



A mechanistic model of small intestinal starch digestion and glucose uptake in the COW

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Accepted Version

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Mills, J. A. N., France, J., Ellis, J. L., Crompton, L. A., Bannink, A., Hanigan, M. D. and Dijkstra, J. (2017) A mechanistic model of small intestinal starch digestion and glucose uptake in the cow. *Journal of Dairy Science*, 100 (6). pp. 4650-4670. ISSN 0022-0302 doi: <https://doi.org/10.3168/jds.2016-12122> Available at <http://centaur.reading.ac.uk/69845/>

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To link to this article DOI: <http://dx.doi.org/10.3168/jds.2016-12122>

Publisher: Elsevier

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ABSTRACT

The high contribution of postruminal starch digestion (up to 50%) to total tract starch digestion on energy-dense, starch-rich diets demands that limitations to small intestinal starch digestion be identified. A mechanistic model of the small intestine is described and evaluated with regard to its ability to simulate observations from abomasal carbohydrate infusions in the dairy cow. The seven state variables represent starch, oligosaccharide, glucose and pancreatic amylase in the intestinal lumen, oligosaccharide and glucose in the unstirred water layer (UWL) at the intestinal wall, and intracellular glucose of the enterocyte. Enzymatic hydrolysis of starch is modelled as a two stage process involving the activity of pancreatic amylase in the lumen and of oligosaccharidase at the brush border of the enterocyte confined within the UWL. Na⁺ dependent glucose transport into the enterocyte is represented along with a facilitative GLUT2 transport system on the basolateral membrane. The small intestine is subdivided into three main sections representing the duodenum, jejunum and ileum for parameterisation. Further sub-sections are defined between which continual digesta flow is represented. The model predicted non-structural carbohydrate disappearance in the small intestine for cattle unadapted to duodenal infusion with $R^2 = 0.92$ and a root mean square prediction error (RMSPE) of 25.4%. Simulation of glucose disappearance for mature Holstein heifers adapted to various levels of duodenal glucose infusion yielded $R^2 = 0.81$ and a RMSPE of 38.6%. Analysis of model behaviour identified limitations to the efficiency of small intestinal starch digestion with high levels of duodenal starch flow. Limitations to individual processes, particularly starch digestion in the proximal section of the intestine, can create asynchrony between starch hydrolysis and glucose uptake capacity.

Key words: starch digestion, small intestine, glucose uptake, mechanistic model

40

41

INTRODUCTION

42 The need to satisfy the energy requirements of high genetic merit dairy cows during early
43 lactation often results in the feeding of substantial quantities of starch rich concentrate. Coupled to
44 this is the use of high starch corn silages as a main or primary forage component in many dairy
45 production systems (Khan et al., 2015). The fate of dietary starch is highly variable and depends
46 on many factors including starch type, processing and interaction with other diet components
47 (Mills et al., 1999a,b; Moharrery et al., 2014; Patton et al., 2012) and maturity of corn at harvest
48 (Hatew et al., 2016; Peyrat et al., 2016). This has significant implications for the productive
49 capacity of the dairy cow (Nocek and Tamminga, 1991).

50 Previously, we developed a model for lactate metabolism in the rumen with a view to
51 address the issue of rumen acidosis (Mills et al, 2014). Whilst starch may be highly degraded by
52 rumen micro-organisms, up to 50% may escape undegraded to the small intestine, in particular
53 with corn, sorghum and legumes (Mills et al., 1999a; Larsen et al., 2009), depending on the ration.
54 The digestion of starch within the small intestine, followed by the absorption of the released
55 glucose, may avoid the inefficiencies of rumen fermentation (Huntington et al., 2006; Reynolds et
56 al., 2014). Digestion of up to 2.5 kg/d of starch in the small intestine of lactating dairy cows has
57 been reported (Reynolds et al., 2014). However, starch reaching the small intestine is by nature
58 less digestible than starch digested in the rumen. As starch flow to the small intestine increases,
59 starch digestibility in the small intestine decreases, and there may be limits to the capacity of the
60 small intestine for enzymatic hydrolysis of starch or glucose uptake by epithelial tissue (Mills et
61 al., 1999b; Huntington et al., 2006; Reynolds et al., 2014). Subsequently, excessive fermentation
62 in the hindgut of starch that escapes digestion in the small intestine may negatively affect fibre

63 digestion and may have negative effects on absorption of microbial lipopolysaccharides (Li et al.,
64 2012). Published data regarding glucose flux across the small intestine in cattle shows highly
65 variable results depending on diet fed or level of postruminal glucose infusion (Huntington and
66 Reynolds, 1986; Reynolds et al., 1988; Reynolds et al., 1991). Patton et al. (2012) compared
67 several models on accuracy of prediction of post-ruminal starch digestion. Even with the large
68 intestine compensating for part of the variation in starch digestion in the small intestine, they still
69 obtained substantial prediction errors of 15% and over 20% of observed means for corn-starch and
70 non-corn starch, respectively. There is hence room for improvement of prediction of intestinal
71 starch digestion. Complementary to improving empirical models (which include fractional rates of
72 passage and digestion; e.g. Patton et al., 2012), is the study of factors which underlie such
73 variation. The objective of the present study is to construct a mechanistic model that can be used
74 to simulate the digestive metabolism of non-structural carbohydrate flowing through the small
75 intestine of the dairy cow.

76

77

THE MODEL

78 The model is based on principles advanced by Mills et al. (1999b) and is illustrated in
79 Figure 1. The level of aggregation adopted to describe the biological processes is similar to that
80 used in previous modelling studies for the rumen and large intestine (Mills et al., 2014). Hence,
81 the model can be considered alone as a tool for small intestinal starch digestion or as an element
82 within a larger model of nutrient digestion and utilisation in the dairy cow. The model consists of
83 three principal sections representing the duodenum, jejunum and ileum between which parameters
84 describing enzyme activity, metabolite transport and intestinal physiology are varied according to
85 literature values. These sections are further subdivided into subsections (2 in duodenum, 15 in

86 jejunum and 30 in ileum; see below for discussion) representing shorter lengths of intestine within
87 which the state variables are represented. Division into subsections facilitates the simulation of
88 digesta flow within each section (i.e., within duodenum, jejunum or ileum) as well as between
89 sections. Equations representative of the model and abbreviations used to define model entities are
90 listed in the Appendix. Associated parameters describing properties of the model and their values
91 are given in Table 1. All pools are expressed in moles, with volume in litres (L) and time in hours
92 (h). The flow equations are described by Michaelis–Menten or mass action forms. To describe
93 non-structural carbohydrates in molar terms, molecular mass of non-polymerised and polymerised
94 glucose is assumed to be 180 and 162 respectively. It is assumed that oligosaccharide resulting
95 from starch hydrolysis contains an average of 5 glucose molecules.

96

97 *Parameterisation*

98 *Intestinal Size and Digesta Passage.* In the absence of other experimental observations, the
99 length of the small intestine is set according to the observations of Gibb et al. (1992) for dairy
100 cows at different stages of lactation. Whilst duodenal length is well characterised within the
101 literature, the proportion of total length attributable to the jejunal and ileal sections is less clear,
102 with little data available especially in the cow. Madge (1975) cites the ratio of duodenal, jejunal
103 and ileal length as 1:4:7. However, most experimental observations for biological activity at these
104 three points relate to measurements taken well within the bounds of the respective sections.
105 Therefore, location-dependent parameters are set according to observed data for the mid
106 duodenum, mid jejunum and terminal ileum (0.9 of ileal length). These parameters are
107 extrapolated in a linear fashion between these three points. The proportions of small intestinal
108 length accounted for by the duodenum, jejunum, and ileum are set at 0.02, 0.35 and 0.63

109 respectively. For the volume and surface area calculations, the small intestine is treated as a
110 cylinder of diameter 5 cm across all sections (Braun et al., 1995).

111 Duodenal nutrient inputs are determined by the composition of infusate or of duodenal
112 nutrient flow reported in the investigations being used for model simulations. Passage of digesta
113 between the intestinal sections and subsections is represented as a fractional rate (k_p) assuming
114 mixing within each section due to myoepithelial contractions (Ruckebusch, 1988). Fractional
115 passage rate between the luminal pools is a function of total mean retention time (**MRT**) for the
116 small intestine. The total MRT in the small intestine is dependent on k_p and intestinal length.
117 Where experimental observations are lacking, k_p is set according to Cant et al. (1999) who
118 observed a rate of 16 m/h in a mature dairy heifer (507 kg). Whilst in reality passage along the
119 small intestine is pulsatile (Ruckebusch, 1988), for simplicity the model assumes a continuous
120 digesta flow between the luminal pools of the intestinal subsections.

121 Small intestinal digesta volume is set at 13% of theoretical lumen volume (12.5 L for a
122 dairy cow with small intestinal length assumed to be 48 m) (Gibb et al., 1992). Water absorption is
123 assigned a fractional rate of 2% of volume per h. This is calculated assuming typical abomasal
124 outflows of 3% dry matter (DM) and ileal outflows of 8% (DM) for a MRT of 2.5 h and hence it is
125 the net result of water absorption from and water influx (with secretions) into the intestinal lumen.

126

127 ***The Intestinal Lumen***

128 *Luminal Starch (Sl)*. Inputs to the luminal starch pool are direct outputs from the
129 equivalent pool in the previous section. The input to the first duodenal section is the output from
130 the rumen, assuming no changes in starch flow occur in omasum and abomasum. The outputs
131 from the luminal starch pool are passage or hydrolysis to luminal oligosaccharide. The potential

132 hydrolysis of the starch is represented by starch digestion turnover time, which is discussed in
133 more detail below under the heading 'Luminal α -Amylase'. Other factors influencing the rate of
134 hydrolysis are the concentration and activity of pancreatic amylase, both of which are described
135 below.

136 *Luminal Oligosaccharide (Ol)*. Inputs to the oligosaccharide pool are directly from
137 abomasal infusions (in the duodenum) or from outflows of the previous section. Outflows are via
138 passage along the intestine or diffusion into the unstirred water layer (UWL). The diffusion
139 coefficient for oligosaccharide ($k_{ol,olou}^{(d)}$) is set at 0.0089 cm²/h by adjusting the glucose diffusion
140 coefficient for the difference in molecular radius (assumed to be proportional to the cube root of
141 molecular weight) of oligosaccharide and glucose.

142 *Luminal Glucose (Gl)*. For typical dietary situations it is assumed that glucose flow from
143 the rumen (via the abomasum) is negligible. Thus in the present model, glucose may only enter the
144 intestinal lumen via infusion at the abomasum. Outputs from the luminal pools are via passage
145 along the intestine or diffusion into the UWL. The diffusion rate constant is discussed in the UWL
146 section.

147 *Luminal Digesta pH*. The pH of the luminal contents can vary considerably between the
148 duodenal and ileal sections of the small intestine. The acidic chyme entering the duodenum (pH
149 2.5) is buffered by pancreatic secretions rich in bicarbonate (pH 8) (Walker et al., 1994
150 Pierzynowski et al., 1988). Armstrong and Beever (1969) reviewed the observations of pH
151 distribution throughout the ruminant small intestine in relation to the pH optima of various
152 carbohydrases and concluded that maximal starch hydrolysis would occur in the proximal
153 jejunum. Whilst most literature data confirm the general increase in luminal pH as digesta moves
154 from the duodenum to the ileum, the actual acidity and rate of change along the tract seems to vary

155 considerably depending on the particular investigation and diet fed. Russell et al. (1981) fed steers
156 either a lucerne only diet or lucerne diets with increasing corn. Duodenal pH (at 10% of intestinal
157 length) was similar for all diets (pH 6.0 – 6.2) although the pH increase towards the ileum was
158 greatest for the diets with least concentrate (ileal pH 7.2). Earlier studies report much lower pH
159 values towards the duodenum. MacRae (1967) observed a pH range at the duodenum of 2.6 – 3.5,
160 with an ileal pH between 8.0 and 8.3. Lennox and Garton (1968) monitored luminal pH in sheep
161 fed grass cubes and observed pH ranges at the proximal jejunum, upper jejunum and distal
162 jejunum of 2.5 – 4.0, 3.9 – 5.0 and 7.2 – 7.9 respectively. Unpublished data from Holstein dairy
163 cows (Reynolds, 2000) indicates a pH of approximately 2.3 at the proximal duodenum and 8.3 at
164 the terminal ileum. Low duodenal pH measurements are obtained where the sampling site is
165 proximal to the pancreatic ducts, as with the results of Russell et al. (1981) and reported by Owens
166 et al. (1986), confirming a substantial rise in luminal pH following pancreatic secretion (> pH
167 6.0). In the model, duodenal luminal pH is set according to observed values and rises in a linear
168 manner to pH at the terminal ileum. Where experimental observations are not available these
169 values are set at 6.0 and 8.0, respectively.

170 *Luminal α -Amylase (Al)*. The first stage of starch digestion in the model is hydrolysis in
171 the intestinal lumen via the action of pancreatic amylase. The product of this reaction is
172 oligosaccharide (see below), and amylase activity is dependent on the quantity secreted and the pH
173 of the intestinal lumen. There is only limited data concerning pancreatic amylase secretion in dairy
174 cows, although various theories as to control mechanisms have been postulated (Fushiki and Iwai,
175 1989; Croom et al., 1992). Detailed representation of neural and hormonal regulatory mechanisms
176 is beyond the scope of the present model. However, nutritional influences particularly concerning
177 interactions with feed carbohydrate are considered. Due to the moderating effect of the rumen on

178 digesta flow, in contrast to non-ruminants, the flow of ruminant pancreatic secretion is relatively
179 constant irrespective of feeding behaviour (Pierzynowski, 1986; Walker and Harmon, 1995).
180 Pancreatic fluid secretion rate in young calves is in the range 0.33 – 0.49 mL/kg body weight
181 (BW)·h⁻¹ (Pierzynowski, 1989; Khorasani et al., 1990). The majority of observations indicate a
182 similar range for mature cattle (0.26 – 0.57 mL/kg BW·h⁻¹), although the influence of diet is
183 pronounced. Pierzynowski et al. (1988) recorded an increase in pancreatic secretion from 0.57 to
184 0.91 mL/kg BW·h⁻¹ when dry cows were fed isoenergetic and isonitrogenous rations with
185 molasses rather than grain as the energy source. Under typical nutritional management there is
186 approximately 12 mg total protein in bovine pancreatic fluid (Walker et al., 1994; Walker and
187 Harmon, 1995) of which less than 2% is amylase (Keller et al., 1958). Harmon (1993) reviewed
188 the literature and concluded that increased postruminal carbohydrate in the form of starch or
189 glucose decreases pancreatic amylase secretion whilst increasing energy intake raises secretion.

190 The model representation allows for increased pancreatic secretions for cows on a high
191 plane of nutrition whilst facilitating a decrease in the concentration of amylase in pancreatic
192 secretion with increased starch or glucose presence. Whilst Russell et al. (1981) observed a
193 doubling in pancreatic amylase activity for homogenised pancreatic tissue in steers fed at either
194 twice or three times maintenance relative to those at maintenance, they did not record fluid
195 secretion rates. In the model, the increase above the basal level of fluid secretion ($v_{Al,PfAl}^{**}$) is
196 sigmoidal and set to yield a doubling in total secretion between 1 and 2 times maintenance feeding
197 (Equation 1.5). The maximum rate of secretion ($v_{Al,PfAl}^*$) is arbitrarily set at 0.8 mL/kg BW·h⁻¹
198 based on the observations described above. The response to duodenal starch delivery (Equation
199 1.6) is inverse to that for energy intake. Minimum concentration of amylase in pancreatic fluid

200 $(v_{Al,PpAl}^{**})$ is set at 10.0 U/mg fluid protein, based on observations by Walker and Harmon (1995).
201 One unit (U) equates to 60 μmol of oligosaccharide released per h (hence $v_{Sl,StOl} = 60 \mu\text{mol/h}$). The
202 maximum concentration of amylase ($v_{Al,PpAl}^{**}$) and the rate of duodenal starch or glucose delivery
203 at which amylase concentration is half maximal ($M_{Al,PfAl}$) are set according to the data of Walker
204 and Harmon (1995) at 22 U/mg fluid protein and 0.21 mol/h, respectively. The affinity of amylase
205 for starch ($M_{Sl,StOl}$) is set at 21.6 mmol/L (Russell et al., 1981) with a modification for starch
206 hydrolysis based on a reference digestion turnover time (T_{St}^*) of 11.8 h for corn starch (Cone,
207 1991). Starch hydrolysis by pancreatic amylase depends on starch source and is set according to
208 the observations of Cone (1991) who used pancreatin in vitro to degrade a range of starches over 4
209 h incubations. The digestion turnover time of starch (T_{St}) is calculated as the reciprocal of the
210 fractional hydrolysis rate constant.

211 The optimum pH for pancreatic amylase activity ($v_{pH,StOl}^{(o)}$) in cattle is 6.9 and activity
212 declines substantially above and below this optimum (Russell et al., 1981). The response of
213 amylase activity to pH is described by a Gaussian type equation (Equation 2.5), with the steepness
214 parameter ($\theta_{Sl,StOl} = 0.6$, $\text{SE} \pm 0.15$, $R^2 = 0.68$) fitted to describe the data of Russell et al. (1981)
215 and Rosenblum et al. (1988).

216

217 ***The Unstirred Water Layer***

218 To facilitate estimates of metabolite concentrations and diffusion parameters, the thickness
219 of the UWL is set at 40 μm based on observations for human jejunal tissue (Levitt et al., 1992). In
220 line with the observations of Lucas (1983), pH of the UWL is independent of luminal pH, with
221 values of 6.1, 6.1, and 7.1 for duodenal, mid-jejunal and terminal ileal sections, respectively.

222 *UWL Oligosaccharide (Ou)*. Input into the UWL oligosaccharide pool is from diffusion
223 across the UWL, whilst outputs are diffusion into the lumen or hydrolysis to glucose via the action
224 of brush border oligosaccharidase. Parameters for enzymatic hydrolysis are discussed below under
225 the heading ‘UWL Oligosaccharidase’, and assumptions for diffusion rates were set out in the
226 previous section ‘The Intestinal Lumen’.

227 *UWL Glucose (Gu)*. Inputs to the UWL glucose pool are from hydrolysis of UWL
228 oligosaccharides, and from diffusion across the UWL (from lumen and from blood). Outputs are
229 diffusion into the lumen and the blood and uptake into the epithelial cells by SGLT1.
230 Pappenheimer and Reiss (1987) promote the concept of solvent drag induced paracellular
231 transport as a significant contributor to glucose absorption in the small intestine of rats.
232 Pappenheimer and Reiss (1987) indicate that where luminal glucose concentrations are greater
233 than 250 mM, paracellular movement of glucose exceeds that for the transcellular route. However,
234 Ferraris et al. (1990) suggest a physiological range in luminal glucose concentration for non-
235 ruminants of 5 – 50 mM, whilst ruminants tend to show even lower concentrations (Bauer, 1996).
236 Therefore, paracellular glucose flux is likely of limited significance in normal feeding situations.
237 This is confirmed by the observations of Krehbiel et al. (1996) who infused 2-deoxyglucose (not
238 transported by sodium dependent glucose co-transporter SGLT1; further explanation follows) into
239 the duodenum of steers and recovered only 7% in the portal vein. However, the absolute
240 significance of paracellular glucose transport is still unknown, especially as glucose may enter the
241 lymphatic drainage and not the portal circulation (Largis and Jacobs, 1971). The influence of
242 localised high glucose concentrations at the UWL may also be significant (Pappenheimer and
243 Reiss, 1987). Meddings and Westergaard (1989) demonstrated that the uptake of luminal glucose
244 in the rat is best described by a carrier system and a diffusion component across the UWL.

245 Therefore, a simple paracellular diffusion component is represented in the model between glucose
246 at the UWL and glucose in the blood, assuming a diffusive surface area based on that for epithelial
247 cell junctions of 4% of total brush border surface area (Krstic, 1979). The diffusion coefficient (
248 $k_{Gu,GuGb}^{(d)}$) is set at 2.412×10^{-2} cm²/h (Levitt et al., 1992).

249 *UWL Oligosaccharidase.* Oligosaccharidase activity is associated entirely with the brush
250 border of the enterocyte (Harmon, 1993) and therefore the hydrolysis of oligosaccharide to
251 glucose only occurs within the UWL. The ability of ruminants to regulate oligosaccharidase
252 activity per unit of intestine appears limited. Reports of adaptive regulation in response to
253 carbohydrate intake (Janes et al., 1985) are likely to be the result of changes in intestinal length as
254 energy intake increases (Harmon, 1993). Therefore, the mean maximum activity is held constant
255 and set according to the data of Kreikemeier et al. (1990) at 0.25, 1.0 and 0.72 U/cm² brush border
256 membrane for the mid-duodenum, mid-jejunum and terminal ileum, respectively. One unit of
257 activity represents 60 μmol glucose produced per hour and therefore $v_{Ou,OuGu}^* = 60$ μmol/h. A
258 similar distribution was reported by Coombe and Siddons (1973). The model employs a Gaussian
259 type equation (Equation 6.7) fitted to the data of Coombe and Siddons (1973) to describe the
260 activity of oligosaccharidase as affected by the UWL pH with a pH optimum of 6.0 and a
261 steepness parameter ($\theta_{Ou,OuGu}$) of 0.19 (SE ±0.0068, R² = 0.98). The affinity for oligosaccharide is
262 set at 4.3 mmol/L, assuming 70% maltase activity (affinity $K_m = 2.3$ mmol/L, mean of Siddons
263 (1968) and Eggermont (1969)) and 30% isomaltase activity ($K_m = 9.1$ mmol/L, Coombe and
264 Siddons (1973)).

265

266 ***Enterocyte Metabolism***

267 In the absence of ruminant data, the cytoplasmic depth of the columnar epithelial cells is
268 set at 25 μm according to observations from rabbits (Stevens, 1992), allowing calculation of
269 enterocyte volume.

270 *Na⁺ Dependent Uptake of Glucose by SGLT1.* The majority of glucose transport into the
271 epithelial cells occurs via a sodium (Na^+) dependent glucose transporter (SGLT1) (Shirazi-
272 Beechey et al., 1995; Dyer et al., 2003) present on the brush border membrane. Hence, transport of
273 glucose into the epithelial cell obeys saturation kinetics, limited by the affinity of SGLT1 for
274 glucose, the maximum uptake rate of glucose per unit of SGLT1 and the quantity of transporter
275 protein present at the membrane. The kinetic properties of SGLT1 have been determined in vitro
276 using brush border membrane vesicles that remove the influence of the unstirred water layer,
277 otherwise present in vivo. Therefore, whilst in vivo estimates of the affinity of SGLT1 for glucose
278 range from 6 to 23 mmol/L (Ferraris et al., 1990), in vitro results using vesicles show the true K_m
279 be between 0.06 and 0.15 mmol/L (Bauer et al., 1997; Zhao et al., 1998). Therefore, $M_{Gu,GuGe}$ is
280 set at 0.1 mmol/L. Maximum glucose uptake rates by SGLT1 per unit of intestinal epithelia vary
281 depending on the luminal glucose presence or degree of adaptation to a particular diet (Hediger
282 and Rhoads, 1994; Bauer, 1996). Diamond and Karasov (1987) demonstrated a 1.5% increase in
283 glucose transporter activity for every 10% increase in dietary sugar level for mice fed isoenergetic
284 diets. Assuming that the signal mechanism for regulation is in the proximal duodenum, $v_{Gu,GuGe}$ is
285 dependent on non-structural carbohydrate entry to the small intestine (Equation 5.9). Since the
286 model's purpose is to simulate the adapted state, and not transition between diets, up-regulation is
287 assumed to have occurred. Bauer et al. (1995) observed a basal transport capacity of 960 mmol
288 glucose/d in steers fed lucerne hay with or without carbohydrate infusion. Therefore, assuming a

289 small intestinal length of 35 m, the mean transport capacity for these steers was $0.73 \mu\text{mol}/\text{cm}^2/\text{h}$
290 intestinal epithelium, a value that has been adopted within the model ($v_{Gu,GuGe}^{**}$). Relatively high
291 basal transporter density allows more efficient utilisation of transient nutrient inputs. Despite
292 evidence suggesting regulation of the maximum SGLT1 capacity to meet or exceed luminal
293 glucose supply in a variety of herbivores (Ferraris et al., 1990), this degree of adaptation in cattle
294 has been questioned (Cant et al., 1999; Lohrenz et al., 2011). It is logical to assume an upper limit
295 to active transport capacity irrespective of luminal glucose delivery. Invariably, the highest
296 transport rates are observed at the pre-ruminant stage of development on milk based diets.
297 Therefore, an absolute mean maximum uptake rate of glucose by SGLT1 ($v_{Gu,GuGe}^*$) is set at 29.2
298 $\mu\text{mol}/\text{cm}^2/\text{h}$ brush border membrane according to data for lambs maintained on a milk-based diet
299 for 5 weeks where transport capacity was 40 times basal levels (Shirazi-Beechey et al., 1991).
300 Based on observations with brush border vesicles from a mid-lactation Holstein cow (Zhao et al.,
301 1998), SGLT1 transport capacity ($v_{Gu,GuGe}^*$, $v_{Gu,GuGe}^{**}$) is distributed at a ratio of 0.66:1.0:0.12,
302 between the mid-duodenum, mid-jejunum and terminal ileum, respectively.

303 *Glucose in the Enterocyte (Ge)*. The inputs to the intracellular glucose pool are from
304 luminal uptake by SGLT1 (Equation 7.2) and from blood by Na^+ independent facilitated diffusion
305 involving GLUT2 transport proteins at the basolateral membrane (Equation 7.3). There are two
306 outputs, one via GLUT2 to the blood (Equation 7.4) and the other via oxidation in the enterocyte
307 (Equation 7.5). Carbohydrate infusion studies frequently report a significant disparity in portal
308 glucose appearance relative to disappearance in the intestine (Kreikemeier and Harmon, 1995).
309 This is due to glucose oxidation by the visceral tissues in order to satisfy the energetic
310 requirements of protein turnover and ion transport. Glucose utilisation by the small intestinal

311 mucosal tissue is estimated to enable a prediction of net glucose flux to the blood. Energetic
312 requirements of the mucosal tissue are calculated assuming that there is a basic energy
313 requirement for protein turnover (0.02 mmol ATP/g mucosa/h) and ion transport (0.031 mmol
314 ATP/g mucosa/h) in the fasted state (Gill et al., 1989). This equates to a combined glucose
315 requirement for protein turnover and ion transport of 0.0043 mmol glucose/g mucosa/h. According
316 to the observations of Kreikemeier et al. (1990), there is 0.15 g mucosa/cm² intestinal epithelium.
317 Added to these requirements are the energy costs associated with Na⁺ dependent transport of
318 amino acids and glucose from the intestinal lumen to the epithelial cytosol. Amino acid uptake is
319 an input into the model based on experimental observations, with a requirement of 0.66 mmol
320 ATP/mmol amino acid as calculated by Gill et al. (1989). For glucose transport, a stoichiometry of
321 3 mol of glucose transported per mole of ATP hydrolysed yields a requirement of 0.33 mmol
322 ATP/mmol glucose (Gill et al., 1989). Therefore, the corresponding glucose requirements in the
323 model for oxidation during glucose transport ($R_{Ge,GuGe}$) and amino acid transport ($R_{Gu,AuAe}$) are
324 0.028 and 0.055 mmol glucose per mmol glucose and amino acid respectively.

325 In non-ruminants, energy requirements for enterocyte metabolism are met primarily by
326 glutamine oxidation (25 – 40% of total CO₂ production), with glucose oxidation only accounting
327 for 6 – 10% of CO₂ production (Windmueller and Spaethe, 1974; Hanson and Parsons, 1977;
328 Windmueller, 1982). However, Okine et al. (1995) observed that between 69 and 76% of energetic
329 requirements are met by glucose for bovine enterocytes in vitro with the remainder met by
330 glutamine oxidation. Therefore, the calculated energy requirements of the mucosa (mol ATP/h)
331 are met assuming a non-limiting supply of glutamine, with glucose oxidation supplying a
332 maximum of 70% of the total requirement for ATP and glucose is the preferred substrate. The fate
333 of metabolised glucose is set at 18% through complete oxidation to CO₂, 36% through lactate,

334 38% through glutamate and 8% through alanine (Okine et al., 1995). For calculation of ATP yield,
335 lactate and alanine are assumed to be lost to the circulation. Yield of ATP from complete
336 oxidation to CO₂ is 36 mol/mol glucose and from glutamate it is 14 mol/mol glutamate (Stryer,
337 1995). Therefore, the mean ATP yield per mol glucose metabolised by the epithelium is set at 11.9
338 mol.

339 *Facilitated Diffusion from the Enterocyte.* There is GLUT2 glucose transport from the
340 epithelial tissue to the blood (Thorens, 1993; Breves and Wolffram, 2006). Zhao et al. (1998)
341 showed that the distribution of GLUT2 in late lactation Holstein cows was similar to that reported
342 for humans (Burant et al., 1991). Northern blot analysis of GLUT2 mRNA demonstrated
343 considerably greater presence in the liver than in the kidney or duodenum (Zhao et al., 1998).
344 However, details of the nutritional management for the cows were not given and data regarding
345 the ability of the epithelial tissue to upregulate GLUT2 transporter presence in the bovine appears
346 to remain unavailable. Thorens (1993) reviewed the studies concerning GLUT2 adaptation to
347 intestinal glucose delivery and concluded that the rate of glucose transport through the basolateral
348 membrane after an acute exposure to increased glucose may result from changes in the intrinsic
349 activity of the transporter, whereas chronic exposure may increase GLUT2 presence. Cheeseman
350 and Harley (1991) showed that from a range of carbohydrate components only glucose and
351 fructose influenced GLUT2 presence in rat small intestine. Kellett and Helliwell (2000) stated that
352 GLUT2 quantity in rat jejunum doubled when luminal glucose concentration was increased from 0
353 to 100 mmol/L. Whilst Kellett and Helliwell (2000) reported a single saturation constant to
354 describe the affinity of GLUT2 for glucose ($K_m = 56$ mmol/L), the kinetics of glucose efflux from
355 the epithelial cell to the blood and uptake from the blood to the cell are in fact asymmetrical.
356 Maenz and Cheeseman (1987), using basolateral membrane vesicles (BLV) from rat jejunum,

357 showed the mean maximum velocity (V_{\max}) for glucose efflux to be 0.20 ± 0.01 nmol glucose/mg
358 BLV protein/sec with a K_m of 23 ± 2 mmol/L glucose, whilst glucose uptake gave a V_{\max} of $1.14 \pm$
359 0.14 nmol glucose/mg BLV/sec and a K_m of 48 ± 5 mmol/L glucose. Such asymmetrical kinetics
360 allows for an efficient glucose delivery system to the blood whilst minimising the reverse flux of
361 glucose from the blood to the epithelial tissue. Hence, $M_{Ge,GeGb}$ and $M_{Gb,GbGe}$ are set at 23 and 48
362 mmol/L, respectively. In the absence of data specific to the dairy cow, basal maximum uptake rate
363 from the cytosol by GLUT2 ($v_{Ge,GeGb}^{**}$) is set at $12.2 \mu\text{mol}/\text{cm}^2/\text{h}$ as measured in isolated rat
364 enterocytes by Cheeseman and Harley (1991) and in line with Lohrenz et al. (2011), the
365 distribution of GLUT2 is equal for each intestinal section. Calculation of basal maximal transport
366 activity assumes $0.15 \text{ g mucosa}/\text{cm}^2$ basolateral membrane (Kreikemeier et al., 1990). The
367 recruitment of additional GLUT2 and hence an increase in $v_{Ge,GeGb}$ is related to an increase in
368 luminal NSC flow (Equation 7.4) with $v_{Ge,GeGb}^*$ set at 4 times basal level (Cheeseman and Maenz,
369 1989) and $M_{NSC,GeGb}$ set at $75 \text{ mmol}/\text{L}$ duodenal NSC flow/h, as for the up-regulation of SGLT1.

370

371 ***The Blood***

372 *Blood Glucose (Gb)*. Plasma glucose concentration represents that of the mesenteric
373 plasma pool in contact with the basolateral membrane of the intestinal epithelium. The
374 concentration of blood glucose is held constant, since a full interpretation of portal drained viscera
375 (PDV) metabolism is beyond the scope of the model. Where observations of blood glucose
376 concentration are not available, estimates of $4 \text{ mmol}/\text{L}$ can be used for cows (Reynolds and
377 Huntington, 1988; Reynolds et al., 1991). The kinetics of glucose flux to and from the epithelial

378 tissue are described in the GLUT2 section whilst the calculation of net glucose flux is explained
379 under Enterocyte Metabolism.

380

381 *Model Summary*

382 The differential equations for the 7 state variables in each of the 47 sub-sections of the
383 small intestine, representing the nutrient pools in the lumen, the UWL and epithelial tissue, are
384 integrated numerically for a given set of initial conditions and parameter values. The model was
385 written in the Advanced Continuous Simulation Language (ACSL) (Mitchell and Gauthier
386 Associates, 1995). A fourth-order fixed-step-length Runge-Kutta method with an integration
387 interval of 0.25 min was used. The results presented were obtained by running the model until a
388 steady state was achieved.

389

390 *Model Evaluation*

391 The model was initially evaluated against its ability to simulate the duodenal glucose
392 infusion study of Cant et al. (1999) involving mature Holstein heifers adapted to the level of
393 glucose supply. Infusion studies utilising adapted animals are rare, the other principal study being
394 that of Bauer et al. (1995). However, Bauer et al. (1995) do not report small intestinal starch or
395 glucose disappearance. To challenge the model further, a data set for small intestinal NSC
396 disappearance was gathered from infusion studies utilising unadapted cattle. These were: the
397 starch, dextrin, and glucose infusions (each at 20, 40 and 60 g/h) of Kreikemeier et al. (1991), the
398 10 and 20 g/h duodenal glucose infusions of Krehbiel et al. (1996), the glucose, starch and dextrin
399 infusions (each at 66 g/h) of Kreikemeier and Harmon (1995), and the duodenal corn starch and
400 ruminal casein infusion of Taniguchi et al. (1995).

401 Walker and Harmon, (1995) showed that 53% of starch hydrolysate (or dextrin) consisted
402 of glucose chain lengths of 7 or less. To accommodate this product within the model scheme, 30%
403 of starch hydrolysate is assumed to enter the luminal oligosaccharide pool directly (approx. 5
404 glucose molecules or less), with the remainder entering the duodenal starch pool.

405 Regression analysis between observed and predicted values was used to demonstrate
406 model performance. Error of prediction is estimated from the calculation of root Mean Square
407 Prediction Error (**rMSPE**) and expressed as a percentage of the observed mean, where:

$$408 \quad \text{MSPE} = \sum (O_i - P_i)^2 / n$$

409 where $i = 1, 2, \dots, n$; n is the number of experimental observations and O_i and P_i are the observed
410 and predicted values (Bibby and Toutenburg, 1977). The MSPE is decomposed into overall bias of
411 prediction, deviation of regression slope from one and the disturbance proportion (Bibby and
412 Toutenburg, 1977).

413 The response of the model to changes in parameter values ($\pm 50\%$, except $\pm 20\%$ for pH)
414 was tested in order to demonstrate sensitivity. The sensitivity and behavioural analyses were
415 performed for a 650 kg dairy cow consuming 22 kg dry matter (DM) at 11 MJ metabolisable
416 energy (ME) per kg DM, with an inflow or infusion of 2.0 kg wheat starch and 2.5 kg of ground
417 corn starch per day. This level of infusion is comparable with the highest observed duodenal
418 starch flows in the literature (McCarthy et al., 1989), with the aim of investigating the rate limiting
419 steps to the digestive and absorptive processes. Results of the sensitivity and behavioural analyses,
420 together with other aspects of model evaluation, are presented in the next section.

421

RESULTS

Comparison between Observations and Predictions

Figure 2 shows the comparison between model predictions and observations of ileal glucose flow in 4 adapted dairy heifers with increasing levels of duodenal glucose infusion (250 – 700 mmol/h) (Cant et al., 1999). There is good agreement between the observed and predicted values ($R^2 = 0.81$) although the small sample gives a high root MSPE (RMSPE) of 38.6%. Figure 2a shows a move away from the line of unity as glucose infusions increase. Figure 2b demonstrates how observed ileal glucose flow reached a plateau beyond 580 mmol/h infusion, whereas the model was unable to simulate this occurrence. It is difficult to ascertain the precise reason for this observed reduction in ileal glucose flow per unit of glucose infusion, although it may relate to the pattern of SGLT1 up-regulation in vivo and the comparatively short periods of adaptation to glucose infusions (3 d). Another explanation could be that the model underestimated the contribution of paracellular diffusion from the digesta to the blood, particularly at high concentrations of luminal glucose. However, the general agreement between observed and predicted results for the lower range of glucose concentrations, more likely to be encountered under normal nutritional management is encouraging.

Figure 3 displays a regression between observed and predicted small intestinal NSC disappearance for cattle unadapted to NSC infusions. The high R^2 (0.92) shows good agreement between the data and a lower RMSPE (25.4%) than for the simulation of glucose infusion in Figure 2a. The trend for under-prediction of luminal NSC removal at higher levels of observed disappearance is surprising and in contrast to results in Figure 2a. The unadapted animals should have a lower capacity to transport glucose and a reduced carbohydrase activity. It is possible that

445 certain internal parameters are more influential than any effect of dietary adaptation (see
446 sensitivity analysis).

447 Figure 4 displays a correlation between the observed net PDV flux of glucose in the same
448 cattle as for Figure 3, and the simulated net release of glucose from the enterocytes to the blood.
449 As expected, the y-intercept indicates a basal level of glucose release to the blood from the small
450 intestine (34 mmol/h), below which no net PDV flux is observed. However, the rate of increase in
451 simulated net glucose flux to the blood is too slow to support the observed increase in PDV
452 glucose flux. This is a clear indication of an overestimate of metabolism of luminal glucose by the
453 model; arterial glucose is a major source of glucose used by enterocytes. This may also be caused
454 by an overestimate of the total contribution of glucose to enterocyte metabolism, especially in the
455 presence of competing substrates such as glutamine. The in vitro measurements for dairy cattle
456 enterocytes that were used for model parameterisation (Okine et al., 1995) may not be
457 immediately applicable in vivo. Indeed, the results of Okine et al. (1995) are somewhat
458 contradictory to the more extensive data for other species, especially non-ruminants (Windmueller
459 and Spaethe, 1974; Windmueller, 1982). Over-estimates of intestinal length will also unduly
460 increase enterocyte glucose demand. Finally, microbial fermentation of glucose within the small
461 intestine was not included in the model, and this may lead to a difference between small intestinal
462 disappearance of starch and portal appearance of glucose. Gilbert et al. (2015) observed more than
463 50% of small intestinal starch disappearance to be due to fermentation in milk-fed calves fed
464 starch in milk replacer twice daily. At present, it is unclear if fermentation of starch has such a
465 significant role in mature cows fed solid feed leading to a much more gradual flow of starch to
466 into the small intestine.

467

468 ***Sensitivity and Behavioural Analyses***

469 *Small Intestinal Starch Flow.* As starch spends more time in the small intestine it is subject
470 to increasing opportunity for enzymatic hydrolysis. Therefore, digestibility increases and Figure
471 5a confirms this type of behaviour in the model. Beyond 4.5 h MRT, the digestion of starch is
472 almost complete. The rate of decline in digestibility below 4.0 h MRT is high (20%/h). This
473 supports the findings summarised from the literature by Reynolds et al. (2014), Mills et al.
474 (1999b) and Nocek and Tamminga (1991) who report a declining small intestinal starch
475 digestibility with increasing postruminal starch flow. Figure 5a indicates that the significance of
476 pancreatic amylase activity as a limit to starch digestion increases as MRT declines. Figure 5b
477 demonstrates the corresponding decline in glucose delivery to the blood. The rate of decline in net
478 blood glucose flux increases as MRT declines because the proportion of flux attributed to glucose
479 oxidation for maintenance of the enterocyte increases. It should be noted that Figure 5b applies
480 primarily to constant starch infusions with increasing MRT. Where digesta flow rate increases in
481 association with starch flow, there is a compensatory elevation in total starch availability for
482 hydrolysis.

483 *Luminal pH.* Figures 6a, b, c highlight the benefits of maintaining duodenal digesta pH
484 between 6 and 6.5. These simulations were run assuming a constant ileal digesta pH due to the
485 extensive buffering capacity in this region of the intestine (Owens et al., 1986). Although the
486 optimal pH for amylase is 6.9, a slightly reduced duodenal pH (6.5) with a gradual rise toward
487 optimum levels at the jejunum resulted in a beneficial effect). Figures 6c, d are evidence of the
488 need for synchronisation between the hydrolysis and uptake processes. Low duodenal pH limits
489 starch hydrolysis initially, and this delay shifts the availability of glucose in the UWL further
490 towards the ileum where SGLT1 capacity is most limiting. Hence, where small intestinal starch

491 flow is high, digesta pH seems to have as much an effect on glucose uptake as it does directly on
492 starch hydrolysis.

493 *Intestinal Size.* The description of small intestinal physiology can substantially affect net
494 glucose flux. It is self-evident that as the proportion of ileum increases relative to the proximal
495 regions of the intestine, the total capacity to transport glucose declines (Figure 7a). However, the
496 relationship between small intestinal length and net blood glucose flux is more complex (Figure
497 7b). As small intestinal length increases from 38 m to 45 m, there is a rise in blood glucose
498 delivery since small intestinal starch digestion and glucose uptake capacity increase. However,
499 further increases in intestinal length result in a sharp reduction in the release of glucose to the
500 blood. This is explained by an increase in mucosal glucose oxidation beyond the supply arising
501 from increased glucose availability due to starch and oligosaccharide hydrolysis. Despite evidence
502 to suggest an overestimation of mucosal glucose requirement (Figure 4), this effect seems
503 physiologically feasible. Indeed, although intestinal length is related to physiological state and
504 body weight, the range for mature Holstein dairy cows has reported to be between 45 and 49 m
505 (Gibb et al., 1992).

506 *Pancreatic Amylase and SGLT1 Activity.* Figure 8a presents the simple linear relationship
507 between the maximum level of pancreatic secretion and starch digestibility. However, when
508 examined against blood glucose flux (Figure 8b), an increasing rate of decline in blood glucose
509 delivery is observed as maximum pancreatic fluid secretion is reduced. Again, the shift in starch
510 hydrolysis and glucose uptake towards the ileum is responsible for this response. The overall low
511 small intestinal digestibility of NSC is highlighted in the analysis of the effects of
512 oligosaccharidase activity on model predictions (Figure 8c). The high rate of starch infusion used
513 for simulation has demonstrated the overall imbalance between the capacity to hydrolyse starch

514 and for the removal of the products of starch hydrolysis. Oligosaccharidase is clearly a limiting
515 factor for the applied level of starch infusion (4.5 kg starch/day), with a doubling in
516 oligosaccharidase density producing a similar rate of increase in blood glucose appearance (Figure
517 8d). A similar impact on net blood glucose flux is achieved by raising maximum SGLT1 activity
518 (Figure 8e). Hence, at this high rate of starch infusion, the processes associated with the unstirred
519 water layer seem limiting as a whole. This implies that the diffusion of NSC, particularly
520 oligosaccharide, across the UWL from the lumen is a crucial rate-limiting step in the recovery of
521 glucose in the blood from digested starch.

522 Those parameters not shown in Figure 8 have little impact on model behaviour within the
523 range tested. The GLUT2 transporter density ($v_{Ge,GeGb}^*$) only changed the net glucose flux when set
524 at less than 40 $\mu\text{mol}/\text{cm}^2/\text{h}$, below which glucose accumulated in the enterocyte. The affinity of
525 SGLT1 for glucose did not alter predictions of glucose uptake for the range tested (0.5 – 1.5
526 μmol). Likewise, the model was largely insensitive to changes in the affinity of oligosaccharidase
527 ($M_{Ou,OuGu}$) (2.2 to 6.8 mmol/L).

528

529

DISCUSSION

530

531 A declining digestibility of starch in the small intestine of lactating dairy cows is evident at
532 higher duodenal starch flows. Reynolds (2006) summarised data and observed a linear decrease of
533 small intestinal digestion with duodenal starch flow (small intestinal starch digestion (g/kg
534 duodenal starch flow) = 785 – 65.6 × duodenal starch flow (kg/d); Reynolds et al., 2014), as did
535 Huntington et al. (2006). There have been several experiments conducted in the literature to
536 elucidate the limits to starch digestion and glucose recovery. The conclusions of these studies have

537 differed depending on the nutritional regime examined together with the physiological state of the
538 cattle and the experimental technique utilised. Owens et al. (1986) reviewed the available data in
539 an attempt to clarify the points of control at which small intestinal starch digestion is limited in the
540 ruminant. They concluded that the enzymatic capacity did not limit intestinal starch digestion,
541 based on observations relating to total small intestinal starch disappearance. However, Owens et
542 al. (1986) also concluded that the influence of processing on particle size and starch granule
543 structure indicated a physical or physio-chemical barrier to carbohydrase activity. This was
544 confirmed by Larsen et al. (2009) who observed that rolling, compared with grinding, was
545 associated with a larger particle size, which reduced accessibility of enzymes and limited starch
546 digestion in the small intestine. The model does not account for the detailed effects of starch
547 particulate size on the rate of starch hydrolysis, unless this is represented by the estimate for the
548 digestion turnover time T_{SI} . In a study specifically aimed to investigate the effect of corn particle
549 size on starch digestion, Rémond et al. (2004) demonstrated a linear decline of 31% in small
550 intestinal starch digestion with mean particle size increasing from 0.7 to 3.7 mm. A similar effect
551 was demonstrated by Offner & Sauvant (2004) who found underprediction of rumen starch
552 digestion from in situ degradation characteristics when mean dietary particle size is less than
553 4mm. As small intestinal starch digestibility was negatively related to resistance to rumen
554 digestion, this would also mean an underpredicted small intestinal starch digestion. This may also
555 be important especially in the case of whole cereal grains for which reduced digestion can be
556 observed. However, a previous attempt at describing the effects of the physical characteristics of
557 starch on both ruminal and intestinal digestion proved largely unsatisfactory as a predictive tool
558 (Ewing and Johnson, 1987) with a lack of available data for parameterisation. In a more recent
559 comparison of starch digestion models, however, Patton et al. (2012) was able to demonstrate

560 potential for improved prediction by taking account of revised starch digestion rates to represent
561 effects of starch source and processing. The ratio of amylose to amylopectin can, in theory at least,
562 be related to starch digestion. Amylopectin may be more rapidly fermented than amylose, since
563 amylopectin has a branched structure which exposes more non-reducing terminal glucose
564 molecules for enzymatic attack than amylose. However, according to Philippeau et al. (1998)
565 ruminal starch degradation was independent of the amylose:amylopectin ratio in starch.

566 Reynolds et al. (2014) and Mills et al. (1999b) both confirmed the observations of Owens
567 et al. (1986) indicating no quantitative upper limit to postruminal starch disappearance. However,
568 both Mills et al. (1999b) and Reynolds et al. (2014) highlight the potential for increasing
569 compensatory large intestinal starch fermentation as total postruminal starch digestion increases.
570 Larsen et al. (2009) demonstrated a higher contribution of the hind gut to (postrumen) starch
571 digestion for legumes rich in protein with a lower total tract starch digestibility as compared to
572 cereals. The inefficiencies of hind gut fermentation can negate the benefits of elevated small
573 intestinal glucose absorption. Therefore, it is important to consider starch hydrolysis and glucose
574 uptake by the small intestine rather than just starch disappearance prior to the ileo-caecal junction
575 (Mills et al., 1999b). There is experimental evidence to suggest that increasing pancreatic
576 secretion does increase small intestinal starch digestion, with the implication that amylase activity
577 can be rate limiting (Castlebury and Preston, 1993; Taniguchi et al., 1995). Taniguchi et al. (1995)
578 stimulated pancreatic secretion with abomasal casein infusion (120 g/d) and increased small
579 intestinal starch digestion by almost 40% at the expense of large intestinal fermentation. Whether
580 these observations are the result of increased proteolytic capacity leading to enhanced access of
581 amylase to the starch granule or simply a result of increased amylolytic capacity is unclear.
582 Indirect support for the importance of amylase activity on starch hydrolysis is found in the study

583 by Nozière et al. (2014) who found a 9% increase in rumen starch degradation when adding
584 exogenous amylase. Although still not support for any effect of amylolytic activity in the small
585 intestine, it is support of the sensitivity of starch hydrolysis for amylase activity. Consistently
586 good empirical relationships are found between rumen and small intestinal starch digestibility
587 (Nocek & Tamminga, 1991; Offner & Sauvant, 2004; Moharrery et al., 2014) but these
588 relationships do not take into account details such as amylase activity. Aiming to account for such
589 details (Mills et al., 1999) warranted the mechanistic approach adopted in the present study.

590 Other studies suggest that, instead of amylase activity, it is the capacity to absorb glucose
591 from hydrolysed starch that limits small intestinal glucose recovery by the cow (Cant et al., 1999).
592 Although Cant et al. (1999) did not infuse starch into the small intestine, they indicate that the
593 ability to upregulate glucose transport capacity was restricted to a level below that for other
594 species, where uptake capacity remains marginally in excess of requirement. Common TMR diets
595 that led to differences in amount of starch available in the small intestine did not regulate glucose
596 transporters (SGLT1 and GLUT2) (Lohrenz et al., 2011), and these authors postulated that it is
597 questionable if providing large amounts of rumen undegradable starch can modulate glucose
598 absorptive capacity in dairy cows under practical conditions. Therefore, absorption of glucose
599 during passage along the small intestine may limit postruminal digestive efficiency, irrespective of
600 enzymatic capacity. In a simulation of small intestinal starch digestion and glucose uptake,
601 Huntington (1997) suggested that the primary limitation to starch disappearance was enzymatic
602 capacity, unless more than 3 kg/d starch passes the duodenum, at which point SGLT1 transport
603 capacity becomes limiting. This 3 kg/d is below the maximum value of observed duodenal starch
604 flows used in reviews on small intestinal starch digestion (e.g., Reynolds et al., 2014). For
605 example, Moharrery et al. (2014) conclude that no limitation occurs up to 2 kg/d of duodenal

606 starch inflow. However, Offner & Sauvant (2004) do report a plateau for small intestinal starch
607 digestion with increasing inflow.

608 The previous simulation studies of Huntington (1997) and Cant et al. (1999) successfully
609 adopted a basal level of aggregation that considered luminal glucose uptake as a single stage
610 process. Hence, apparent affinity constants for glucose uptake by SGLT1 were uncorrected for the
611 effects of the UWL surrounding the microvilli. There are restrictions imposed with this
612 methodology since the potential for diffusion across the UWL and oligosaccharide hydrolysis
613 cannot be accounted for as independent limits to glucose uptake. The reductionist approach taken
614 in the present model allows for estimation of rate limiting processes and provides a tool for the
615 identification of novel methods of nutritional manipulation aimed at elevating the recovery of
616 duodenal starch as absorbed glucose. After careful evaluation, the model has potential for practical
617 application, especially when considered as part of a feed evaluation system incorporating a
618 mechanistic rumen model predicted duodenal starch inflow (e.g. Dijkstra et al., 1992). The ability
619 to vary starch hydrolysis rate and account explicitly for changing levels of pancreatic secretion are
620 particular strengths of the model, ultimately resulting in a predictive tool for starch digestion
621 rather than just glucose uptake under controlled conditions. Wider application of the model may
622 necessitate a more complete description of hydrolysis rates of different starches following
623 different degrees of processing (Theurer, 1986).

624 The behavioural analysis has shown that at high duodenal starch flows, most of the
625 parameters representing maximum rates of transport or hydrolysis are limiting to some extent.
626 This goes some way to explaining the apparent contradiction between published studies
627 identifying single rate limiting factors. This also corresponds with indications in literature that
628 there are limitations to small intestinal starch digestion (e.g. Offner & Sauvant, 2004). For

629 example, when the model is used to simulate the 3 levels of abomasal starch infusion by
630 Kreikemeier et al. (1991), the effect of luminal starch flow can be clearly seen as a determinant of
631 the pattern of starch appearance and disappearance along the small intestine. Figure 9a shows the
632 elevation in luminal oligosaccharide flow distal from 50% of small intestinal length with the two
633 highest rates of starch infusion (250, 375 mmol/h). The decline in oligosaccharide flow is much
634 slower for the highest levels of starch infusion. As a result, the oligosaccharide load through the
635 ileum is increased disproportionately. Since oligosaccharidase is relatively active in the ileal
636 section, substantial quantities of glucose are produced in the UWL. The flow of glucose in the
637 duodenum and jejunum section at low levels of starch entering the duodenum is small (Fig 9b).
638 However, the ability of the brush border SGLT1 in the distal section of the ileum is limited. At
639 high levels of starch entering the duodenum, a significant amount of glucose formed from
640 oligosaccharides in the UWL diffuses to the lumen where it is lost via passage to the large
641 intestine (up to 25 mmol/h, representing up to 10% of starch infused into the abomasum; Figure
642 9b). This type of analysis can be useful in diet evaluation, to prevent unnecessary outflow of
643 glucose, oligosaccharides or starch to the large intestine, and to evaluate how appropriate actions
644 can be taken such as reducing rumen escape starch, decreasing passage rate or increase the
645 intrinsic digestibility of starch through feed processing. Recently, Gilbert et al. (2015) found
646 evidence for a limitation of starch hydrolysis by maltase activity in the brush border of the small
647 intestine of milk-fed calves. Similar limitations may occur in the adult ruminant. Simulation
648 results in Figure 9 reiterate the need to coordinate enzymatic starch hydrolysis and intestinal
649 glucose uptake, to maximise the efficiency of postruminal starch digestion for a given diet.

650 The present mechanistic model of small intestinal starch digestion and trans-epithelial
651 glucose transport in dairy cattle provides insight into control points for maximising postruminal

652 carbohydrate digestive efficiency, and may be a useful tool for starch evaluation. Some limitations
653 and weaknesses of the model that affect its prediction accuracy and precision have been identified.
654 Firstly, data on metabolism at the brush border in high-producing dairy cattle were largely lacking,
655 and this part of the model was partly parameterized using in vitro data and data from non-
656 ruminants. The relative lack of data at brush border level indicates that this is an area where
657 further research is required to improve the model. Moreover, microbial fermentation of starch in
658 the small intestine was not represented. Secondly, the total contribution of glucose to enterocyte
659 metabolism appears to be overestimated, especially in the presence of competing substrates such
660 as glutamine. Thirdly, the model underestimated the contribution of paracellular diffusion of
661 glucose from the digesta to the blood, particularly at high concentrations of luminal glucose.
662 Finally, the model does not account for the detailed effects of starch particulate size on the rate of
663 starch hydrolysis, unless this is represented by the estimate for the digestion turnover time as an
664 input to the model.

665 The model predictions, combined with observations from the literature, would suggest a
666 series of rate limiting steps is involved with small intestinal starch metabolism. The ideal situation
667 is one where pancreatic fluid secretion is plentiful, the concentration of amylase in the fluid is
668 high, oligosaccharide transport and hydrolysis are both non-limiting, with SGLT1 activity in
669 excess of requirement. This situation will only exist for low levels of duodenal starch delivery.
670 Beyond this, limitations will be imposed. The model has shown that these limits can be localised
671 within a section of the small intestine (Figure 9a). However, spatial asynchrony between
672 carbohydrase activity and transport capacity can amplify the effect of such localised limitations
673 (Figure 9b) to affect the overall efficiency of small intestinal starch hydrolysis and glucose
674 absorption.

675

676

CONCLUSIONS

677

678 This research suggests that there is no single factor that has the potential to limit starch
679 digestion in the small intestine under conditions evaluated. The balance between small and large
680 intestinal starch digestion is determined by a complex interplay of hydrolysis and transport
681 processes, that themselves are dependent on diet and physiological state. The mechanistic
682 approach adopted in the model provides for a more comprehensive quantitative understanding
683 than has previously been available. Examination of the processes, ranging from duodenal starch
684 delivery to glucose transport out of the enterocyte and NSC flow to the large intestine, as one
685 complete system is particularly beneficial. Such an approach avoids undue emphasis on certain
686 potentially rate limiting steps and the oversight of others. The mechanistic model of small
687 intestinal starch digestion and transepithelial glucose transport is a useful tool for feed evaluation
688 especially where substantial quantities of starch flow undegraded from the rumen. The model also
689 provides an insight into control points for maximising postruminal carbohydrate digestive
690 efficiency.

691

692

ACKNOWLEDGEMENTS

693

694 The authors thank Dr C. K. Reynolds, University of Reading, for providing helpful
695 suggestions and other intellectual input during the conduct of this research. Funding was obtained
696 in part from the Canada Research Chairs program (National Science and Engineering Council,

697 Ottawa). The contribution of André Bannink and Jan Dijkstra was partly funded by the Product
698 Board Animal Feed (Zoetermeer, the Netherlands).

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Table 1. Parameter values*†.

Transaction	M_{ijk}	$M_{NSC_{jk}}$	v_{ijk}^*	v_{ijk}^{**}	$v_{ijk}^{(o)}$	θ_{ijk}	$k_{ijk}^{(d)}$	Y_{ijk}	T_{SI}^*
GbGe	0.048	0.075	4.88E-5	1.22E-5					
GbGu							0.0242		
GeGb	0.023	0.075	4.88E-5	1.22E-5		5.0			
GlGu							0.0242		
GuGb							0.0242		
GuGe	1.0E-4	0.075	2.92E-4	7.3E-7		5.0			
GuGl							0.0242		
OIOu							0.0089		
OuGu	0.0045		6.0E-5		6.0	0.19		0.9	
OuOl							0.0089		
PfAl	0.21		22000	5000		3.0			
PfPp								11.9	
PpAl	2.5		8.0E-4	3.0E-4		15.0			
SIOl	0.0216		6.0E-5		6.9	0.6			11.8

940 *See Appendix for explanation of notation

941 † N.B. Parameters relating to small intestinal physiology are considered as input parameters,
 942 subject to variation dependent on animal type, and are not displayed here but are discussed in the
 943 text.

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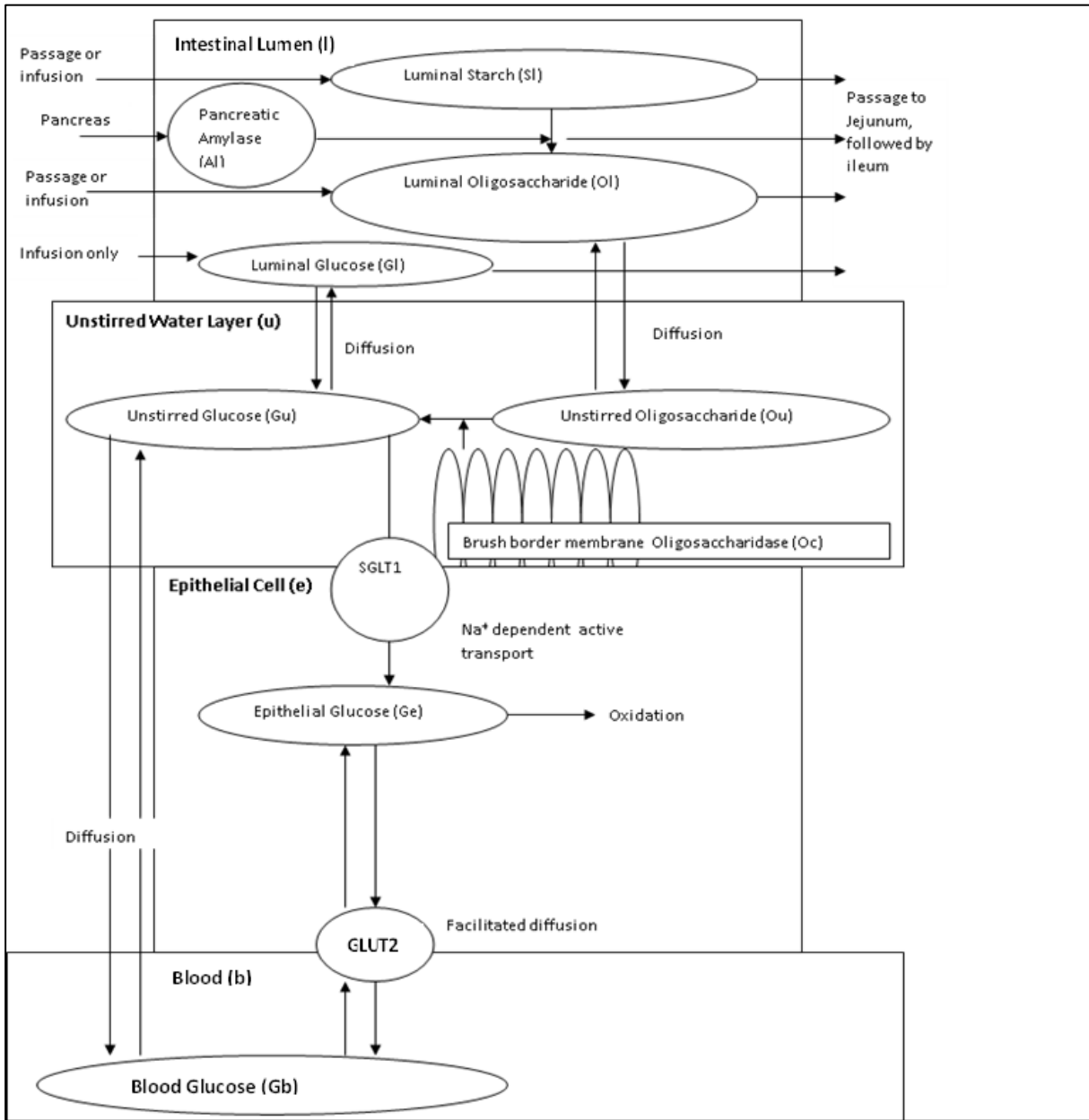
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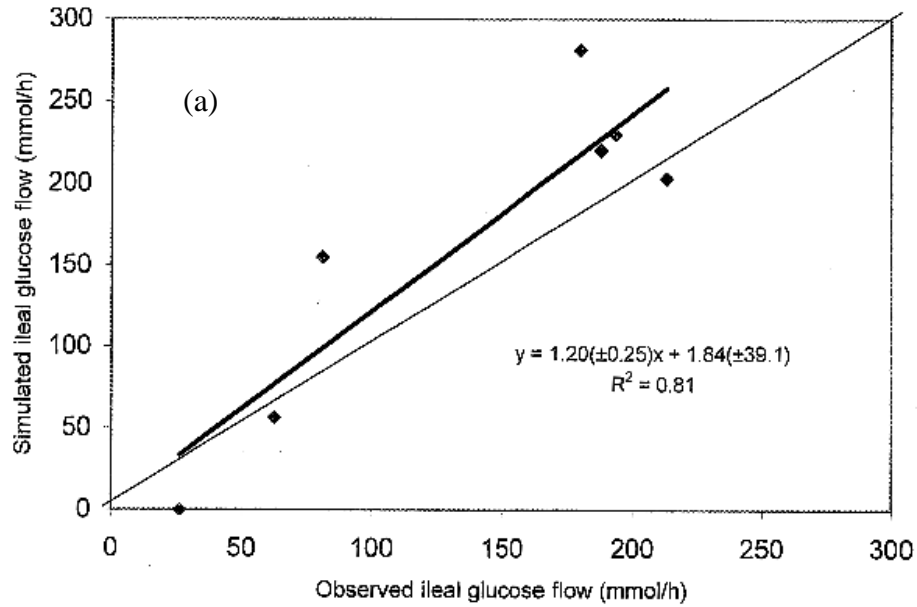
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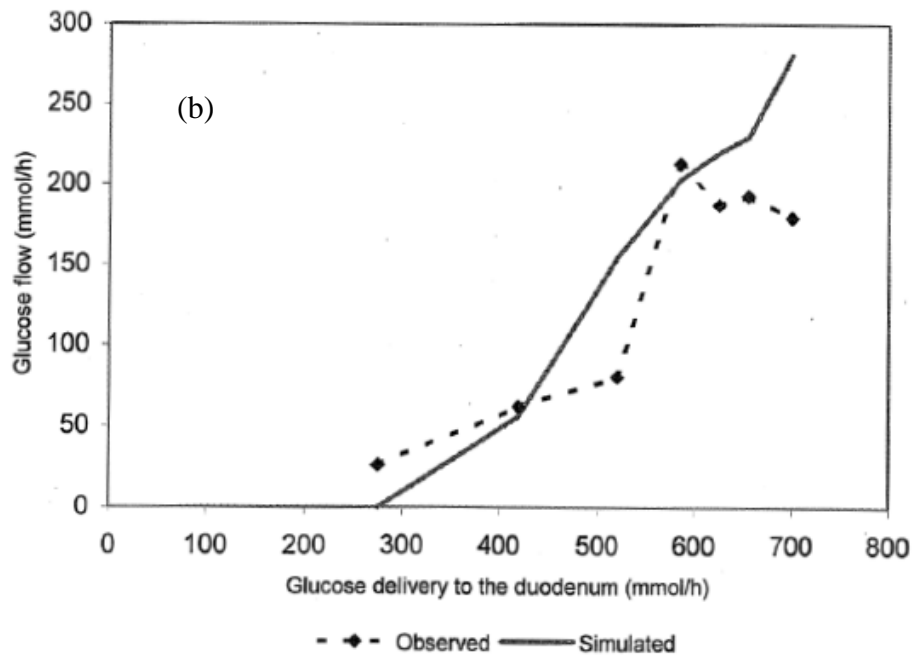


963 **Figure 1.** Diagrammatic representation of one sub-section in the small intestine model.

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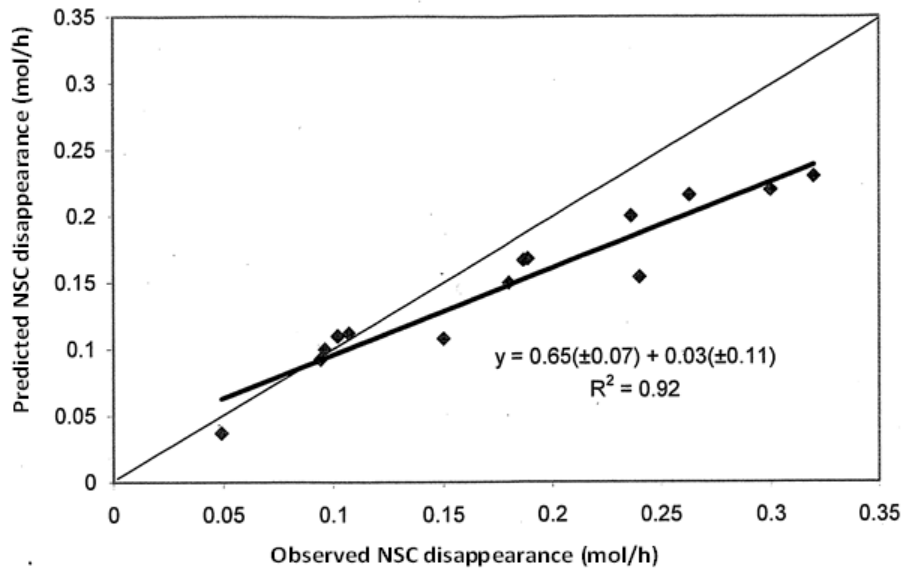
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967 **Figure 2.** Observed and simulated ileal glucose flow in dairy heifers infused with glucose at the
 968 duodenum. (a) regression [root Mean Square Prediction Error (RMSPE) = 38.6% of observed
 969 mean, bias of prediction = 20.7% of MSPE, error due to regression = 39.3%, disturbance
 970 proportion = 40.0%], (b) flows.

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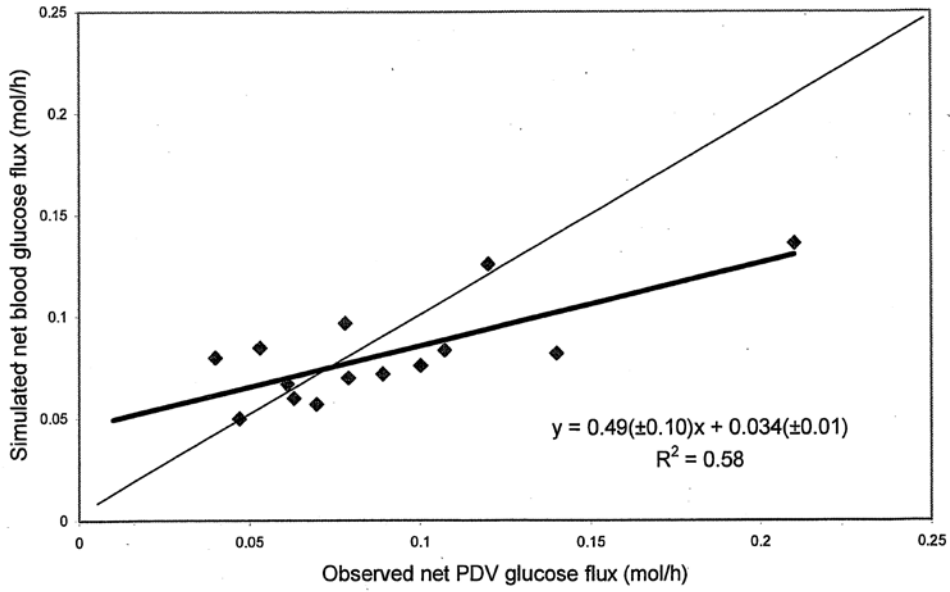
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 973 **Figure 3.** Regression of observed and predicted non-structural carbohydrate (NSC) disappearance
 974 in the small intestine. root Mean Square Prediction Error (RMSPE)= 25.4%, bias of prediction =
 975 45.2% of MSPE, error due to regression = 25.1%, disturbance proportion = 29.7%.

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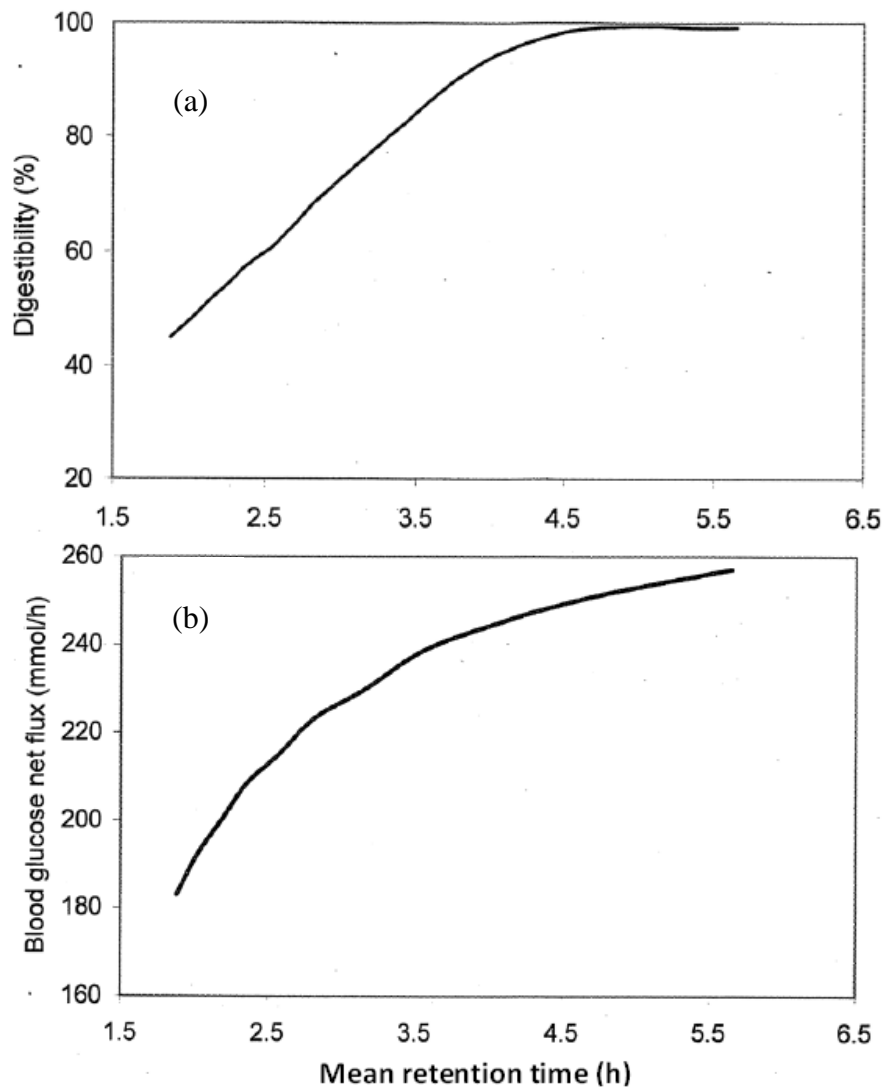
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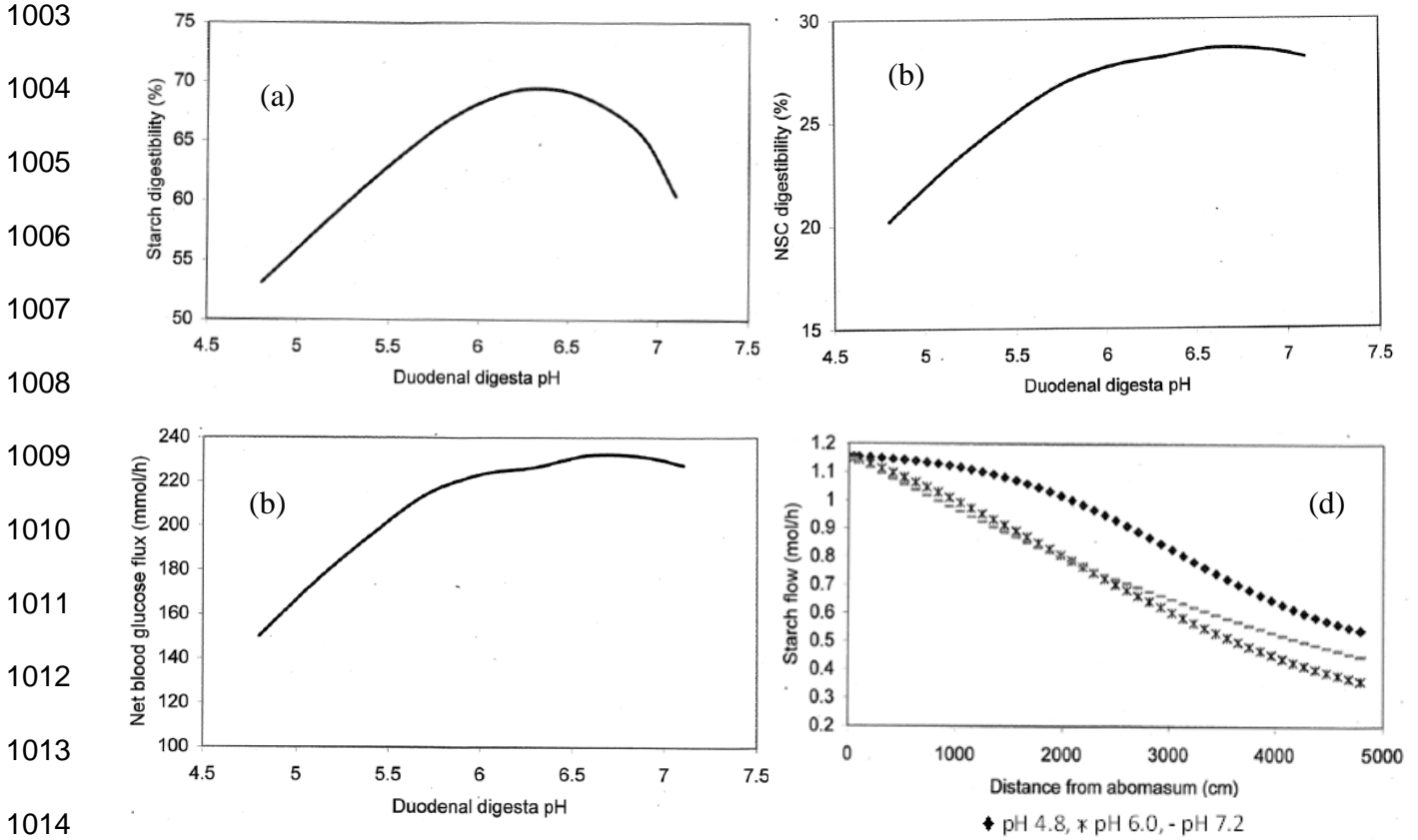


980
981 **Figure 4.** A comparison of observed net portal drained visceral (PDV) glucose flux and simulated
982 net glucose release from the small intestinal epithelial tissue.
983



1000 **Figure 5.** Small intestinal starch flow. (a) simulated relationship between starch digestibility and
1001 mean retention time (MRT), (b) simulated net blood glucose flux and MRT.

1002



1015 **Figure 6.** Luminal pH. (a) simulated starch digestibility over a range of duodenal digesta pH, (b)
 1016 simulated non-structural carbohydrate (NSC) digestibility over a range of duodenal digesta pH, (c)
 1017 duodenal digesta pH and net blood glucose flux, (d) duodenal digesta pH and starch flow down the
 1018 small intestine.

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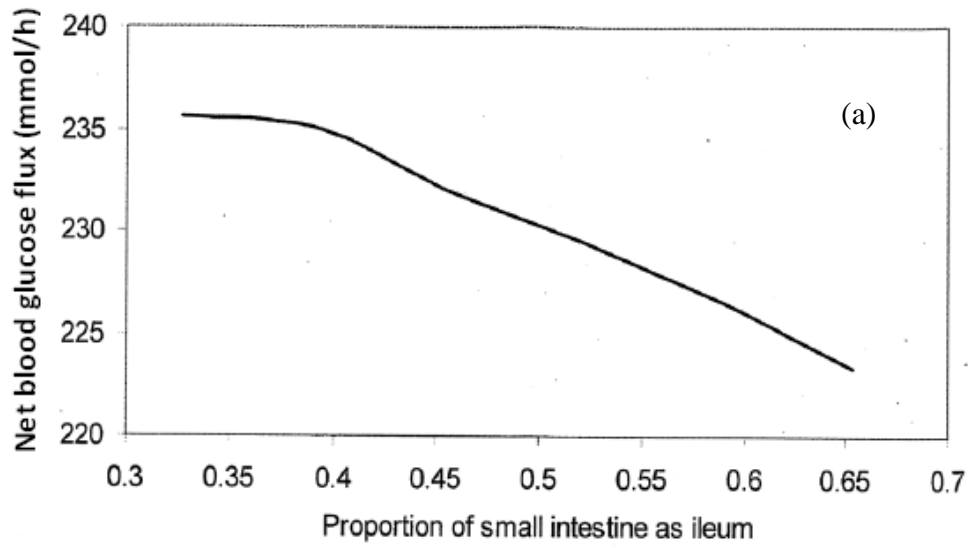
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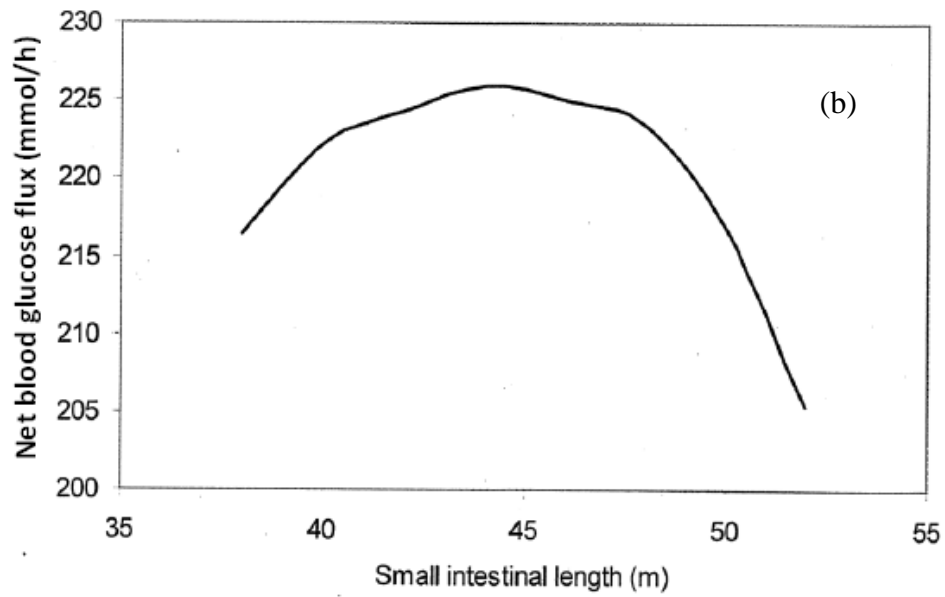
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1038 **Figure 7.** Intestinal physiology. (a) net blood glucose flux and proportion of small intestine

1039 defined as ileum, (b) small intestinal length and net blood glucose flux.

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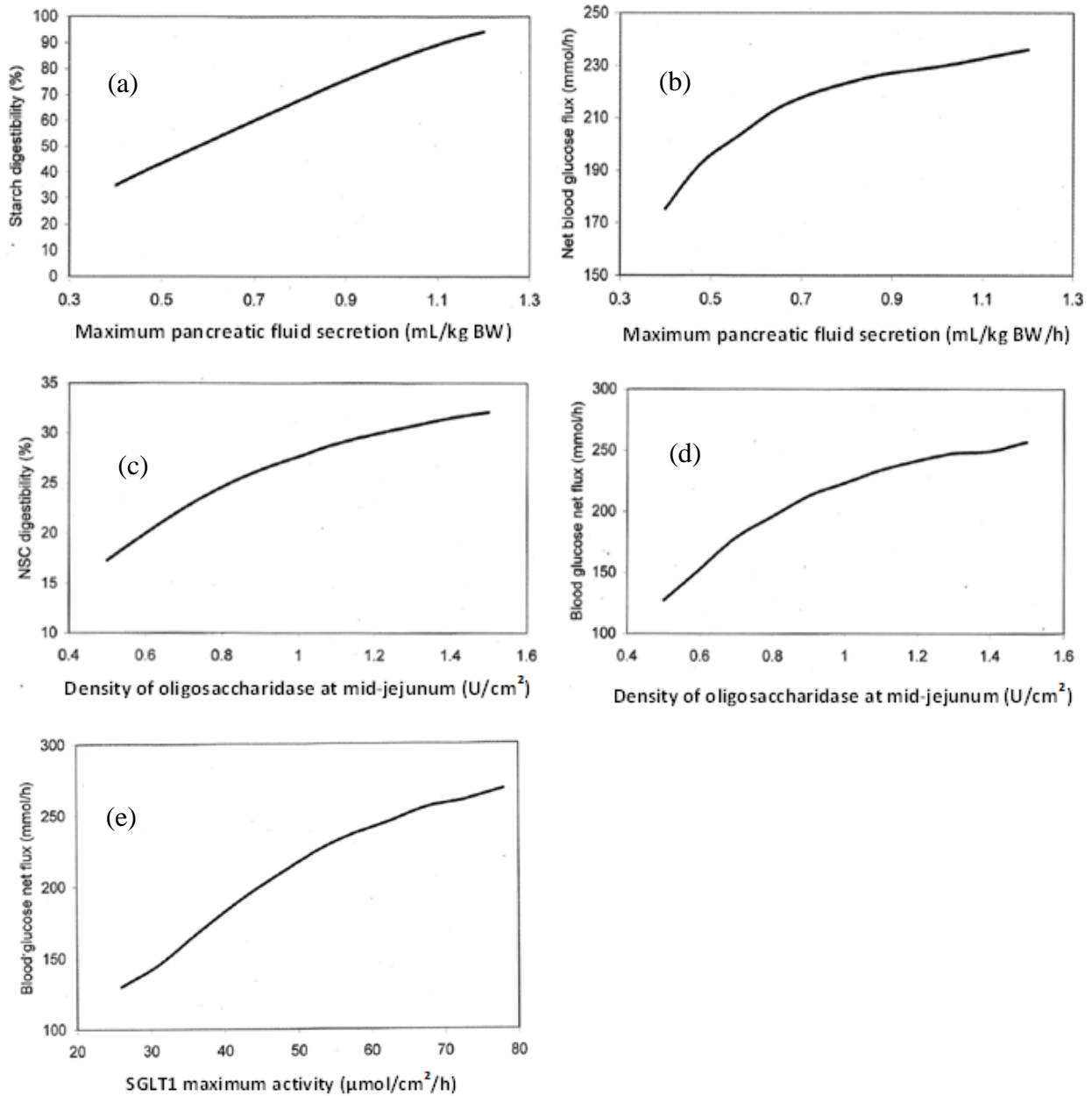
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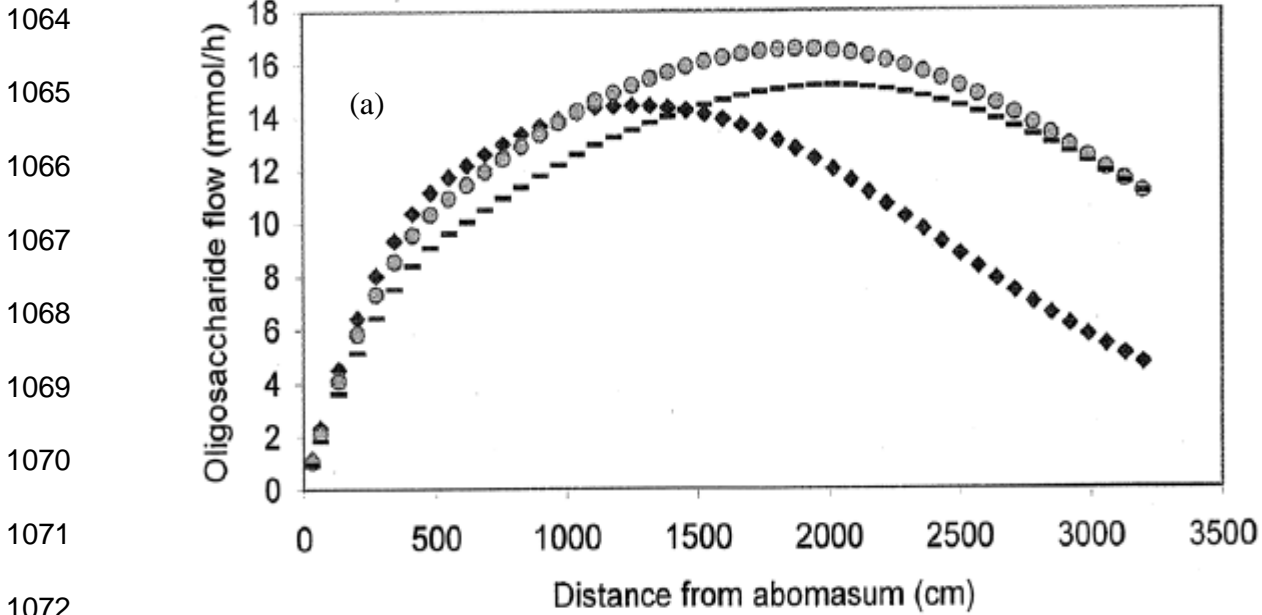
1059 **Figure 8.** Pancreatic amylase and SGLT1 activity. (a) maximum rate of pancreatic fluid secretion

1060 and starch digestibility, (b) maximum rate of pancreatic fluid secretion and net blood glucose flux,

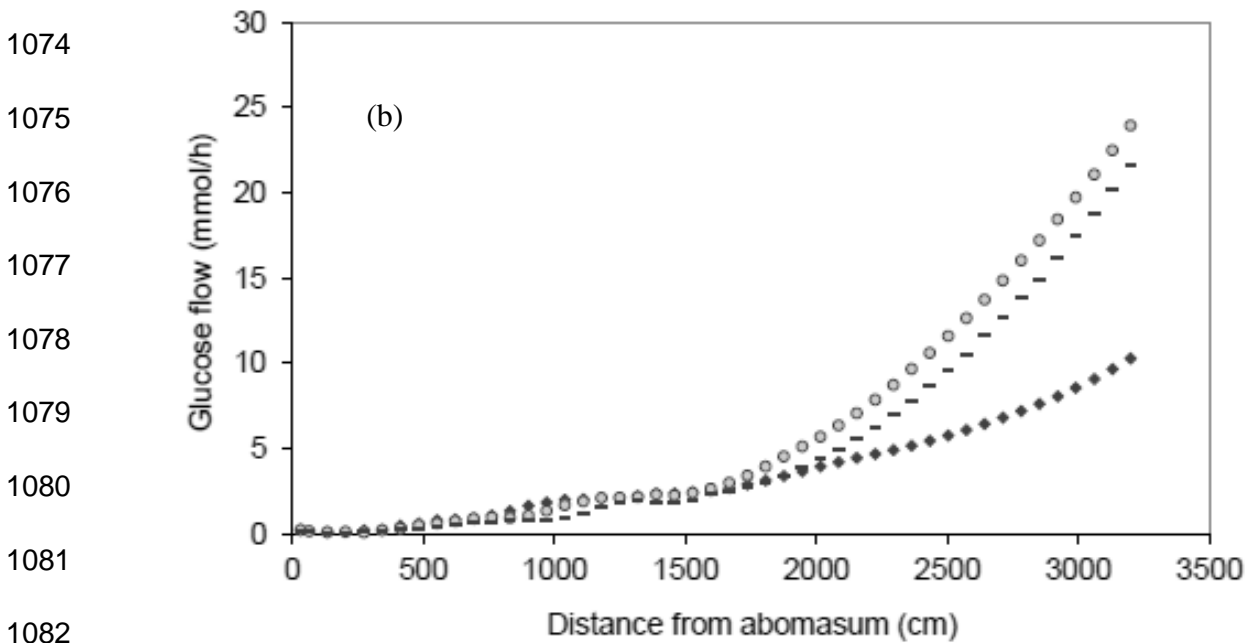
1061 (c) non-structural carbohydrate (NSC) digestibility and maximum oligosaccharidase activity, (d)

1062 response of blood glucose net flux to density of oligosaccharidase at the unstirred water layer, (e)

1063 simulated net blood glucose flux and SGLT1 maximum activity.



◆ 125 mmol/h ○ 250 mmol/h - 375 mmol/h



◆ 125 mmol/h ○ 250mmol/h - 375 mmol/h

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1085 **Figure 9.** Simulated flow during passage along the small intestine in steers infused with

1086 increasing levels of starch into the abomasum. (a) oligosaccharide, (b) glucose.

1087 **Appendix: Mathematical Model Statements***

1088

1089 **Duodenal Lumen**

1090 *Amylase in Lumen, Q_{Al} mol*

1091 *Concentration:*

1092
$$C_{Al} = Q_{Al} / V_{Lu} \quad (1.1)$$

1093 *Inputs:*

1094
$$P_{Al,PfAl} = Y_{Pp,PfPp} Y_{Al,PpAl} U_{Pf,PfAl} \quad (1.2)$$

1095 *Outputs:*

1096
$$U_{Al,AlEx} = k_p Q_{Al} \quad (1.3)$$

1097 *Differential equation:*

1098
$$\frac{dQ_{Al}}{dt} = P_{Al,PfAl} - U_{Al,AlEx} \quad (1.4)$$

1099 *Auxiliary equations:*

$$U_{Pf,PfAl} = \left(v_{Al,PfAl}^{**} + \left(v_{Al,PfAl}^* - v_{Al,Pf}^{**} \right) / \left(1 + \left(M_{Al,PfAl} / MEIM \right)^{\theta_{Al,PfAl}} \right) \right) BW \quad (1.5)$$

1100
$$Y_{Al,PpAl} = v_{Al,PpAl}^{**} + \left(v_{Al,PpAl}^* - v_{Al,PpAl}^{**} \right) / \left(1 + \left(SI_{Flow} / M_{Al,PpAl} \right)^{\theta_{Al,PpAl}} \right) \quad (1.6)$$

$$v_{pH,SIoI} = \exp \left(-\theta_{SI,SIoI} \left(pH_{Lu} - v_{pH,SIoI}^{(o)} \right)^2 \right) \quad (2.5)$$

1101 *Starch in the Lumen, Q_{Sl} mol*

1102 *Concentration:*

1103
$$C_{Sl} = Q_{Sl} / V_{Lu} \quad (2.1)$$

1104 *Input:*

1105
$$D_{Sl} = \text{driving variable} \quad (2.2)$$

1106 *Outputs:*

$$U_{Sl,SlEx} = k_p Q_{Sl} \quad (2.3)$$

1107

$$U_{Sl,SlOl} = (v_{Sl,SlOl} v_{pH,SlOl} Q_{Al}) / \left(1 + \left(M_{Sl,SlOl} \left(T_{Sl} / T_{Sl}^* \right) \right) / (C_{Sl}) \right) \quad (2.4)$$

1108 *Auxiliary equation:*

$$v_{pH,SlOl} = \exp \left(-\theta_{Sl,SlOl} \left(pH_{Lu} - v_{pH,SlOl}^{(o)} \right)^2 \right) \quad (2.5)$$

1110 *Differential equation:*

$$\frac{dQ_{Sl}}{dt} = D_{Sl} - U_{Sl,SlEx} - U_{Sl,SlOl} \quad (2.6)$$

1112 *Glucose in Lumen, Q_{Gl} mol*

1113 *Concentration:*

$$C_{Gl} = Q_{Gl} / V_{Lu} \quad (3.1)$$

1115 *Inputs:*

$$D_{Gl} = \text{driving variable} \quad (3.2)$$

1116

$$P_{Gl,GuGl} = U_{Gu,GuGl} \quad (3.3)$$

1117 *Outputs:*

$$U_{Gl,GlGu} = k_{Gl,GlGu}^{(d)} S_{Lu} C_{Gl} \quad (3.4)$$

1118

$$U_{Gl,GlEx} = k_p Q_{Gl} \quad (3.5)$$

1119 *Differential equation:*

$$\frac{dQ_{Gl}}{dt} = D_{Gl} + P_{Gl,GuGl} - U_{Gl,GlGu} - U_{Gl,GlEx} \quad (3.6)$$

1121 *Oligosaccharide in Lumen, Q_{Ol} mol*

1122 *Concentration:*

$$C_{Ol} = Q_{Ol} / V_{Lu} \quad (4.1)$$

1124 *Inputs:*

$$D_{Ol} = \text{driving variable} \quad (4.2)$$

1125 $P_{Ol,SlOl} = U_{Sl,SlOl} \quad (4.3)$

$$P_{Ol,OuOl} = U_{Ou,OuOl} \quad (4.4)$$

1126 *Outputs:*

1127 $U_{Ol,OlOu} = S_{Lu} k_{Ol,OlOu}^{(d)} C_{Ol} \quad (4.5)$

$$U_{Ol,OlEx} = k_p Q_{Ol} \quad (4.6)$$

1128 *Differential equation:*

1129 $\frac{dQ_{Ol}}{dt} = D_{Ol} + P_{Ol,SlOl} + P_{Ol,OuOl} - U_{Ol,OlOu} - U_{Ol,OlEx} \quad (4.7)$

1130 **Duodenal Unstirred Water Layer (UWL)**

1131 *Glucose in UWL, Q_{Gu} mol*

1132 *Concentration:*

1133 $C_{Gu} = Q_{Gu} / V_{Wl} \quad (5.1)$

1134 *Inputs:*

$$P_{Gu,GlGu} = U_{Gl,GlGu} \quad (5.2)$$

1135 $P_{Gu,OuGu} = Y_{Gu,OuGu} U_{Ou,OuGu} \quad (5.3)$

$$P_{Gu,GbGu} = U_{Gb,GbGu} \quad (5.4)$$

1136 *Outputs:*

$$U_{Gu,GuGl} = S_{Lu} k_{Gu,GuGl}^{(d)} C_{Gu} \quad (5.5)$$

1137 $U_{Gu,GuGb} = S_{Lu} k_{Gu,GuGb}^{(d)} C_{Gu} \quad (5.6)$

$$U_{Gu,GuGe} = (v_{Gu,GuGe} S_{Lu}) / (1 + M_{Gu,GuGe} / C_{Gu}) \quad (5.7)$$

1138 *Differential equation:*

1139 $\frac{dQ_{Gu}}{dt} = P_{Gu,GlGu} + P_{Gu,OuGu} + P_{Gu,GbGu} - U_{Gu,GuGl} - U_{Gu,GuGb} - U_{Gu,GuGe} \quad (5.8)$

1140 *Auxillary equations:*

$$v_{Gu,GuGe} = v_{Gu,GuGe}^{**} + \left((v_{Gu,GuGe}^* - v_{Gu,GuGe}^{**}) / \left(1 + (M_{NSC,GuGe} / NSC_{Flow})^{\theta_{Gu,GuGe}} \right) \right) \quad (5.9)$$

1141 $NSC_{Flow} = k_p (Q_{Gl} + Q_{Ol} + Q_{Sl}) \quad (5.10)$

$$SI_{Flow} = k_p Q_{Sl} \quad (5.11)$$

1142 *Oligosaccharide in UWL, Q_{Ou} mol*

1143 *Concentration:*

1144 $C_{Ou} = Q_{Ou} / V_{Wl} \quad (6.1)$

1145 *Input:*

1146 $P_{Ou,OlOu} = U_{Ol,OlOu} \quad (6.2)$

1147 *Outputs:*

1148 $U_{Ou,OuGu} = (v_{Ou,GuDu} S_{Lu}) / (1 + M_{Ou,OuGu} / C_{Ou}) \quad (6.3)$

$$U_{Ou,OuOl} = S_{Lu} k_{Ou,OuOl}^{(d)} C_{Ou} \quad (6.4)$$

1149 *Differential equation:*

1150 $\frac{dQ_{Ou}}{dt} = P_{Ou,OlOu} - U_{Ou,OuGu} - U_{Ou,OuOl} \quad (6.5)$

1151 *Auxiliary equations:*

$$v_{Ou,GuDu} = v_{Ou,OuGu}^* Q_{Oc} v_{pH,OuGu} \quad (6.6)$$

1152 $v_{pH,OuGu} = \exp\left(-\theta_{Ou,OuGu} \left(pH_{Wl} - v_{pH,OuGu}^{(o)}\right)^2\right) \quad (6.7)$

1153 **Duodenal Enterocyte**

1154 *Glucose in enterocyte, Q_{Ge} mol*

1155 *Concentration:*

1156 $C_{Ge} = Q_{Ge} / V_{En} \quad (7.1)$

1157 *Inputs:*

1158 $P_{Ge,GuGe} = U_{Gu,GuGe} \quad (7.2)$

$$P_{Ge,GbGe} = U_{Gb,GbGe} \quad (7.3)$$

1159 *Outputs:*

1160
$$U_{Ge,GeGb} = [(v_{Ge,GeGb}^{**} S_{Lu}) + ((v_{Ge,GeGb}^* - v_{Ge,GeGb}^{**}) S_{Lu}) / (1 + (M_{NSC,GeGb} / NSC_{Flow})^{\theta_{Ge,GeGb}})] / (1 + M_{Ge,GeGb} / C_{Ge}) \quad (7.4)$$

$$U_{Ge,GeOx} = R_{Ge,GuGe} U_{Gu,GuGe} + R_{Gu,AuAe} U_{Au,AuAe} + 0.0043 S_{Lu} \quad (7.5)$$

1161 *Differential equation:*

1162
$$\frac{dQ_{Ge}}{dt} = P_{Ge,GuGe} + P_{Ge,GbGe} - U_{Ge,GeGb} - U_{Ge,GeOx} \quad (7.6)$$

1163 **Blood**

1164 *Blood Glucose, Q_{Gb} mol*

$$U_{Gb,GbGu} = k_{Gb,GbGu}^{(d)} C_{Gb} S_{Lu} \quad (8.1)$$

1165
$$U_{Gb,GbGe} = [(v_{Ge,GeGb}^{**} S_{Lu}) + ((v_{Ge,GeGb}^* - v_{Ge,GeGb}^{**}) S_{Lu}) / (1 + (M_{NSC,GeGb} / NSC_{Flow})^{\theta_{Ge,GeGb}})] / (1 + M_{Gb,GbGe} / C_{Gb}) \quad (8.2)$$

1166 * Displayed is model code representing one small intestinal sub-section, the first proximal

1167 duodenum. The model is repeated for each subsection hereafter until the terminal ileum is reached.

1168

Table A1. Definition of symbols for entities and processes represented in the model

Symbol	Entity
Ae	Amino acids in enterocyte
Al	Amylase in lumen
Au	Amino acids in UWL
BW	Liveweight
Du	Duodenum
Ex	Exit to next subsection
Gb	Glucose in blood
Ge	Glucose in enterocyte
Gl	Glucose in lumen
Gu	Glucose in UWL
Il	Ileum
Je	Jejunum
Lu	Lumen
MEIM	Metabolisable energy intake (MEI) to MEI at maintenance ratio
NSC	Non-structural carbohydrate
Oc	Oligosaccharidase
Ol	Oligosaccharide in lumen
Ou	Oligosaccharide in UWL
Pf	Pancreatic fluid
Pp	Pancreatic protein

Sl	Starch in lumen
Wl	Water layer (unstirred)

1169

Table A2. General notation used in the model

Symbol	Entity
C_i	Concentration of i , mol/L
D_i	Entry of i from passage or infusion (driving variable), mol/h
$k_{i,jk}^{(d)}$	Diffusion constant for i in $j - k$ transaction, /h
k_p	Fractional rate of passage constant, /h
$M_{i,jk}$	Michaelis-Menten affinity constant with respect to i for $j - k$ transaction, mol/L
$P_{i,jk}$	Rate of production of i in $j - k$ transaction, mol/h
pH_i	pH of region i
Q_i	Quantity of i , mol
$R_{i,jk}$	Requirement for i in $j - k$ transaction, mol/mol
S_i	Surface area of i , cm ²
T_i	Digestion turnover time of substrate i , h
T_i^*	Maximum level with respect to i , h
$\theta_{i,jk}$	Steepness parameter associated with i for $j - k$ transaction
$U_{i,jk}$	Rate of utilization of i by $j - k$ transaction, mol /h
V_i	Effective volume of i , L or kg
$v_{i,jk}$	Velocity for $j - k$ transaction with respect to i , mol/h
$v_{i,jk}^*$	Maximum level with respect to i for $j - k$ transaction, mol/h
$v_{i,jk}^{**}$	Minimum level with respect to i for $j - k$ transaction, mol/h

$v_{i,jk}^{(o)}$ Optimum level of i for $j - k$ transaction

$Y_{i,jk}$ Yield of i in $j - k$ transaction, mol/mol

1170