384.
- Serdaroğlu, M., & Öztürk, B. (2017). Use of olive oil-in-water gelled emulsions in model turkey
   breast emulsions. *IOP Conference Series: Earth and Environmental Science*, 85, 012071.

Shao, Y., & Tang, C.-H. (2016). Gel-like pea protein Pickering emulsions at pH3.0 as a potential 1089 1090 intestine-targeted and sustained-release delivery system for β-carotene. Food Research International, 79, 64-72. 1091 Sok Line, V. L., Remondetto, G. E., & Subirade, M. (2005). Cold gelation of β-lactoglobulin oil-in-1092 water emulsions. Food Hydrocolloids, 19, 269–278. 1093 Soukoulis, C., Cambier, S., Hoffmann, L., & Bohn, T. (2016). Chemical stability and bioaccessibility 1094 1095 of  $\beta$ -carotene encapsulated in sodium alginate o/w emulsions: Impact of Ca 2+ mediated 1096 gelation. Food Hydrocolloids, 57, 301-310. Soukoulis, C., Tsevdou, M., Andre, C. M., Cambier, S., Yonekura, L., Taoukis, P. S., & Hoffmann, L. 1097 1098 (2017). Modulation of chemical stability and *in vitro* bioaccessibility of beta-carotene loaded 1099 in kappa-carrageenan oil-in-gel emulsions. Food Chemistry, 220, 208-218. 1100 Su, J., Guo, Q., Chen, Y., Dong, W., Mao, L., Gao, Y., & Yuan, F. (2020). Characterization and formation mechanism of lutein pickering emulsion gels stabilized by  $\beta$ -lactoglobulin-gum 1101 1102 arabic composite colloidal nanoparticles. Food Hydrocolloids, 98, 105276. Suzawa, E., & Kaneda, I. (2010). Rheological properties of agar microgel suspensions prepared using 1103 1104 water-in-oil emulsions. Journal of biorheology, 24, 70-76. Tang, C.-H., Luo, L.-J., Liu, F., & Chen, Z. (2013). Transglutaminase-set soy globulin-stabilized 1105 emulsion gels: Influence of soy β-conglycinin/glycinin ratio on properties, microstructure and 1106 1107 gelling mechanism. Food Research International, 51, 804-812. 1108 Tang, C.-h., Yang, M., Liu, F., & Chen, Z. (2013). A novel process to efficiently form transglutaminase-set soy protein isolate-stabilized emulsion gels. LWT - Food Science and 1109 1110 Technology, 53, 15-21. 1111 Tang, C. H., Chen, L., & Foegeding, E. A. (2011). Mechanical and water-holding properties and microstructures of soy protein isolate emulsion gels induced by CaCl2, glucono-delta-lactone 1112 (GDL), and transglutaminase: influence of thermal treatments before and/or after 1113 emulsification. Journal of Agricultural and Food Chemistryistry, 59, 4071-4077. 1114 Tolano-Villaverde, I. J., Torres-Arreola, W., Ocaño-Higuera, V. M., & Marquez-Rios, E. (2015). 1115 1116 Thermal gelation of myofibrillar proteins from aquatic organisms. CyTA - Journal of Food, 1–7. 1117 Torres, O., Tena, N. M., Murray, B., & Sarkar, A. (2017). Novel starch based emulsion gels and 1118 1119 emulsion microgel particles: Design, structure and rheology. Carbohydrate Polymers, 178, 1120 86-94. Van Vliet, T. (1988). Rheological properties of filled gels. Influence of filler matrix interaction. 1121 Colloid and Polymer Science, 266, 518-524. 1122 1123 Wakita, K., & Imura, T. (2018). High internal phase emulsion gels stabilized by natural casein peptides. Journal of Oleo Science, 67, 1579-1584. 1124 Wan, Z., Sun, Y., Ma, L., Yang, X., Guo, J., & Yin, S. (2017). Responsive emulsion gels with tunable 1125 properties formed by self-assembled nanofibrils of natural saponin glycyrrhizic acid for oil 1126 1127 structuring. Journal of Agricultural and Food Chemistryistry, 65, 2394–2405. Wang, L. J., Yin, S. W., Wu, L. Y., Qi, J. R., Guo, J., & Yang, X. Q. (2016). Fabrication and 1128 1129 characterization of Pickering emulsions and oil gels stabilized by highly charged zein/chitosan complex particles (ZCCPs). Food Chemistry, 213, 462-469. 1130 Wang, Q., Jiang, J., & Xiong, Y. L. (2019). Genipin-Aided Protein Cross-linking to Modify Structural 1131 1132 and Rheological Properties of Emulsion-Filled Hempseed Protein Hydrogels. Journal of Agricultural and Food Chemistry, 67, 12895–12903. 1133 Wang, S., Zhang, Y., Chen, L., Xu, X., Zhou, G., Li, Z., & Feng, X. (2018). Dose-dependent effects 1134 of rosmarinic acid on formation of oxidatively stressed myofibrillar protein emulsion gel at 1135 different NaCl concentrations. Food Chemistry, 243, 50-57. 1136 1137 Wang, X., Luo, K., Liu, S., Adhikari, B., & Chen, J. (2019). Improvement of gelation properties of soy protein isolate emulsion induced by calcium cooperated with magnesium. Journal of 1138 Food Engineering, 244, 32–39. 1139 1140 Wang, X., Xiong, Y. L., & Sato, H. (2017). Rheological enhancement of pork myofibrillar protein-1141 lipid emulsion composite gels via glucose oxidase oxidation/transglutaminase cross-linking pathway. Journal of Agricultural and Food Chemistryistry, 65, 8451-8458. 1142

1143	Wang, X. F., Luo, K. Y., Liu, S. T., Zeng, M. M., Adhikari, B., He, Z. Y., & Chen, J. (2018). Textural
1144	and rheological properties of soy protein isolate tofu-type emulsion gels influence of soybean
1145	variety and coagulant type. <i>Food Biophysics</i> , 13, 324–332.
1146	Wang, Z., Neves, M. A., Kobayashi, I., Uemura, K., & Nakajima, M. (2013). Preparation,
1147	characterization, and gastrointestinal digestibility of oil-in-water emulsion-agar gels. Biosci
1148	Biotechnol Biochem, 77, 467–474.
1149	Wouters, A. G., & Delcour, J. A. (2019). Cereal protein based nanoparticles as agents stabilizing air-
1150	water and oil-water interfaces in food systems. Current Opinion in Food Science, 25, 19-27.
1151	Wu, M., Xiong, Y. L., & Chen, J. (2011). Role of disulphide linkages between protein-coated lipid
1152	droplets and the protein matrix in the rheological properties of porcine myofibrillar protein-
1153	peanut oil emulsion composite gels. <i>Meat Science</i> , 88, 384–390.
1154	Wu M Xiong Y L. Chen I Tang X & Zhou G (2009) Rheological and microstructural
1155	properties of porcine myofibrillar protein-linid emulsion composite gels. <i>Journal of Food</i>
1156	Science 74 E207_217
1157	Vi 7 Lin W McClements D I & Zou I (2010) Pheological structural and microstructural
1157	AI, Z., Elu, W., McClements, D. J., & Zou, E. (2019). Kneological, subclutal, and inclosubclutal
1120	properties of ethanor induced cold-set whey protein emuision gets. Effect of on content. <i>Food</i>
1159	Chemistry, 291, 22–29.
1160	Xiao, J., Wang, X. a., Gonzalez, A. J. P., & Huang, Q. (2016). Kafirin nanoparticles-stabilized
1161	Pickering emulsions: Microstructure and rheological behavior. Food Hydrocolloids, 54, 30–
1162	<i>39.</i>
1163	Xu, Y. T., Liu, T. X., & Tang, C. H. (2019). Novel pickering high internal phase emulsion gels
1164	stabilized solely by soy $\beta$ -conglycinin. Food Hydrocolloids, 88, 21–30.
1165	Xu, W., Huang, L., Jin, W., Ge, P., Shah, B. R., Zhu, D., & Jing, J. (2019). Encapsulation and release
1166	behavior of curcumin based on nanoemulsions-filled alginate hydrogel beads. International
1167	Journal of Biological Macromolecules, 134, 210–215.
1168	Yang, X., Gong, T., Li, D., Li, A., Sun, L., & Guo, Y. (2019). Preparation of high viscoelastic
1169	emulsion gels based on the synergistic gelation mechanism of xanthan and konjac
1170	glucomannan. Carbohydrate Polymers, 226, 115278.
1171	Yang, X., Gong, T., Lu, Yh., Li, A., Sun, L., & Guo, Y. (2020). Compatibility of sodium alginate
1172	and konjac glucomannan and their applications in fabricating low-fat mayonnaise-like
1173	emulsion gels. Carbohydrate Polymers, 229, 115468.
1174	Yang, X., Li, A., Yu, W., Li, X., Sun, L., Xue, J., & Guo, Y. (2020). Structuring oil-in-water emulsion
1175	by forming egg yolk/alginate complexes: Their potential application in fabricating low-fat
1176	mayonnaise-like emulsion gels and redispersible solid emulsions. International Journal of
1177	Biological Macromolecules, 147, 595–606.
1178	Ye, A., & Taylor, S. (2009). Characterization of cold-set gels produced from heated emulsions
1179	stabilized by whey protein. International Dairy Journal, 19, 721–727.
1180	Yonekura L. & Nagao A (2007) Intestinal absorption of dietary carotenoids <i>Molecular Nutrition</i>
1181	& Food Research 51 107–115
1182	Zhang C & Zhang H (2018) Formation and stability of core-shell nanofibers by electrospinning of
1183	gel-like corn oil-in-water emulsions stabilized by gelatin <i>Journal of Agricultural and Food</i>
118/	Chemistryistry 66 11681_11690
1104	Zhao X Zou V E Shao I I Chen X Han M V & Xu X I (2017) Comparison of the
1105	Acidia and Alkalina Transmant on Emulsion Composite Cal Properties of the Proteins
1100	Actuic and Alkanne Treatment on Emulsion Composite Ger Properties of the Proteins
1107	East Drassing and Drassing 41 a12894
1100	<i>Food Processing and Preservation</i> , 41, e12664.
1189	Zneng, H., Mao, L., Cui, M., Liu, J., & Gao, Y. (2020). Development of food-grade bigets based on k-
1190	carrageenan hydrogel and monoglyceride oleogels as carriers for p-carotene: Roles of oleogel
1191	iraction. Food Hydrocollolas, 103, 103855.
1192	Znou, F., Sun, W., & Zhao, M. (2015). Controlled formation of emulsion gels stabilized by salted
1193	myofibrillar protein under malondialdehyde (MDA)-induced oxidative stress. <i>Journal of</i>
1194	Agricultural and Food Chemistryistry, 63, 3766–3777.
1195	Zhuang, X., Jiang, X., Zhou, H., Han, M., Liu, Y., Bai, Y., Cu, X. L., Zhou, G. H. (2019). The effect
1196	of insoluble dietary fiber on myofibrillar protein emulsion gels: Oil particle size and protein
1197	network microstructure. LWT, 101, 534–542.

- Zhang, Z. P., Zhang, R. J., Zou, L. Q., Chen, L., Ahmed, Y., Bishri, W. A., Balamash, K., &
   McClements, D. J. (2016). Encapsulation of curcumin in polysaccharide-based hydrogel
   beads. *Food Hydrocolloids*, 58, 160–170.
- Zou, Y., Guo, J., Yin, S. W., Wang, J. M., & Yang, X. Q. (2015). Pickering emulsion gels prepared by
   hydrogen-bonded zein/tannic acid complex colloidal particles. *Journal of Agricultural and Food Chemistryistry*, 63, 7405–7414.
- Zou, Y., Yang, X. Q., & Scholten, E. (2018). Rheological behavior of emulsion gels stabilized by
   zein/tannic acid complex particles. *Food Hydrocolloids*, 77, 363–371.

## 1206 **Table 1**

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1207 Selected examples of materials and methods used to prepare bulk emulsion gels.

Materials	Methods	Matrix	Emulsifier/oil category and content	Structure	References
Protein	Heat treatment	Myofibrillar protein (MP)	MP/soybean oil ( $\varphi = 0.25$ ), peanut oil	O/W	(Feng, et al., 2017; Wang, et al., 2018; Wu, Xiong,
			(5%) or lard and peanut oil (0–15%)		& Chen, 2011; Wu, Xiong, Chen, Tang, & Zhou,
					2009)
		Myofibrillar protein	SPI/canola oil (10%)	O/W	(Jiang & Xiong, 2013)
		Chicken protein isolate (CPI)	CPI/pork backfat (20%)	O/W	(Zhao, Zou, Shao, Chen, Han, & Xu, 2017)
		Whey protein isolate (WPI)	WPI, glycerol monopalmitate, Tween	O/W	(Chen & Dickinson, 1998a; Chen & Dickinson,
			20, DGML, DGMO, or lecithin/canola		1999a; Chen & Dickinson, 1999b; Chen, Dickinson,
			oil (20%), soybean oil (20%),		Langton, & Hermansson, 2000; Guo, Bellissimo, &
			sunflower oil ( $\varphi = 0.05-0.25$ ) or		Rousseau, 2017; Guo, Ye, Lad, Dalgleish, & Singh,
			triolein ( $\phi = 0.3$ )		2013; Gwartney, Larick, & Foegeding, 2004)
	One-step cold-set	Cold-soluble gelatin	No emulsifier/olive oil (52%)	O/W	(Pintado, Ruiz-Capillas, Jimenez-Colmenero,
					Carmona, & Herrero, 2015)
	Cold-set after heat	Gelatin	No emulsifier/sunflower oil ( $\varphi = 0.3$ )	O/W	(Sato, Moraes, & Cunha, 2014)
	treatment				

	Gelatin	WPI/sunflower oil or fat (50% in	O/W emulsion	(Oliver, Berndsen, van Aken, & Scholten, 2015;
		emulsions) or medium-chain	filled	Oliver, Scholten, & van Aken, 2015; Sala, van Vliet,
		triglycerides (40% in emulsions)		Cohen Stuart, Aken, & van de Velde, 2009)
Enzyme treatment	Myofibrillar protein	SPI or MP/canola oil (10% or 25%)	O/W	(Jiang & Xiong, 2013; Wang, Xiong, & Sato, 2017)
(TGase)				
	Soy protein isolate (SPI)	SPI/soy oil ( $\phi = 0.2-0.6$ )	O/W	(Tang, Luo, Liu, & Chen, 2013; Tang, Yang, Liu, &
				Chen, 2013; Tang, Chen, & Foegeding, 2011)
	Bovine serum albumin	Bovine serum albumin/n-tetradecane	O/W	(Kang, Kim, Shin, Woo, & Moon, 2003)
		(φ = 0.45)		
	Sodium caseinate	Sodium caseinate/olive oil (52%) or	O/W	(Lim, Kim, Choi, & Moon, 2015; Pintado, Ruiz-
		sunflower oil (45%)		Capillas, Jimenez-Colmenero, Carmona, & Herrero,
				2015)
	Sodium caseinate	PGPR/perilla oil (80% in W <sub>1</sub> /O)	W <sub>1</sub> /O/W <sub>2</sub>	(Freire, Bou, Cofrades, & Jimenez-Colmenero,
				2017)
	Gelatin	Sodium caseinate/perilla oil (80% in	W <sub>1</sub> /O/W <sub>2</sub>	(Flaiz, Freire, Cofrades, Mateos, Weiss, Jimenez-
		W <sub>1</sub> /O)		Colmenero, et al., 2016)
Acidification	Soy protein	Soy protein/soy oil (40% or $\varphi = 0.2-$	O/W	(Fang Li & Hua, 2013;Li, Kong, Zhang, & Hua,
treatment		0.3)		2012; Tang, Chen, & Foegeding, 2011)
(GDL/citric acid)				

	Whey protein isolate	WPI/sunflower oil (20%) or milk fat	O/W	(Mao, Roos, & Miao, 2014; Ye & Taylor, 2009)
		(20-30%)		
	Sodium caseinate	Sodium caseinate/sunflower oil (10%),	O/W	(Chen & Dickinson, 2000; Dickinson & Merino,
		vegetable fat (30%) or n-tetradecane ( $\phi$		2002; Kiokias & Bot, 2005; Montes de Oca-Ávalos,
		= 0.3)		Huck-Iriart, Candal, & Herrera, 2016)
	Micellar casein isolate	WPI or casein/sunflower oil or fat	O/W emulsion	(Oliver, Scholten, & van Aken, 2015; Oliver, Wieck,
		(50% in emulsions) or milk fat (5–25%	filled	& Scholten, 2016)
		in emulsions)		
	Whey protein isolate	WPI, Tween 20, or	O/W emulsion	(Oliver, Scholten, & van Aken, 2015; Sala, van
		lactoferrin/sunflower oil or fat (50% in	filled	Vliet, Cohen Stuart, Aken, & van de Velde, 2009)
		emulsions) or medium-chain		
		triglycerides (40% in emulsions)		
Addition of ions	Soy protein	Soy protein/soy oil ( $\varphi = 0.2$ )	O/W	(Tang, Chen, & Foegeding, 2011)
	Whey protein isolate or $\beta$ -	Proteins/sunflower oil (30%) or milk	O/W	(Liang, Leung Sok Line, Remondetto, & Subirade,
	lactoglobulin	fat (20–30%)		2010; Sok Line, Remondetto, & Subirade, 2005; Ye
				& Taylor, 2009)
Malondialdehyde	Myofibrillar protein	MP/soybean oil (20%)	O/W	(Zhou, Sun, & Zhao, 2015)
(MDA)				

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Protein/protein	Heat treatment	Micelle casein/whey protein isolate	Proteins/sunflower oil (5–15%)	O/W	(Balakrishnan, Nguyen, Schmitt, Nicolai, &
					Chassenieux, 2017)
	Enzyme treatment	Potato protein/zein	Potato protein and zein/olive oil ( $\varphi$ =	O/W	(Glusac, Davidesko-Vardi, Isaschar-Ovdat,
	(Tyrosinase)		0.4)		Kukavica, & Fishman, 2018)
Polysaccharide	Cold-set after heat	к-Carrageenan	Polysorbate 80 or no	O/W or bigels	(Poyato, Astiasarán, Barriuso, & Ansorena, 2015;
	treatment		emulsifier/sunflower oil (40%) or corn		Zheng, Mao, Cui, Liu, & Gao, 2020)
			oil with monoglycerides (25-75%)		
		Gellan gum	Tween 80/sunflower oil (10–30%)	O/W	(Lorenzo, Zaritzky, & Califano, 2013)
		Agar	Polyglycerol esters of fatty	O/W emulsion	(Kim, Gohtani, Matsuno, & Yamano, 1999; Wang,
			acids/soybean oil (20% in emulsions)	filled	Neves, Kobayashi, Uemura, & Nakajima, 2013)
			or corn oil ( $\phi = 0.3$ in emulsions)		
		κ-Carrageenan	WPI or Tween 20/medium-chain	O/W emulsion	(Sala, van Vliet, Cohen Stuart, Aken, & van de
			triglycerides (40% in emulsions)	filled	Velde, 2009)
	Addition of ions	Alginate	No emulsifier/sunflower oil ( $\phi = 0.3$ or	O/W	(Herrero, Ruiz-Capillas, Pintado, Carmona, &
			52%) or chia oil (40%)		Jiménez-Colmenero, 2018; Pintado, Ruiz-Capillas,
					Jimenez-Colmenero, Carmona, & Herrero, 2015;
					Sato, Moraes, & Cunha, 2014)

	Self-assembly	Inulin	Soy lecithin/olive oil (21–38%)	O/W	(Paradiso, Giarnetti, Summo, Pasqualone, Minervini,
	(crystallisation)				& Caponio, 2015)
Polysaccharide/	Self-assembly	Alginate/konjac glucomannan	Egg yolk or Tween 80/rapeseed oil	O/W emulsion	(Yang, Gong, Lu, Li, Sun, & Guo, 2020)
polysaccharide	(compatibility)		(10–60% in emulsions)	filled	
		Xanthan/konjac glucomannan	Tween 80/rapeseed oil (20%)	O/W emulsion	(Yang, Gong, Li, Li, Sun, & Guo, 2019)
				filled	
Protein/	Cold-set after heat	Whey protein isolate/xanthan gum	Span 80 and Tween 60/babacu seed oil	O/W emulsion	(Geremias-Andrade, Souki, Moraes, & Pinho, 2017)
polysaccharide	treatment		(2.8%) and tristearin (1.2%)	filled	
		Soy protein isolate/xanthan gum	Span 80 and Tween 80/tristearin	O/W emulsion	(Brito-Oliveira, Bispo, Moraes, Campanella, &
			(4.5%)	filled	Pinho, 2017)
		Gelatin/Agar	WPI/sunflower oil (40% in emulsions)	O/W emulsion	(Devezeaux de Lavergne, Tournier, Bertrand, Salles,
				filled	van de Velde, & Stieger, 2016)
	Enzyme treatment	Soy protein isolate/sugar beet	Tween 20, SPI, SBP or SPI and	O/W	(Feng, Jia, Zhu, Liu, Li, & Yin, 2019; Hou, Guo,
	(TGase or laccase)	pectin (SBP)	SBP/corn oil (15%) or medium-chain		Wang, & Yang, 2016)
			triglyceride oil (10%)		
	Acidification	Whey protein isolate/carrageenan	WPI/canola oil (50%)	O/W	(Lam & Nickerson, 2014)
	treatment (GDL)				
	Heat treatment and	Gelatin/alginate	No emulsifier/sunflower oil ( $\phi = 0.3$ )	O/W	(Sato, Moraes, & Cunha, 2014)
	addition of ions				

		Whey protein concentrate/Persian	No emulsifier/milk fat (1%)	O/W	(Khalesi, Emadzadeh, Kadkhodaee, & Fang, 2019)
		gum			
	Self-assembly	Egg yolk protein/alginate (pH <	Egg yolk protein/rapeseed oil (30%)	O/W emulsion	(Yang, et al., 2020)
	(electrostatic	pKa of proteins)		filled	
	attraction)				
Organic	Self-assembly	Sapoin glycyrrhizic acid	No emulsifier/sunflower oil, algal oil,	O/W	(Wan, Sun, Ma, Yang, Guo, & Yin, 2017)
compounds			and flaxseed oil (40%)		
		$\beta$ -Sitosterol/ $\gamma$ -Oryzanol	No emulsifier/sunflower oil (40–90%)	W/O	(Bot, den Adel, Regkos, Sawalha, Venema, & Flöter,
					2011; Sawalha, den Adel, Venema, Bot, Floter, &
					van der Linden, 2012)
		Sapoin glycyrrhizic acid	PGPR in W <sub>1</sub> /O and no emulsifier in	W <sub>1</sub> /O/W <sub>2</sub>	(Ma, Wan, & Yang, 2017)
			double emulsions/sunflower oil (70%		
			in W <sub>1</sub> /O)		

## 1208 **Table 2**

# 1209 Selected examples of materials and methods used to prepare emulsion gel particles.

Materials	Methods	Matrix	Emulsifier/oil category and content	Structure	References
Polysaccharide	External gelation (ionic	Alginate	No emulsifier or WPI/canola oil (1–10%),	O/W	(Benavides, Cortes, Parada, & Franco, 2016;
	gelation)		safflower oil (10%), thyme essential oil (1%),		Ching, Bansal, & Bhandari, 2016; Corstens, et
			or D-limonene		al., 2017; Lević, et al., 2015)
		к-Carrageenan or	Tween 80/corn oil (10%) in emulsions	O/W emulsion	(Zhang, et al., 2016)
		alginate		filled	
Protein/polysaccharide	External gelation (ionic	Pectin/WPI	WPI/oily solution of vitamin A (20%)	O/W	(Ruffin, Schmit, Lafitte, Dollat, & Chambin,
	gelation)				2014)
		Alginate/WPI	WPI/sunflower oil (0.5–20%)	O/W	(Feng, Yue, Wusigale, Ni, & Liang, 2018)
		Alginate/lupin	Lupin protein/linseed oil ( $\varphi = 0.15-0.69$ )	O/W	(Piornos, Burgos-Díaz, Morales, Rubilar, &
		protein			Acevedo, 2017)

## 1210 **Table 3**

# 1211 Selected examples of materials and methods used to prepare fluid emulsion gels.

Materials	Methods	Matrix	Emulsifier/oil category and content	Structure	References
Protein	Pickering emulsion/self-	/	Soy glycinin nanoparticles/soy oil ( $\phi = 0.1, 0.3, \text{ or } 0.5$ )	O/W	(Liu & Tang, 2016; Luo, Liu, & Tang, 2013;
	support		or unknown oil ( $\varphi = 0.1-0.89$ )		Xu, Liu, & Tang, 2019)
		/	Pea protein isolate/soy oil ( $\varphi = 0.2-0.6$ )	O/W	(Shao & Tang, 2016)
		/	WPI/camellia oil ( $\phi = 0.75$ )	O/W	(Liu, Gao, McClements, Zhou, Wu, & Zou,
					2019)
		/	Casein peptides/unknown oil (61% and 77%)	O/W	(Wakita & Imura, 2018)
		Sarcoplasmic	Sarcoplasmic protein/canola oil (50%)	O/W	(Hemung, Benjakul, & Yongsawatdigul, 2013)
		protein			
		/	Wheat gluten/corn oil (60%)	Oil-in-glycerol	(Liu, Chen, Guo, Yin, & Yang, 2016)
	Electrospinning	Gelatin	Gelatin/corn oil ( $\varphi = 0.2-0.8$ )	O/W	(Zhang & Zhang, 2018)
	emulsion/self-support				
Polysaccharide	Pickering emulsion/self-	Starch granule	Octenylsuccinate quinoa starch OSQS/corn oil (30-	O/W	(Li, Zhang, Li, Fu, & Huang, 2020)
	support		60%)		
	Disrupted gel systems	Starch	OSA modified starch/sunflower oil (40%)	O/W emulsion	(Torres, Tena, Murray, & Sarkar, 2017)

	(homogenization)			filled	
	Disrupted gel systems	Sodium	Tween 80/canola oil (5%)	O/W	(Soukoulis, Cambier, Hoffmann, & Bohn,
	(mechanical shearing)	alginate			2016)
Protein/polysacc	Pickering emulsion/self-	/	Zein and gum arabic complex (ZGAPs)/medium-chain	O/W	(Dai, Sun, Wei, Mao, & Gao, 2018)
haride	support		triglyceride oil ( $\varphi = 0.1-0.7$ )		
		1	Zein and chitosan complex (ZCCPs)/algal oil (20-70%)	O/W	(Wang, Yin, Wu, Qi, Guo, & Yang, 2016)
		/	Zein and tannic acid complex particles (ZTP)/corn oil ( $\boldsymbol{\phi}$	O/W	(Zou, Guo, Yin, Wang, & Yang, 2015; Zou,
			= 0.5)		Yang, & Scholten, 2018)
		1	$\beta$ -lactoglobulin and gum arabic complex/medium-chain	O/W	(Su, et al., 2020)
			triglyceride oil ( $\varphi = 0.3-0.7$ )		
	Disrupted gel systems	WPI/alginate	WPI/olive oil (5–25%)	O/W	(Leon, Medina, Park, & Aguilera, 2018)
	(mechanical shearing)				
1212					

### 1 Figure legends

- Fig. 1. Structures of two idealized models of emulsion gels: (A) emulsion droplet-filled gels, and (B)
  emulsion droplet-aggregated gels (Dickinson, 2012).
- 4 Fig. 2. Schematic presentation of two kinds of fillers in emulsion gels: (A) active fillers (droplets

5 covered by black line), and (B) inactive fillers (droplets covered by white line).

- 6 Fig. 3. Visual appearances of alginate-based (A) bulk emulsion gels, (B) emulsion gel particles, and
- 7 (C) fluid emulsion gels. Preparing alginate-based emulsion gels includes two steps: first preparing
- 8 emulsions with 1 wt% sodium alginate and 0.5 wt% Tween 80 in water phase and sunflower oil at 40
- 9 wt% and then turning emulsions into gels. For the preparation of bulk emulsion gel, 0.5 wt% CaCl<sub>2</sub>
- 10 was added to the emulsion, and the samples were allowed to gel for 6 h in stand. For the production of
- emulsion gel particles, the emulsion was dropped into a 2 wt% CaCl<sub>2</sub> solution, and the samples were
- 12 allowed to gel in the CaCl<sub>2</sub> solution for 6 h with mild magnetic stirring. For producing fluid emulsion
- 13 gel,  $0.5 \text{ wt\% CaCl}_2$  was added to the emulsion, and the mixture was sheared under constant paddle
- 14 stirring at 600 rpm for 6 h.











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## **Highlights**

- Preparation methods differ according to the emulsion gel types, gelling agents, and purposes. •
- Different emulsion gels have different morphological properties and structure-property • relationships.
- Structures of matrix and emulsion droplets can affect mechanical and release properties of • bulk emulsion gels.
- Uses of emulsion gels as fat replacers and delivery systems were discussed. •

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