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Glucagon-like peptide-1 regulation by food proteins and protein hydrolysates

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Abstract

Glucagon-like peptide-1 (GLP-1) is an enterohormone with a key role in several processes controlling body homeostasis, including glucose homeostasis and food intake regulation. It is secreted by the intestinal cells in response to nutrients, such as glucose, fat and amino acids. In the present review, we analyse the effect of protein on GLP-1 secretion and clearance. We review the literature on the GLP-1 secretory effects of protein and protein hydrolysates, and the mechanisms through which they exert these effects. We also review the studies on protein from different sources that has inhibitory effects on dipeptidyl peptidase-4 (DPP4), the enzyme responsible for GLP-1 inactivation, with particular emphasis on specific sources and treatments, and the gaps there still are in knowledge. There is evidence that the protein source and the hydrolytic processing applied to them can influence the effects on GLP-1 signalling. The gastrointestinal digestion of proteins, for example, significantly changes their effectiveness at modulating this enterohormone secretion in both *in vivo* and *in vitro* studies. Nevertheless, little information is available regarding human studies and more research is required to understand their potential as regulators of glucose homeostasis.

Key words: Enterohormones: Glucagon-like peptide-1: Dietary protein: Hydrolysates: Secretagogues

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Introduction

The gastrointestinal tract is responsible for the digestion and absorption of nutrients, and acts as a barrier against luminal pathogens. Moreover, the gastrointestinal tract cooperates in controlling the metabolism through hormones secreted from enteroendocrine cells, which are the body's largest endocrine organ⁽¹⁾. Enteroendocrine cells are capable of responding to luminal content because their apical side has chemosensing machinery such as taste receptors (TASR), G protein-coupled receptors (GPCR), specific transporters and channels. Their secretory products are stored in characterised secretory vesicles, before being secreted through the basolateral membrane by exocytosis^(2,3). When luminal content moves through the gastrointestinal tract, specific macronutrients stimulate the chemosensing machinery, which leads to the modulation of gut hormone release. Gut hormones exert their effect via vagal nerve or endocrine/paracrine signalling, through the interaction of specific receptors expressed in different tissues of the body. These hormones, which are mainly glucagon-like peptide-1 (GLP-1), cholecystokinin (CCK), peptide YY (PYY), gastric inhibitory polypeptide (GIP) and ghrelin, influence the functioning of the digestive tract, but also modulate insulin secretion from the pancreas, the energy storage of adipose

tissue and neuronal signalling in appetite centres in the brain to mediate the regulation of food intake by terminating hunger and inducing satiety.

Since dietary compounds modulate enterohormone secretion, and given the central role of enterohormones in body homeostasis, such an interaction could have beneficial health implications⁽⁴⁾. In this context, protein and protein hydrolysates are currently being studied to determine their effects on GLP-1 modulation, either through secretion or clearance, which may influence the processes regulated by this hormone such as regulation of glycaemia homeostasis and food intake control. The nutrient-sensing machinery of carbohydrates and lipids is better understood than the detection and pathways followed by protein digestion. The main reasons for this gap in knowledge is the redundant signalling in the gut for the different protein digestion products and the complexity of protein digests⁽⁵⁾. Here we review the literature on this subject in order to determine if the evidence supports differential effects of food proteins on GLP-1 profile. We will introduce the relevance of GLP-1 signalling on health. Then we will focus on the effects on GLP-1 secretion of proteins and its hydrolysates, and the suggested mechanisms. Finally, we will briefly review the use of protein hydrolysates as dipeptidyl peptidase-4 (DPP4) inhibitors. We compile a

Abbreviations: CaSR, Ca-sensing receptor; DPP4, dipeptidyl peptidase-4; GIP, gastric inhibitory polypeptide; GLP-1, glucagon-like peptide-1; HSGH, halibut skin gelatin hydrolysate; OGTT, oral glucose tolerance test; SGLT-1, Na-dependent GLUT-1; T2DM, type 2 diabetes mellitus; PepT1, peptide transporter 1; TSGH, tilapia skin gelatin hydrolysate.

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66 significant number of scientific studies to highlight the impor-
67 tance of the different protein sources, the hydrolysis conditions
68 applied to them, and the resulting digestion products.

69 Relevance of glucagon-like peptide-1 signalling in health

70 There is evidence to suggest that specific enterohormones
71 administered at physiological concentrations can influence the
72 appetite of rodents and human subjects (for a review, see
73 Murphy & Bloom⁽⁶⁾). Likewise, the effects of gut hormones on
74 food intake and body weight have been observed in bariatric
75 surgery (such as Roux-en-Y gastric bypass), which induces a
76 huge increase in GLP-1 and peptide YY (PYY) secretion and
77 is used to treat obesity⁽⁷⁾. Therefore, the modulation of entero-
78 hormone signalling may be an important target in the prevention
79 of obesity and related/associated pathologies. Moreover, endog-
80 enous gut hormones regulate appetite physiologically, unlike
81 the drugs that are currently available, which mainly influence
82 the central neurotransmitter systems. Therefore, gut hormone-
83 based therapies might lead to fewer side effects⁽⁶⁾.

84 Furthermore, modulation of endogenous incretin hormones
85 (GLP-1 and GIP) could be an interesting strategy for preventing
86 and/or managing type 2 diabetes mellitus (T2DM)⁽⁸⁾. T2DM is
87 the most common endocrine disorder, characterised by insulin
88 resistance and impaired insulin secretion, and it is one of the fast-
89 est growing non-communicable diseases in the world⁽⁹⁾. The
90 main goal in the treatment of T2DM is to keep blood glucose lev-
91 els within the normal physiological range. In this regard, GLP-1
92 and GIP are therapeutically interesting peptides because they
93 are important mediators of glycaemic homeostasis, as they are
94 responsible for approximately 50–70 % of the total insulin
95 secreted after glucose intake⁽¹⁰⁾. GLP-1, together with GIP, is
96 responsible for the incretin effect, since it binds to GLP-1 recep-
97 tor in β -cells in the pancreas leading to an increase in intra-
98 cellular Ca and a subsequent insulin secretion in response to glu-
99 cose⁽¹¹⁾. It has also been shown that GLP-1 enhances markers of
100 proliferation and differentiation, and decreases markers of apop-
101 tosis in the pancreas of Zucker diabetic rats^(12,13). Furthermore,
102 GLP-1 improves the glycaemic profile by inhibiting glucagon
103 secretion and improves glucose disposal in peripheral tissues⁽¹⁰⁾.
104 In that way, for patients with T2DM, a non-pharmacological
105 therapeutic approach could be achieved by targeting these
106 incretins (GLP-1 and GIP) through protein- and protein hydroly-
107 sate-based strategies. This approach would be mainly focused
108 on increasing GLP-1 levels rather than stimulating GIP because
109 in these patients the responsiveness of their β -cells to GIP action
110 is decreased⁽¹⁴⁾. Furthermore, only GLP-1 exerts an appetite-sup-
111 pressing effect, while GIP does not seem to do the same⁽¹⁰⁾.
112 Accordingly, many incretin-based therapies focus on using
113 GLP-1 analogues, promoting endogenous GLP-1 secretion or
114 using DPP4 inhibitors.

115 DPP4 is a ubiquitous aminodipeptidase that exists essentially
116 as a membrane-anchored cell-surface enzyme⁽¹⁵⁾. It is expressed
117 throughout the body tissues, such as kidneys, the gastrointestinal
118 tract, liver, pancreas, and the endothelial and epithelial cells on
119 the vascular bed. Its soluble form is found in plasma and there-
120 fore it is in close proximity with hormones circulating in the

blood^(16,17). The main activity of DPP4 is to remove N-terminal
121 dipeptides from polypeptides⁽¹⁸⁾, which preferably have a pro-
122 line or alanine in the second position from the N-terminal.
123 Some of the main DPP4 substrates are GLP-1 and the other incre-
124 tin hormone GIP, which are peptides with N-terminal Tyr-Ala
125 and His-Ala, respectively⁽¹⁹⁾. The intact GLP-1 is rapidly hydro-
126 lysed by DPP4 into a shorter, inactive form, once it reaches the
127 plasma. GLP-1 has a half-life of 1–2 min⁽¹⁸⁾. Only 25 % of the
128 active GLP-1 reaches the portal circulation and subsequently
129 the liver, where a further 40–50 % is digested by the DPP4 in hep-
130 atocytes. This means that only 15 % of the secreted GLP-1 enters
131 the systemic circulation and may reach other tissues, such as the
132 pancreas or the brain⁽²⁰⁾. Therefore, DPP4 is responsible for inac-
133 tivating more than 80 % of the secreted GLP-1⁽¹⁸⁾. Studies focus
134 not only in the development of DPP4-inhibitory drugs, but also
135 on peptides derived from food sources with DPP4-inhibitory
136 capacity. 137

138 Although pharmacological compounds are being studied⁽²¹⁾,
139 natural compounds might be used to prevent the develop-
140 ment of overweight- and obesity-related problems from early
141 preclinical stages through interaction with the enteroendo-
142 crine system⁽²²⁾. 142

143 Dietary regulation of glucagon-like peptide-1 secretion

144 Nutrient ingestion is the primary physiological stimulus for
145 inducing GLP-1 secretion by L cells, located in the ileum and
146 colon in the human gastrointestinal tract. GLP-1 secretion occurs
147 in a biphasic pattern, which consists of a rapid release in 15–
148 30 min after a meal, followed by a second minor peak that occurs
149 in 60–120 min. Enteroendocrine cells have been shown to
150 respond to carbohydrates, lipids and proteins. 150

151 Glucose and fat have been reported to be strong GLP-1-
152 secretagogues after they have been ingested⁽²³⁾, or directly
153 administered into the intestine^(24,25) or into perfused ileal
154 segments⁽²⁶⁾. In the murine model, glucose-stimulated GLP-1
155 release is blocked using Na-dependent GLUT-1 (SGLT-1) knock-
156 out mice and SGLT-1 inhibitors^(27,28), which suggests that glucose
157 metabolism uses glucose transport via SGLT-1 to induce GLP-1
158 secretion. It has also been proposed that sweet taste receptors
159 (T1R2, T1R3) are involved in the glucose-sensing mechanism,
160 but there is still some controversy about whether this is so^(29,30).
161 On the other hand, it has been reported that G-protein-coupled
162 receptors (GPCR) are activated by dietary fat to stimulate GLP-1
163 release, including GPR40 and GPR120 by medium-chain fatty
164 acids, long-chain fatty acids and long-chain unsaturated FA; and
165 GPR41 and GPR43 by SCFA (for reviews, see Hirasawa *et al.*⁽³¹⁾
166 and Reimann⁽³²⁾). 166

167 Other food components could also modulate GLP-1 secre-
168 tion. Flavonoid structures, present in several vegetables, also
169 stimulate GLP-1 secretion⁽³³⁾. In both *ex vivo*⁽³⁴⁾ and rat
170 models⁽³⁵⁾, these compounds have been shown to improve
171 the metabolic status altered by a cafeteria diet treatment⁽³⁶⁾. 171

172 Effects of proteins on glucagon-like peptide-1 secretion

173 Dietary proteins undergo digestion by gastric (pepsin) and pan-
174 creatic (chymotrypsin and trypsin) proteases and membrane 174

175 digestion by peptidases associated with the brush-border mem- 233
 176 brane of enterocytes. The different digestive proteases cleave the 234
 177 peptide bonds at preferential positions. The primary endprod- 235
 178 ucts are dipeptides and tripeptides, which will enter the cell 236
 179 through peptide transporters. Free amino acids are also released 237
 180 after luminal protein digestion and after peptide hydrolysis 238
 181 within the intestinal cells, and then exit across the basolateral 239
 182 membrane via specific amino acid transporters. 240

183 GLP-1 release is activated by luminal intestinal chemosen- 241
 184 sors, which could be reached by peptides of different sizes, 242
 185 mixed with free amino acids. 243

186 Studies in human, animal and enteroendocrine cells have 244
 187 shown increased GLP-1 secretion by free amino acids such as 245
 188 L-phenylalanine, L-alanine and L-glutamine^(37,38) and L-asparagine⁽³⁹⁾. 246
 189 The effect of glutamine has been confirmed in healthy, obese 247
 190 and diabetic human subjects^(40,41). Tolhurst *et al.*⁽⁴²⁾ demon- 248
 191 strated this effect in isolated mouse L cells and reported that 249
 192 the mechanisms were associated with an increase in cyclic AMP 250
 193 (cAMP) and cytosolic Ca²⁺ levels. They also found evidence to 251
 194 suggest that electrogenic Na-coupled amino acid uptake is 252
 195 responsible for initiating membrane depolarisation and voltage 253
 196 gated Ca²⁺, while a second pathway increases intracellular 254
 197 cAMP levels. Young *et al.*⁽⁴³⁾ also reported similar results with 255
 198 L-proline, L-serine, L-alanine, L-glycine, L-histidine, L-cysteine and 256
 199 L-methionine in the STC-1 cell line. 257

200 When analysing the effects of protein on GLP-1 release, many 258
 201 studies focus on the effects of protein hydrolysates, produced by 259
 202 the hydrolysis of food protein with commercial enzymes (sum- 260
 203 marised in Tables 1–3). Sometimes, especially in *in vitro* studies, 261
 204 these are digestive enzymes that simulate intestinal digestion. 262
 205 However, many different hydrolysates are obtained through 263
 206 treatment with enzymes other than pepsin, chymotrypsin or 264
 207 trypsin. Protein hydrolysis can have two main benefits: (1) pro- 265
 208 tein will be more quickly digested after intake; and (2) bioactive 266
 209 peptides^(44–52) might be released. Thus, the degree of protein 267
 210 digestion may impact the capability of protein to stimulate 268
 211 GLP-1 release, as discussed below. 269

212 *In vitro* studies on the STC-1 cell line showed a clear stimu- 270
 213 lation by whole dairy proteins (whey, casein, α -lactalbumin, 271
 214 β -lactoglobulin)^(53–55). Moreover, the stimulation of GLP-1 by 272
 215 whey protein β -lactoglobulin in STC-1 cells was partially lost 273
 216 when treated with trypsin (β -lactoglobulin 7.3-fold increase and 274
 217 hydrolysates 2–5.8-fold increase, all *v. vehicle* control), and 275
 218 totally lost when digested with chymotrypsin for 60 min or 276
 219 more⁽⁵⁴⁾. In the same cell line, the stimulatory effects of whey 277
 220 protein on GLP-1 were lost after extensive hydrolysis with micro- 278
 221 bial (not described) enzymes, or after a simulated gastrointestinal 279
 222 digestion that included a 90-min treatment with pepsin and a 280
 223 150-min treatment with Corolase PP⁽⁵⁶⁾. Another study showed 281
 224 that treating whey or casein with trypsin or DPP4 for 30 min 282
 225 did not lead to any loss of GLP-1-stimulatory properties⁽⁵³⁾. 283

226 In humans, dairy protein is one of the most studied protein 284
 227 sources involving GLP-1 secretion. Intraduodenal infusion of 285
 228 whey protein hydrolysate has stimulated plasma GLP-1 in lean 286
 229 and obese subjects⁽⁵⁷⁾, reduced glucose concentration and sup- 287
 230 pressed energy intake⁽⁵⁸⁾ compared with saline. In these studies, 288
 231 hydrolysed, rather than intact, whey protein was selected 289
 232 because it more closely resembles partially digested protein. 290

233 Also in patients with T2DM, a whey preload increased GLP-1 233
 234 secretion, lowered plasma glucose levels and increased the 234
 235 insulin response^(59,60) compared with water and sucralose, 235
 236 respectively. It has been shown that whey, casein and casein 236
 237 hydrolysates increase GLP-1 secretion^(61–63). However, there is 237
 238 no agreement about whether there are any differences between 238
 239 their effect on GLP-1 secretion. Hall *et al.*⁽⁶²⁾ showed that 120 min 239
 240 after being ingested, whey protein induced a 2-fold increase in 240
 241 postprandial GLP-1 levels compared with casein protein. On the 241
 242 other hand, when comparing whey, casein and their hydroly- 242
 243 sates, Calbet & Holst⁽⁶³⁾ showed that the release of GLP-1 was 243
 244 not influenced by the source or hydrolysis process. Also, a com- 244
 245 mercially available whey protein hydrolysate showed a higher 245
 246 GLP-1 release 30 min after an oral glucose tolerance test 246
 247 (OGTT) than did casein glycomacropeptide (CGMP), but 247
 248 not compared with whey isolate or α -lactalbumin-enriched 248
 249 whey⁽⁶⁴⁾ (incremental AUC_{30min} median; 593 (hydrolysate), 249
 250 270 (CGMP); $P=0.045$). Thus, the studies performed with whey 250
 251 and whey hydrolysates do not show any differences in the effects 251
 252 of the two sources in terms of GLP-1 secretion. Calbet & Holst⁽⁶³⁾ 252
 253 suggested that this is because the dairy protein hydrolyses rap- 253
 254 idly in the intestine and there is a subsequent rise in peripheral 254
 255 amino acids independent of the fractionation. 255

256 Other protein sources have also been shown to stimulate 256
 257 GLP-1 release *in vivo*. A similar rise in rat plasma GLP-1 levels, 257
 258 comparable with that caused by dairy protein, has been 258
 259 observed after pea-protein meals⁽⁶⁵⁾. Furthermore, also in rats, 259
 260 pea protein and pea-protein hydrolysate have been shown to 260
 261 similarly stimulate GLP-1 release, although the hydrolysate 261
 262 showed stronger eating-inhibitory properties⁽⁶⁶⁾ (total energy 262
 263 intake: 63 \pm 6 kJ, 46 \pm 3 kJ, 67 \pm 5 kJ after pea protein, the hydroly- 263
 264 sate and the control, respectively). *In vitro* studies with STC-1 264
 265 cells showed that intact pea protein increases GLP-1 release. 265
 266 On the other hand, various pea-protein hydrolysates obtained 266
 267 by enzymic hydrolysis with subtilisin were tested, and only 267
 268 one of them maintained its GLP-1-secretory capacity⁽⁵³⁾. 268

269 Cereal protein has also been shown to stimulate GLP-1. Maize 269
 270 protein zein (a major maize protein) hydrolysate attenuated gly- 270
 271 caemia in rats under the intraperitoneal glucose tolerance test, 271
 272 associated with enhanced secretions of GLP-1 and GIP⁽⁶⁷⁾ com- 272
 273 pared with water. *In vitro* (GLUTag cells), zein hydrolysate was 273
 274 shown to stimulate GLP-1 release more than egg albumin, coun- 274
 275 try bean and meat hydrolysates⁽⁶⁸⁾. However, the type of 275
 276 hydrolysis was different in the various sources, so the effect of 276
 277 the protein source *per se* cannot be concluded from this paper. 277
 278 The stimulation of GLP-1 secretion by maize zein hydrolysate in 278
 279 GLUTag cells is not affected by treatment with pepsin/pancreatin 279
 280 for 60 min, although it is reduced after pronase treatment⁽⁶⁷⁾ 280
 281 compared with the positive control, KCl 70 mM. The authors sug- 281
 282 gested that the hydrolysate is not further cleaved by pepsin treat- 282
 283 ment (the degree of hydrolysis was only 8.6 %). 283

284 Oral administration of rice protein hydrolysates also 284
 285 increased total GLP-1 in plasma, and improved glycaemic 285
 286 response in rats⁽⁶⁹⁾ (the control used was 2 g/kg of glucose sol- 286
 287 ution). In the same study, rice protein hydrolysates (degree of 287
 288 hydrolysis 5–10 %) stimulated GLP-1 in GLUTag cells, with the 288
 289 potency depending on the enzyme and the time of digestion⁽⁶⁹⁾ 289
 290 compared with the blank treatment. The effect of the whole rice 290

Table 1. Stimulation of glucagon-like peptide-1 (GLP-1) secretion by protein and protein hydrolysates in humans

Protein	Hydrolysis conditions	Subjects	n*	Protein dose	Secretion	Increment v.	Reference
Turkey	Intact protein	Healthy subjects	8	Ingestion of 352 g	↑	Fat isoenergetic meal	(23)
Whey	N.D.	Obese and lean men	12	Intraduodenal infusion of 24 g	↑	Saline	(57)
Whey	N.D.	Healthy men	16	Intraduodenal infusion of 48 g	↑	Saline	(58)
Whey	N.D.	Healthy men	16	Intraduodenal infusion 8 g	↑	Saline	(58)
Whey	N.D.	Healthy men	16	Intraduodenal infusion 24 g	↑	Saline	(58)
Whey	N.D.	Healthy men	16	Intraduodenal infusion 48 g	↑	Saline	(58)
Whey	Intact protein	T2DM subjects of both sexes	21	Ingestion of 17 g	↑	Sucralose	(59)
Whey	Intact protein	T2DM subjects of both sexes	15	Ingestion of 50 g	↑	Water	(60)
Casein	Intact protein	Overweight to obese men and women	24	Ingestion of 30 g	↑	Time 0	(61)
Whey	N.D.	Healthy men and women	9	Ingestion of 48 g	↑	Casein	(62)
Whey	Alcalase/53 °C/pH 8.0/–†	Healthy men	6	Stomach infusion of 36 g	↑	Time 0	(63)
Whey	+ Neutrased/53 °C/pH 7.0/–†	Healthy men	6	Stomach infusion of 36 g	↑	Time 0	(63)
Casein	Intact protein	Healthy men and women	9	Ingestion of 48 g	↑	Casein	(62)
Casein	Alcalase/53 °C/pH 8.0/–†	Healthy men	6	Stomach infusion of 36 g	↑	Time 0	(63)
Casein	+ Neutrased/53 °C/pH 7.0/–†	Healthy men	6	Stomach infusion of 36 g	↑	Time 0	(63)
Whey	Intact protein	Healthy men and women	9	Ingestion of 48 g	↑	Casein	(62)
Whey	N.D.	T2DM subjects of both sexes	11	Ingestion of 45 g	↑	CGMP-enhanced whey	(64)

↑, GLP-1 secretion is incremented v. the control, specified in each row; N.D., hydrolysis conditions not described; T2DM, type 2 diabetes mellitus; CGMP, casein glycomacropeptide.

* Number of subjects per group.

† Time not known.

AQ5



Table 2. Stimulation of glucagon-like peptide-1 (GLP-1) secretion by protein and protein hydrolysates *in vitro**

Protein	Hydrolysis conditions	Cell line	Protein treatment	Secretion	Increment v.	Reference
Egg albumin	N.D.	STC-1	2 h	↑	KRB	(26)
Meat			2.5–20 mg/ml			
Meat	N.D.	Small-intestinal cultures	2 h	↑	Saline with 0.1 % BSA	(37)
			5.0–50 mg/ml			
			2 h	–		
			0.5 mg/ml			
Milk	N.D.	Small-intestinal cultures	2 h	↑		
Vegetables			5.0 mg/ml			
Casein	Intact protein	STC-1	2 h	↑	Hanks' buffered salt solution	(53)
Codfish			1.0 mg/ml			
Egg						
Pea (DPS†)						
Wheat						
Whey						
Ovomucoid	Intact protein	STC-1	2 h	–		
Pea (Pisane†)			1.0 mg/ml			
Pea (SM†)						
Soyabean						
Casein hydrolysate	N.D.	STC-1	2 h	–		
			1.0 mg/ml			
Egg	N.D.	STC-1	2 h	↑		
			1.0 mg/ml			
Pea	Subtilisin/–†/pH 8.0/–†	STC-1	2 h	–		
	+ PSE/–†/pH 6.0/–†		1.0 mg/ml§			
Pea (HP90†)	N.D.	STC-1	2 h	–		
Wheat			1.0 mg/ml			
Yoghurt whey	Intact protein	STC-1 pGIP/neo	3 h	↑	HEPES	(54)
	Intact protein	STC-1 pGIP/neo	3 h	↓		
			50–100 mg/ml			
Cheese whey	Intact protein	STC-1 pGIP/neo	3 h	–		
			5.0–10, 100 mg/ml			
			3 h	↑		
			25–50 mg/ml			
β-Lactoglobulin	Intact protein	STC-1 pGIP/neo	3 h	↑		
			0.63–10 mg/ml			
			3 h	–		
			0.31 mg/ml			
	Chymotrypsin/37 °C/pH 7.4/30 min	STC-1 pGIP/neo	3 h	↑		
	Trypsin/37 °C/pH 7.4/30–150 min		10 mg/ml			
	Chymotrypsin/37 °C/pH 7.4/60–150 min	STC-1 pGIP/neo	3 h	–		
			10 mg/ml			
α-Lactalbumin	Intact protein	STC-1 pGIP/neo	3 h	–		
			0.31–0.63 mg/ml			
			3 h	↑		
			1.3–10 mg/ml			

Regulation of glucagon-like peptide-1

Table 2. (Continued)

Protein	Hydrolysis conditions	Cell line	Protein treatment	Secretion	Increment v.	Reference
Casein	Intact protein	STC-1 pGIP/neo	3 h 0.31–10 mg/ml	↑	HEPES	(55)
α-Casein	Intact protein	STC-1 pGIP/neo	3 h	↑		
β-Casein	Intact protein	STC-1 pGIP/neo	0.16–5.0 mg/ml	–		
κ-Casein	Intact protein	STC-1 pGIP/neo	3 h 0.16–0.31 mg/ml	–		
			3 h 0.63–5.0 mg/ml	↑		
α-Casein	Chymotrypsin/37 °C/pH 7.4/30–150 min	STC-1 pGIP/neo	3 h 5.0 mg/ml	–		
	Trypsin/37 °C/pH 7.4/30–150 min	STC-1 pGIP/neo	3 h	↑		
	Pepsin/37 °C/pH 2.3/30–150 min	STC-1 pGIP/neo	5.0 mg/ml	–		
β-Casein	Chymotrypsin/37 °C/pH 7.4/30–150 min	STC-1 pGIP/neo	3 h	–		
	Trypsin/37 °C/pH 7.4/30–150 min	STC-1 pGIP/neo	5.0 mg/ml	–		
	Pepsin/37 °C/pH 2.3/30–150 min	STC-1 pGIP/neo	3 h 5.0 mg/ml	↑		
Whey	Intact protein	STC-1	4 h 10.0 mg/ml	↑	KRB with 10 mm-glucose	(56)
	Pepsin/37 °C/pH 2/90 min	STC-1	4 h 10 mg/ml	–		
	+ Corolase PP/37 °C/pH 7.5/150 min	STC-1	4 h	–		
Whey DH32	N.D.	STC-1	4 h	–		
	Pepsin/37 °C/pH 2/90 min	STC-1	10 mg/ml	–		
	+ Corolase PP/37 °C/pH 7.5/150 min	STC-1	4 h	–		
Whey DH45	N.D.	STC-1	4 h	–		
	Pepsin/37 °C/pH 2/90 min	STC-1	10 mg/ml	–		
	+ Corolase PP/37 °C/pH 7.5/150 min	STC-1	4 h	–		
Maize zein	Papain/55 °C/pH 7.2/60 min	GLUtag	1 h	↑	HEPES	(67)
	Papain/55 °C/pH 7.2/60 min	GLUtag	10 mg/ml	–		
	+ Pepsin/37 °C/pH 1.85/60 min	GLUtag	1 h	–		
	+ Pancreatin + trypsin/37 °C/pH 8.2/120 min	GLUtag	1 h	–		
	Papain/55 °C/pH 7.2	GLUtag	2.0 mg/ml	–		
	+ Pronase/37 °C/pH 7.0	GLUtag	1 h	↑		
Maize zein	Papain/55 °C/pH 7.0/60 min	GLUtag	5.0–20 mg/ml	–	HEPES	(68)
		GLUtag	1 h	–		
Egg albumin	N.D.	GLUtag	5.0–20 mg/ml	–		
BSA	Intact protein	GLUtag	1 h	–		
Meat	N.D.	GLUtag	5.0 mg/ml	–		
Bean	Pepsin/37C/pH 1.9/10 min	GLUtag	5.0 mg/ml	–		



Table 2. (Continued)

Protein	Hydrolysis conditions	Cell line	Protein treatment	Secretion	Increment v.	Reference	
Rice endosperm	Papain/55 °C/pH 7.2/60 min	GLUTag	1 h	↑	HEPES	(69)	
	Pepsin/37 °C/pH 1.85/30 min Pepsin/37 °C/pH 1.85/60 min	GLUTag	10 mg/ml 1 h	–			
Rice bran	Papain/55 °C/pH 7.2/60 min	GLUTag	1 h	–			
	Pepsin/37 °C/pH 1.85/30 min Pepsin/37 °C/pH 1.85/60 min	GLUTag	10 mg/ml 1 h	↑			
Wheat (770 Da fraction)	N.D.	GLUTag	2 h	–	Saline	(70)	
			0.1–0.25 mg/ml 2 h	↑			
Wheat (7740 Da fraction)	N.D.	GLUTag	0.5–1.0 mg/ml 2 h	–			
			1.0 mg/ml				
Wheat gluten	N.D.	GLUTag	1 h	–	HEPES	(71)	
			5 mg/ml 1 h	↑			
α-Lactalbumin	N.D.	GLUTag	10 mg/ml 1 h	–			
			5 mg/ml 1 h	↑			
Wheat gluten	N.D. + Pepsin/37 °C/pH 1.85/30–60 min	GLUTag	10 mg/ml 1 h	↑			
			10 mg/ml				
α-Lactalbumin	N.D. + Pancreatin/37 °C/pH 8.2/60–120 min	GLUTag	1 h	↑			
			10 mg/ml				
Cuttlefish viscera	N.D. + Pepsin/37 °C/pH 1.85/30–60 min	GLUTag	10 mg/ml				
Cuttlefish viscera	Pancreatin/37 °C/pH 8.2/60–120 min Intact protein + Salivary fluid H ₁₁ /50 °C/pH 8.0/4 h + Salivary fluid H ₁₁ /50 °C/pH 8.0/4 h + Salivary fluid H ₁₁ /50 °C/pH 8.0/4 h + Salivary fluid H ₁₁ /50 °C/pH 8.0/4 h + Salivary fluid Pepsin/37 °C/pH 2.5–3/120 min	STC-1	2 h 13 mg/ml	–	Baseline	(72)	
		STC-1	2 h 13 mg/ml	↑	UCVP + salivary fluid		
		STC-1	2 h 13 mg/ml	–	UCVP + salivary fluid		
		STC-1	2 h 13 mg/ml	–	UCVP + IVD		

Regulation of glucagon-like peptide-1

Table 2. (Continued)

Protein	Hydrolysis conditions	Cell line	Protein treatment	Secretion	Increment v.	Reference
	H \parallel /50 °C/pH 8.0/4 h + Salivary fluid + Pepsin/37 °C/pH 2.5–3/120 min H \parallel /50 °C/pH 8.0/4 h + Salivary fluid + Pepsin/37 °C/pH 2.5–3/120 min + Pancreatin/37 °C/pH 7.0/120 min H \parallel /50 °C/pH 8.0/4 h + Salivary fluid + Pepsin/37 °C/pH 2.5–3/120 min + Pancreatin/37 °C/pH 7.0/120 min	STC-1	2 h 13 mg/ml	↓	UCVP + IVD	
	Intact protein + Salivary fluid	STC-1	2 h 13 mg/ml	–	UCVP + IVD	
Bovine Hb	Salivary fluid + Pepsin/37 °C/pH 2.5–3.0/60–120 min Salivary fluid + Pepsin/37 °C/pH 2.5–3.0/120 min + Pancreatin/37 °C/pH 7.0/30–120 min Salivary fluid + Pepsin/37 °C/pH 2.5–3.0/120 min + Pancreatin/37 °C/pH 7.0/120 min	STC-1	2 h 13 mg/ml	↑	HEPES	(73)
Bovine Hb	Salivary fluid + Pepsin/37 °C/pH 2.5–3.0/120 min + Pancreatin/37 °C/pH 7.0/30–120 min Salivary fluid + Pepsin/37 °C/pH 2.5–3.0/120 min + Pancreatin/37 °C/pH 7.0/120 min	STC-1	2 h 13 mg/ml	↑	HEPES	
Meat	N.D.	GLUTag	2 h 1.0–50 mg/ml	↑	Baseline	(74)

Table 2. (Continued)

Protein	Hydrolysis conditions	Cell line	Protein treatment	Secretion	Increment v.	Reference
Salmon skin gelatin	Alcalase/50 °C/pH 7.0/4 h	GLUTag	2 h	↑	Glucose 2 mM	(76)
	Alcalase + Flavourzyme/50 °C/pH 7.0/4 h		2.5 mg/ml			
	Promod/50 °C/pH 7.0/4 h	GLUTag	2 h	–		
			2.5 mg/ml			
Salmon trimmings	Alcalase + Flavourzyme/50 °C/pH 7.0/4 h + Pepsin/37 °C/pH 2.0/90 min + Corolase PP/37 °C/pH 7.0/150 min	GLUTag	2 h	–		
	Alcalase/50 °C/pH 7.0/4 h		2.5 mg/ml			
	Alcalase + Flavourzyme/50 °C/pH 7.0/4 h	GLUTag	2 h	↑		
			2.5 mg/ml			
	Promod/50 °C/pH 7.0/4 h	GLUTag	2 h	↓		
			2.5 mg/ml			
Meat	Alcalase + Flavourzyme/50 °C/pH 7.0/4 h + Pepsin/37 °C/pH 2.0/90 min + Corolase PP/37 °C/pH 7.0/150 min	GLUTag	2 h	↑	KRB with 0.2 % BSA	(82)
	N.D.	NCI-H716	20 mg/ml			
Chicken feet	Neutrase/25 °C/pH 7.0/24 h	STC-1	2 h	↑	HEPES	(99)
			5 mg/ml			
		Ileum explants	1 h	↑	KRB with 10 mM-glucose	
			15 mg/ml			

↑ GLP-1 secretion is incremented v. the control, specified in each row; –, GLP-1 secretion is not altered v. the control, specified in each row; ↓ GLP-1 secretion is reduced v. the control, specified in each row; BSA, bovine serum albumin; Corolase PP, a porcine pancreatic enzyme preparation; DH32, 32 % degree of hydrolysis; DH45, 45 % degree of hydrolysis; DPS, Dutch Protein Services; H, hydrolysis; HEPES, 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid; IVD, *in vitro* digestion with pepsin and pancreatin, always indicates the same hydrolysis conditions as the protein that is compared with; KRB, Krebs–Ringer modified buffer; N.D., hydrolysis conditions not described; PSE, proline-specific endoprotease; UCVP, undigested cuttlefish viscera protein.

* The salivary fluid does not contain enzymes.

† Pea protein origin: DPS, from Dutch Protein Services; Pisane, from Cosucra; SM, from Nutralys; HP90, from Triballat.

AQ6 ‡ Temperature or time not known.

AQ7 § This pea hydrolysate did not stimulate GLP-1 secretion; nor did the 10 kDa permeate. Nevertheless, the supernatant fraction obtained after centrifugation increased GLP-1 secretion compared with the control.

|| Hydrolysis with cuttlefish hepatopancreas digestive proteases.

¶ Hydrolysis with cuttlefish smooth hound intestine digestive proteases.

Table 3. Stimulation of glucagon-like peptide-1 (GLP-1) secretion by protein and protein hydrolysates in animals

Protein	Hydrolysis conditions	Species	n*	Protein dose	Secretion	Increment v.	Reference
Egg albumin	N.D.	Wistar male rats	7–9	Jejuno-ileum administration of 25 mg/ml Jejuno-ileum administration of 50 mg/ml Colon administration of 25 mg/ml Colon administration of 50 mg/ml	↑	Saline	(26)
Salmon skin gelatin	Flavourzyme/50 °C/pH 7.0/4 h	Sprague–Dawley male rats†	12	5 weeks Oral administration 300 mg/d	↑	Water	(47)
Porcine skin gelatin	Flavourzyme/50 °C/pH 7.0/6 h	Sprague–Dawley male rats	12	6 weeks Oral administration 300 mg/d	–	Water	(48)
	Flavourzyme/50 °C/pH 7.0/6 h	Sprague–Dawley male rats†		6 weeks Oral administration 300 mg/d	↑		
Halibut skin gelatin	Flavourzyme/50 °C/pH 7.0/4 h	Sprague–Dawley male rats†	11	4 weeks 750 mg/kg/d	↑	Water	(49)
Tilapia skin gelatin	Flavourzyme/50 °C/pH 7.0/6 h	Sprague–Dawley male rats		4 weeks 750 mg/kg/d	–		
Whey	Intact protein	SPF Wistar male rats	9	Oral administration of about 3 g/kg BW	–	Sucrose	(65)
Pea	Intact protein	Sprague–Dawley male rats	10	Intragastric infusion of 136 mg/ml	–	Saline	(66)
Pea	N.D.						
Maize zein	Papain/55 °C/pH 7.2/60 min	Sprague–Dawley male rats	7–10	Oral administration§ of 2 g/kg BW	↑	Water	(67)
Meat	N.D.	Sprague–Dawley male rats	7–10	Oral administration§ of 2 g/kg BW	–		
Maize zein	Papain/55 °C/pH 7.2/60 min	Goto-Kakizaki male rats	6–7	Oral administration‡ of 2 g/kg BW	↑		
Whey	Papain/55 °C/pH 7.2/60 min	Goto-Kakizaki male rats	6–7	Oral administration‡ of 2 g/kg BW	–		
Maize zein	Papain/55 °C/pH 7.0/60 min	Sprague–Dawley male rats	6–9	Duodenal administration of 100–250 mg/ml Ileal administration of 100 mg/ml Ileal administration of 250 mg/ml Duodenal loop administration of 300 mg/ml Jejunal loop administration of 300 mg/ml Ileal loop administration of 300 mg/ml	↑ – ↑ –	Water Time 0	(68)
Rice endosperm	Pepsin/37 °C/pH 1.85/30 min	Sprague–Dawley male rats	4–6	Oral administration of 2 g/kg BW Oral administration‡ of 0.1–1.0 g/kg BW Oral administration‡ of 2 g/kg BW Oral administration§ of 1–2 g/kg BW	↑ – ↑ ↑	Water	(69)
Rice bran	Pepsin/37 °C/pH 1.85/30 min	Sprague–Dawley male rats	4–6	Oral administration of 2 g/kg BW Oral administration‡ of 0.1–1.0 g/kg BW Oral administration‡ of 2 g/kg BW	↑ – ↑		
Wheat (770 Da fraction)	N.D.	Sprague–Dawley male rats	8	Oral administration§ of 2 g/kg BW	↑	Saline	(70)
Wheat gluten	N.D.	Wistar/ST male rats	5–7	Oral administration of 1 g/kg BW	–	Water	(71)
α-Lactalbumin							
Meat	N.D.	Wistar male rats	6	Duodenal infusion of 50 mg/ml	↑	Baseline	(75)
Lysozyme	Alcalase/60 °C/pH 8.0/6 h	ZDF male rats	9	Oral administration of 1 g/kg BW	–	Untreated rats	(100)
Maize zein	Papain/55 °C/pH 7.0/60 min	Sprague–Dawley male rats	6–8	Ileal administration§ of 250 mg/ml	↑	Water	(101)
Meat	N.D.						

N.D., hydrolysis conditions not described; ↑, GLP-1 secretion is incremented v. the control, specified in each row; –, GLP-1 secretion is not altered v. the control, specified in each row; SPF, specific pathogen-free; BW, body weight; ZDF, Zucker diabetic fatty.

* Number of animals per group.

† Sprague–Dawley streptozotocin-induced diabetic rats.

‡ Changes in plasma GLP-1 after oral administration of the protein under the oral glucose tolerance test.

§ Changes in plasma GLP-1 after oral administration of the protein under the intraperitoneal glucose tolerance test.

291 protein was not assessed. The authors found that GLP-1 secre- 347
 292 tion was weaker after 60 min digests with pepsin in rice endo- 348
 293 sperm protein hydrolysates than after 30 min digests, which 349
 294 suggests that oligo- or larger peptides, rather than small peptides 350
 295 or free amino acids, might be responsible for this stimulation. 351
 296 The results for wheat protein were just the opposite. In GLUTag 352
 297 cells, a low-molecular fraction of wheat protein hydrolysate 353
 298 enhanced GLP-1 secretion while a high-molecular fraction did 354
 299 not⁽⁷⁰⁾. The low-molecular fraction of wheat protein hydrolysate 355
 300 had a glucose-lowering effect mediated by GLP-1 in rats⁽⁷⁰⁾ after 356
 301 an oral administration compared with 0.9 % NaCl. Also, in 357
 302 another study in a distal enteroendocrine cell model (GLUTag 358
 303 cells), the effect of wheat hydrolysate on the stimulation of 359
 304 GLP-1 secretion was largely enhanced by pepsin/pancreatin 360
 305 digestion relative to the blank⁽⁷¹⁾. 361

306 For other protein sources, *in vitro* studies also showed that 362
 307 GLP-1-secreting activity of digested protein was greater than that 363
 308 of the original source. In a study performed with cuttlefish (*Sepia* 364
 309 *officinalis*) viscera, a hydrolysate (obtained from digestion with 365
 310 cuttlefish hepato-pancreatic enzymes) was found to exert GLP-1- 366
 311 secreting action while the undigested protein did not⁽⁷²⁾. These 367
 312 results were found with the samples solubilised in saliva, but 368
 313 they were subjected to further *in vitro* simulated gastrointestinal 369
 314 digestion (including treatment with pepsin and pancreatin). 370
 315 Results showed that gastrointestinal digestion increased the 371
 316 GLP-1-secretory effects of both the hydrolysate and the initially 372
 317 undigested protein, leading to no differences between the 373
 318 hydrolysate and the non-hydrolysate gastrointestinally digested 374
 319 samples. Also, intestinal digested bovine Hb protein had a 375
 320 greater effect on GLP-1 release than partially digested protein 376
 321 (saliva and gastric digest) in STC-1 cells⁽⁷³⁾. 377

322 Taken together, all these studies prove that several protein 378
 323 sources increase GLP-1 secretion, which is associated to benefits 379
 324 such as food intake or glucose homeostasis regulation. *In vivo* 380
 325 studies do not fully clarify whether previous hydrolysis of the 381
 326 protein sources with commercial enzymes leads to stronger 382
 327 GLP-1-secreting effects. *In vitro* data show that many protein 383
 328 sources, including purified proteins, activate GLP-1 release. 384
 329 However, digestion as it might physiologically happen upon 385
 330 protein intake might stimulate or reduce the effect of the undi- 386
 331 gested protein, depending on the original source. This suggests 387
 332 that some high-molecular-weight peptides might reach enter- 388
 333 oendocrine cells and activate GLP-1 secretion, while in other 389
 334 cases the lower-molecular-weight peptides or the amino acids 390
 335 released after digestion are responsible for the secretion. 391

336 *Mechanisms involved in the effects of protein as* 337 *glucagon-like peptide-1 secretagogue*

338 The mechanisms through which the proteins and peptides 396
 339 released after protein hydrolysis (either 'synthetic' or simulated 397
 340 digestion) act as secretagogues are still not fully understood, but 398
 341 several pathways have been shown to be involved. Studies on 399
 342 the mechanisms through which protein and protein hydrolysates 400
 343 stimulate GLP-1 secretion are carried out using *in vitro* (i.e. enter- 401
 344 oendocrine cell lines such as STC-1 and GLUTag) and *ex vivo* 402
 345 (i.e. perfused intestine and intestinal explants) models, and also 403
 346 primary cultures. 404

Many of the studies that focus on the mechanisms that stimu- 347
 late GLP-1 secretion use commercial meat peptones, that is meat 348
 hydrolysates produced by the digestion of meat with proteolytic 349
 enzymes which lead to a complex mixture of partially metabol- 350
 ised proteins. 351

With this protein source, it seems that one key player in the 352
 oligopeptide stimulation of GLP-1 release is peptide transporter 353
 1 (PepT1) (Fig. 1). Meat peptone was shown to stimulate GLP-1 354
 secretion in mouse colonic primary culture through PepT1-de- 355
 pendent uptake, followed by an increase in intracellular Ca, 356
 and activation of Ca-sensing receptor (CaSR)⁽⁷⁴⁾. Very recently 357
 Modvig *et al.*⁽⁷⁵⁾ used isolated perfused rat small intestine to 358
 study GLP-1 secretion stimulated by meat peptone. The sensory 359
 mechanisms underlying the response depended on di-/tripep- 360
 tide uptake through PepT1 and subsequent basolateral activa- 361
 tion of the amino acid-sensing receptor (CaSR) (Fig. 2). CaSR 362
 might also be activated by free amino acids taken up from the 363
 intestinal lumen by different amino acid transporters⁽⁷⁵⁾. 364

It has been pointed out that it is difficult to determine the 365
 PepT1-dependent oligopeptide-sensing pathway in GLUTag 366
 and STC-1 cell lines, because the expression of endogenous 367
 PepT1 is lower than in native L cells⁽⁷⁴⁾. Therefore, the effects 368
 of peptones observed in both cell lines may be due to the free 369
 amino acids that some of these peptones contain, as has been 370
 suggested in an *in vitro* study on the effects of salmon hydroly- 371
 sate⁽⁷⁶⁾ carried out in GLUTag cells. However, other studies on 372
 these cell lines do not share this view. As mentioned above, 373
 GLP-1 secretion is activated by dairy proteins⁽⁵³⁻⁵⁵⁾, low-molecu- 374
 lar-weight wheat (with less than 1 % free amino acids)⁽⁷⁰⁾, intact 375
 pea-protein⁽⁵³⁾ or peptin-resistant zein hydrolysate⁽⁶⁷⁾. Furthermore, 376
 three synthetic peptide sequences (ANVST, TKAVEH and KAAT) 377
 were reported to be able to enhance GLP-1 secretion in 378
 STC-1 cells⁽⁷⁷⁾. The authors concluded that the incretin effect 379
 of proteins is associated with the amino acid profile, but the 380
 specific amino acid motif that triggers GLP-1 secretion stimu- 381
 lation was not determined. Thus, receptor or peptide trans- 382
 porters other than PepT1 expressed in STC-1 and GLUTag 383
 cells might be involved in the peptide stimulation of GLP-1. 384
 For instance, one of the mediators suggested was the G protein- 385
 coupled receptor family C group 6 subtype A (GPCR6A)⁽⁷⁰⁾ 386
 (Fig. 3). 387

Protein hydrolysates are also detected by the umami receptor 388
 (T1R1-T1R3 heterodimer)⁽⁷⁸⁾ (Fig. 4) and G protein-coupled 389
 receptor 92/93 (GPR92/93)⁽⁷⁹⁾, which leads to the release of 390
 the gut-derived satiety factor cholecystokinin. There is no direct 391
 evidence of umami stimulation and GLP-1 secretion, but the 392
 T1R1 receptors were co-expressed with GLP-1-expressing 393
 STC-1 cells⁽⁸⁰⁾, which suggests that umami receptors play a role 394
 in GLP-1 signalling. 395

An increase in intracellular Ca has been reported to be a path- 396
 way activated by protein hydrolysates to mediate GLP-1 secre- 397
 tion. Pais *et al.*⁽³⁷⁾ reported that meat peptone-stimulated GLP-1 398
 secretion from primary L cells was also associated with Ca 399
 influx through voltage gate Ca channels (Fig. 3). In NCI-H716 400
 human enteroendocrine cells, tetrapeptides, but not single 401
 amino acids or any of the dipeptides, tripeptides and pentapep- 402
 tides tested, were found to induce a robust and selective [Ca²⁺]_i 403
 response associated with increased secretion of GLP-1⁽⁸¹⁾. 404

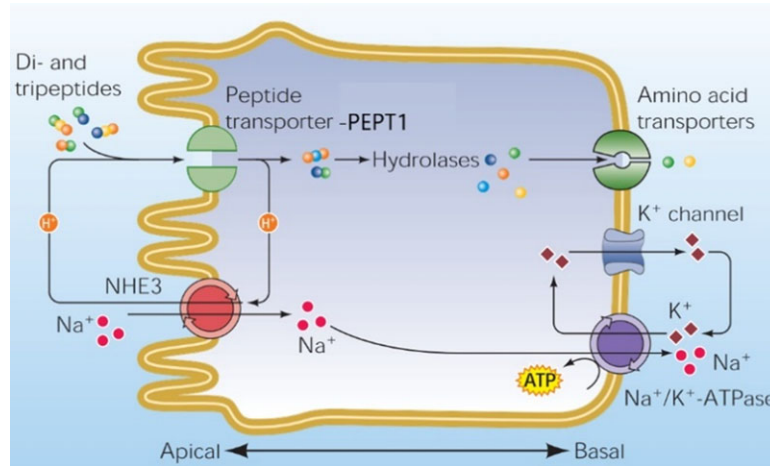
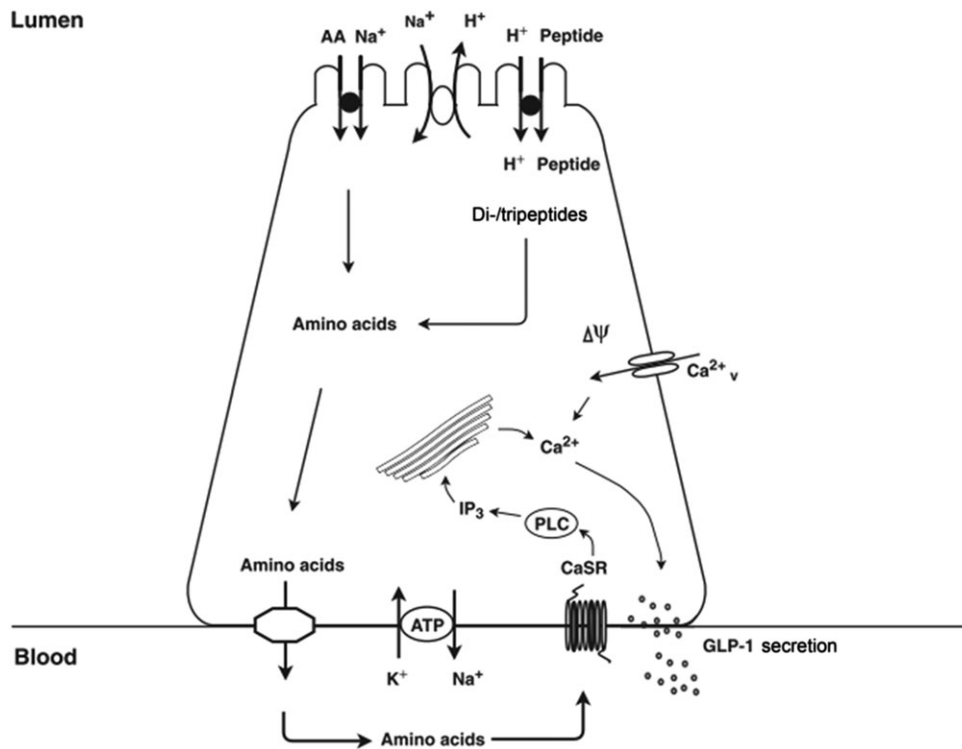


Fig. 1. The intestinal transporter form PEPT1 (SLC15A1) is located in apical membranes with a functional coupling to the apical Na⁺/H⁺ antiporter (NHE3) for pH recovery from the peptide-transport-induced intracellular acid load. Adapted from Daniel *et al.*⁽¹⁰³⁾.



AQ8 Fig. 2. Illustration of the endocrine L cell and the proposed mechanisms by which peptide stimulates glucagon-like peptide-1 (GLP-1) release. Di-/tripeptides are taken up by PepT1 and are degraded by cytosolic peptidases to their respective amino acids (AA). Intracellular amino acids are then transported to the interstitial side through basolateral amino acid transporters, wherefrom they stimulate the L cells by activating amino acid sensors, like calcium-sensing receptor (CaSR), situated on the basolateral membrane. IP₃, inositol trisphosphate; PLC, phospholipase C. Adapted from Modvig *et al.*⁽⁷⁵⁾.

405 Moreover, these effects were not observed in either STC-1 or in
 406 GLUTag rodent cells. Interestingly, in the same paper, the
 407 authors showed that casein protein hydrolysate elicited an
 408 increase in GLP-1 without modulating intracellular Ca.

409 It has been suggested that GLP-1 secretion is mediated by
 410 other intracellular pathways such as extracellular signal-
 411 regulated kinase 1/2 (ERK1/2), mitogen-activated protein kinase
 412 (MAPK) and p38 MAPK, activated by peptones and mixtures of
 413 essential amino acids in NCI-H716 cells⁽⁸²⁾.

414 Altogether, the studies show that which signalling path-
 415 ways are involved in GLP-1 secretion by different peptide mix-
 416 tures will depend on the peptide length, the sequences and/or
 417 the amino acid composition, and whether there are free amino
 418 acids in the mixture. Furthermore, the model studied has to be
 419 carefully considered since there are differences in the expres-
 420 sion of key genes (such as pepT-1) and some effects might
 421 depend on the vectoriality of the system (the capacity to differ-
 422 entiate basolateral and apical processes).

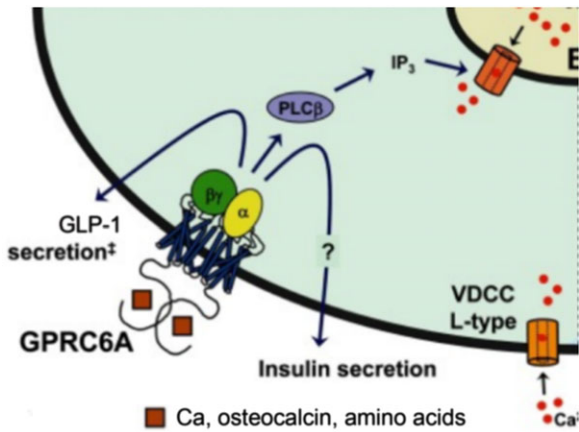


Fig. 3. Signalling through G protein-coupled receptor family C group 6 subtype A (GPCR6A) in β - or gut cells. GPCR6A can be directly activated by amino acids and use calcium as an allosteric regulator. IP₃, inositol triphosphate; PLC β , phospholipase C β ; GLP-1, glucagon-like peptide-1; VDCC, voltage-dependent calcium channel. ‡ Described in enterocyte L cells of the small intestine. Adapted from Wauson *et al.*⁽¹⁰⁴⁾.

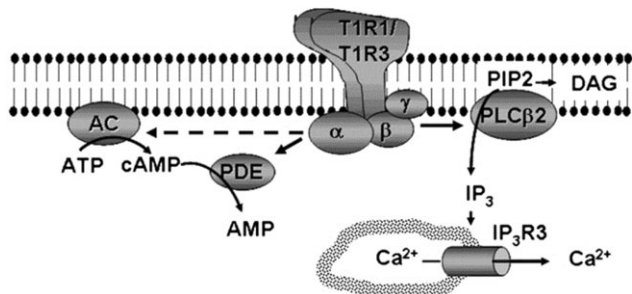


Fig. 4. The T1R1/T1R3 heterodimer is coupled to a heteromeric G protein, where the Gbc subunit appears to mediate the predominant leg of the signalling pathway. Ligand-binding activates Gbc, which results in activation of phospholipase C β 2 (PLC β 2), which produces inositol triphosphate (IP₃) and diacylglycerol (DAG). IP₃ activates IP₃ receptor type 3 (IP₃R3) which results in the release of Ca²⁺ from intracellular stores. AC, adenylyl cyclase; cAMP, cyclic AMP; PDE, phosphodiesterase; PIP2, phosphatidylinositol 4,5-bisphosphate. Adapted from Kinnamon⁽¹⁰⁵⁾.

423 Protein bioactivity on glucagon-like peptide-1 clearance

424 Like the studies on the effects of protein on GLP-1 secretion,
425 most of the studies on the effects of protein on DPP4 inhibition
426 are performed with protein hydrolysates. Over the past few
427 years, bioactive peptides have shown their potential as DPP4
428 inhibitors, a research area that is currently expanding. *In vitro*
429 simulated gastrointestinal digestion has been reported to pro-
430 duce DPP4-inhibitory protein hydrolysates^(83,84). Also, hydrolysis
431 with a range of enzymes is used to release DPP4-inhibitory pep-
432 tides^(69,85-90). Thus, a wide range of protein sources has been
433 used to obtain hydrolysates, for which DPP4-inhibitory activity
434 has been screened mainly *in vitro*.

435 Research has shown that the amino acid sequence plays a
436 much greater role in DPP4-inhibitory activity than other physico-
437 chemical parameters such as length, isoelectric point, hydropho-
438 bicity and net charge^(91,92). DPP4 preferentially cleaves
439 substrates that bear proline or alanine at their P₁ position
440 (Xaa-Pro and Xaa-Ala; where Xaa represents any amino acid)

and also acts on substrates that bear other residues, such as gly- 441
cine, serine, valine and leucine⁽⁹³⁾. Hydrophobic and basic res- 442
idues at the P₂ position enhance the affinity for cleavage 443
compared with acidic residues⁽⁹⁴⁾. The presence of tryptophan 444
residue at the N-terminal position increases the susceptibility 445
to cleavage. Although the residues at the N-terminal position 446
may have a major impact by inhibiting DPP4, the authors pointed 447
out that the C-terminal amino acid also affects the potency of 448
DPP4 because it is involved in the interaction with the enzyme⁽⁹⁵⁾. 449

To date, some studies have been carried out on the *in vivo* 450
DPP4-inhibitory effects of the hydrolysates and peptides from 451
dietary proteins. Peptides derived from milk and bean proteins, 452
which have been shown to inhibit the activity of DPP4 *in vitro*, 453
were also found to have glycaemic effects in mice^(96,97) as plasma 454
glucose levels decreased after an OGTT. A β -casein-derived pep- 455
tide LPQNIPPL found in Gouda-type cheese with *in vitro* DPP4- 456
inhibitory effects has also been tested with animal models. Oral 457
administration of this octapeptide resulted in 1.8-fold lower 458
postprandial glucose AUC; however, insulin plasma levels did 459
not differ⁽⁹⁸⁾. In these studies, the authors did not measure 460
plasma DPP4 activity, so it is not known whether the lower blood 461
glucose was caused by inhibition of DPP4 activity. Chicken feet 462
hydrolysates with DPP4-inhibitory activity *in vitro* improved 463
hyperglycaemia in diet and aged models of glucose homeostasis 464
impairment⁽⁹⁹⁾. 465

As well as hydrolysates from milk and bean protein, *in vivo* 466
models hydrolysate from the egg protein lysosyme has also 467
shown a 25 % reduction in blood serum DPP4 activity and a trend 468
towards higher serum GLP-1 levels after 90 min in diabetic rats 469
undergoing chronic treatment⁽¹⁰⁰⁾. Streptozotocin-induced dia- 470
betic rats were used to evaluate the effects of porcine skin gelatin 471
hydrolysates⁽⁴⁸⁾, Atlantic salmon skin gelatin⁽⁴⁷⁾, and halibut and 472
tilapia skin gelatin⁽⁴⁹⁾. In all these studies, diabetic animals 473
showed reduced blood glucose levels during OGTT, increased 474
plasma insulin and active GLP-1 levels, and reduced plasma 475
DPP4 activity after a chronic treatment with these proteins com- 476
pared with water. Diabetic rats treated for 42 d with a daily dose 477
of 300 mg/kg of porcine skin gelatin showed their plasma glu- 478
cose AUC reduced from 30 000 to 28 000 mg \times min/dl (1665 479
to 1554 mmol \times min/l), insulin levels increased 2-fold, active 480
GLP-1 levels reduced from 15 to 13.5 μ M and DPP4 activity 481
reduced by half⁽⁴⁸⁾. In another study in which the animals were 482
treated for 35 d with a daily dose of 300 mg/kg of Atlantic salmon 483
skin gelatin hydrolysate, blood glucose levels were reduced to 484
less than 200 mg/dl (11.1 mmol/l) during OGTT, insulin levels 485
increased 3-fold, active GLP-1 levels increased 1.6-fold and 486
DPP4 activity was reduced from 115.5 to 82.6 % (lower than 487
in normal rats)⁽⁴⁷⁾. When these animals received a 30 d treatment 488
involving a daily dose of 750 mg/kg of halibut (HSGH) or tilapia 489
skin gelatin hydrolysate (TSGH) the plasma glucose was lower 490
than 200 mg/dl (11.1 mmol/l) in the TSGH-treated group. When 491
TSGH was administered, insulin levels were 1.56 g/l, higher than 492
that of HSGH (1.14 g/l) and the diabetic control group (0.43 g/l). 493
The active GLP-1 plasma levels of the diabetic control rats 494
(5.14 μ M) were lower than those for TSGH-treated group 495
(13.32 μ M) and for HSGH-treated group (7.37 μ M) and the DPP4 496
activity reduced from 115.5 in the diabetic group to 86.6 and 497
71.6 % in the HSGH- and TSGH-treated groups, respectively⁽⁴⁹⁾. 498

Moreover, rodents receiving halibut and tilapia skin gelatin hydrolysates also showed increased total GLP-1 levels. Therefore, the findings of this study suggest that these hydrolysates exert their anti-hyperglycaemic effect via dual actions of DPP4 inhibition and GLP-1 secretion enhancement. Similarly, the ileal administration of zein protein hydrolysate to rats was found to potentiate the incretin effect when administered before an intraperitoneal glucose tolerance test, resulting in decreased glucose concentration, increased insulin levels, decreased plasma DPP4 activity, and increased total and active GLP-1 secretion compared with water⁽¹⁰¹⁾. Rice-derived peptides were likewise found to act via dual action. Oral administration increased plasma GLP-1 levels compared with water during an intraperitoneal glucose tolerance test, and ileal administration reduced plasma DPP4 activity and increased the ratio of active GLP-1 to total GLP-1⁽⁶⁹⁾ in rats. *In vitro* studies also showed dual mechanisms for protein hydrolysates; both enhanced GLP-1 secretion and inhibited DPP4, as has been shown for the cuttlefish (*Sepia officinalis*) viscera protein hydrolysate and bovine Hb hydrolysate^(72,77), whey proteins⁽⁵⁶⁾ and chicken feet hydrolysate⁽⁹⁹⁾. Therefore, these two mechanisms might also take part *in vivo* for some protein sources, leading to an increase in active GLP-1 and improve glycaemia.

Human studies, although limited, offer some evidence that food-derived peptides, mostly from dairy protein, act as DPP4 inhibitors⁽¹⁰²⁾. It was shown that a whey preload, consumed before the breakfast meal, reduced glucose levels by 28 % and increased insulin and total GLP-1 levels by 105 and 141 %, respectively, compared with water. Nevertheless, no significant differences in plasma DPP4 activity were found. This could be interpreted as whey protein acting as an endogenous inhibitor of DPP4 in the proximal small intestine, but not in the plasma (intestinal DPP4 activity was not assessed)⁽⁶⁰⁾. Further studies are needed to examine the potential of casein- and whey-derived peptides, as well as peptides derived from other sources, to act with DPP4 inhibitors in human subjects.

Conclusions

Food proteins target the enteroendocrine system. They directly enhance GLP-1 release from enteroendocrine cells. Current studies suggest that the source of the protein might lead to differences in GLP-1 secretion, although there is not enough literature to enable the different proteins to be compared. The effect of gastrointestinal digestion can also enhance or decrease GLP-1-secreting capacity depending on the protein type. Thus, it is important to consider this digestion when discussing the effects of protein on GLP-1 secretion *in vitro*. In addition, peptides with DPP4-inhibitory effects can be released during the digestion process, which could modulate the life span of target enterohormones. However, whether this hydrolysis remains important after intestinal digestion *in vivo* remains to be clarified. Thus, the use of protein/protein hydrolysates to ameliorate situations of glucose derangements is promising, but more research, specifically human studies, is required to define the most effective sources/treatments.

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